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IKZF1 Deletions with COBL Breakpoints Are Not Driven by RAG-Mediated Recombination Events in Acute Lymphoblastic Leukemia



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Abstract

IKZF1 deletion ($\Delta IKZF1$) is an important predictor of relapse in both childhood and adult B-cell precursor acute lymphoblastic leukemia (B-ALL). Previously, we revealed that *COBL* is a hotspot for breakpoints in leukemia and could promote *IKZF1* deletions. Through an international collaboration, we provide a detailed genetic and clinical picture of B-ALL with *COBL* rearrangements (*COBL-r*). Patients with B-ALL and *IKZF1* deletion ($n = 133$) were included. *IKZF1* 1-8 were associated with large alterations within chromosome 7: monosomy 7 (18%),

isochromosome 7q (10%), 7p loss (19%), and interstitial deletions (53%). The latter included *COBL-r*, which were found in 12% of the *IKZF1* 1-8 cohort. Patients with *COBL-r* are mostly classified as intermediate cytogenetic risk and frequently harbor *ETV6*, *PAX5*, *CDKN2A/B* deletions. Overall, 56% of breakpoints were located within *COBL* intron 5. Cryptic recombination signal sequence motifs were broadly distributed within the sequence of *COBL*, and no enrichment for the breakpoint cluster region was found. In summary, a diverse spectrum of alterations characterizes Δ *IKZF1* and they also include deletion breakpoints within *COBL*. We confirmed that *COBL* is a hotspot associated with Δ *IKZF1*, but these rearrangements are not driven by RAG-mediated recombination.

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Introduction

B-cell precursor acute lymphoblastic leukemia (B-ALL) comprises multiple subtypes defined by structural and numerical chromosomal alterations. These initiating lesions include aneuploidy and chromosomal rearrangements, leading to expression of ectopic fusion proteins and/or to deregulation of gene expression. During the evolution of B-ALL, a series of secondary genomic alterations, such as DNA copy number alterations (CNAs) and sequence mutations, usually emerges [1]. These secondary events commonly involve deletions of genes that encode regulators of cell-cycle (*CDKN2A* and *CDKN2B*) and B-cell development (*PAX5* and *IKZF1*) [2].

IKZF1 deletions occur in ~15% patients with B-ALL and are more recurrent in the *BCR-ABL1* and Philadelphia-like subgroups, affecting more than 70% and 40% of the patients, respectively [3]. *IKZF1* is located within chromosome band 7p12 and codifies the transcription factor IKAROS. It has two main domains: a DNA-binding zinc finger domain (encoded by exons 4-6) and a zinc finger dimerization domain (encoded by exon 8). Some intragenic *IKZF1* deletions impair its DNA-binding domain resulting in a dominant-negative effect (e.g. Δ 4-7), while deletions of the whole gene (Δ 1-8), as well as deletions of exons 1-2 (e.g. Δ 2-3) lead to the gene haploinsufficiency. *IKZF1* deletions have been associated with dismal prognosis in both pediatric and adult B-ALL [4,5], despite the type of deletion [6]. Interestingly, sequencing of *IKZF1* breakpoints suggested that aberrant recombination activating gene (RAG)-mediated recombination is responsible for these deletions [7]. Due to their clinical relevance, the identification of *IKZF1* deletions has become an essential prognostic biomarker over the last decade. Hence, aiming to detect these deletions, we developed a rapid and efficient polymerase chain reaction (PCR)-based method [8]. Recently, by delineating the genetic landscape of *IKZF1* Δ 1-8, we identified *COBL* (*COBL* cordon-bleu *WH2* repeat protein) gene as an originating hotspot for *IKZF1* deletions [9]. Here we established an international collaboration to enable a detailed genetic and clinical characterization of B-ALL with *COBL* rearrangements (*COBL-r*).

Material and Methods

Patients

This study included 146 patients diagnosed with B-ALL and *IKZF1* deletions. This patient cohort comprised two independent groups. The first one included 133 patients with 1) *IKZF1* complete deletion, Δ 1-8 ($n = 104$); or 2) intragenic deletion of *IKZF1* including exon 8, Δ n-8 ($n = 29$). The second group referred to 13 patients already diagnosed with *IKZF1* deletions and *COBL-r* at collaborating centers [8–10].

Patient samples and information were obtained through the cooperative efforts of nine centers worldwide, and the project was conducted in accordance with the declaration of Helsinki.

Detection of *IKZF1* Deletions and CNAs

IKZF1 deletions and CNAs in *EBF1*, *JAK2*, *CDKN2A*, *CDKN2B*, *PAX5*, *ETV6*, *BTG1*, *RBI*, and the pseudoautosomal region 1-PAR1 (*SHOX*, *CRLF2*, *C2F2RA*, *IL3RA*, and *P2RY8*) were determined by either multiplex ligation-dependent probe amplification (MLPA), or single nucleotide polymorphism (SNP) array analyses. The SALSA MLPA P335-(A3-B2) and/or SALSA MLPA P202-B1 (MRC Holland) were used for MLPA experiments, and the data analyses were performed using Coffalyser software. The CytoScan HD Array (Affymetrix) assessed the occurrence of CNAs in the Brazilian (partially), British, and French cohorts. Data were analyzed with Chromosome Analysis Suite software, version 3.2 (Applied Biosystems), and the GRCh38/hg38 build of the Human Genome Assembly.

Characterization of *COBL* Breakpoints

We developed customized MLPA assays for the screening of CNAs within chromosome 7. The probes' design and assay conditions have been previously described [9]. To validate our SNP array and MLPA findings, we performed either multiplexed long-distance PCR (M-PCR) or long-distance inverse (LDI)-PCR. These PCR approaches allowed us to confirm *COBL-r* and to determinate the breakpoints at nucleotide level.

Patient Data and Survival Analyses

For the comparison of laboratory and clinical data between patients with or without *COBL-r*, we used the Pearson χ^2 test. Overall survival (OS) was defined as the time in months from the date of diagnosis to death or to the last follow-up assessment for patients alive. Kaplan-Meier method was used to estimate OS rates of patients according to *COBL* status with differences compared by the log-rank test. Statistical analyses were performed using R software, version 3.5.2, and P values < .05 were considered statistically significant.

Analysis of Activation-Induced Deaminase (AID) or RAG Recognition Signal Sequences in Recombined *COBL* Alleles

An agnostic search for motifs located within fragments spanning 50 bp from the breakpoint junctions of *COBL-r* was performed using MEME. The limit of output motifs was set to 5, and width ranged from 2 to 15 bp. The FIMO (Find Individual Motif Occurrences) tool was used for the analysis of WGCW and CG sequences, and the recombination signal sequence (RSS) consensus sequence was used to search for cryptic RSS sequences [11,12].

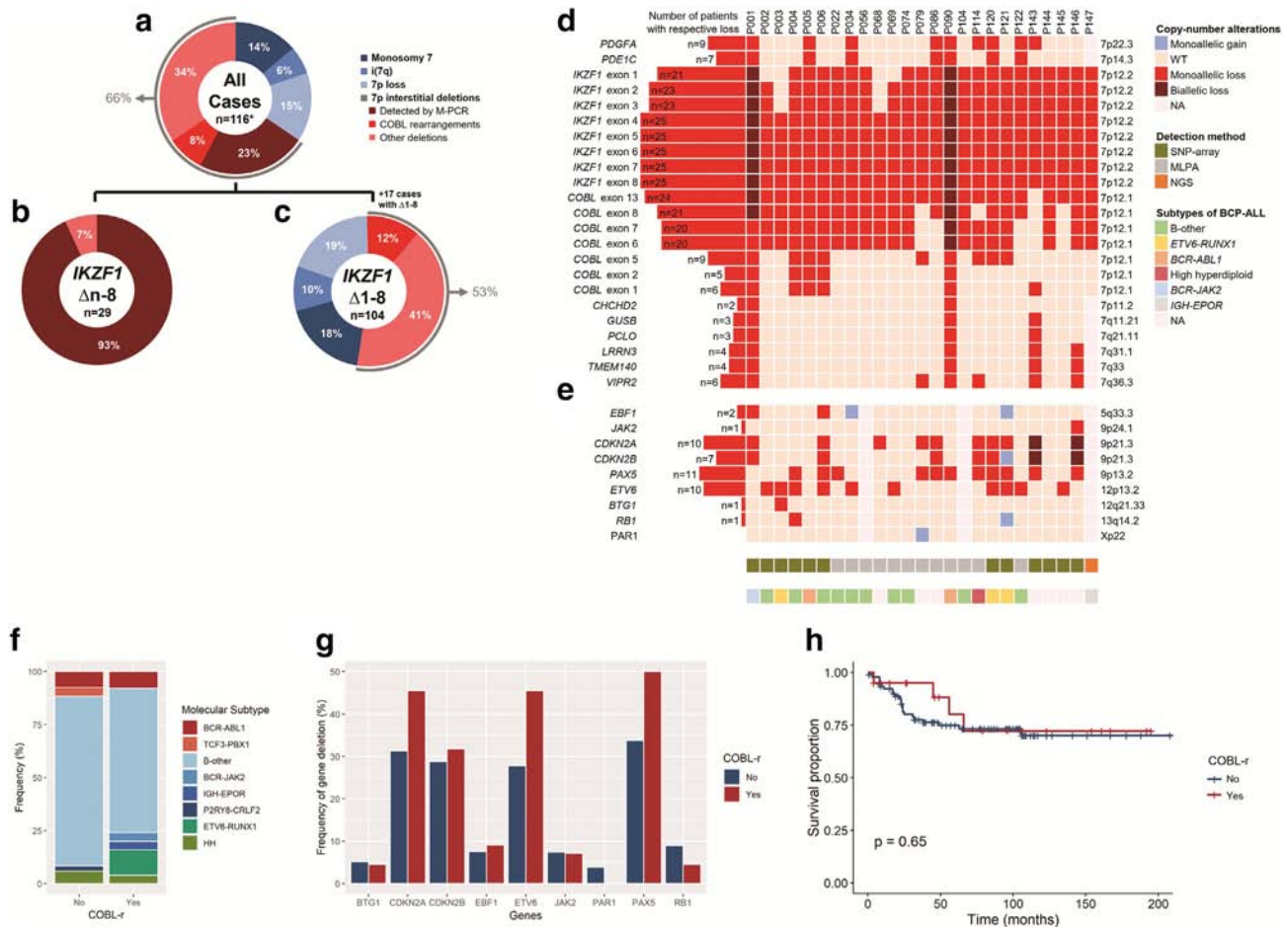


Figure 1. Genetic alterations associated with *IKZF1* deletions in B-ALL. (A) Patients with *IKZF1* deletions spanning its exon 8 presented four types of aberrations on chromosome 7: monosomy 7; isochromosome 7q, i(7q); 7p loss; and interstitial deletions within 7p arm. *One cohort of 17 cases comprising only complete deletions of *IKZF1* was excluded from this analysis because they would lead to a biased estimative of CNAs' frequencies among patients with *IKZF1* deletions overall. Then, patients were divided into two groups: (B) intragenic deletions of *IKZF1* spanning its exon 8 ($n = 29$) and (C) complete deletions of *IKZF1* ($n = 104$). (D) The *COBL* rearrangements were identified in 25 patients. Most of the deletions spanned between *IKZF1* exon 1 and *COBL* intron 5, although they presented variable size. (E) These patients had additional alterations, such as *ETV6*, *PAX5*, *CDKN2A*, and *CDKN2B* deletions. NA, not available. (F) Distribution of molecular subtypes of B-ALL among patients with or without *COBL*-r. Legend colors illustrate alterations associated with high risk (red), intermediate risk (blue), and good risk (green). HH, high hyperdiploidy. (G) The frequency of deletions in genes associated with B-ALL was compared according to *COBL* status. *PAX5*, *CDKN2A*, *CDKN2B*, and *ETV6* deletions were recurrently observed; however, the frequency of these CNAs was similar for all these genes, as indicated by P values above each bar. *PAR1*, pseudoautosomal region 1. (H) No significant difference was observed when comparing the overall survival of B-ALL patients with *COBL*-r vs. *COBL* wild type ($P = .65$).

Results

Most of the deletions encompassing *IKZF1* exon 8 were classified as whole-gene deletions ($\Delta 1-8$: 78% vs. $\Delta n-8$: 22%). The CNAs found in the cohort were associated with monosomy 7 (14%; $n = 16$); isochromosome 7q (6%; $n = 7$); 7p loss (15%; $n = 17$), and interstitial deletions within 7p (66%; $n = 76$) (Figure 1A). Among the $\Delta n-8$ group, most of the alterations (93%) were *IKZF1* deletions detectable by the M-PCR, i.e., $\Delta 2-8$ or $\Delta 4-8$ (Figure 1B). Alternatively, 47% of the $\Delta 1-8$ group presented with aberrations involving loss of the whole 7p, such as monosomy 7 (18%; $n = 19$), isochromosome 7q (10%; $n = 10$), and 7p loss (19%; $n = 20$). The remaining 55 patients (53%) harbored interstitial deletions; *COBL*-r were found in 12% of the $\Delta 1-8$ cohort (Figure 1C).

We identified *COBL*-r in 25 B-ALL cases with *IKZF1* deletions (Figure 1D). *COBL*-r were detected by MLPA screening in 133 patients with B-ALL and *IKZF1* deletions ($n = 12$) or SNP array/NGS investigation performed at collaborating centers ($n = 13$), as

described in the methodology. The loss of genes located within 9p locus — *PAX5* ($n = 11$), *CDKN2A* ($n = 10$), and *CDKN2B* ($n = 7$) — and *ETV6* deletions ($n = 10$) were the most recurrent additional alterations among these patients (Figure 1E). Demographic and laboratory data for the 25 patients with *COBL*-r showed they included 17 males and 8 females who were mainly children and adolescents ($n = 21$), with a median age at diagnosis of 5.5 years (range 1-59 years) and a median white blood cell count of $7.5 \times 10^9/l$ (range $1.5-459.6 \times 10^9/l$) (Table 1). The patients were treated on diverse therapy protocols (Table 2). Seven patients relapsed within a median of 5.5 years (range 1.4-16.3 years), and six of them experienced isolated disease recurrence in the bone marrow ($n = 5$) or testes ($n = 1$). One patient relapsed at both sites: bone marrow and central nervous system. The median follow-up was 5.1 years (range 0.3-16.3 years), and the five deaths (mainly in patients with early relapse) occurred at a median 4.2 years (range 0.3-5.5 years) following diagnosis. In addition, the comparison between patients

Table 1. Demographic and Laboratory Characteristics of B-ALL Cases with COBL Rearrangements.*

Patient	Age (Years)	Gender	WBC ($\times 10^9/l$)	Blasts at BM	5' Breakpoint	3' Breakpoint	Detection Method	Karyotype
P004	1	F	96.000	95%	ELMO1 intron 14	Upstream COBL	SNP array	46,XX,+2,t(3;16)(p21;p13),del(4)(q22),-7,der(12)t(7;12)(q11;p13),-13,+16,-18,-19,+mar1,+mar2
P034	1	M	3.400	98%	RAD50 intron 5	COBL intron 5	LDI-PCR	NA
P086	1	M	5.000	95%	7pter	COBL intron 7	MLPA/LDI-PCR	NA
P120	1	F	459.600	80%	7p14	COBL intron 5	SNP array/M-PCR	NA
P056	2	F	1.600	98%	SPATA intron 1	COBL intron 5	LDI-PCR	46,XX
P122	2	M	13.400	40%	7pter	COBL intron 7	MLPA	NA
P145	2	F	92.000	98%	TCRGC2 intron 1	COBL intron 13	SNP array	47,XX,del(7)(p11),+10,add(12)(p13),del(12)(p11.2)[4]
P022	4	F	7.500	85%	7p12	COBL intron 2-5	MLPA	46,XX
P003	5	M	10.000	95%	IKZF1 intron 3	COBL intron 5	SNP array/M-PCR	46,XY
P074	5	M	16.400	78%	7p12.1	COBL intron 5 [‡]	LDI-PCR	NA
P079	5	F	5.700	88%	7p	COBL intron 8-13	MLPA	NA
P121	5	F	7.470	50%	7p12.2	COBL intron 5	SNP array/M-PCR	NA
P144	5	M	31.000	NA	7p14.2	COBL intron 5	SNP array	47,XY,+X,del(16)(q13),i(17)(q10),ider(21)(q10)dup(21)(q?) [3]/ 47,idem,add(7)(p1)[3]
P002	6	M	115.000	97%	7p12.2	COBL intron 5	SNP array/M-PCR	46,XY,del(6)(q21q25),der(12)del(12)(p1p13)t(12;17)(p13;q11),der(17)t(12;17)
P069	10	F	204.000	89%	IKZF1 intron 1	COBL intron 5	LDI-PCR	NA
P146	11	M	1.500	90%	SPATA intron 2	COBL intron 5	SNP array	48,XY,+X,t(6;20)(p1;q1),t(7;9)(p1;p2),i(9)(q10),+12,der(21)dup(21)(q?) [9]
P068	12	M	NA	NA	IKZF1 intron 3	COBL intron 5	RNA-seq/LDI-PCR	48,XY,+X,del(4)(q25),-7,del(9)del(9)t(4;9)(q25;p13),+21,+mar[cp8]/
P104	13	M	1.680	84%	7p14	COBL intron 5	MLPA	46,XY,del(7)(p14),der(9;12)(q10;q10),+mar[6]
P090	15	M	4.500	83%	7p22	COBL intron 5	MLPA	NA
P114	16	M	55.000	45%	7p	COBL intron 2-5	MLPA	NA
P143	16	M	6.200	96%	COBL intron 1 [†]	7p12.1	SNP array	46,XY,idel(9)(p13)[1]/ 45,idem,-7[3]/ 45,idem,add(2)(p25),-7[9]
P147	Adolescent	M	NA	NA	SUN3 intron 5	COBL intron 5	NGS	NA
P005	21	M	2.820	90%	7pter	Upstream COBL	SNP array	46,XY,t(9;22)(q34;q11)
P006	38	M	3.040	67%	VWC2 intron 3	Upstream COBL	SNP array	46,XY
P001	59	M	58.000	75%	7p12	COBL intron 7 [§]	SNP array	45,XY,-7,t(9;22;15)(p24;q11;q21),add(9)(p13)

BM, bone marrow; LDI-PCR, long-distance inverse PCR; M-PCR, multiplex PCR; MLPA, multiplex ligation-dependent probe amplification; NA, not available; WBC, white blood cell count.

* In this case, blast percent was estimated from peripheral blood.

† Patient had COBL rearrangement and 7p12.3::GRB10 intron 5 deletion.

‡ Patient had an inversion within COBL intron 5.

§ Patient had COBL rearrangement and monosomy 7.

Table 2. Clinical Characteristics and Outcome of B-ALL Cases with COBL Rearrangements

Patient	Clinical Trial	CNS Disease	MRD Day 33	MRD Day 78	PR	CMR Day 33	Relapse	Outcome	Follow-Up (Months)
P147	AALL0232_1	NA	NA	NA	NA	NA	NA	NA	NA
P068	AALL0331/UKALLR3	No	Negative	Negative	NA	Yes	BM	Dead	45
P034	AIEOP-BFM ALL2000	No	Negative	Negative	Good	Yes	No	1st CR	96
P022	AIEOP-BFM ALL2000	No	Negative	Negative	Good	Yes	No	1st CR	79
P114	AIEOP-BFM ALL2000	No	NA	NA	Good	Yes	NA	Dead	NA
P120	AIEOP-BFM ALL2000	No	Positive	Positive	Poor	No	No	Dead	4
P104	ALL IC 2009	Yes	Negative	Negative	Good	Yes	No	1st CR	38
P145	UKALL2003	No	Negative	Negative	NA	Yes	No	1st CR	105
P146	UKALL2003	No	Negative	NA	NA	Yes	BM, CNS	Dead	66
P144	UKALL97	No	NA	NA	NA	Yes	No	1st CR	26
P143	UKALL97	No	NA	NA	NA	Yes	No	1st CR	167
P003	CAALLF01	No	Negative	NA	Good	Yes	NA	1st CR	4
P079	COALL 05-92	No	NA	NA	NA	Yes	No	1st CR	161
P086	COALL 06-97	No	Negative	NA	NA	Yes	Testes	2nd CR	195
P074	COALL 06-97	No	Negative	NA	NA	Yes	BM	2nd CR	154
P069	COALL 07-03	No	Positive	NA	NA	No	No	1st CR	123
P004	EORTC 58081	No	Negative	Negative	Good	Yes	No	1st CR	49
P090	EsPhALL	No	Positive	NA	Good	Yes	Isolated BM	Dead	56
P006	FRALLE 93	No	Positive	Positive	Poor	Yes	Yes	Relapse	192
P002	FRALLE BT	No	Positive	Negative	Poor	Yes	No	1st CR	15
P122	GBTLLALL99	No	NA	NA	Good	Yes	NA	Alive	NA
P121	GBTLLALL99	No	Negative	Negative	Good	Yes	No	Alive	NA
P005	GRAAPH	No	Positive	Positive	Not evaluable	Yes	No	1st CR	27
P056	NA	No	Positive	NA	Good	Yes	BM	Lost follow-up	17
P001	NA	No	Positive	NA	Good	Yes	No	SCT; alive	9

BM, bone marrow; CNS, central nervous system; CMR, complete morphological remission; MRD, minimal residual disease; NA, not available; PR, prednisone response; SCT, stem-cell transplantation; WBC, white blood cell count. MRD-negative status was defined as <0.01% leukemic cells in bone marrow and peripheral blood.

with *COBL*-r vs. *COBL* wild-type revealed that both groups presented similar laboratory and clinical characteristics (Table 3). Among the B-ALL cytogenetic abnormalities, *TCF3-PBX1* and *ETV6-RUNX1* were exclusively found in patients without or with *COBL*-r, respectively (Figure 1F). Although we did not observe any significant difference in the frequency of additional gene deletions when comparing patients with vs. without *COBL*-r, it is worthy of note that *CDKN2A*, *ETV6*, and *PAX5* deletions were more frequent in patients with *COBL*-r (Figure 1G). Follow-up data were available for 111 B-ALL patients with ($n = 20$) or without *COBL*-r ($n = 91$). The OS of patients with *COBL*-r was similar to those with *IKZF1* deletion only (hazard ratio, 1.278; 95% CI, 0.35-4.68; $P = .646$) (Figure 1H).

The breakpoints of the *COBL*-r were determined at nucleotide level in 11 of the 25 cases, and 56% of breakpoints were located within *COBL* intron 5 (Figure 2B). To address the possible causes of these breakpoints, we first performed an agnostic motif search. This analysis identified the motif CASWGTGG (E-value = 0.87) within all 22 breakpoint sequences of *COBL*-r (Figure 2A; Supplementary Table S1). CASWGTG is similar to the heptamer of the RAG RSS, which is composed of a heptamer (5'-CACAGTG-3') and a nonamer (5'-ACAAAAACC-3') sequence, interspaced by 12 or 23 random nucleotides associated to RAG-type rearrangements. Since the nonamer was absent in our first analysis, we then investigated the presence of complete motifs associated with the occurrence of rearrangements in leukemia: cryptic RSS sequence (RAG-type fusions), WGCW (AID-type fusions), and CG sequences. WGCW and CG sequences were not found; however, we identified cryptic RSS in 5 out of 22 breakpoint regions, although none of them were spanning breakpoints within *COBL* (Supplementary Table S2).

Additionally, we performed a robust analysis for the identification of cryptic RSSs within the whole sequence of *COBL* (Figure 2, C-D). The results revealed a broad distribution of this motif throughout the gene and no enrichment for the breakpoint cluster located in intron 5 of *COBL*.

Discussion

In this study, among the Δn -8 group, the majority of the alterations were *IKZF1* deletions which had already been detected by the M-PCR method [8]. Since the remaining Δn -8 samples harbored deletions restricted to *IKZF1*, either they had DNA fusions outside the breakpoint cluster region of *IKZF1* or M-PCR failed to detect them. Among the $\Delta 1$ -8 group, *COBL*-r were found in 12% of the patients. This result revealed that *COBL*-r are more frequent among $\Delta 1$ -8 but rarely related to Δn -8. Although *COBL*-r were not detected within the Δn -8 group for the cases included in the current proposal, we found these deletions in patients with *COBL*-r from a previous study [9]. These patients had *IKZF1-COBL* fusions, which involved *IKZF1* intron 1 ($n = 1$) or intron 3 ($n = 2$) and *COBL* intron 5.

Confirming the idea that *COBL* represents a genomic hotspot for *IKZF1* deletions in B-ALL [9], we identified *COBL*-r in 25 B-ALL cases with *IKZF1* deletions. Although either good (*ETV6-RUNX1*, high hyperdiploid) or high-risk (*BCR-ABL1*) cytogenetic groups have been observed in some of these patients, most of them had the so-called “B-other” subclassification with either normal or other abnormal (*BCR-JAK2*, *IGH-EPOR*) cytogenetic profile. In the current genetic risk stratification, many cases with *COBL*-r would then be classified as intermediate cytogenetic risk. We found that these patients also presented other secondary abnormalities commonly identified in B-ALL, such as *PAX5* and *CDKN2A/B* and *ETV6* deletions. This combination is of special interest because these patients could benefit from the newly proposed combined risk stratification strategies, such as *IKZF1*^{plus} or the UKALL-CNA classifiers [13–15]. Considering that patients with B-ALL share similar laboratory and clinical characteristics regardless of *COBL* involvement, *IKZF1* deletions may play a major role on risk stratification for these patients.

Regarding the breakpoints of the *COBL*-r, it is remarkable that 56% of them were located within *COBL* intron 5, although this region represents only 16% (47,770 out of 300,587 bp) of the entire gene. Based on this observation, we formulated two hypotheses to explain the presence of a hotspot for breakpoints within *COBL*: 1) the production of a truncated *COBL* protein, encoded by exons 1-5 only, could have a role on leukemogenesis, or 2) there is a breakage mechanism involving *COBL* intron 5, thus enriching this area for gene rearrangements.

COBL protein has three Wiskott-Aldrich syndrome protein homology 2 domains for actin binding. It shows substantial expression in neurons and muscle cells, although levels are low in blood [16]. *COBL* functions as an actin nucleator, controlling neuronal morphology and development [17]. Considering that *COBL* does not play a direct role in lymphoid development, the enrichment of *COBL*-r in B-ALL is more likely to be related to mechanisms controlling DNA breakage and promotion of genetic fusions.

Usually, genetic rearrangements in lymphoid malignancies are caused by either AID or the RAG complex. Therefore, we searched for motifs located within the breakpoint junctions of *COBL*-r, which could potentially provide a rational explanation for the observed

Table 3. Demographic and Laboratory Data of Patients with B-ALL *

	<i>COBL</i> Rearrangement		<i>P</i> Value
	No	Yes	
	<i>n</i> = 121	<i>n</i> = 25	
	<i>n</i> (%)	<i>n</i> (%)	
Gender			0.473
Male	48 (39.7)	8 (32.0)	
Female	73 (60.3)	17 (68.0)	
Age at diagnosis			0.701
<1 year	2 (1.7)	0 (0.0)	
1-9 years	49 (41.5)	12 (48.0)	
≥10 years	67 (56.8)	13 (52.0)	
WBC ($\times 10^9/L$)			0.948
<50	85 (70.2)	16 (69.6)	
≥50	36 (29.8)	7 (30.4)	
NCI risk			0.797
Standard	69 (58.0)	14 (60.9)	
High	50 (42.0)	9 (39.1)	
CNS disease			0.741
No	96 (94.1)	23 (95.8)	
Yes	6 (5.9)	1 (4.2)	
Prednisone response			0.143
Good	66 (91.7)	11 (78.6)	
Poor	6 (8.3)	3 (21.4)	
Relapse			0.607
No	75 (70.8)	13 (65.0)	
Yes	31 (29.2)	7 (35.0)	
Outcome			0.604
Alive	94 (77.7)	20 (83.3)	
Dead	27 (22.3)	4 (16.7)	

WBC, white blood cell count; NCI, National Cancer Institute of US; CNS, central nervous system.
* Pearson χ^2 calculation.

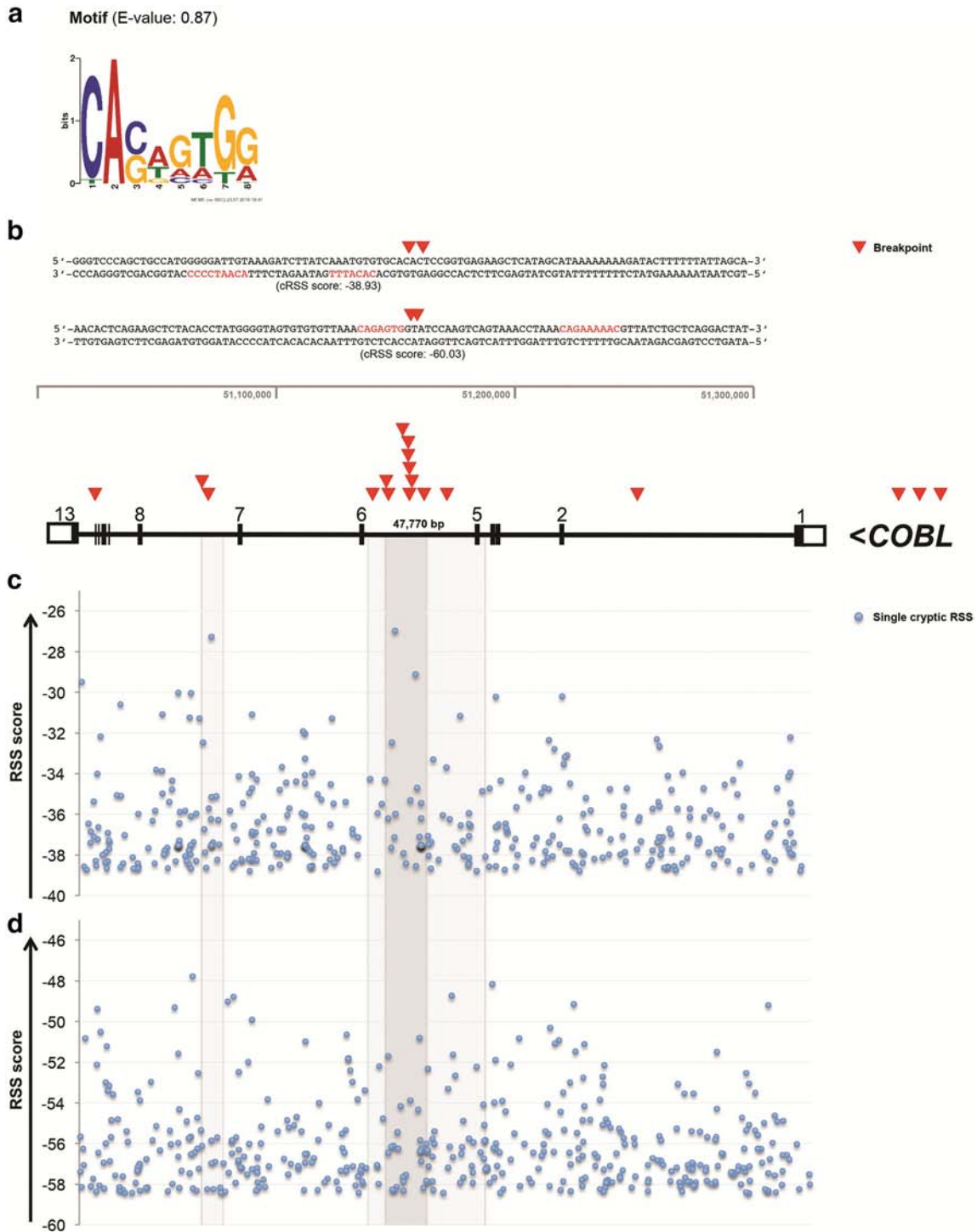


Figure 2. Identification of motifs within the breakpoint sequences. (A) An agnostic motif search using MEME identified the sequence CASWGTGG (E-value = 0.87) among 22 breakpoint sequences. (B) The map of 19 deletion breakpoints (red triangles) within *COBL* revealed a hotspot located at intron 5. Three breakpoints were detected within a downstream region of 7p12.1. The sequences highlight two breakpoint clusters located at *COBL* intron 5. The mapped cryptic recombination signal sequences were not statistically significant. The cryptic recombination signal sequences (cRSS) with a spacer of 12-bp (c) and 23-bp (d) were mapped along *COBL* gene. The highest RIC scores represent cRSS (blue dots) associated with RSS functionality. The gray area highlights the *COBL* intron 5 and breakpoint cluster regions.

chromosomal rearrangements. RSSs are recognized by RAG enzymes during V(D)J recombination, and previous studies have located cryptic RSS immediately internal to the breakpoints of intragenic deletions of *IKZF1*⁷. Although the mutual motif within sequences

spanning the breakpoints was similar to the heptamer sequence, our results do not support the idea that aberrant RAG-mediated recombination is the mechanism responsible for *IKZF1* and *COBL* codeletions.

In summary, our results highlight *COBL* as a hotspot for interstitial deletions within chromosome 7, especially for deletions including the *IKZF1* gene. Most of the *COBL-r* arose within *COBL* intron 5, leading to complete deletion of *IKZF1*; nevertheless, we also observed fusions between both genes. The analysis of the breakpoint sequences revealed a common motif resembling the heptamer of recombination signal sequence, but the analysis of the whole consensus sequence did not provide evidence for RAG-mediated recombination. Lastly, our study demonstrates that patients with *IKZF1* deletions are associated with worse outcome regardless of *COBL-r*.

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Conflict of Interest

The authors declare no conflict of interest.

Author Contribution

Conception and design: B. A. L., C. M., R. M., M. E. Experiments, data analysis, and interpretation: B. A. L., C. M., T. C. B., C. P. P., N. D., M. B., U. S., C. P., N. C. V., R. M., M. E. Provision of samples, data collection, and assembly: B. A. L., M. B. M., N. D., C. J. H., U. S., M. H., M. S. P. O., G. C., R. S., C. N. A., G. T., S. G., S. B. Writing and/or revision of the manuscript: B. A. L., R. M., M. B. M., C. J. H., G. C., R. S., C. N. A., M. E. Final manuscript approval: all authors.

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