The \textit{Traf2DNxBCL2-tg} Mouse Model of Chronic Lymphocytic Leukemia/Small Lymphocytic Lymphoma Recapitulates the Biased IGHV Gene Usage, Stereotypy, and Antigen-Specific HCDR3 Selection of Its Human Counterpart

Gema Perez-Chacon\textsuperscript{1,2}† and Juan M. Zapata\textsuperscript{1,2*}

\textsuperscript{1} Instituto de Investigaciones Biomédicas “Alberto Sols”, CSIC-UAM, Madrid, Spain, \textsuperscript{2} Instituto de Investigación Hospital Universitario La Paz (IDIPAZ), Madrid, Spain

Chronic lymphocytic leukemia (CLL)/Small lymphocytic lymphoma (SLL) is a heterogeneous disease consisting of at least two separate subtypes, based on the mutation status of the immunoglobulin heavy chain variable gene (IGHV) sequence. Exposure to antigens seems to play a role in malignant transformation and in the selection and expansion of more aggressive CLL clones. Furthermore, a biased usage of particular IGHV gene subgroups and the existence of stereotyped B-cell receptors (BCRs) are distinctive characteristics of human CLL. We have previously described that \textit{Traf2DN/BCL2} double-transgenic (tg, +/+ ) mice develop CLL/SLL with high incidence with aging. In this model, TNF-Receptor Associated Factor (TRAF)-2 deficiency cooperates with B cell lymphoma (BCL)-2 in promoting CLL/SLL in mice by specifically enforcing marginal zone (MZ) B cell differentiation and rendering B cells independent of BAFF for survival. In this report, we have performed the sequencing of the IGHV-D-J rearrangements of B cell clones from the \textit{Traf2DN/BCL2-tg+/+} mice with CLL/SLL. The results indicate that these mice develop oligoclonal and monoclonal B cell expansions. Allotransplantation of the oligoclonal populations into immunodeficient mice resulted in the preferential expansion of one of the parental clones. The analysis of the IGHV sequences indicated that 15% were mutated (M) and 85% unmutated (UM). Furthermore, while the \textit{Traf2DN/BCL2-tg−/−} (wild-type), \textit{−/+} (BCL2 single-tg) and \textit{+/+} (Traf2DN/DN single-tg) littermates showed the expression of various IGHV gene subgroups, the CLL/SLL expanded clones from the \textit{Traf2DN/BCL2-tg+/+} (double-transgenic) mice showed a more restricted IGHV gene subgroup usage and an overrepresentation of particular IGHV genes. In addition, the HCDR3-encoded protein sequence indicates the existence of stereotyped immunoglobulin (Ig) in the BCRs and strong similarities with BCR recognizing autoantigens and pathogen-associated antigens. Altogether, these results
highlight the remarkable similarities between the CLL/SLL developed by the Traf2DN/BCL2-tg+/- mice and its human counterpart.

Keywords: TRAF2, BCL2, chronic lymphocytic leukemia, CLL, small lymphocytic lymphoma, IGHV, BCR stereotypy

INTRODUCTION

Chronic lymphocytic leukemia (CLL) is the most common adult leukemia in the Western world. CLL and small lymphocytic lymphoma (SLL) are two manifestations of the same B cell neoplasia and are characterized by the accumulation of slowly proliferating CD5+CD23+ B lymphocytes with dysregulated apoptosis (1–3).

It is well established that CLL is a heterogeneous disease consisting of at least two separate subtypes, based on phenotypic and clinical behavior. Approximately 55% of CLL patients have mutated (M) immunoglobulin heavy chain variable (IGHV) genes (4–6), which have a better prognosis than patients with unmutated (UM)- IGHV genes (6–8). According to phenotypic analysis and gene expression profiling both M- and UM-CLL are antigen-experienced B cells (9, 10). The differences in clinical outcome and biological characteristics between CLL patients with M- and UM-IGHV genes could be related to distinct differences in mutation incidence and distribution reflecting specific underlying mutagenic mechanisms between these two groups (11). As a result, M- and UM-CLL show differences in BCR reactivity profile (12) and signaling (13).

In addition, CLL can also be classified according to the expression of stereotyped HCDR3, which are found in a 41% of CLL patients (14, 15). Indeed, the remarkable similarity of HCDR3 regions within sets of patients strongly supports the notion that B cell receptor (BCR) recognition of particular antigens is a driving force in clonal selection, expansion and evolution in CLL [reviewed in (16, 17)].

CLL cells are low proliferating cells, mostly quiescent and with dysregulated apoptosis. Only a small percentage are proliferating cells, which makes difficult their expansion in immunodeficient mice. Besides, human CLL cell xenotransplantation may result in the expansion of B cell clones that do not recapitulate the IGHV-D-J rearrangements of the parental clone [reviewed in (18)]. In addition, it has been shown that donor T cells are required to support CLL implantation (19). However, proliferating T cells could result in a graft versus host disease that hampers the utility of CLL xenotransplanted mice.

Mouse models of CLL are useful tools for the study of CLL etiology and as preclinical platforms for new drug testing. Several CLL mouse models are currently available, which recapitulate key aspects of the human disease [reviewed in (20)]. However, a majority of these CLL mouse models, including the profusely studied Eµ-T Cell Leukemia-1 (Eµ-TCL-1)-tg mice [reviewed in (21)], only produce UM-CLL clones, thus implying that M-CLL etiology is not properly represented in these mice.

We previously described that B cell-specific Traf2DN/BCL2-double-tg (+/+ ) mice develop CLL/SLL with high incidence (22, 23). In this mouse model, expression of Traf2DN causes the depletion of endogenous TRAF2, resulting in unbridled BAFF signaling and constitutive NFκB2 activation, causing the expansion of marginal zone (MZ) B cells (24). BCL2 overexpression, which is a CLL trademark (25), would provide MZ B cells with non-redundant and complementary protection against apoptosis that predisposes these cells to CLL/SLL.

In this report we show that the CLL/SLL arising in the Traf2DN/BCL2-tg+/- mice consists of expanded M- and UM-CLL/SLL clones. Expanded clones show a biased IGHV gene usage, stereotypy and express HCDR3 that are similar to those recognizing autoantigens and pathogen antigens, thus closely resembling human CLL.

MATERIALS AND METHODS

Transgenic Mice

Lymphocyte-specific Traf2DN-tg expressing a 1D4-epitope–tagged TRAF2 deletion mutant lacking the N-terminal 240 amino acids (AA) encompassing the RING and zinc finger domains (Traf2DN) (26) and B cell-specific BCL2-tg mice mimicking the t(14;18)(q32;21) translocation involving BCL2 and Igh found in human follicular lymphoma (27) have been previously described. Traf2DN-tg (FVB/N) and BCL2-tg (BALB/c) heterozygous mice were bred to produce F1 litters with progeny of the four possible genotypes ((wild-type-/-; Traf2DN-tg (single-positive, */+); BCL2-tg (single-positive, */+); and Traf2DN/BCL2 (double-positive, */++)) expressed on FVB/N x BALB/c mixed background as previously described (22). Analysis of the transgenic mouse genotypes was performed by polymerase chain reaction (PCR) using primers specific for Traf2 (F) 5'-GACCAGAGCAAGATTGAGGC-3' and (R) 5'-GCACATAGGAATTTTGCC-3' and BCL2 (F) 5'-TTAGAGGTCTTTACCTGGCCTG-3' and (R) 5'-ACCTTAGAGACGGTGACC-3'. The animal protocols were approved by the Bioethics Committee of the hosting institution. Mice showing symptoms of distress and pain (heavy breath, weight loss, distended belly, respiratory distress, lethargy, etc) were euthanized. All transgenic mice in the study were heterozygotes for each transgene.

Isolation of Mononuclear Cells

Spleens, lymph nodes and blood from Traf2DN/BCL2-tg mice of the different genotypes were collected and mononuclear cells were bred to produce F1 litters with progeny of the four possible genotypes ((wild-type-/-; Traf2DN-tg (single-positive, */+); BCL2-tg (single-positive, */+); and Traf2DN/BCL2 (double-positive, */++)) expressed on FVB/N x BALB/c mixed background as previously described (22). Analysis of the transgenic mouse genotypes was performed by polymerase chain reaction (PCR) using primers specific for Traf2 (F) 5'-GACCAGAGCAAGATTGAGGC-3' and (R) 5'-GCACATAGGAATTTTGCC-3' and BCL2 (F) 5'-TTAGAGGTCTTTACCTGGCCTG-3' and (R) 5'-ACCTTAGAGACGGTGACC-3'. The animal protocols were approved by the Bioethics Committee of the hosting institution. Mice showing symptoms of distress and pain (heavy breath, weight loss, distended belly, respiratory distress, lethargy, etc) were euthanized. All transgenic mice in the study were heterozygotes for each transgene.

Flow Cytometry Analysis

Mononuclear cells were incubated with 50 μg/ml human γ-globulin for 10 minutes at 4°C. Then, 10⁶ cells were incubated with a combination of FITC-, PE-, or APC-conjugated antibodies against mouse CD45R/B220, CD21, CD23, IgM, IgD, CD5, and CD43
(all from BD Biosciences). After 30 minutes of incubation at 4°C, cells were washed with PBS and analyzed by flow cytometry in a FACScanto II cytometer and the FACSDiVa 6.1.2 (BD Biosciences) flow cytometry analysis software.

**Immunohistochemistry**

Tissues and organs from transgenic mice were fixed in 10% formalin (Sigma-Aldrich) or in Bouin’s solution (Sigma-Aldrich) for bone marrow analysis and embedded in paraffin. Tissue sections (5 μm) were deparaffinized and then stained with hematoxylin and eosin, dehydrated, and mounted in DPX (Fluka). Blood smears were stained with Wright-Giemsa (Sigma-Aldrich).

**Immunoglobulin IGHV-D-J Sequence Analysis**

Tissues and cells from Traf2DNxBCL2 mice representative of all different genotypic combinations (+/+; +/-; -/+ and -/-) were extracted and the plasmids were extracted using the Wizard® Minipreps DNA Purification System (Promega). Products were sequenced in a capillary sequencer by GATC Biosciences using the IMGT/V-QUEST analysis tool (29). Since our mice are FVB/N x BALB/c F1 hybrids and the GL of these strains are found in the IMGT repertoire IG database, to discriminate between SS and SHM, we considered unmutated (UM). Isoelectric point (pI) of HCDR3 was calculated using the Compute pI/Mw tool (ExPaSy Bioinformatics Resource Portal, http://web.expasy.org/compute_pi/). HCDR3 analysis was carried out comparing the sequence in the protein BLAST database.

**Statistics**

IBM SPSS statistics v.26 (SPSS, Chicago, IL) and Graph Pad Prism 5 were used for statistical analysis. Statistical significance for HCDR3 length and isoelectric point (pI) was determined using the t-Student test. Pearson Chi-Square and likelihood ratio tests with Monte Carlo correction were applied for assessing the significance of the IGHV-D-J subgroups distribution among genotypes. Proportion test was used to determine the significance of IGHV gene expression frequency.

**RESULTS**

**Characteristics of BCRs Expressed by CLL/SLL B Cells From the Traf2DNxBCL2-tg+/+ Mice**

As stated above, Traf2DNxBCL2-tg+/+ mice develop CLL/SLL with high incidence as they age (22). In most mice, SLL arises first, involving splenomegaly, lymphadenopathy and infiltration of different tissues and organs, later progressing to CLL (22, 23). An example of the histology features of the bone marrow, blood, spleen and lung of a representative Traf2DNxBCL2-tg+/+ mouse with CLL/SLL is shown in Figure 1A. In addition, flow cytometry analysis of the B cell populations in this mouse (Figure 1B) identified two B cell populations. One majority population, with larger cells based on their forward scatter (FSC) profile and expressing low levels of B220, IgD, CD21 and CD23 and high levels of IgM (Figure 1B), corresponds to the CLL/SLL expanded cells (blue). These cells were CD43high and CD5low or null (not shown). The other population (FSCsmall) is composed of seemingly normal B2 cells expressing B220high, IgMlow, IgDhigh, CD21high, and CD23middle (green) (Figure 1B). These cells were CD43null and CD5null (not shown). The expanded CLL/SLL population is found in blood, spleen and in pleural effusion (Figure 1B).

To ascertain the BCR characteristics of these CLL/SLL cells, we have analyzed the sequences of the HCDR3 of these mice. Table 1 shows the HCDR3 characteristics and frequency of the expanded clones isolated from Traf2DNxBCL2-tg+/+ mice with CLL/SLL. Based on the HCDR3 sequences, these mice develop oligoclonal (mice: #13, #16, #65, #72 and #74) and monoclonal (mice: #29, #40, #45, #50 and #51) B cell expansions (Table 1). Interestingly, when spleen and blood were compared, we have examples of mice with identical expanded clones in both sources (mice: #16, #40 and #65) but also a mouse (#55) with different clones in spleen and blood.
In addition, blood lymphocytes or splenocytes (40-60 x 10^6) from representative Traf2DNxBCL2-tg+/+ mice (#55, #72 and #74) were allotransplanted into immunodeficient SCID/NOD mice. Animals were euthanized when they develop any sign of illness (distended belly, respiratory distress, lethargy, etc). As shown in Table 1, only one of the expanded CLL/SLL clones found in each of the parental mice was selectively expanded in the immunodeficient allotransplanted mice.

**IGHV-D-J Subgroups and Gene Usage by the Expanded CLL/SLL Cells From the Traf2DNxBCL2-tg+/+ Mice**

CLL clones from human CLL patients express mostly IgM and have a biased usage of IGHV genes compared to normal B cells [reviewed in (30)]. These characteristics are also shared by the Eµ-TCL-1-tg (28, 31) and the MDR^-/- and miR-15a/16-1^-/- (32) mouse models of CLL. Thus, to ascertain whether B cells from the Traf2DNxBCL2-tg^+/+ mice with CLL/SLL have similar characteristics, we have analyzed the Ig isotypes and IGHV-D-J rearrangements expressed by B cells from these mice and compared them with those found in mice representing all other genotype combinations. For this purpose, Traf2DN-tg (FVB/N background) and BCL2-tg (BALB/c background) mice were crossed to produce F1 litters with mice harboring the different transgene combinations, Traf2DNxBCL2-tg^-/-, +/-, -/+ and +/+ . The analyses were performed when the Traf2DNxBCL2-tg^+/+ mice developed CLL/SLL, using for comparison age- and sex-matched mice representing all genotypes and genders. As shown in Figure 2A, retrotranscription and amplification of the mRNAs encoding for IgM, IgG and IgA shows that B cells from Traf2DNxBCL2-tg^+/+ mice developed CLL/SLL, using for comparison age- and sex-matched mice representing all genotypes and genders. As shown in **Figure 2A**, retrotranscription and amplification of the mRNAs encoding for IgM, IgG and IgA shows that B cells from Traf2DNxBCL2-tg^+/+ mice with CLL/SLL almost exclusively express IgM, while all three Igs (M, G and A) mRNAs could be readily detected in B cells from representative mice of all the other genotypes. The relative expression of IgM, IgG and IgA in the Traf2DNxBCL2-tg with the different genotypes and in the expanded CLL/SLL clones is shown in **Figure 2B**.

Next, we studied whether a biased IGHV gene usage was also a feature of the CLL/SLL developed by the Traf2DNxBCL2-tg^+/+ mice. The IGHV, IGHD and IGHJ genes and the HCDR3 sequences expressed in the expanded B cell clones from the
| Animal no. | Age (months) | Sex | Tissue | IGHV Family | IMGT IGHV gene | IGHD gene | IGHJ gene | SHM % | SHM status | Frequency % | HCDR3 length | HCDR3 | pI |
|-----------|--------------|-----|--------|-------------|----------------|-----------|-----------|------|-----------|-------------|--------------|-------|-----|
| 13        | 13           | F   | Spleen | VH14        | IGHV14-2*02 F | VHS7m.a2psi.88 | D2 | DSP2.9 | JH4  | 1.1 UM | 5/13 38 | GRDDGGYYAMYD | 12 | 3.93 |
| 16        | 12           | M   | Spleen | VH1        | IGHV1-85*01 F | VHJ558.88.194 | D1 | DFL16.1 | JH3  | 0.35 UM | 6/10 60 | ASYAFAY | 7 | 5.57 |
| 29    | 20           | F   | Spleen | VH1        | IGHV1-80*01 F | VHJ558.83.189 | D2 | DSP2.2 | JH4  | 2.1 M | 8/10 80 | ASPSYDYYYYAMYD | 15 | 3.56 |
| 45        | 20           | F   | Spleen | VH1        | IGHV1-9*01 F | VHJ558.b9 | D2 | DSP2.2 | JH4  | 0.7 UM | 8/10 80 | ASRGTYDFFFF | 11 | 4.03 |
| 50        | 18           | M   | Spleen | VH1        | IGHV1-74*01 F | VHJ558.b9 | D4 | DQ52   | JH4  | 0.7 UM | 10/10 100 | AGSTRMIMIMMYD | 13 | 4.21 |
| 55 P      | 16           | F   | Spleen | VH1        | IGHV1-74*01 F | VHJ558.b9 | D2 | DSP2.4 | JH4  | 0.0 UM | 10/10 100 | AGSTRMIMIMMYD | 10 | 3.56 |
| 65 F1     | -            | M   | Spleen | VH1        | IGHV1-77*01 F | VHJ558.80.186 | D1 | DFL16.1e | JH2  | 1.4 UM | 5/10 50 | AVVYDGGYAMYD | 15 | 3.56 |
| 72 P      | 15           | F   | Spleen | VH1        | IGHV14-2*02 F | VHS7m.a2psi.88 | D2 | DSP2.9 | JH4  | 0.7 UM | 4/14 29 | GRDDGGYYAMYD | 12 | 3.93 |
| 72 F1     | -            | F   | Spleen | VH1        | IGHV1S130*01 F | Unknown | D2 | DSP2.2 | JH2  | 0.35 UM | 4/14 29 | ARRNWDFEDY | 11 | 4.56 |
| 74 P      | 15           | F   | Spleen | VH1        | IGHV1S130*01 F | Unknown | D1 | DFL16.1 | JH2  | 0.0 UM | 6/10 60 | ASGPDFDY | 8 | 3.56 |

Table shows the mouse ID number, the tissue source of the mRNA sample, the age and the sex of the mice. The immunoglobulin IGHV, IGHD and IGHJ subgroups and genes found recombined in each CLL B cell clone are indicated, according to IMGT/V-QUEST and Vbase2 analysis tools. SHM status indicates whether the IGHV region is unmutated (UM; ≤2% difference from the GL sequence) or mutated (M; >2% difference from the GL sequence) after correcting for SSP as described in Supplementary Materials and Methods. The frequency and % of occurrence of the B cell clones isolated from the indicated tissues of each mouse is also shown. All clones encoded a productive Ig and the HCDR3 sequence is also provided. Basic (red) and acid (green) AAs are highlighted and stereotyped HCDR3 are shown in bold. The length and isoelectric point (pI) of the HCDR3 sequence are shown. Additional information pertaining to these CLL clones is provided in Supplementary Table 4.
mice with -/-, +/-, -/+ genotypes and the whole list of clones
In addition, similar information from the
the analysis of the
IGHV14 (10%) subgroup genes. A similar picture emerges from
followed by IGHV2 (13%), IGHV5 (13%), IGHV10 (10%) and
subgroups, with a larger representation of IGHV1 (37%)
Figure 1
frontiers in Immunology | www.frontiersin.org April 2021 | Volume 12 | Article 6276026
[45x249]mRNA from the indicated mice was extracted and retrotranscribed into
cDNAs from the various genotypes (−/−, n = 5; −/+ , n = 3; +/−, n = 4; +/+ ,
spleen, n = 8, blood, n = 3) were analyzed as indicated in A and the resulting
bands were quantified. The results show the percentage of each Ig (M, G and
A) found in mice of the various genotypes (average ± SD). Statistical
significance: **p < 0.005; ***p < 0.0005).

FIGURE 2 | Immunoglobulin subtypes found in Traf2DNxBCL2-tg mice representative of all different genotypic combinations (−/−; −/+; +/− and +/+).
(A) mRNA from the indicated mice was extracted and retrotranscribed into
cDNAs using random primers. Then, PCR was performed using specific
primers for the IGHV-D-J region of IgM, IgG or IgA, as described in Materials
and Methods. The amplified PCR fragments were analyzed in 2% agarose
gels and staining with SYBR Safe and UV light. (B) IgM, IgG and IgA cDNAs
from mice of the various genotypes (−/−, −/+ , +/−, +/+ ) were analyzed in the same way as in A and the results show the percentage of each Ig (M, G and
A) found in mice of the various genotypes (average ± SD). Statistical
significance: **p < 0.005; ***p < 0.0005).

Traf2DNxBCL2-tg mice with CLL/SLL are shown in Table 1. In addition, similar information from the Traf2DNxBCL2-tg mice with −/−, −/+ , +/− genotypes and the whole list of clones isolated from the Traf2DNxBCL2-tg mice are shown in
Supplementary Tables 1-4, respectively. A schematic representation of the IGHV, IGHD and IGHJ subgroups expressed in B cells from mice of each genotype and those
used by the expanded B cell clones of the Traf2DNxBCL2-tg mice with CLL/SLL are shown in Figure 3 and Supplementary Figure 1. B cell clones isolated from the Traf2DNxBCL2-tg (wild-type) mice demonstrated the usage of various IGHV subgroups, with a larger representation of IGHV1 (37%) followed by IGHV2 (13%), IGHV5 (13%), IGHV10 (10%) and
IGHV14 (10%) subgroup genes. A similar picture emerges from the analysis of the Traf2DNxBCL2-tg mice of −/− and −/+ genotypes, also showing the usage of various IGHV subgroups, with IGHV1 being the most prominently used in all of them, consistent with

The larger representation of this subgroup in the murine GL
reertoire (33). In contrast, IGHV1 (51%), IGHV5 (19%),
IGHV14 (14%) and IGHV3 (9%) are the subgroup genes most
conspicuously used by B cells from the Traf2DNxBCL2-tg mice and also by the expanded CLL/SLL clones (Figure 3 and
Supplementary Table 5). Interestingly, the IGHV subgroup expression frequency observed in other CLL mouse models is
seemingly different to that of the Traf2DNxBCL2-tg expanded
CLL/SLL clones, with the exception of IGHV1, which is the most expressed IGHV gene subgroup in all of them (Supplementary Table 5) (see discussion).

To determine whether this restricted IGHV gene subgroup usage was a distinctive characteristic of the Traf2DNxBCL2-tg CLL/SLL model or was instead a general feature that could also be found in the expanded B cell clones of other types of B cell malignancies, we analyzed the IGHV subgroup repertoire used
by the mature non-Hodgkin lymphomas (NHL) developed by the Traf3xBCL2-tg mice (34). These mice are also F1 hybrids of FBV/N x BALB/c background, and therefore are genetically equivalent to the Traf2DNxBCL2-tg mice. As shown in
Supplementary Figure 2, in B cells from the Traf3xBCL2-tg mice the IGHV subgroup usage (IGHV1 (43%), IGHV5 (13%), IGHV14 (11.8%), IGHV2 (10.5%) is similar to that found in the wild-type (Traf2DNxBCL2-tg) mice. The expanded B cell clones from the Traf3xBCL2-tg mice that have developed post-germinal center (GC) NHL malignancies used more frequently genes from the IGHV1 subgroup genes (56%), similar to the Traf2DNxBCL2-tg CLL/SLL clones, but the usage of IGHV5 (12.5%) and IGHV14 (6.3%) genes is much
reduced compared to the latter. Expression of IGHV2 genes is also found in the Traf3xBCL2-tg mice, while it is absent in the Traf2DNxBCL2-tg mice.

Regarding the usage of the IGHD genes, B cell clones of all genotypes preferentially used IGHD2 subgroup gene members (Figure 3 and Supplementary Figure 1) and no statistical
significance was observed in the IGHD subgroup distribution among the various Traf2DNxBCL2 genotypes (p = 0.275;
LR = 0.327). In contrast, there is a favored usage of the IGHD4 gene by the Traf2DNxBCL2-tg mice (all clones, 58%; expanded CLL clones, 63%) compared to the mice with the other Traf2DNxBCL2 genotype combinations (p = 0.024;
LR = 0.024) (Figure 3) and also compared to the average IGHD4 usage in mice (21.5%) (35).
Next, we assessed whether Traf2DNxBCL2-tg CLL/SLL clones show any preferential usage of particular IGHV genes similar to what has been described in human CLL [reviewed in
(30)] and the Eμ-TCL-1-tg mice (28). Indeed, as shown in Table 2,
we observed that 3 genes are overrepresented in the Traf2DNxBCL2-tg CLL/SLL clones compared to the B cells from mice of all other genotypes and in the Traf3xBCL2-tg
mice. Thus, VH7183.a47.76 (IGHV5) is found in 25% of the 
Traf2
DNx
BCL2
-tg+/+ CLL/SLL clones. It is found recombined to a 
wide variety of IGHD and IGHJ genes producing distinct HCDR3 
sequences (Table 1). The expression of this gene is also 
overrepresented in B cell clones from the Traf2DNxBCL2-tg-/+ 
(18%) and the Traf2DNxBCL2-tg-/+ (14.3%) mice compared to the 
1.86% rearrangement frequency found in the BALB/c strain (33). 
Proportion test shows that these differences are statistically 
significant (P>0.0001) (Table 2). The expression of these gene in 
the Traf3xBCL2-tg-/+ B cell lymphoma clones is not significantly 
different to the expression in the normal population. Another 
gene, VHJ558.b9 (IGHV1) is found in 15% of the Traf2DNxBCL2-
tg-/+ CLL/SLL clones (13.3% of all Traf2DNxBCL2-tg-/+ B cell 
clones), a statistically significant difference (P<0.0001) compared 
to the 1.18% rearrangement frequency found in the C57BL/6 
strain (33). It is also overrepresented in the Traf3xBCL2-tg-/+ B 
cell clones but not in the expanded B cell lymphoma clones. 
Finally, VHSM7.a2psi.88 (IGHV14) is rarely or not found in 
BALB/c and C57BL/6 strains (33), but it was found in 15% of 
the expanded CLL/SLL clones and in 8.9% of all Traf2DNxBCL2-
tg-/+ B cell clones (P<0.0001) (Table 2). Interestingly, this 
VHSM7.a2psi.88 was found recombined to DSP2.9 and IGHJ4 
genes in expanded clones from 3 different Traf2DNxBCL2-tg-/+ 
mice with CLL/SLL, producing an identical HCDR3 sequence 
(see below).

Altogether, these results suggest that a preferential usage of 
IGHV subgroups and genes by the expanded CLL/SLL clones 
from the Traf2DNxBCL2-tg-/+ mice is occurring, similar to what 
has been previously observed in human CLL patients and in the 
Eµ-TCL-1-tg mouse model of CLL.

FIGURE 3 | Analysis of the IGHV, IGHD and IGHJ genes subgroup usage in Traf2DNxBCL2-tg. Circle diagrams representing the percentage of the IGHV, IGHD and 
IGHJ subgroup usage of mice representative of all different genotypic combinations (-/-; +/-; -/+ and +/+), including those found in the Traf2DNxBCL2-tg-/+ expanded 
CLL/SLL clones are shown.

Perez-Chacon and Zapata Mouse CLL Recapitulates Human CLL
Table 2: A restricted set of IGHV genes predominates in the expanded Traf2DNxBCL2-tg+/+ CLL/SLL clones.

| Mouse tg-line | Genotype | IGHV5 VH7183.a47,76 | P | IGHV1 VH558.b9 | P | IGHV14 VHS71.a2psi88 | P |
|---------------|----------|----------------------|---|----------------|---|----------------------|---|
| Traf2DNxBCL2  | +/-      | 4.3                  | 0.12 | 1.4     | 0.86 | nf                   | 0.9 |
|               | +/       | 4                    | 0.48 | 4     | 0.004 | nf                   | 0.87 |
|               | +/-      | 14.3                 | <0.0001 | 6     | 0.014 | nf                   | 0.85 |
|               | +/-      | 18                   | <0.0001 | 13    | <0.0001 | 2     | <0.0001 |
|               | +/-      | 25                   | <0.0001 | 15    | <0.0001 | 15    | <0.0001 |
| Traf3xBCL2    | +/-      | 2.5                  | 0.5 | 8.1     | <0.0001 | nf                   | 0.69 |
|               | +/-      | 6.2                  | 0.18 | nf     | 0.66 | nf                   | 0.9 |
|               | +/-      | 1.9                  | -    | 1.2     | -     | <0.1                  | -  |

The IGHV subgroup and its frequency (%) in the B cell clones isolated from the Traf2DNxBCL2-tg and Traf3xBCL2-tg mice of the indicated genotypes, as well as the expression frequency (%) of these genes in mice (33) is shown. Statistical significance was calculated using the chi-squared test, and significant results are highlighted in bold. Clones from different tissues of the same mouse are only counted once. Clones found in parental and F1 mice are only counted once. +/- indicated the expanded B cell clones (CLL/SLL in the Traf2DNxBCL2-tg+/+ mice and mature non-Hodgkin lymphoma in the Traf3xBCL2-tg+/+ mice). nf, not found.

Analysis of the IGHV Somatic Hypermutation Status and HCDR3 Features of the CLL/SLL Clones From the Traf2DNxBCL2-tg+/+ Mice

Patients with CLL segregate into two groups based on the number of SHMs in the rearranged IGHV genes of the transformed clones. Approximately 55% of CLL patients have transformed B cells with mutations in IGHV genes (M) (4–6). The rest of the patients have UM IGHV CLL clones, which correlates with poor disease prognosis (6–8).

To determine the frequency of M vs. UM IGHV regions in the expanded CLL/SLL clones of the Traf2DNxBCL2-tg+/+ mice, we first compared the IGHV sequence of the transformed clones with the available GL sequences stored in the IMGT1 repertoire IG database (mostly based on the C57BL/6, with scattered presence of 129/Sv and BALB/c lines GL sequences (33)). The results obtained using the IMGT/V-QUEST analysis tool showed that many sequences have considerable variations with their respective IGHV genes. A similar result was obtained when the IGHV sequences from the B cell clones isolated from mice with the other genotypes (+/-, +/-, +/+ and all +/-) was compared. Since our mice are FVB/N X BALB/c hybrids and because it has been shown that the IGHV GL repertoire is conserved in most of the corresponding IGHV gene from different mouse lines (33, 36), these variations might reflect the reported differences among strains and the absence of IGHV GL sequences from the FVB/N mouse line in the IMGT repertoire IG database.

To determine whether some of these variations with the GL sequences may be the result of SSSs, we have performed a clustal W sequence comparison of the IGHV region of Traf2DNxBCL2-tg B cell clones with identical IGHV alleles, irrespective of their genotype. Indeed, we have observed the existence of nucleotide mismatches compared to the IMGT referenced IGHV gene that are conserved in most of the corresponding IGHV gene from different Traf2DNxBCL2-tg individuals and genotypes. Since SHM randomly introduces any of the 4 nucleotides in a given spot, a mismatch of the same nucleotide in the same position of several identical alleles compared to the GL sequence strongly suggest the existence of a polymorphism. The number of SSSs found in the IGHV rearranged sequences from the Traf2DNxBCL2-tg B cell clones ranges from 1 (0.35% of the IGHV sequence) to 24 (9.375%), averaging a 2.54% SSSs, consistent with previously reported IGHV GL differences among mouse strains (36). The criteria for discriminating between SSP and SHM and examples of the IGHV gene sequence comparisons are shown in supplementary materials and methods and Supplementary Figure 3, respectively.

The estimation of SHM events (%) according to the criteria described above found in the expanded CLL/SLL B cell clones is shown in Table 1. In addition, the percentage of similarity of the IGHV region of the analyzed B cell clones with the GL and the SSS and SHM estimated events for the B cell clones from all the Traf2DNxBCL2-tg genotypes is shown in Supplementary Tables 1–4. A standard 2% difference with the GL was applied to categorize UM or M IGHV clones (4, 28, 37). As shown in Supplementary Table 6, the CLL/SLL expanded clones from the Traf2DNxBCL2-tg+/+ mice were 85% UM and 15% M (identical clones found in a different tissue of the same mouse as well as identical clones found in parental and allotransplanted F1 mice were only counted once). Similar UM and M percentages were found in Traf2DNxBCL2-tg-/- (78.6% UM vs. 21.4% M), and Traf2DNxBCL2-tg+/- (all clones) (80% UM vs. 20% M) mice and a larger population of UM B cell clones was also found in Traf2DNxBCL2-tg+/- (60% UM vs. 40% M). In contrast, Traf2DNxBCL2-tg+/+ B cell clones are split in half (46% UM vs. 54% M). This result is consistent with the fact that Traf2-deficiency causes the expansion of MZ B cells (24, 38), which are mostly UM (39). In addition, Bcl2 overexpression has been shown to reduce the SHM rate (40).

It has been reported in human CLL patients that UM- and M-CLL clones have a biased usage of IGHV subgroups. Thus, IGHV1 genes predominate in the rearrangements of UM-CLL cells while IGHV3 and IGHV4 genes are more frequently found in M-CLL cells (5, 6, 41). A larger percentage of IGHV1 genes are also found in UM-CLL clones from the Eµ-TCL-1-tg mice (28) and the MDR+/- and mir15a/16-1-/- mice (32). A comparison between the IGHV, IGHD and IGHJ subgroup usage between M- and UM clones from the Traf2DNxBCL2-tg+/+ is shown in Figure 4. Our analyses showed that UM-B cell clones from the Traf2DNxBCL2-tg+/+ used more frequently IGHV1 (46%), IGHV14 (20%), IGHV5 (17%) and IGHV3 (11%), while M-B

Frontiers in Immunology | www.frontiersin.org April 2021 | Volume 12 | Article 627602

8 Perez-Chacon and Zapata Mouse CLL Recapitulates Human CLL
cell clones used IGHV1 (70%), IGHV5 (20%) and IGHV11 (10%). A similar trend was observed in UM- and M-CLL/SLL clones, although the reduced number of M-CLL/SLL clones (n=3) avoid any conclusion. In addition, IGHD2 and IGHJ4 genes were overrepresented in both UM- and M-Traf2DNxBCL2-tg/+ B cell clones and in expanded CLL/SLL clones (Figure 4).

Other HCDR3 features, such as length, charge and AA sequence are intrinsic HCDR3 characteristics. HCDR3 sequences vary in their AA composition, charge and length depending on how the IGHV, IGHD and IGHJ genes recombine (5, 42, 43). A summary of the analysis of these HCDR3 features in the expanded Traf2DNxBCL2-tg/+ B cell clones and of B cells from the other genotypes is shown in Supplementary Table 6.

The analysis of the HCDR3 AA sequence of the expanded Traf2DNxBCL2-tg/+ CLL/SLL clones shows an average length of 11.6 ± 2.9 AAs, which is similar to the HCDR3 length in all other genotypes (-/-, 11.92 ± 2.7; +/-, 10.71 ± 2.6; -/+, 12.11 ± 3.3; +/-, 11.44 ± 2.8) and in accordance with the HCDR3 average length in mice (11.5 ± 1.9 AAs) (44). Moreover, no remarkable differences were observed between the HCDR3 average length of M vs. UM clones in any of the genotypes, including the expanded CLL/SLL clones (Supplementary Table 6). Although long HCDR3 have been proposed to be a characteristic of UM-CLL HCDR3 in humans (5, 42), long HCDR3 were found in both M- and UM-clones from mice of the different genotypes, including the expanded CLL/SLL clones. It is noteworthy that the HCDR3 average length of the Traf2DNxBCL2-tg/+ UM-CLL/SLL clones (11.18 ± 2.9 AAs) is similar to the HCDR3 average length of the UM-CLL clones from the Eµ-TCL-1-tg mice [11.6 ± 2.3 AAs (28) and 10.6 ± 2.4 AAs (31)] and from the IgH-TEµ mice [11.4 ± 2.32 (45)]. The average length of the UM-CLL/SLL clones developed by the MDR/− and miR15a/16-1/− is slightly longer [12.87 ± 2.17 AAs (32)], but compared to that of the Traf2DNxBCL2-tg/+ UM-CLL/SLL clones these differences did not reach statistical significance (P = 0.073).

The average isoelectric point (pI) of the HCDR3 expressed in the analyzed B cell clones from the Traf2DNxBCL2-tg mice of the distinct genotypes is also shown in Supplementary Table 6. The Traf2DNxBCL2-tg/+ expanded CLL/SLL clones have the most acidic HCDR3 (4.54 ± 1.31) compared to that of Traf2DNxBCL2-tg mice with the other genotypes. Indeed, only one Traf2DNxBCL2-tg/+ CLL/SLL clone had a HCDR3 with a basic pI. Aspartic acid and arginine are the most frequently found acidic and basic AAs, respectively. In addition, tyrosine is frequently overrepresented, with some HCDR3 containing as much as 44% of tyrosine, compared to the average 25% frequency for this AA found in the mouse HCDR3 (44). Of note is that the average pI of the HCDR3 of UM-CLL/SLL clones from the Traf2DNxBCL2-tg/+ is significantly more acidic than the average pI of the HCDR3 from the Eµ-TCL-1-tg UM-CLL clones (4.5 ± 1.4 vs. 5.9 ± 1.9) (P=0.02).
Identical HCDR3 Are Expressed in Traf2DNxBCL2-tg+/+ CLL/SLL Clones From Distinct Mice

A distinctive characteristic of human CLL is the expression of structurally identical or highly similar HCDR3 between unrelated individuals producing structurally similar BCRs (14, 15, 46). This occurrence is known as HCDR3 stereotypy and points out toward the role of antigens in the clonal selection and pathogenesis of the disease [reviewed in (17, 30, 47)]. Stereotyped HCDR3 rearrangements account for 41% of human CLL clones (15). BCR stereotypes are also found in CLL clones from the Eμ-TCL-1-tg (28, 31) and from the MDR−/− and mir-15a/16-1−/− (32) mice, among others.

Our results also demonstrate the existence of two identical HCDR3 sequences in the Traf2DNxBCL2-tg+/+ expanded CLL/SLL clones. One is found in mice #50 and #55 (ASGYDYAMYD) and the other in mice #13, #65 and #72 (GRDDGYYYAMYD) (Table 1), accounting for the 25% of the CLL/SLL clones. These Traf2DNxBCL2-tg+/+ CLL/SLL stereotyped HCDR3 sequences are found in UM clones, in agreement with the findings in stereotyped HCDR3 sequences from human CLL (17) and from the above mentioned CLL mouse models (28, 32), that are also found in UM-CLL clones. In addition, it is worth noting that other stereotyped sequences are also found in seemingly not expanded clones from the Traf2DNxBCL2-tg+/+ mice (Supplementary Table 7). These stereotyped sequences might belong to low represented CLL/SLL clones. Including these low represented clones, we found 18 stereotyped HCDR3 account (31% of all Traf2DNxBCL2-tg+/+ B cell clones), of which 16 of them are UM and 2 are M. Interestingly, stereotyped HCDR3 sequences were also found in a few B cell clones from the Traf2DNxBCL2-tg−/+ and −/− mice (Supplementary Table 7). Of note is that some of these stereotyped sequences are found in clones expressing distinct IGHV genes, although always from the same IGHV subgroup. In addition, Supplementary Figure 4 shows a Clustal W analysis of the IGHV regions from representative clones producing identical HCDR3. Even though these IGHV regions are UM, they differ in their SHM pattern.

It is also worth mentioning that HCDR3 expressed in CLL clones from the Eμ-TCL-1-tg (28), the MDR−/− and mir-15a/16-1−/− (32) mice, among others. Our results further demonstrate that the expression of both transgenes is necessary to trigger CLL/SLL in these mice (22). In addition, the comparison of B cell clones isolated from Traf2DNxBCL2-tg−/+ mice with all possible transgene combinations reveal a more restricted set of IGHV subgroup and IGHV gene usage by the expanded Traf2DNxBCL2-tg−/+ (wild-type), −/+ (expressing only TRAF2DN), −/− (expressing only BCL2) and −/+ (expressing both TRAF2DN and BCL2). Our results further demonstrate that monoclonal and oligoclonal B expansions are only observed in the Traf2DNxBCL2-tg+/+ mice that developed CLL/SLL, thus confirming that the expression of both transgenes is necessary to trigger CLL/SLL in these mice (22).

The comparison of the BCR HCDR3 sequences of the expanded Traf2DNxBCL2-tg+/+ CLL/SLL clones with similar sequences found in public databases showed high homology with HCDR3 recognizing autoantigens, such as phosphatidylcholine (82 and 75% homology), cardiolipin (86% homology), dsDNA (80% homology), as well as to pathogen antigens, such as hepatitis C virus E2 protein (81% homology), CMV glycoprotein B (76% homology), Bordetella (75% homology) and Vaccinia protein A3 (80% homology) (Table 3 and Supplementary Table 9). Many of these antigens were already described as being recognized by the BCR of human and mouse CLL clones, including phosphatidylcholine (28), cardiolipin (55, 56), dsDNA (55, 57) and CMV (58, 59), further supporting the role of these types of antigens in the etiology of Traf2DNxBCL2-tg+/+ CLL/SLL.

DISCUSSION
Characteristics of the HCDR3 of the Traf2DNxBCL2-tg Mice

The results presented herein underscore the similarities between the CLL/SLL developed by the Traf2DNxBCL2-tg+/+ mice and the CLL developed by human patients. This includes a biased usage of IGHV genes, the existence of CLL/SLL clones with stereotyped HCDR3 and the expansion of CLL/SLL clones with HCDR3 similar to those recognizing autoantigens and bacteria antigenic determinants (1, 30, 60, 61). Furthermore, compared to other CLL mouse models, such as the Eμ-TCL-1-tg (28) and the MDR−/− and the mir-15a/16-1−/− (32) mice, only generate UM-CLL clones, the CLL/SLL developed by the Traf2DNxBCL2-tg+/+ mice produce both UM- and M-CLL/SLL clones, similar to human CLL, albeit a vast majority of them are UM.

In this report we have compared the IGHV, IGHD and IGHJ gene usage and the HCDR3 sequences expressed in B cells from mice representing all the different genotypes obtained by crossing heterozygous Traf2DN-tg and BCL2-tg mice, that is, Traf2DNxBCL2-tg−/+ (wild-type), −/+ (expressing only TRAF2DN), −/− (expressing only BCL2) and −/+ (expressing both TRAF2DN and BCL2). Our results further demonstrate that monoclonal and oligoclonal B expansions are only observed in the Traf2DNxBCL2-tg+/+ mice that developed CLL/SLL, thus confirming that the expression of both transgenes is necessary to trigger CLL/SLL in these mice (22). In addition, the comparison of B cell clones isolated from Traf2DNxBCL2-tg−/+ mice with all possible transgene combinations reveal a more restricted set of IGHV subgroup and IGHV gene usage by the expanded Traf2DNxBCL2-tg−/+ (wild-type), −/+ (expressing only TRAF2DN), −/− (expressing only BCL2) and −/+ (expressing both TRAF2DN and BCL2). Our results further demonstrate that monoclonal and oligoclonal B expansions are only observed in the Traf2DNxBCL2-tg+/+ mice that developed CLL/SLL, thus confirming that the expression of both transgenes is necessary to trigger CLL/SLL in these mice (22). In addition, the comparison of B cell clones isolated from Traf2DNxBCL2-tg−/+ mice with all possible transgene combinations reveal a more restricted set of IGHV subgroup and IGHV gene usage by the expanded Traf2DNxBCL2-tg−/+ (wild-type), −/+ (expressing only TRAF2DN), −/− (expressing only BCL2) and −/+ (expressing both TRAF2DN and BCL2). Our results further demonstrate that monoclonal and oligoclonal B expansions are only observed in the Traf2DNxBCL2-tg+/+ mice that developed CLL/SLL, thus confirming that the expression of both transgenes is necessary to trigger CLL/SLL in these mice (22).

Putative Antigens Recognized by the HCDR3 Sequences of the CLL/SLL Clones From the Traf2DNxBCL2-tg+/+ by Comparison With Those in Public Databases

As stated above, CLL cells frequently express BCR recognizing autoantigens and pathogen-associated antigens [reviewed in (16, 17)] that are involved in the clonal selection and progression of the disease (31, 50–54).
worth noting that Traf2DNxBCL2-tg+/+ CLL/SLL clones have a MZ origin (24) and that the IGHV2 gene subgroup is underrepresented in transformed B cells of a MZ origin (62, 63).

As described above, the analysis of the IGHV subgroup usage of the expanded Traf2DNxBCL2-tg+/+ CLL/SLL clones indicates a preponderance of certain gene subgroups (IGHV1 > IGHV5 > IGHV14 > IGHV3) (Supplementary Table 5). In contrast, the expression of IGHV subgroups found in other CLL mouse models described in the literature is seemingly different. This includes the Eµ-TCL-1-tg [IGHV1 > IGHV11 = IGHV12 > IGHV4, calculated from (28)], the MDR−/− and miR-15a/16-1−/− [IGHV1 > IGHV11 > IGHV12, calculated from (32)] and the IgH-TEL-tg [IGHV1 = IGHV11, calculated from (45)] mice (Supplementary Table 5). Although IGHV1 is most frequently found in all these mouse models, the preferential use of IGHV5 and IGHV14 by the Traf2DNxBCL2-tg+/+ CLL/SLL clones instead of the use of IGHV11 and IGHV12 seen in the other CLL mouse models might indicate that the CLL/SLL developed by the Traf2DNxBCL2-tg+/+ mice arises from a B cell subset different to that of the other CLL mouse models (see below). These differences might also underlie the reported differences in IGHV-D-J usage by mice of different strains (33, 36). In this regard, differences in IGHV subgroup usage have been also observed in CLL from distinct human populations [(64, 65) and references therein]. However, it is noteworthy that mouse IGHV5 and IGHV11 belong to the IGHV clan whose human counterpart is IGHV3, mouse IGHV1 and IGHV14 share clan with human IGHV1, and mouse IGHV3 and IGHV12 are in the same clan than human IGHV4 (66).

Although some mouse CLL clones may have longer HCDR3 than normal B cells, as it has been shown in sets of UM-CLL in humans (5, 42), a comparison of the HCDR3 average length of the Traf2DNxBCL2-tg+/+ UM-CLL/SLL clones and those from the Eµ-TCL-1-tg, the MDR−/−, the miR-15a/16-1−/− and the IgH-TEL-tg mice showed no significant differences among them and, in all cases, it was similar to the HCDR3 average length of normal mouse B cells. However, even though this result suggests that this feature of human UM-CLL is not shared by its mouse counterparts, an analysis of a large cohort of CLL samples from 2662 patients have shown that the stereotyped HCDR3 sequences seem to cluster in discrete groups of 9, 13, 20 and 22 AAs (14), thus suggesting that long HCDR3 are not a general feature of human UM-CLL but rather of some stereotype subtypes. Thus, due to the limited number of mouse CLL

| TABLE 3 | HCDR3 sequence alignments of the Traf2DNxBCL2-tg+/+ CLL/SLL clones and their putative target antigens. |

A clustal W alignment and dendrogram comparing the HCDR3 from the expanded Traf2DNxBCL2-tg+/+ CLL/SLL clones is shown. The ID number of the mouse, the source of the tissue, the frequency of occurrence for each clone and the SHM status of the clone (M or UM) are indicated. The putative antigens recognized by the CLL/SLL HCDR3 were determined using NCBI protein Blast (non-redundant sequences restricted to Mus musculus, taxid. 10090) and selecting the antigen recognized by antibodies encoding HCDR3 with the highest similarities to the HCDR3 expressed by the CLL/SLL clones (the antigen candidate and the % of HCDR3 similarities is indicated). Those HCDR3 with ≥ 75% similarities are highlighted. The GenBank accession code for the referred antibodies is provided. Alignment was performed using the clustal W muscle tree method UPGM https://www.ebi.ac.uk/Tools/phylogeny/simple_phylogeny/.
HCDR3 sequences available, it remains an open question whether a similar distribution could be observed in mouse CLL.

It is important to note that the Traf2DNxBCL2-tg mice are in a FVB/NxBALB/c mixed background. This is relevant considering that the vast majority of the IGHV sequences available at the IMGT repertoire IG database are from C57BL/6 mice and that IGHV from FVB/N mice are not represented in this database. As stated above, there is a large sequence GL IGHV variation between mouse strains (33, 36). Therefore, a direct comparison of the IGHV sequences from the Traf2DNxBCL2-tg mice with those GL stored at IMGT would not be representative of the actual percentage of variation of the analyzed IGHV sequences with the GL. To provide a more precise analysis, we have compared all available sequences of the same IGHV gene from the Traf2DNxBCL2-tg (irrespective of their genotype) and Traf3xBCL2-tg mice (also in a FVB/NxBALB/c mixed background), as well as with available FVB/N IGHV sequences available in public databases. These comparisons allowed discriminating what differences with the GL sequence were more likely SSIP or SHM. Our results indicate that a majority of the IGHV sequences of the Traf2DNxBCL2-tg-/+, +/- and ++ have ≤2% differences with the GL and are categorized as UM. However, it is important to state that these results are an estimation of SHM events and that a comparison with the GL of the mouse strain analyzed is required for an accurate assessment of SHM events.

**Insights Into the Cellular Origins on Mouse CLL**

CLL ontogeny is still a matter of intense study and discussion (2, 67, 68). This also applies to the identification of the cellular source of mouse CLL, notwithstanding our deeper knowledge on mouse B cell ontogeny and differentiation compared to that of humans. Questions still remain even on whether human and CLL arises from a single or multiple cell types.

MZ B cells are IgM+ cells responding to T-independent antigens. They have a limited IGHV-D-J repertoire usage often producing polyclonal BCR recognizing autoantigens and pathogen antigens (69). MZ B cells are mostly UM but they can go through extra-germinal center SHMs producing also M-MZ B cells (70). Our studies on the mechanisms causing CLL/SLL development in the Traf2DNxBCL2-tg+/+ mice showed that B cell-specific TRAF2DN expression caused proteasome-dependent degradation of endogenous TRAF2, thus rendering B cell-specific Traf2DN-tg mice into bona fide B cell-specific Traf2-deficient mice (24). Confirming previous results (38, 71), we showed that the lack of functional TRAF2 enforces MZ B cell accumulation and releases B cells from the need of BAFF for survival (24). BCL2 overexpression, a defining characteristic of human CLL cells (25) would provide in this model a necessary additional level of protection against apoptosis, likely through a similar mechanism to that described in human CLL (72). Altogether, our results would be consistent with a role for Traf2-deficiency and BCL2 overexpression in promoting MZ B cells expansion and predisposing MZ B cells to CLL/SLL transformation (24).

A role for MZ B cells as the source of the CLL/SLL arising in the Traf2DNxBCL2-tg+/+ mice might explain why in this mouse model SLL arises first, later progressing to CLL (22). This would be in line with the ability of MZ B cells to move into circulation (73). Moreover, since the CLL/SLL developed by the Traf2DNxBCL2-tg+/+ mice may express or not CD5 on their surface (22), this could reflect that MZ B cells were at different activation stage at the time of transformation [reviewed in (2)].

On the other hand, various lines of evidence suggest that the CLL developed by other mouse models might arise from a different B cell type. In this regard, there is evidence pointing out to a B1a cell origin for UM-CLL developed by some of the available CLL mouse models [reviewed in (2)]. First, the preferential usage of IGHV1 and IGHV11 genes by the Eμ-TCL-1-tg (28, 51, 74) and by the MDR−/− and miR-15a/16-1−/− mice (32) is similar to the preferential IGHV subgroup usage of mouse splenic B1a cells (75). Second, the most frequently expressed clones in B1a cells (both, peritoneal and splenic) have HCDR3 with the sequences MRYGNYWYFDV, MRYSYNYWYFDV, MRYGSYWYFDV and MRYGSSYWYFDV (75) found in BCRs that are reactive to phosphatidylcholine (28). These HCDR3 are commonly found in expanded CLL clones from the Eμ-TCL-1-tg mice (28) the MDR−/− and miR-15a/16-1−/− mice (32) and the IgH-TEmice (45) (Supplementary Table 8). Third, Hayakawa and coworkers (52) have shown that allotransplantation of B1 cells, but not of other B cell subtypes, from the Eμ-TCL1-tg mice resulted in CLL with a biased repertoire, including stereotyped BCRs, thus recapitulating the CLL developed by the Eμ-TCL1-tg mice.

Of note is that both MZ B cells and B1 cells have been proposed as a possible source for CLL cells [reviewed in (2)]. However, there is conflicting evidence for MZ B cells as the source of human CLL (2, 67) and we are still lacking clear evidence on the existence of a human counterpart of mouse B1 cells. Therefore, despite the high similarities of the CLL developed by humans and the available mouse CLL models, including the Traf2DNxBCL2-tg+/+ mice, additional research is needed to elucidate whether mouse and human CLL have a similar ontogeny and cell type origin.

**Possible Role of Autoantigens and Pathogen Antigens in the CLL/SLL Developed by the Traf2DNxBCL2-tg+/+ Mice**

Although CLL cells relying on antigen-independent, cell-autonomous BCR signaling have been described (76), there is ample evidence for the role of autoantigen-stimulated BCR in CLL clonal selection, expansion and clonal evolution (31, 50–54). Our results showing the similarities of the HCDR3 expressed by the expanded Traf2DNxBCL2-tg+/+ CLL/SLL clones to those recognizing autoantigens and pathogens suggest that antigen-stimulation would also drive disease progression in our CLL/SLL mouse model, similarly to what has been demonstrated in the Eμ-TCL1-tg mice (31, 51).

Stereotyped HCDR3 sequences are mostly found in UM-CLL clones in humans and produce BCRs that frequently recognized
autoantigens [reviewed in (16, 17)]. In agreement with these findings, the identical HCDR3 found in the Traf2DNxBCL2-tg+/+ mice were also UM-CLL/SLL clones. Moreover, we found several Traf2DNxBCL2-tg+/+ UM-CLL/SLL clones expressing HCDR3 highly similar to HCDR3 recognizing autoantigens (phosphatidylcholine) and pathogen antigens (CMV, hepatitis C virus, and lipoteichoic acid). However, HCDR3 with similar antigen specificities were also found in M-CLL/SLL clones, recognizing autoantigens, such as cardiolipin and dsDNA, and pathogen antigens (Bordetella) (Table 3). In this regard, Herve and coworkers (12) have shown that both M- and UM-CLL clones derived from self-reactive B cell precursors and our data would be in agreement with those results.

Finally, the presence of B cell clones with similar HCDR3 sequence in mice with different genotypes (Traf2DNxBCL2-tg+/+; +/+ and +/+) suggests that all mice are exposed to similar antigens and have similar immune responses to them. Exposure to the same antigens should be expected considering that mice in this study are littermates and are housed together. The fact that only the Traf2DNxBCL2-tg+/+ mice develop CLL/SLL highlights the need of both Traf2 deficiency and BCL2 overexpression for promoting CLL development in this mouse model and underlines a role for autoantigen- and pathogen antigens-specific HCDR3 in driving disease progression.

DATA AVAILABILITY STATEMENT
The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT
The animal study was reviewed and approved by Bioethics Committee of the Consejo Superior de Investigaciones Cientificas (CSIC).

REFERENCES
1. Chiorazzi N, Rai KR, Ferrarini M. Chronic lymphocytic leukemia. N Engl J Med (2005) 352(8):804–15. doi: 10.1056/NEJMra041720
2. Chiorazzi N, Ferrarini M. Cellular origin(s) of chronic lymphocytic leukemia: cautionary notes and additional considerations and possibilities. Blood (2011) 117(6):1781–91. doi: 10.1182/blood-2010-07-155663
3. Kipps TJ, Stevenson FK, Wu CJ, Croce CM, Packham G, Wierda WG, et al. Chronic lymphocytic leukaemia. Nat Rev Dis Primers (2017) 3:16096. doi: 10.1038/nrdp.2016.96
4. Schroeder HW Jr, Dighiero G. The pathogenesis of chronic lymphocytic leukemia: analysis of the antibody repertoire. Immunol Today (1994) 15 (6):288–94. doi: 10.1016/0167-5699(94)90009-4
5. Fais F, Ghiotto F, Hashimoto S, Sellars B, Valetto A, Allen SL, et al. Chronic lymphocytic leukemia B cells express restricted sets of mutated and unmutated antigen receptors. J Clin Invest (1998) 102(8):1515–25. doi: 10.1172/JCI3009
6. Hamblin TJ, Davis Z, Gardiner A, Oscar DG, Stevenson FK. Unmutated Ig VH genes are associated with a more aggressive form of chronic lymphocytic leukemia. Blood (1999) 94(6):1848–54. doi: 10.1182/blood.V94.6.1848.1848_1854
7. Damle RN, Wasil T, Fais F, Ghiotto F, Valetto A, Allen SL, et al. Ig V gene mutation status and CD38 expression as novel prognostic indicators in chronic lymphocytic leukemia. Blood (1999) 94(6):1840–7. doi: 10.1182/blood.V94.6.1840
8. Maloum K, Davi F, Merle-Beral H, Pritsch O, Magnac C, Vuillier F, et al. Expression of unmutated VH genes is a detrimental prognostic factor in chronic lymphocytic leukemia. Blood (2000) 96(1):377–9. doi: 10.1182/blood.V96.1.377.013k56f_377_379
9. Damle RN, Ghiotto F, Valetto A, Albesiano E, Fais F, Yan XI, et al. B-cell chronic lymphocytic leukemia cells express a surface membrane phenotype of activated, antigen-experienced B lymphocytes. Blood (2002) 99(11):4087–93. doi: 10.1182/blood.V99.11.4087
10. Klein U, Tu Y, Stolovitzky GA, Mattioli M, Cattoretti G, Husson H, et al. Gene expression profiling of B cell chronic lymphocytic leukemia reveals a homogeneous phenotype related to memory B cells. J Exp Med (2001) 194 (11):1625–38. doi: 10.1084/jem.194.11.1625

AUTHOR CONTRIBUTIONS
GP-C designed, performed, and analyzed the experiments and helped writing the paper. JZ designed and analyzed the experiments and wrote the paper. All authors contributed to the article and approved the submitted version.

FUNDING
This work was supported by grants from the Agencia Estatal de Investigacion (PID2019-110405RB-100/AEI/10.13039/501100011033) and from the Instituto de Salud Carlos III (PI16/000895). We acknowledge support of the publication fee by the CSIC Open Access Publication Support Initiative through its Unit of Information Resources for Research (URICI). The cost of this publication was paid in part by FEDER funds.

ACKNOWLEDGMENTS
We are indebted to Laura Barrios (Scientific Calculation Center, SGAI, CSIC) for statistical analyses. We are thankful to Maria G. Gonzalez-Bueno for excellent technical support. We are grateful to the personnel of the Animal facility and Genomics facilities at Instituto de Investigaciones Biomedicas “Alberto Sols”. We also thank Dr. Paloma Perez-Aciego (Fundacion LAIR, Madrid) for kindly providing reagents and helpful discussions. Pablo Carr, Andrea de Andrés and Blanca Jimenez are acknowledged for helpful technical assistance.

SUPPLEMENTARY MATERIAL
The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2021.627602/full#supplementary-material
23. Kress CL, Konopleva M, Martinez-Garcia V, Krajewska M, Lefebvre S, Hyer R, et al. TNFR-associated factor (TRAF) domain and Bcl-2 cooperate to induce small B cell lymphoma. *J Clin Invest* (2005) 115(6):1636–43. doi: 10.1172/JCI24387

24. Perez-Chacon G, Llobet D, Pardo C, Pindado J, Choi Y, Reed JC, et al. TNFR-associated factor (TRAF) domain and Bcl-2 cooperate to induce small B cell lymphoma. *J Clin Invest* (2005) 115(6):1636–43. doi: 10.1172/JCI24387

25. Robertson LE, Plunkett W, McConnell K, Keating MJ, McDonnell TJ. Bcl-2 deficiency in B lymphocytes predisposes to chronic lymphocytic leukaemia. *Proc Natl Acad Sci USA* (2004) 101(47):16600–5. doi: 10.1073/pnas.0407541101

26. Lee SY, Reichlin A, Santana A, Sokol KA, Nussenzieg MC, Choi Y, TRAF2 is essential for JNK but not NF-κB activation and regulates lymphocyte proliferation and survival. *Immunity* (1997) 7:703–13. doi: 10.1016/S1074-7613(00)80390-8

27. Katsumata M, Siegel RM, Louie DC, Miyashita T, Tsujimoto Y, Nowell PC, et al. Differential effects of Bcl-2 on B and T lymphocytes in transgenic mice. *Proc Natl Acad Sci USA* (1994) 91:11376–80. doi: 10.1073/pnas.91.23.11376

28. Yan XJ, Alisalam E, Zanesi N, Yancopoulos G, Sawyer A, Romano E, et al. B cell receptors in TCEL1 transgenic mice resemble those of aggressive, treatment-resistant human chronic lymphocytic leukemia. *Proc Natl Acad Sci USA* (2006) 103(31):11713–8. doi: 10.1073/pnas.0604564103

29. Brochet X, Lefranc MP, Giudicelli V. IMGT/V-QUEST: the highly customized and integrated system for IG and TR standardized V-J and V-D-J sequence analysis. *Nucleic Acids Res* (2008) 36(Web Server issue):W503–8. doi: 10.1093/nar/gkn316

30. Karan-Djusevic T, Pavlovic S. Somatic Hypermutational Status and Gene Repertoire of Immunoglobulin Rearrangements in Chronic Lymphocytic Leukemia. In: I Gheorghita, editor. *Lymphocyte Updates - Cancer, Autoimmunity and Infection*. Rijeka, Croatia: InTech (2017). p. 49–78. (Chapter 3).

31. Iacovelli S, Hug E, Benardo S, Dheuhen-von Minden M, Gobessi S, Rinaldi A, et al. Two types of BCR interactions are positively selected during leukemia development in the Emu-TCL1 transgenic mouse model of BLL. *Blood* (2015) 125(10):1578–88. doi: 10.1182/blood-2014-07-587790

32. Iken U, Lia M, Crespo M, Siegel R, Shen Q, Mo T, et al. The DLEu2/mir-15a/16-1 cluster controls B cell proliferation and its deletion leads to chronic lymphocytic leukemia. *Cancer Cell* (2010) 17(1):28–40. doi: 10.1016/j.ccr.2009.11.019

33. Collins AM, Wang Y, Roskin KM, Marquis CP, Jackson KJ. The mouse antibody chain repertoire is germline-focused and highly variable between inbred strains. Philosophical transactions of the Royal Society of London Series B. * Biol Sci* (2015) 370(1676):20140236. doi: 10.1098/rstb.2014.0236

34. Perez-Chacon G, Amdros M, Vallesio-Cremades MT, Lefebvre S, Reed JC, Zapata JM. Dysregulated TRAF3 and BCL2 Expression Promotes Multiple Classes of Mature Non-hodgkin B Cell Lymphoma in Mice. *Front Immunol* (2019) 10:3114. doi: 10.3389/fimmu.2019.03114

35. Rettig TA, Ward C, Bye BA, Pecaut MJ, Chapes SK. Characterization of the naïve murine antibody repertoire using unamplified high-throughput sequencing. *PLoS One* (2018) 13(1):e0190982. doi: 10.1371/journal.pone.0190982

36. Watson CT, Kos JT, Gibson WS, Newman L, Deikus G, Busse CE, et al. A comparison of immunoglobulin IGHV, IGHD and IGJH genes in wild-derived and classical inbred mouse strains. *Immunol Cell Biol* (2019) 97(10):888–901. doi: 10.1111/icmb.12288

37. Tobin G, Thunberg U, Laurell A, Karlsson S, Aleskog A, Willander K, et al. Patients with chronic lymphocytic leukaemia with mutated VH genes presenting with Binet stage B or C form a subgroup with a poor outcome. *Haematologica* (2005) 90(4):465–9.

38. Gardam S, Siero F, Basten A, Mackay F, Brink R. TRAF2 and TRAF3 signal adapters act cooperatively to control the maturation and survival signals delivered to B cells by the BAFF receptor. *Immunity* (2008) 28(3):391–401. doi: 10.1016/j.immuni.2008.01.009

39. Martin F, Kearney JF. Marginal zone B cells. *Nat Rev Immunol* (2002) 2(5):323–35. doi: 10.1038/nri799

40. Mastache EF, Lindroth K, Fernandez C, Gonzalez-Fernandez A. Somatic hypermutation of Ig genes is affected differently by failures in apoptosis caused by disruption of Fas (lpr mutation) or by overexpression of Bcl-2. *Scandinavian J Immunol* (2006) 63(6):420–9. doi: 10.1111/j.1365-3083.2006.01758.x

41. Maurer K, Zahrich D, Gorgun G, Li A, Zhou J, Ansen S, et al. Immunoglobulin gene segment usage, location and immunogenicity in mutated and unmutated chronic lymphocytic leukaemia. *Br J Haematol* (2005) 129(4):499–510. doi: 10.1016/j.bjha.2005.05.0480

42. Johnson TA, Rassenti LZ, Kipps TJ, Ig VH genes expressed in B cell chronic lymphocytic leukaemia exhibit distinctive molecular features. *J Immunol* (1997) 159(1):235–46.

43. Widhoff GF, Kipps TJ. Normal B cells express 51p-encoded Ig heavy chains that are distinct from those expressed by chronic lymphocytic leukaemia B cells. *J Immunol* (2001) 166(1):95–102. doi: 10.4049/jimmunol.166.1.95

44. Shi B, Ma L, He X, Wang X, Wang P, Zhou L, et al. Comparative analysis of human and mouse immunoglobulin variable heavy regions from IMGT/LIGM-DB with IMGT/HIGH-VDJ. *Theor Biol Med Model* (2014) 11(1):30. doi: 10.1186/1742-4682-11-30

45. Pal Singh S, De Bruijn MJW, de Almeida MP, Meijers RWJ, Nitschke L, Pal S, et al. Identi fi cation of Distinct Unmutated Chronic Lymphocytic Leukemia Subsets in Mice Based on Their T Cell Dependency. *Front Immunol* (2019) 8:1996. doi: 10.3389/fimmu.2018.01996

46. Agathangelidis A, Darzentas N, Hadzidimitriou A, Brochet X, Murray F, Yan XJ, et al. Stereotyped B cell receptors in one-third of chronic lymphocytic
leukemia: a molecular classification with implications for targeted therapies. *Blood* (2012) 119(19):4647–75. doi: 10.1182/blood-2011-11-393694

47. Agathangelidou A, Vardi A, Baliakas P, Stamatopoulos K. Stereotyped B-cell receptors in chronic lymphocytic leukemia. *Leuk Lymphoma* (2014) 55 (10):2252–61. doi: 10.3109/10428194.2013.879715

48. Colombo M, Bagnara D, Reverberi D, Matis S, Cardillo M, Massara R, et al. Tracking CLL-biased stereotyped immunoglobulin gene rearrangements in normal B cell subsets using a high-throughput immunogenetic approach. *Mol Med* (2020) 26(1):25. doi: 10.1186/s12849-020-00151-9

49. Muggen AF, de Jong M, Wolvers-Tettero ILM, Kallemeijn MJ, Teodosio C, et al. Production of autoantibodies by CD5-expressing B lymphocytes from chronic lymphocytic leukemia B cells. *Blood* (2013) 121 (23):4708–17. doi: 10.1182/blood-2012-08-447904

50. Chen SS, Batliwalla F, Holodick NE, Yan XJ, Yancopoulos S, Croce CM, et al. Chronic lymphocytic leukemia requires BCL2 to sequester prodeath BIM, explaining sensitivity to BCL2 antagonist ABT-737. *Crit Rev Immunol.* (2018) 38(2):145–58. doi: 10.1615/CritRevImmunol.2018024985

51. Jimenez de Oya N, De Giovanni M, Fioravanti J, Ubelhart R, Di Lucia P, et al. The authors declare that the research was conducted in the normal BCR repertoire in healthy individuals increases with age. *Immun Ageing* (2019) 16:22. doi: 10.1186/s12979-019-0163-x

52. Zwick C, Fadle N, Regitz E, Kemele M, Stilgenbauer S, Buhler A, et al. Autoantigenic targets of B-cell receptors derived from chronic lymphocytic leukemias bind to and induce proliferation of leukemic cells. *Blood* (2013) 121(23):4708–17. doi: 10.1182/blood-2012-08-447904

53. Hayakawa K, Formica AM, Brill-Dashoff J, Shinton SA, Ichikawa D, Zhou Y, et al. Early generated B1 B cells with restricted BCRs become chronic lymphocytic leukemia with continued c-Myc and low Bmf expression. *J Exp Med* (2016) 213(13):3007–24. doi: 10.1084/jem.20160712

54. Broker BM, Klajman A, Youinou P, Jouquan J, Worman CP, Murphy J, et al. Marginal zone B cells emerge from innate-like antibody-producing lymphocytes. *Nat Rev Immunol* (2013) 13(2):118–32. doi: 10.1038/nri3383

55. Hendricks J, Bos NA, Kroese FGM. Heterogeneity of Memory Marginal Zone B Cells. *Crit Rev Immunol.* (2018) 38(2):145–58. doi: 10.1615/CritRevImmunol.2018024985

56. Grech AP, Amesbury M, Chan T, Gardam S, Bastein A, Brink R. TRAF2 differentially regulates the canonical and noncanonical pathways of NF-κB activation in mature B cells. *Immunity* (2004) 21(5):629–42. doi: 10.1016/j.immuni.2004.09.011

57. Hendricks J, Bos NA, Kroese FGM. Heterogeneity of Memory Marginal Zone B Cells. *Crit Rev Immunol.* (2018) 38(2):145–58. doi: 10.1615/CritRevImmunol.2018024985

58. Bichi R, Shinton SA, Martin ES, Koval A, Calin GA, Cesari R, et al. Human chronic lymphocytic leukemia: a molecular classification with implications for targeted therapies. *Blood* (2012) 119(19):4647–75. doi: 10.1182/blood-2011-11-393694

59. Ten Hacken E, Gounari M, Ghia P, Burger JA. The importance of B cell receptor isotypes and stereotypes in chronic lymphocytic leukemia. *Leukemia* (2019) 33(2):287–98. doi: 10.1038/s41375-018-0303-x

60. Traver-Glehne A, Davi F, Ben Simon E, Callet-Bauchu E, Felman P, Basegio L, et al. Analysis of VH genes in marginal zone lymphoma reveals marked heterogeneity between splenic and nodal tumors and suggests the existence of clonal selection. *Haematologica* (2005) 90(4):470–8.