Chronic lymphocytic leukemia patients with IGH translocations are characterized by a distinct genetic landscape with prognostic implications

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Abbreviations

CLL: chronic lymphocytic leukemia; COSMIC: Catalogue of Somatic Mutations in Cancer; DLBCL: diffuse large B-cell lymphoma; ExAC: Exome Aggregation Consortium; FISH: Fluorescence in situ hybridization; FL: follicular lymphoma; GATK: Genome Analysis Toolkit; ICGC: International Cancer Genome Consortium; IGH: immunoglobulin heavy chain; IGHR-CLLs: CLL patients with IGH rearrangements;

IGHV: immunoglobulin heavy-chain variable; IGV: Integrative Genomics Viewer; IWCLL: International Workshop on CLL; MACS: magnetically activated cell sorting; NGS: next-generation sequencing; NHL: non-Hodgkin lymphoma; OS: Overall survival; PCAWG: Pan-Cancer Analysis of Whole Genomes.; TFT: Time to first treatment; VAF: variant allele frequency; VCF: variant calling files.

Novelty and Impact

In this work, we assess for the first time the genetic landscape of CLLs with *IGH* rearrangements by targeted NGS, characterizing recurrently mutated genes with prognostic implications and demonstrating that these entities exhibit an intermediate mutational profile between CLL and non-Hodgkin lymphoma. Moreover, our findings showed that the incorporation of NGS and the IGH-probe in the CLL-FISH panel used in clinical routine could be extremely useful, especially for elucidating prognosis in 'normal FISH' cases.

ABSTRACT

Chromosome 14q32 rearrangements/translocations involving the immunoglobulin heavy chain (IGH) are rarely detected in chronic lymphocytic leukemia (CLL). The prognostic significance of the IGH translocation is controversial and its mutational profile remains unknown. Here, we present for the first time a comprehensive next-generation sequencing (NGS) analysis of 46 CLL patients with IGH rearrangement (IGHR-CLLs) and we demonstrate that IGHR-CLLs have a distinct mutational profile with recurrent mutations in NOTCH1, IGLL5, POT1, BCL2, FBXW7, ZMYM3, MGA, BRAF and HIST1H1E genes. Interestingly, BCL2 and FBXW7 mutations were significantly associated with this subgroup and almost half of BCL2, IGLL5 and HISTH1E mutations reported were previously identified in non-Hodgkin lymphomas (NHL). Notably, IGH/BCL2 rearrangements were associated with a lower mutation frequency and carried BCL2 and IGLL5 mutations, while the other IGHR-CLLs had mutations in genes related to poor prognosis (NOTCH1, SF3B1 and TP53) and shorter time to first treatment (TFT). Moreover, IGHR-CLLs patients showed a shorter TFT than CLL patients carrying 13q-, normal FISH and +12 CLL, being this prognosis particularly poor when NOTCH1, SF3B1, TP53, BIRC3 and BRAF were also mutated. The presence of these mutations not only was an independent risk factor within IGHR-CLLs, but also refined the prognosis of low-risk cytogenetic patients (13q-/normal FISH). Hence, our study demonstrates that IGHR-CLLs have a distinct mutational profile from the majority of CLLs and highlights the relevance of incorporating NGS and the status of IGH by FISH analysis to refine the risk-stratification CLL model.

INTRODUCTION

Chronic lymphocytic leukemia (CLL) is a disease that displays extreme clinical heterogeneity, clearly reflecting the marked biological diversity, which has led to the identification of a plethora of prognostic markers¹⁻⁴. Chromosomal abnormalities are the hallmark of the disease and their correlation to the clinical course has contributed to patients risk stratification since the 2000s⁵. In the last years, CLL molecular and cellular biology has been enriched by seminal insights that have led to a better understanding of CLL pathogenesis² and, consequently, to the identification of molecular markers whose evaluation is well-established in clinical routine, such as the *IGHV* mutational status or *TP53* gene abnormalities. The integration of these markers together with the new relevant genetic alterations reported in next-generation sequencing (NGS) studies, specifically those of *NOTCH1*, *SF3B1* and *BIRC3* genes, could be used to refine Döhner hierarchical cytogenetic model^{2,6-12}.

Although more than 80% of CLL patients carry cytogenetic alterations, chromosome 14q32 rearrangements/translocations involving the immunoglobulin heavy chain gene (*IGH*) was considered a rare aberration affecting fewer than 4% of CLL patients^{5,13}. Nevertheless, with the emergence of new molecular approaches and large-scale genomic studies in CLL, a higher incidence of *IGH* rearrangements has been reported in the recent years (5-15%) ¹⁴⁻¹⁶. This cytogenetic abnormality contributes to CLL pathogenesis by deregulating the *IGH*-partner genes^{17,18} and their prognostic significance is still controversial. Previous studies have shown that patients carrying 14q32 rearrangements (IGH*R*-CLLs) have an intermediate-adverse outcome¹⁹⁻²¹, particularly when compared with favorable and intermediate-risk cytogenetics^{22,23}. However, some studies have reported that patients carrying 14q32 rearrangements with *BCL2* have a better clinical course^{24,25}.

CLL patients with *IGH* rearrangements are still poorly characterized at the molecular level, partly due to the low incidence of cases, the *IGHR* co-occurrence with other cytogenetic alterations, and the difficulty of distinguishing between *IGHR*-CLLs and

forms of non-Hodgkin lymphoma (NHL)²⁶. Furthermore, the *IGH* probe is not included in the classic four-probe CLL FISH panel for the 13q14, 12p11.1-q11, 11q22 and 17p13 regions used in routine clinical practice²⁶, which is partially responsible for this subgroup passing unnoticed. In this study, we characterize the genetic landscape of CLL patients with 14q32/*IGH* translocations for the first time, demonstrating that *IGHR*-CLLs have a distinct mutational profile from other classic cytogenetic groups of CLLs, dependent on whether *BCL2* is involved or not in the *IGH* rearrangement, and as well as the presence of certain mutations. Taken together, our results improve our understanding of the molecular underpinnings of this cytogenetic CLL subgroup, allowing us to refine the prognosis of *IGHR*-CLL patients.

METHODS

Patients

The study was based on 862 CLL patients, diagnosed according to the International Workshop on CLL (IWCLL) criteria^{27,28}. All of them were screened for *IGH* translocation and positive cases for *IGH* rearrangement were individually reviewed to rule out the possibility that they represented a different lymphoproliferative disorder (see **Supplementary Methods**). Samples and clinical data were collected from 16 Spanish institutions.

Mutational analysis was performed in 233 untreated CLL patients: 46 with 14q32/IGH rearrangements and 187 as the control group. Patients in the control group were selected according to sample and clinical data availability and absence of treatment and were representative of the disease in terms of demographic and clinical characteristics (Supplementary Table S1). Patients risk classification criteria is described in Supplementary Methods and a diagram of the patients included in the different outcome analyses is shown in Supplementary Fig. S1.

In the IGHR-CLL group, the median time between diagnosis and *IGH* rearrangement detection was 1 month (range, 0-117 months), and the median follow-up was 57 months (range, 1-157 months). Within IGHR-CLLs, 31/46 (67.4%) received treatment after FISH

test, with a median TFT of 19 months (range: 7-30). Most of them (93.5%) received conventional chemoimmunotherapy and 2 patients were treated with ibrutinib.

The study was approved by the local ethical committee (*Comité Ético de Investigación Clínica*, *Hospital Universitario de Salamanca*). Written informed consent was obtained from all participants before they entered the study.

Fluorescence in situ hybridization (FISH)

Interphase FISH was performed on peripheral blood or bone marrow samples using the following commercially probes: ATM, CEP12, D13S319 and TP53 (Vysis, Abbott Laboratories, IL, USA). Dual color break-apart FISH probes were performed for IGH/BCL2 and IGH/BCL6 translocations. The methods used for FISH analysis have been described elsewhere²⁹. Signal screening was carried out in at least 200 cells with well-delineated fluorescent spots. In all cases, a score of $\geq 10\%$ was considered positive, based on the cut-off value used by our laboratory.

Next-generation sequencing

NGS studies were performed in 233 cases and in the same sample as the FISH test. Genomic DNA was isolated from peripheral blood or bone marrow by magnetically activated cell sorting (MACS) CD19+ B-lymphocytes. B cell purity was greater than 98% by flow cytometry, as previously described in our group³⁰. The Agilent SureSelect^{QXT} Target Enrichment system for Illumina Multiplexed Sequencing (Agilent Technologies, Santa Clara, CA, USA) was used to produce libraries of exonic regions from 54 genes CLL-related as well as from *BCL2*, *IGLL5* and *NOTCH1* UTR regions (**Supplementary Methods**). Genes included in the custom-designed panel^{31,32} are involved in CLL pathogenesis and the UTR regions were considered due to the previous identification of *IGLL5*, *BCL2* and *NOTCH1* UTRs somatic mutations in CLL^{8,33,34}. (**Supplementary Table S2**). Paired-end sequencing (151-bp reads) was run on the Illumina NextSeq instrument (Illumina, San Diego, CA, USA).

Data analysis

Raw data quality control was performed with FastQC (v0.11.8) and Picard tools (v2.2.4) to collect sequencing metrics. Demultiplexed files (FASTQ) were aligned to the reference genome (GRCh37/hg19 genome), read duplicates were marked with SAMTools (v1.3.1) and post-alignment was performed with GATK (v3.5). Coverage for each region was assessed using BEDTools (v.2.26.0). A minimum quality score of Q30 was required for ensuring high-quality sequencing results. Finally, somatic variant calling, and annotation were performed using an in-house pipeline, based on VarScan (v2.4) and ANNOVAR (v.2017Jul16), respectively.

Median coverage of target regions was 600 reads/base, with at least 100X in 97% of them. To validate variants detected with VAF <5% using the custom panel, samples were conducted to resequencing using different amplicon-based approaches (Illumina Nextera XT/454 Roche³⁰) with read depth above 1000X, allowing to report variants down to 2% (**Supplementary Methods**).

Data was then filtered according to the severity of the consequence, considering variants that lead to an amino acid change in the protein sequence (missense, nonsense, frameshift) and those in the splice site and UTRs. To discard single nucleotide polymorphisms (SNPs), minor allelic frequencies (MAFs) were consulted in several databases (dbSNP, 1000 genomes, ExAC and our in-house database) and only variants with a MAF of <0.01 were selected for further analysis. In addition, variants with a VAF between 40-60% or greater than 90% were manually reviewed prioritizing variants described in *in silico* tools (Polymorphism Phenotyping v2 (PolyPhen-2), Sorting Intolerant From Tolerant (SIFT) and ClinVar) as deleterious, damaging, pathogenic or likely pathogenic.

Aligned reads were manually reviewed with the Integrative Genomics Viewer (IGV) to confirm and interpret variant calls and reduce the risk of false positives. Variants described in the Catalogue of Somatic Mutations in Cancer database (COSMIC82 database) or mutations in driver genes previously described in seminal papers were rescued for the analysis (CLL and non-Hodgkin lymphoma)^{7,8,33,35-37}. Manually screening

in VarSome and International Cancer Genome Consortium (ICGC) Databases was performed for assessing the functional impact of mutations.

Statistical analysis

Statistical analyses were performed using IBM SPSS v23.0 for Windows (IBM Corp., Armonk, NY, USA) and SDM-PSI v6.21 software for the false discovery rate (FDR) correction in multiple comparisons. Continuous variables were analyzed with the Mann-Whitney U test, while the chi-square and Fisher's exact tests were used to assess associations between categorical variables. Overall survival (OS) and time to first therapy (TFT) were calculated from the date FISH test was performed to the date of death, first treatment or last follow-up (considering disease-unrelated deaths as competing events). Statistically significant variables related to OS and TFT were estimated by the Kaplan-Meier method, using the log-rank test to compare the curves of each group. Univariate and multivariate analyses of the OS and TFT employed the Cox regression method. Results were considered statistically significant for values of P<0.05. FDR was used to correct P-values for multiple hypothesis testing when appropriate, by applying the Benjamini and Hochberg method³⁸. Adjusted P-values (Q-values) were considered significant when Q<0.1.

RESULTS

CLL patients with *IGH* translocations have a distinct mutational profile with high mutation frequencies in *NOTCH1*, *BCL2*, *FBXW7*, *ZMYM3* and *MGA*

NGS analysis of the 233 CLL patients revealed that 75% of cases had at least one mutation in any of the 54 genes included in the targeted-NGS approach, and the median frequency of mutations per patient was 2 (range: 0-7). The most frequently mutated genes were *NOTCH1* (19.3%), *IGLL5* (15%), *SF3B1* (10.7%), *TP53* (10%), *ATM* (9%), *POT1* (8.5%), *RPS15* (6.9%), *CHD2* (6%), *NFKBIE* (5.1%), *BIRC3* (5.1%) and *XPO1* (4.3%).

Regarding the 46 IGH*R*-CLLs, we identified a total of 109 mutations located in 35 genes. The median frequency of mutations per patient was 2 (range: 0-6), and 82% of patients (38/46) harbored at least one mutation. Moreover, 61% of patients (28/46) presented more than one mutated gene. The most frequently mutated genes in this cohort were *NOTCH1* (30.4%), *IGLL5* (17.4%), *SF3B1* (13%), *POT1* (13%), *TP53*, *BCL2*, *FBXW7*, *ZMYM3* and *MGA* (8.7% each) followed by *BRAF*, *EGR2* and *RPS15* (6.5% each) (**Fig. 1A**; **Supplementary Table S3**). Other genes such as *ATM* (4.3%) or *CHD2* and *MYD88* (2.2% each) were mutated at low frequencies.

The comparison between the mutational profiles of IGHRs-CLLs and the control group showed higher mutation frequencies in *NOTCH1*, *BCL2*, *FBXW7*, *ZMYM3* and *MGA* within IGHR-CLLs, especially those of *BCL2* and *FBXW7* (*Q*=0.048, *Q*=0.06, respectively) (**Fig. 1A**; **Supplementary Table S3**).

Furthermore, 61% of IGH*R*-CLLs (28/46) carried additional FISH alterations (**Fig 1B**). Their mutational profile was analyzed with respect to the presence of IGH*R* together with 13q, 11q, 17p deletion or trisomy 12, and only *TP53* mutations were significantly associated with 17p or 11q deletion in IGH*R*-CLLs (*Q*=0.048). We observed that the mutational profile of patients with IGH*R* as a sole aberration (18/46) was similar to that of the entire IGH*R*-CLL cohort: *NOTCH1* (33.3%), *IGLL5* (27.8%), *SF3B1* (16.7%), *BCL2*, *ZMYM3*, *MGA* and *FUBP1* (11.1% each) followed by *FBXW7* and *BRAF* (5.6% each). All mutation frequencies are shown in **Supplementary Table S4**.

Interestingly, we reported a higher incidence of *IGLL5*, *BCL2* and *HIST1H1E* mutations in this subgroup compared to the described in previous large-scale CLL studies^{7,8} (**Fig. 1B**). IGH*R*-CLL patients showed *IGLL5* mutations targeting the signal peptide domain (4/10) and the 5'UTR region (3/10), *BCL2* mutations affecting the 5'UTR region (2/6) and the exon 2 (4/6), and *HIST1H1E* mutations located in the exon 1 (**Fig 2**). According to the ICGC Database, most of the coding mutations in *IGLL5* (6/7), *BCL2* (3/4) and *HIST1H1E* (1/2) identified in our study, had functional impact in the gene function (**Table 1**). In addition, six out of 17 mutations detected in the aforementioned three genes were

previously described in non-Hodgkin lymphomas (as reported in the COSMIC and ICGC database and whole-exome and whole-genome data from NHL patients^{35-37,39,40}).

Moreover, five of the mutations reported in IGLL5 and BCL2 were located in the 5'UTR of the gene. Specifically, the novel BCL2 recurrent mutation identified in the 5'UTR region (genomic position chr18:60985900) was exclusively found in IGHR-CLLs when compared with the control group (P=0.048) (**Supplementary Fig. S2**).

Detailed lists of the mutations detected in the IGHR-CLLs and the control group are shown in **Supplementary Tables S5, S6 and S7**.

CLL patients with IGH/BCL2 exhibit a lower mutation frequency and a different mutational profile than patients with other IGH translocations

We next sought to assess whether the mutational landscape changes depend on the *IGH* translocation partner, e.g. *BCL2* and *BCL6*. In our study, 13/46 patients (28%) carried *IGH/BCL2* translocation (**Figure 1B**) and 2/46 (4.3%) harbored an *IGH/BCL6* rearrangement (ID 8 and 20). Due to the small number of *BCL6* rearrangements, we performed further analysis comparing *IGH/BCL2* vs. the rest of IGH*R* cases.

In the IGHR patients, fewer CLLs with IGH/BCL2 translocation had mutations in at least one gene compared with the subgroup with other IGH translocations (7/13, 54% vs. 31/33, 94%; P=0.001). The median mutation frequency per patient was significantly lower in the group with IGH/BCL2 compared with that without it (1 vs. 2, P=0.030).

The most frequently mutated genes in the *IGH/BCL2* group were *BCL2* (23%), *IGLL5* (23%), *HIST1H1E* (15%) and *NOTCH1* (15%), whereas for all other IGHR-CLLs, the most frequently mutated genes were *NOTCH1* (36%), *SF3B1* (18%), *POT1* (18%), *TP53* (12%), and *FBXW7* (12%) (**Supplementary Fig. S3A**). It is worth mentioning that neither *TP53* nor *SF3B1* mutations, widely associated with poor prognosis, were detected in CLL patients with an *IGH/BCL2* translocation, reflecting a different mutational profile from all other IGHR-CLLs. The mutational analysis of 9 *IGH/BCL2* cases previously reported in a WES/WGS study of CLL⁸ also showed the presence of mutations in *BCL2*,

and *NOTCH1*, and the absence of poor-prognosis genes such as *TP53* or *SF3B1* (**Supplementary Fig. S3B**). They have been previously reported in WES of However, no statistically significant associations were detected in our analysis, probably due to the small number of cases (**Supplementary Table S8**).

In *IGH/BCL2* cases that also harbored *BCL2* mutations, we observed that 60-87% of the cells carried the rearrangement, while *BCL2* mutations VAFs range from 11% to 40%, suggesting that somatic mutations occurred later in time than the rearrangement (**Table 1**).

Patients carrying IGH translocations exhibit an intermediate-adverse outcome

We also analyzed the clinical and biological characteristics of IGHR-CLLs within the entire cohort (n=862) (**Table 2**). Patients carrying this cytogenetic alteration showed a higher incidence of poor prognosis markers such as Binet stage B or C (Q=0.039), high β 2-microglobulin (Q=0.0007) and lactate dehydrogenase levels (Q=0.054), unmutated IGHV) (Q=0.054) and need for treatment (Q=0.007). In addition, two IGHR-CLLs developed Richter syndrome during follow-up (patient IDs: 18 and 35). Regarding the presence of additional cytogenetic alterations, 34.8% of IGHR-CLL patients (16/46) carried trisomy 12, showing significant co-occurrence of the two events (trisomy 12 and IGHR) (Q=0.0007). By contrast, the presence of the 13q deletion in IGHR-CLLs was significantly less frequent than in CLLs without IGH rearrangements (Q=0.0014) (**Table 2**).

Within IGHR-CLLs, 31/46 (67.4%) received treatment after FISH test, with a median TFT of 19 months (95% CI, 7-30 months). Patients with an *IGH* translocation showed shorter TFT than the 13q- and normal FISH subgroups (median: 19 vs. 120 and 184 months; p<0.0001, p<0.0001), and longer TFT than the 11q- and 17p- subgroups (19 vs. 5, 6 months; p=0.042, p=0.31). The median TFT of the +12 subgroup was slightly higher than that of IGHR-CLLs (28 vs. 19 months; p=0.37). In terms of OS, we observed similar trends (**Supplementary Fig. S4A**). Differences in outcome among the cytogenetic subgroups were consistent with the prevalence of unfavorable clinical and biological

features in IGHR-CLLs, suggesting that this subgroup exhibits an intermediate-adverse prognosis. In addition, the clinical comparison between IGHR-CLLs and control CLLs selected for the mutational analysis (n=233) showed quite similar results to the presented in this section, also demonstrating that control group was representative of the entire cohort (Supplementary Table S1) (Supplementary Fig. S4B).

In our entire cohort (n=862), 31% of patients showed no alterations using 13q14/D13S319, 12p11.1-q11/CEP12, 11q22/ATM, 17p13/P53 probes. However, it is worth mentioning that 6.7% of patients who would be classified as normal FISH in our cohort using the four-probe CLL FISH panel customarily used in routine clinical practice, actually carried an *IGH* rearrangement. The presence of this cytogenetic alteration had a negative effect on the time to first treatment within this group of patients: CLL patients with IGHR as the sole FISH abnormality had a significantly shorter TFT than those without any FISH aberration (23 vs. 120 month, P = 0.01) (**Fig. 3A**).

The presence of the IGH/BCL2 translocation was associated with mutated IGHV (P=0.001), and patients with this alteration showed a longer TFT than those with another IGHR (56 vs. 4 months, P=0.05). By contrast, the presence of IGH/BCL2 rearrangement was not associated to any additional cytogenetic alteration (13q-, Q=0.822; +12, Q=0.822; 11q-/17p-, Q=0.822) and there was no significant difference in terms of OS between patients with IGH/BCL2 and patients with other IGH rearrangements (P=0.433) (**Supplementary Fig. S5**).

Genetic mutations refine the prognosis of IGHR and low-risk cytogenetic CLL patients

IGHR-CLL untreated patients with at least one mutated gene showed a shorter TFT than IGHR-CLLs without gene mutations (10 months vs. median TFT not reached, P=0.026) (**Fig. 3B**). These differences were more significant among recurrent gene mutations previously associated with worse prognosis (*NOTCH1*, *SF3B1*, *TP53*, *BIRC3* and *BRAF*) (2 vs. 88 months, P<0.0001) (**Fig. 3C**). Specifically, TFT was shorter in IGHR patients with *TP53* mutations (0 vs. 23 months, P<0.0001) as well as with *BRAF* mutations (2 vs.

23 months, P = 0.042) (**Supplementary Fig. S6**). In contrast, the presence of *IGLL5* or *BCL2* mutations showed a better impact in terms of TFT, as IGH*R*-CLL patients with mutated *IGLL5* or *BCL2* showed a longer TFT than those without mutations in any of these genes (median TFT not reached vs. 9 months, P = 0.001) (**Fig. 3D**).

In the univariate analysis, other variables associated with a shorter TFT were Binet's stage B/C (P=0.001), splenomegaly (P=0.025), unmutated IGHV status (P=0.013), TP53 disruption/mutation (P=0.003) and the absence of IGLL5/BCL2 mutations (P=0.008). Only the presence of mutations in NOTCH1, SF3B1, TP53, BIRC3 and BRAF was significantly related to a shorter TFT within IGHR-CLL patients in the multivariate analysis (HR=0.255, 95% CI=0.07-0.9, P=0.030) (Table 3).

Since the presence of mutations in these five genes has a prognostic impact within IGHR-CLL patients as well as in the control group (median TFT not reached vs. 12 months, P<0.0001) (Supplementary Fig. S7), we propose an integrated mutational and cytogenetic model to account for our observations in the studied cohort (187 control and 46 IGHR-CLLs). Low-risk patients in our control series (13q-/normal FISH, n=134) segregated into two groups according to the presence of mutations in NOTCH1, SF3B1, TP53, BIRC3 and BRAF (median TFT not reached vs. 24 months, P<0.0001) (Fig. 3E). These mutations also contributed to a worse outcome in intermediate-risk patients (IGHR /+12, n=72) (56 vs. 2 months, P<0.0001). However, the small number of cases with these mutations was insufficient to demonstrate a statistically significant difference in clinical impact in the high-risk cytogenetics subgroup (11q-/17p-, n=27) (4 vs. 0 months, P=0.580). The median TFT of those intermediate-risk patients with mutations was similar to that of patients with high-risk cytogenetic alterations (2 vs. 5 months; P=0.548), and the TFT of low-risk patients with mutations was not significantly different from that of intermediate-risk patients without mutations in any of the five genes (24 vs. 56, P=0.210). Therefore, by including NOTCH1, SF3B1, TP53, BIRC3 and BRAF mutations in the cytogenetic model, approximately 27.6% (37/134) of low-risk patients were reclassified into an intermediate-risk subgroup, and 51% (35/72) of intermediate-risk patients were reclassified into a high-risk subgroup (Fig. 3E).

DISCUSSION

The identification of novel recurrent mutations in CLL has provided a more comprehensive perspective on the genomic landscape and the biological mechanisms underlying the clinical heterogeneity of the disease^{2-4,7,8}. Previous studies have shown that CLLs carrying *IGH* rearrangement could have a worse outcome than low-risk cytogenetic CLL patients^{22,23}. However, their clinical course and molecular characteristics are not well defined¹⁹⁻²⁴. Here, we adopted a targeted NGS approach to assess for the first time the mutational profile of 46 *IGHR*-CLL patients.

Overall, the mutational analysis revealed that IGHR-CLL patients had a high incidence of mutations, not only in well-known CLL drivers such as *NOTCH1*, *SF3B1*, *POT1*, *TP53* and *FBXW7*—previously described in unselected large CLL cohorts—, but also in less commonly mutated genes such as *BCL2*, *FBXW7*, *ZMYM3* and *MGA*^{7,8,33}, being *BCL2* and *FBXW7* significantly associated with the *IGH* translocation (**Fig 1A**). Although we observed the co-occurrence between IGHR and trisomy 12 previously described, we demonstrated that IGHR-CLLs mutational profile did not depend on the presence of additional cytogenetic aberrations: CLLs with only IGHR also exhibited a high mutation frequency in genes well-known associated with trisomy 12 such as *NOTCH1*⁸, as well as in the majority of recurrently mutated genes in the entire IGHR-CLL cohort (**Fig. 1B**).

Strikingly, several IGHR-CLLs showed mutations in *IGLL5*, *BCL2* and *HIST1H1E*, mainly those with *IGH/BCL2* translocations (**Supplementary Fig. S3A**). Although these gene mutations have been detected at low frequencies in other CLL cohorts^{7,8}, they have been extensively reported in other hematological malignances such as diffuse large B-cell lymphoma (DLBCL) and follicular lymphoma (FL)^{35-37,39-44} (**Table 1**). Considering that our series of IGHR-CLL patients is well characterized at the immunophenotypic and clinical levels, the presence of mutations previously described in lymphomas suggests that patients with *IGH* translocation are a cytogenetic subgroup with a mutational profile distinct from the other CLLs and, probably, with different genetic mechanisms underlying their disease pathogenesis. Here, we demonstrated that IGHR-CLLs had an intermed iate

genetic landscape between those of CLL and non-Hodgkin lymphoma, and we suggest that the mutational analysis of patients with *IGH* translocations such as *IGH/BCL2* could help distinguish between CLL and NHL cases.

In this work, we have reported mutations in coding and non-coding regions of BCL2 and IGLL5 and, specifically, we identified a novel recurrent 5'UTR mutation in BCL2. Although this mutation has not been previously described, proximal mutations have been detected in other CLL and NHL studies^{8,34,44}. Moreover, the vast majority of BCL2 mutations reported in Puente et al. were detected in cases harboring IGH/BCL2 translocations⁸, which is consistent with our results (**Supplementary Fig. S3**). Regarding IGLL5, a previous study identified 5'UTR and coding mutations in low-risk IGHVmutated CLLs as well as in the presence of rearrangements³³. These statements are consistent with our results, as most of patients harboring mutations in both genes exhibited mutated *IGHV* and were associated with longer time to first treatment (**Fig. 3D**). In addition, data compiled in the ICGC repository suggested a functional impact of the BCL2 and IGLL5 coding mutations in the gene function. However, the functional impact of UTR somatic mutations has not been well-established yet. Puente et al. demonstrated the negative impact of the 3'UTR mutation in NOTCH18 and recent findings from the Pan-Cancer Analysis of WholeGenomes (PCAWG) Consortium identified novel driver candidates, including mutations in UTR regions, with a potential role in CLL pathogenesis⁴⁵. Nevertheless, further investigation is needed in order to determine the importance of BCL2 and IGLL5 non-coding mutations in CLL.

The clinical impact of *IGH* translocations is currently under discussion¹⁹⁻²⁵. The median TFT was shorter in IGHR-CLLs than that of patients with low-risk cytogenetic alterations, but similar to that of patients with trisomy 12 (**Supplementary Fig. S4A**), indicating that *IGH* translocations could be associated with an intermediate-adverse outcome. Indeed, 6,7% of CLL patients who would be considered 'normal FISH' using the customary four-probe CLL FISH panel in our study, carried the *IGH* translocation and also had a worse prognosis than CLLs lacking IGHR (**Fig. 3A**), thus highlighting the

value of including the IGH probe in the CLL FISH panel to improve patient outcome prediction²⁵.

Previous studies have shown that patients with a IGH/BCL2 translocation had a favorable clinical course, similar to that of patients with low-risk chromosomal alterations, whereas patients with other IGH rearrangements had a similar prognosis to the high-risk subgroups^{24,25}. In our study, patients with IGH/BCL2 not only were associated with IGHV-M and longer TFT (Supplementary Fig. S5A), but also exhibited lower mutation rate compared with other IGH translocations that may contribute to understand why these entities have a better prognosis than the rest of IGHR. Regarding the mutational profile of IGH/BCL2 translocations, a previous large-scale CLL study showed mutations in BCL2, IGLL5 and NOTCH1 within 9 IGH/BCL2 cases, which strongly supports our findings (Supplementary Fig. S3B). On the other hand, IGHR-CLLs without IGH/BCL2 rearrangement presented higher mutation frequencies in genes related to bad prognosis, such as NOTCH1, SF3B1, TP53, BRAF and RPS15 (Fig. 1B; Supplementary Fig. S3A). The high frequency of these mutations may reflect a genomic instability in IGHR-CLLs without IGH/BCL2, which could be also influenced by the role of the translocated partner in the rearrangement. Furthermore, two IGHR-CLLs developed Richter transformation to DLBCL. One of them harbored IGH/BCL2 rearrangement together with trisomy 12 and NOTCH1, and the other patient had an IGHR with unknown partner, NOTCH1 and TP53 mutations. These observations are in line with previous findings regarding the molecular pathways frequently altered at transformation^{46,47}. Altogether, these molecular characteristics could be the underlying mechanisms of the IGHR-CLLs poorer outcome²⁵.

In our cohort, patients harboring *NOTCH1*, *SF3B1*, *TP53*, *BIRC3* or *BRAF* mutations experienced an adverse clinical course (**Fig. 3C**), which is consistent with previous studies^{6-8,11,48}. Within IGH*R*-CLLs, the presence of these mutations contributed to shorter TFT being identified as an independent adverse prognostic factor (**Table 3**). Specifically, IGH*R*-CLL patients harboring *BRAF* mutations exhibited an adverse outcome (**Supplementary Fig. S6A**), which corroborates previous results showing that patients carrying these mutations display an aggressive disease ^{49,50}.

Therefore, the present study proposes an integrated mutational and cytogenetic model for CLL prediction that includes IGHR and BRAF mutational status as novel components with respect to previous prognostic models^{9,10}. The presence of mutations in any of the aforementioned five genes caused a significant shift to a more aggressive outcome in low (13q-/normal FISH) and intermediate-risk (+12/IGHR) CLLs, refining their prognosis and providing information that could help in therapeutic decisions. Interestingly, low-risk patients with mutations in NOTCH1, SF3B1, TP53, BIRC3 or BRAF still had a significantly better outcome than did intermediate-risk patients with any of those mutations (Fig. 3E). These results may indicate that the co-occurrence of cytogenetic abnormalities and gene mutations could have different clinical impacts, depending on the type of the genetic alterations involved.

In conclusion, our study revealed significant differences in the mutational profile and the frequencies of CLL-mutated genes in patients with *IGH* rearrangements. The distribution of genetic mutations differed within the *IGHR*-CLL subgroup: patients with *IGH/BCL2* translocation had higher frequencies of *BCL2* and *IGLL5* mutations than those without the translocation. Conversely, patients with other *IGHR* showed higher mutation frequencies of genes related to bad prognosis (*NOTCH1*, *SF3B1*, *TP53*, *BIRC3* and *BRAF*) than did those with the *IGH/BCL2*. Notably, the presence of those somatic mutations enables us to refine not only the prognosis of *IGHR*-CLLs but also the outcome of low-risk cytogenetic patients. Thus, this mutational analysis improves our understanding of the molecular heterogeneity of CLL patients and could help improve prognostic stratification of CLLs.

Conflict of interest

The authors declare no competing interests

Data Accessibility

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Ethics statement

The present study was approved by the local ethics committee (Comité Ético de Investigación Clínica, Hospital Universitario de Salamanca). Written informed consent was obtained from all participants before they entered the study.

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Author Contribution

CPC, AERV and JMHR designed research. CPC performed the research and statistical analyses, analyzed the data, and drafted the manuscript. MHS analyzed the data and critically reviewed the manuscript. TG selected the samples and individually reviewed all IGHR-CLL cases by FISH and immunophenotypic studies. MQA contributed to the analysis and interpretation of the results and critically reviewed the manuscript. MMI and JMHS proceeded the samples and performed the next-generation sequencing studies. MJV, AGC, CA, MVP, SA, MS, ARM, JD, JRDV, JAQ, and JAHR provided the patients' data. RB and MHS designed the custom-CLL panel and the sequencing studies. AERV and JMHR performed research and critically reviewed and approved the final version of the manuscript.

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Figure legends

Figure 1. Mutational profile of CLL patients with *IGH* rearrangements. A) Mutational frequencies and associations in the CLL cohort according to the presence of *IGH* rearrangements. Significant p/q-values are annotated with asterisks (N=233). B) Each column represents a patient; each row corresponds to a genomic alteration. Patients are clustered according to the *IGHR* (*IGH/BCL2* translocation is indicated in light blue; other *IGH* translocations are shown in dark blue). Missense, frameshift, nonsense, splicing and UTR mutations are reported in red, green, yellow, pink and brown, respectively. The presence of a cytogenetic alteration is shown in gray and the *IGHV* unmutated status is represented in purple (N=46).

Figure 2. Schematic representation of *BCL2*, *IGLL5* and *HIST1H1E* mutations. Positions of coding mutations are indicated according to the aminoacid change at the protein level; positions of UTR mutations are indicated according to the nucleotide change in the DNA sequence (GRCh37/hg19 genome); with respect to the UTR regions, only *BCL2* and *IGLL5* 5'UTR regions were covered in the sequencing analysis (see Supplementary Table S2). Number of cases are denoted by circles in each mutation line and the color of the circles indicates the mutation subtype (missense, frameshift and nonsense). Mutations identified in the COSMIC database in non-Hodgkin lympho mas (NHL) are represented with red lines; mutations reported in the COSMIC database in CLL are indicated with blue lines; all other mutations are shown in black. Abbreviation: Aa, aminoacid.

Figure 3. Clinical impact of IGHR and genetic mutations in CLL patients. A) Kaplan-Meier analysis of TFT according to the presence of *IGH* translocation in CLLs with normal FISH (N=268). Kaplan-Meier analysis of TFT in IGHR-CLL patients with B) any mutation, C) *NOTCH1*, *SF3B1*, *TP53*, *BIRC3* or *BRAF* mutations and D) *IGLL5/BCL2* mutations (N=46). E) Kaplan-meier analysis of TFT in the three risk stratifications subgroups according to the presence of mutations in *NOTCH1*, *SF3B1*, *BIRC3*, *TP53* and *BRAF* genes. In low-risk patients, the presence of mutations in some of

these five genes is significantly associated with shorter TFT (median not reached vs. 24 months, P<0.0001) as well as in the intermediate-risk subgroup (56 vs. 2 months, P<0.0001).

Zable 1.	IGLL5, BCL2 a	nd <i>HISTH1E</i> n	nutations identified in I	GHR-CLLs	(N=46)

Peti	tra	GH/BCL2 anslocation % of cells)	Gene	DNA ^a /cDNA change: AA change	VAF, %	Function	ID COSMIC ^b	Previously described	SIFT/ PolyPhen-2 pathogenicity prediction	Reported as somatic in VarSome	Functional impact ^b ICGC
	0	NO	BCL2	C60985900G	12.74	5' UTR	-	-	-	YES	-
	1	YES (85)	BCL2	C60985900A	33.87	5' UTR	-	-	-	YES	-
لن	1	YES (85)	BCL2	c.G405T:p.E135D	30.15	exonic	-	-	T/P	YES	UNKNOWN
4:	2	YES (60)	BCL2	c.G140A:p.G47D	11.39	exonic	COSM220809	DLBCL ³⁵	T/P	YES	YES
	5	YES (87)	BCL2	c.G589A:p.G197S	40.1*	exonic	COSM5947452	DLBCL/FL ³⁶	-/B	YES	YES
	5	YES (87)	BCL2	c.C175T:p.P59S	37.82*	exonic	COSM4170930	DLBCL/FL ³⁶	T/B	YES	YES
2	8	NO	IGLL5	c.G26C:p.G9A	51.09	exonic	COSM5713869	DLBCL	T/B	YES	-
	5	NO	IGLL5	c.G312T:p.K104N	9.42	exonic	-	-	D/P	YES	YES
3	2	NO	IGLL5	c.G72A:p.W24X	42.86	exonic	-	-	-	-	YES
	6	NO	IGLL5	C23230223-	26.12	5' UTR	-	CLL^{33}	-	-	-
	6	NO	IGLL5	G23230229C	26.51	5' UTR	-	-	-	YES	-
1	1	YES (85)	IGLL5	c.G88A:p.G30S	33.93	exonic	-	-	D/P	YES	YES
1	5	YES (87)	IGLL5	c.T167G:p.V56G	19.88	exonic	-	CLL^8	T/B	YES	YES
3.	5	YES (87)	IGLL5	c.C182T:p.S61F	19.88	exonic	COSM3357314	CLL8/DLBCL	T/B	YES	YES
	5	YES (87)	IGLL5	A23230172C	17.58	5' UTR	-	CLL^8	-	-	-
4.	3	YES (41)	IGLL5	c.G94A:p.A32T	43.48	exonic	COSM5949859	CLL	D/B	YES	YES
()	4	YES (77)	HIST1H1E	c.G515C:p.S172T	42.33	exonic	-	-	D/D	-	-
3′	7	YES (98)	HIST1H1E	c.C500T:p.A167V	41.74	exonic	COSM1292261	FL ³⁹ /CLL ⁷	T/B	YES	YES

^{*}C nfirmed as somatic in the matched CD19- cell fraction.

ositions of UTR mutations are indicated according to the nucleotide change in the DNA sequence (GRCh37/hg19 genome)(reference transcripts: see Suppl. Table S5).

b Hoematopoietic and lymphoid tissue

Ab reviations: AA=aminoacid; VAF=variant allele frequency; DLBCL=diffuse large B cell lymphoma; FL=follicular lymphoma; CLL=chronic lymphocytic leukemia; r=Tolerable; B=Benign; D=Damaging; P=Pathogenic.

UNKNOWN: reported in ICGC database with unknown functional impact in the gene. "-" indicates the variant has not been previously reported in the databases or seminal parters.

Accept

IGH-translocation no IGH-translocation P Characteristic $\boldsymbol{\varrho}$ (N=46)(N=816)Median age at diagnosis, years (range) 0.112^{b} 69 (43-89) 66 (25-97) 0.542 Gender Male, % 0.814 63 63.8 0.698^{c} Median time from diagnosis to FISH, months (range) 1 (0-117) 1 (0-253) 0.568^{b} 0.808 Binet B or C, % 38.6 0.014^{c} 0.039 22.2 Med an YBCa count, ·109/L (range) 17.6 (2.3-196) 17.8 (2.4-964) 0.721^{b} 0.841 Median lymphocytes count, ·10⁹/L (range) 0.874 11.6 (0.6-186) 12.2 (0.8-960) 0.874^{b} Median platelet count, ·10⁹/L (range) 0.808 172 (55-295) 187 (2-587) 0.456^{b} Median l'emoglobin level, g/dL (range) 14.1 (6.6-16.5) 14.2 (4.4-18.9) 0.577^{b} 0.808 High pa-microglobulin level, % 67.4 36.3 <0.0001^c 0.0007 • Higher te dehydrogenase level, % 27.3 15.7 0.027^{c} 0.054 Hepatomegaly, % 7.1 6.9 0.824^{c} 0.852 Sr lenomegaly, % 15.9 16.8 0.852^{c} 0.852 7.9 11.1 0.595^{c} 0.757 Richter transformation 4.3 1.7 0.148^{c} 0.259 Unmutated, % 60.6 44.9 0.025^{c} 0.054 13q deletion, % 26.1 0.0014 43.1 0.0003^{c} trisomy 12, % 34.8 14.5 <0.0001c 0.0007

4.3

6.5

67.4

57 (1-157)

10.9

4.3

44.0

133 (106-159)

 0.426^{c}

 0.334^{c}

 0.002^{c}

 0.155^{b}

0.596

0.520 **0.007**

0.543

Table 2. Clinical and biological characteristics of CLL patients depending on the presence of *IGH* rearrangements (N=862).

Median follow-up, months (range)

11q deletion, %

1-1-1ion, %

Need for treatment, %

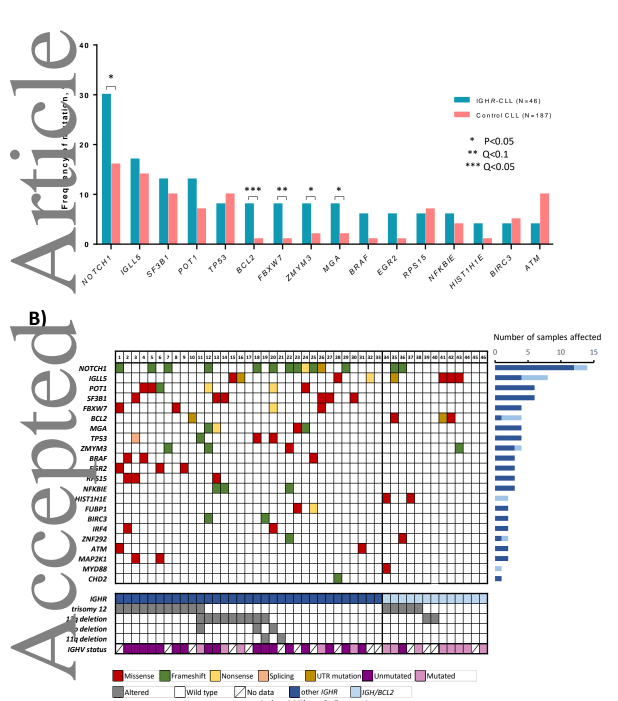
a VI C: hite blood cells

bMann Whitney U test

Table 3. Univariate and multivariate analysis for Time to First Treatment (TFT) in IGHR-CLL patients (N=46).

		Univariate 95% CI				Multivariate 95% CI			
		HR	Lower	Upper	P	HR	Lower	Upper	P
	Male Gender	0.675	0.33	1.37	0.276				
	Binet B/C	0.257	0.12	0.56	0.001	0.558	0.22	1.43	0.221
	CD38 positivity	0.810	0.36	1.82	0.614				
	GHV-unmutated	0.325	0.13	0.79	0.013	0.566	0.20	1.64	0.29
	LDH high	0.726	0.32	1.66	0.448				
	2M high	0.558	0.25	1.27	0.163				
	Hepatomegaly	0.340	0.10	1.16	0.085				
•	Splenomegaly	0.368	0.15	0.88	0.025	0.403	0.14	1.19	0.099
<u> </u>	symptoms	0.483	0.18	1.27	0.141				
	11q deletion	0.421	0.09	1.82	0.246				
-	GH/BCL2 translocation absence	0.443	0.18	1.09	0.076				
'	GLL5/BCL2 mutations absence	0.139	0.03	0.60	0.008	0.821	0.14	4.76	0.828
	TP53 disruption/mutation	0.143	0.04	0.51	0.003	0.304	0.07	1.28	0.105
1	<i>BRAF</i> mutations	0.325	0.09	1.13	0.076				
	NOTCH1/SF3B1/TP53/BIRC3/BRAF mutations	0.204	0.08	0.47	0.0002	0.255	0.07	0.88	0.030
	Presence of mutation	0.238	0.05	10.0	0.051				

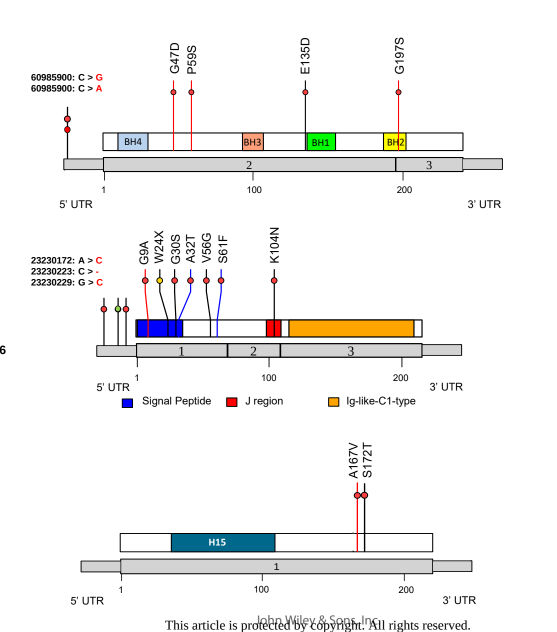
β2M, β2-microglobulin level; LDH, lactate dehydrogenase level.



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BCL2

IGLL5



Missense CLL in COSMIC NHL in COSMIC Frameshift Nonsense No COSMIC Id

Domains (239 aa)

Exons & UTR regions

aa position

Domains (214 aa)

Exons & UTR regions

aa position

Domains (219 aa)

Exons & UTR regions

aa position

