CRLF2 and IKZF1 abnormalities in Mexican children with acute lymphoblastic leukemia and recurrent gene fusions: exploring surrogate markers of signaling pathways

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Abstract

The gene fusions BCR-ABL1, TCF3-PBX1, and ETV6-RUNX1 are recurrent in B-cell acute lymphoblastic leukemia (B-ALL) and are found with low frequency in coexistence with CRLF2 (cytokine receptor-like factor 2) rearrangements and overexpression. There is limited information regarding the CRLF2 abnormalities and dominant-negative IKZF1 isoforms associated with surrogate markers of Jak2, ABL, and Ras signaling pathways. To assess this, we evaluated 24 Mexican children with B-ALL positive for recurrent gene fusions at diagnosis. We found CRLF2 rearrangements and/or overexpression, dominant-negative IKZF1 isoforms, and surrogate phosphorlated markers of signaling pathways coexisting with recurrent gene fusions. All the BCR-ABL1 patients expressed CRLF2 and were positive for pCrkl (ABL); most of them were also positive for pStat5 (Jak2/Stat5) and negative for pErk (Ras). TCF3-PBX1 patients with CRLF2 abnormalities were positive for pStat5, most of them were also positive for pCrkl, and two patients were also positive for pErk. One patient with ETV6-RUNX1 and intracellular CRLF2 protein expressed pCrkl. In some cases, the activated signaling pathways were reverted in vitro by specific inhibitors. We further analyzed a TCF3-PBX1 patient at relapse, identifying a clone with the recurrent gene fusion, P2RY8-CRLF2, rearrangement, and phosphorylation of the three surrogate markers that we studied. These results agree with the previous reports regarding resistance to treatment observed in patients with recurrent gene fusions and coexisting CRLF2 gene abnormalities. A marker phosphorylation signature was identified in BCR-ABL1 and TCF3-PBX1 patients. To obtain useful information for the assessment of treatment in B-ALL patients with recurrent gene fusions, we suggest that they should be evaluated at diagnosis for CRLF2 gene abnormalities and dominant-negative IKZF1 isoforms, in addition to the analyses of activation and inhibition of signaling pathways.

Keywords: CRLF2; IKZF1; activated signaling pathways; acute lymphoblastic leukemia; primary rearrangements; Mexican children; TCF3-PBX1 concurrent with P2RY8-CRLF2

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Introduction

B-cell acute lymphoblastic leukemia (B-ALL) is a complex disease with many gene abnormalities and disturbed functional processes; thus, a plethora of genetic lesions and heterogeneous abnormal signaling pathways can be detected [1,2]. The fusions BCR-ABL1, TCF3-PBX1, ETV6-RUNX1, and KMT2A-variant are recurrent genetic abnormalities with biological significance in B-ALL [3]. These gene fusions are initial leukemogenic events that confer self-renewal properties to hematopoietic stem cells or lymphoid progenitors [1].

The novel BCR-ABL1-like B-ALL subtype is recognized as a high-risk subgroup characterized by alterations in kinases or their receptors, resulting in the constitutive activation of signaling pathways. Recurrent gene fusions and BCR-ABL1-like are mutually exclusive [4]. BCR-ABL1-like patients present CRLF2 (cytokine receptor-like factor 2) gene rearrangements in 60% of the cases. The most frequent rearrangements are the P2RY8-CRLF2 fusion, generated by deletion of the pseudoautosomal region PAR1 and present in 25.8% of the BCR-ABL1-like patients, and the IGH-CRLF2 rearrangement, produced by t(X:14)(p11;q32) or t(Y:14)(p11;q32) and observed in 3.5% of BCR-ABL1-like patients [5]. Both abnormalities result in the binding of transcriptional control elements to the coding sequence of CRLF2, leading to overexpression of the encoded protein [2,6,7]. Another hallmark of the BCR-ABL1-like subgroup is the high frequency of abnormalities in the IKZF1 gene (IKZF1a), which are mainly deletions or dominant-negative isoforms [4].

The coexistence of CRLF2 rearrangements and overexpression with recurrent B-ALL abnormalities, such as ETV6-RUNX1, BCR-ABL1, iAMP21, or hyperdiploidy, has been reported. CRLF2 overexpression occurs more frequently in children with hyperdiploidy (19%), and P2RY8-CRLF2 fusion is present in patients with iAMP21 (20%) [8–11]. Although less frequent, the coexistence of CRLF2 rearrangements with the BCR-ABL1 fusion has been observed in adults [10,12–14]. In children, CRLF2 rearrangements have been described coexisting with ETV6-RUNX1, KMT2A-AFF1, and only in one patient with TCF3-PBX1 [14–17]. Notably, CRLF2 rearrangements concurrent with BCR-ABL1 confer to patients’ resistance to the ABL pathway inhibitors; furthermore, some of these patients have been diagnosed as BCR-ABL1-like [18]. Interestingly, the coexistence of recurrent gene fusions with CRLF2 rearrangements in the same blast has been observed mostly in patients of Hispanic and Mexican American origin [18,19].

The IKZF1 gene encodes the transcription factor Ikaros; importantly, in leukemic cells, one of its targets is the CRLF2 promoter [20]. The most frequent somatic abnormality in B-ALL is IKZF1 deletion, which is present in 83% of BCR-ABL1 patients [21]. IKZF1 deletions and point mutations are also frequent among BCR-ABL1-like patients and present in 68% of cases [22]. In contrast, IKZF1a is less common in cases with gene fusions such as TCF3-PBX1 (1–4%) [23,24] and ETV6-RUNX1 (3–6%) [24–26].

As mentioned above, CRLF2 rearrangements, concurrent with other recurrent gene fusions, have been rarely described. It is possible to identify recurrent genetic fusions in combination, either in the same cell or in separate cells of the same patient [14,18,27]. These observations prompted us to describe a group of Mexican children with B-ALL and recurrent gene fusions, searching for CRLF2 rearrangements and overexpression, dominant-negative IKZF1 isoforms (Ik6, Ik8, and IKZF1a), and surrogate markers for Jak2, ABL, and Ras signaling pathways. All these features were selected as they are characteristic of BCR-ABL1-like patients [22].

Materials and methods

Patients

One hundred and thirty-eight bone marrow samples from B-ALL children at diagnosis were collected at the Instituto Nacional de Pediatría and Hospital Infantil de México Federico Gómez in Mexico City, Mexico. Diagnosis of B-ALL was based on morphological characteristics of bone marrow smears together with the immunophenotype analysis of leukemic cells by flow cytometry. Mononuclear cells from bone marrow samples were isolated and used for obtaining suitable biologic material for the genetic and flow cytometry analyses. The detection of gene fusions was performed by reverse transcription polymerase chain reaction (RT-PCR) using a diagnostic kit for 28 leukemia gene fusions (HemaVision RT-PCR, Risskov, Denmark). Genetic and flow cytometry studies were performed in patients with fusions when RNA and/or cells were available. Informed consent was obtained from the guardians of patients according to the Declaration of Helsinki. The project was approved and it followed the guidelines of the Research and Institutional Ethics Committees of the participant Institutions.

Determination of CRLF2 transcript expression

RNA was extracted from bone marrow mononuclear cells using the RNeasy kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. cDNA was obtained by standard methods (Invitrogen, Waltham,
Relative quantification of CRLF2 transcripts was determined by real-time RT-PCR, using a LightCycler 2.0 Instrument (Roche Applied Science, Mannheim, Germany) with TaqMan gene expression probes from the Universal Probe Library System (Roche Applied Science). The primer sets that we used for CRLF2 (NM_022148) were forward primer 5′-AGCGACTGTCAGGTTGA-3′ and reverse primer 5′-AATTTGGACACCTTGGTGGG-3′; the probe we used was #53. Quantification of transcripts was calculated by the ΔΔCt method using the GUSβ gene as an endogenous control for data normalization. The primer sets for GUSβ (NM_000181.3) were forward primer 5′-CGCCCTGCCTATCTGTATTC-3′ and reverse primer 5′-TCCCCACAGGGAGTGTGTGTAG-3′; the probe was #57. Gene expression was analyzed in duplicate. The CRLF2 high expression cut-off value was determined in an independent cohort of 70 ALL patients without recurrent primary alterations. The normalized numerical values of relative expression were divided into high or low CRLF2 expression groups (quartile 1 – quartile 2 versus quartile 3 – quartile 4) and the cut-off value (14.3) was determined by GraphPad Prism 8 software (GraphPad, San Diego, CA, USA) [20,28]. Statistical analysis with analysis of variance (ANOVA) showed that CRLF2 expression in the two groups was significantly different (p = 0.05). It must be noted that in the mentioned cohort the first patient above the cut-off value was positive for IGH-CRLF2, and no patient with this rearrangement was found below this value.

Analysis of the P2RY8-CRLF2 deletion

The deletion was detected by RT-PCR based on the methods described by Palmi et al [29]. To detect the deletion in samples with minor cell subclones, the PCR product of the first reaction was reamplified using the same set of primers.

Analysis of Ik6 and Ik8 transcripts

The transcripts of the dominant-negative isoforms Ik6 and Ik8 were detected by nested RT-PCR, based on the methods described by Iacobucci et al [30,31]. The expected sizes of the PCR products of the Ik8 and Ik6 isoforms are 390 and 255 base pairs, respectively.

Determination of CRLF2 protein expression by flow cytometry

In patients from whom cells were available, flow cytometry studies were performed. CRLF2 protein expression was assessed using an allophtocyocyanin (APC)-conjugated monoclonal antibody that recognizes the extracellular domain (Becton Dickinson, Franklin Lakes, NJ, USA) [32]. Bone marrow cells were stained with cell surface monoclonal antibodies: CD45 Amcyan, CD34 PECy7, CD19 PerCP, and CD10 PE (Becton Dickinson). To identify cell surface CRLF2 or intracellular CRLF2, two different staining conditions were used: CRLF2 antibody incubation in fresh cells and CRLF2 antibody incubation after fixation, and permeabilization with BD Cytofix/Cytoperm® (Becton Dickinson). The antibody concentration and conditions for the assay were performed as per the manufacturer’s recommendations. Flow cytometry was performed using a BD FACSVersace Cell Analyzer System and data were analyzed by FlowJo vX software (Becton, Dickinson and Company, Ashland, OR, USA).

Identification of kinase alterations and inhibition assay by phosphoflow cytometry

To identify Jak2/Stat5, ABL, and Ras pathway alterations in ALL patient samples, fresh cells from bone marrow were stimulated with 200 ng/ml thymic stromal lymphopoietin (TSLP) (PeproTech, Cranbury, NJ, USA) for 30 min at 37 °C, and stained with the following cell surface monoclonal antibodies: CD45 Amcyan, CD34 PECy7, CD19 PerCP, and CD10 PE (Becton Dickinson, USA) [32]. Then, the cells were fixed with BD Phosflow Fix Buffer I, permeabilized with BD Phosflow Perm Buffer III (Becton Dickinson), and incubated with monoclonal antibodies to identify phosphorylated targets of each pathway: Stat5 (pY694) Pacific Blue to identify Jak2, CrkL (pY207) AF488 for ABL, and Erk 1/2 (pT202/pY204) AF488 for Ras. In cases with an adequate number of cells, the abnormal pathway activity was selectively inhibited in vitro for 30 min with ruxolitinib (5 nM) (Selleckchem, Houston, TX, USA) to inhibit Jak2/Stat5 pathway, and with imatinib or dasatinib (5 μM) (Selleckchem) to prevent ABL abnormal activation. The antibody concentration and conditions for each assay were performed based on the manufacturer’s recommendations. Flow cytometry was performed using a BD FACSVersace Cell Analyzer System and data were analyzed by FlowJo vX software.

FISH studies in a patient with TCF3–PBX1 at relapse

In this patient, a previous conventional cytogenetic analysis with GTG bands was performed to determine the t(1;19)(q25;p13). The LSI 1ptel/1p36/1q35 and ToTelVysion mix #14 including p and q subtelomeres of chromosome 19 by Vysis-Abbott (Abbott Park, IL, USA) probes were used to detect the t(1;19) by
fluorescence in situ hybridization (FISH). The P2RY8-CRLF2 fusion was detected by single-sequence double-color probes (Cytocell, Cambridge, UK).

Results

CRLF2 rearrangements

One hundred and thirty-eight bone marrow samples were collected from B-ALL patients at diagnosis; 24 patients presented recurrent gene fusions (Table 1). We analyzed CRLF2 expression, the P2RY8-CRLF2 fusion, and cell surface and intracellular CRLF2 protein in these patients. Of these 24 patients, 5 were positive for the P2RY8-CRLF2 fusion, 8 presented CRLF2 overexpression, CRLF2 protein was detected on the cell surface in 11, and 14 cases had intracellular CRLF2 protein; 3 of these patients were not analyzed for cell surface protein. A total of 19 of 24 patients had the P2RY8-CRLF2 fusion, CRLF2 overexpression, or CRLF2 protein expression, and the following recurrent gene fusions: 6 of 6 BCR-ABL1-positive patients, 9 of 12 cases with TCF3-PBX1, and 4 of 6 patients with ETV6-RUNX1 (Table 1).

P2RY8-CRLF2 fusion

None of the patients with BCR-ABL1 presented the P2RY8-CRLF2 fusion; however, 4 of 12 patients with TCF3-PBX1 and 1 of 6 cases with ETV6-RUNX1 were positive.

CRLF2 expression

CRLF2 overexpression was found in 4 of 6 patients with BCR-ABL1, 3 of 12 with TCF3-PBX1, and 1 of 6 with ETV6-RUNX1 (Table 1 and Figure 1A). Thus, CRLF2 overexpression was more frequent in patients with BCR-ABL1 than in those with other fusions (*p = 0.0178, one-way ANOVA test, GraphPad Prism 8 software).

CRLF2 protein

The CRLF2 protein was determined in patients with available samples (16/24, 66%) (Table 1). Interestingly, most patients presented CRLF2 protein in blasts, on the cell surface, and within cells (Table 1 and Figure 1A).

Table 1. CRLF2 abnormalities in children with B-ALL and recurrent gene fusions.

| Patients | Primary abnormality | P2RY8-CRLF2 | CRLF2 gene expression | CRLF2 protein (%) | On cell surface | Intracellular | Ik6 and Ik8 Isoforms |
|----------|---------------------|-------------|-----------------------|-------------------|----------------|-------------|---------------------|
| L1       | Neg                 | High        | 1.54                  | Neg               | ND             | ND          | Ik6, Ik8            |
| L2       | Neg                 | High        | 4.18                  | High              | 26             | Neg         |                     |
| L3       | Neg                 | High        | ND                    | ND                | ND             | Neg         | Ik6                |
| L4       | Neg                 | High        | ND                    | ND                | Neg            | Ik6, Ik8    |                     |
| L5       | Neg                 | Low         | 13.6                  | High              | 69.5           | Ik6         |                     |
| L6       | Neg                 | Low         | ND                    | ND                | 100            | Neg         | Ik6                |
| L7       | Pos                 | High        | 1.7                   | Low               | 64.8           | Neg         |                     |
| L8       | Neg                 | High        | 2                     | Low               | Neg            | Ik6, Ik8    |                     |
| L9       | Neg                 | Low         | 3                     | 34                | Neg            | Ik6         |                     |
| L10      | Pos                 | Low         | 23.2                  | 90.8              | Neg            | Ik6         |                     |
| L11      | Pos                 | ND          | 1.7                   | 91.5              | Neg            | Ik6         |                     |
| L12      | Neg                 | Low         | 10.7                  | 46.7              | Neg            | Ik6         |                     |
| L13      | Neg                 | Low         | 3.5                   | 63.2              | Ik6, Ik8       |             |                     |
| L14      | Pos                 | Low         | 92                    | ND                | Neg            | Ik6         |                     |
| L15      | Neg                 | High        | ND                    | Neg               | Neg            | Ik6         |                     |
| L16      | Neg                 | Low         | Neg                   | ND                | Neg            | Ik6         |                     |
| L17      | Neg                 | Low         | ND                    | ND                | Neg            | Ik6         |                     |
| L18      | Neg                 | Low         | ND                    | ND                | Neg            | Ik6, Ik8    |                     |
| L19      | Pos                 | Low         | ND                    | ND                | Neg            | Ik6, Ik8    |                     |
| L20      | Neg                 | High        | ND                    | ND                | Neg            | Ik6         |                     |
| L21      | Neg                 | Low         | 91.7                  | 98.8              | Neg            | Ik6, Ik8    |                     |
| L22      | Neg                 | Low         | ND                    | 100               | Neg            | Ik6, Ik8    |                     |
| L23      | Neg                 | Low         | ND                    | ND                | Neg            | Ik6, Ik8    |                     |
| L24      | Neg                 | Low         | ND                    | ND                | Neg            | Ik6, Ik8    |                     |

Cells in gray represent positive results.
ND, not determined; Neg, negative; Pos, positive.

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In BCR-ABL1 patients, cell surface CRLF2 protein was present in a lower percentage of blasts (1.54–13.6%) than the intracellular protein (26–100%). Similar results were observed in the TCF3-PBX1 patients; cell surface CRLF2 protein was present in 1.7–23.2% of blasts and the intracellular protein was present in a high proportion (34–92%). Within the ETV6-RUNX1 group, patient L21 presented a high percentage of blasts positive for both CRLF2 cell surface and intracellular protein (91.7 and 98.8%) and patient L22 showed 100% of blasts with intracellular CRLF2 protein.

All the patients positive for cell surface and intracellular CRLF2 protein were analyzed for CRLF2 expression and presented high or low expression levels (Table 1).

Surrogate markers of signaling pathways

Fifteen patients were analyzed for surrogate markers of Jak2, ABL, and Ras signaling pathways (Table 2). All BCR-ABL1 patients analyzed for these surrogate markers were positive for CRLF2 protein. Most of the BCR-ABL1 patients (3/4) were positive for Stat5 phosphorylation. As expected, all these patients were positive for phosphorylation of the main target of the ABL pathway, CrkL, and were negative for Erk1/2, the Ras pathway’s main target.

Nine patients in the TCF3-PBX1 group were analyzed; eight were positive for CRLF2 protein and for Stat5 phosphorylation. Unexpectedly, seven of nine cases were positive for CrkL and two were positive for Erk 1/2 (Table 2 and Figure 2A). Interestingly, one TCF3-PBX1 patient (L15) negative for CRLF2 protein was negative for the analyzed surrogate markers. In one patient with TCF3-PBX1 (L14), the Stat5 phosphorylation was reverted in vitro using the specific inhibitor ruxolitinib (Table 2 and supplementary material, Figure S1).

Two patients with ETV6-RUNX1, who expressed CRLF2 protein, were analyzed for surrogate markers of the ABL pathway; patient L21 was positive for CrkL phosphorylation, reverted in vitro with imatinib and dasatinib (Table 2 and supplementary material, Figure S1). None of the patients of this subtype were positive for Stat5 phosphorylation.

IKZF1 dominant-negative isoform transcripts

We found a high frequency of dominant-negative IKZF1 isoform transcripts in all the patients (13/24, 54.1%). The Ik6 and Ik8 isoforms were observed...
particularly in four of six BCR-ABL1 cases, either in
the same patient or independently. The isoforms
were present in 6 of 12 cases with TCF3-PBX1 and
3 of 6 cases with ETV6-RUNX1 (Table 1).

Patients’ evolution
Patients’ follow-up time is less than 3 years. At
present, three of the patients with CRLF2 rearrange-
ments or overexpression and phosphorylated surrogate
markers have developed early relapses and died; one
of them was BCR-ABL1 positive (L2) and two showed
TCF3-PBX1 fusion (L7 and L13). It was possible to
study a 2-year-old male diagnosed with B-ALL at
relapse, who was positive for TCF3-PBX1. The sample
at diagnosis was not available to evaluate CRLF2
rearrangements, overexpression, or signaling path-
ways. The patient was treated based on the St. Jude
Total Therapy XIII-B protocol and relapsed 2.4 years
later presenting the same fusion as at diagnosis. We
determined by FISH in metaphases the coexistence of
TCF3-PBX1 and P2RY8-CRLF2 fusions in the same
cell at relapse (see supplementary material, Figure S2).
The patient was positive for CRLF2 surface protein
(15%) and phosphorylation of Stat5 (93%), CrkL
(90%), and Erk 1/2 (66%). Ruxolitinib and dasatinib
inhibited two signaling pathways; Ras signaling inhi-
bition was not determined (see supplementary material,
Figure S2).

Discussion
We identified 24 B-ALL Mexican children positive for
recurrent gene fusions: BCR-ABL1, TCF3-PBX1, or
ETV6-RUNX1. These patients were assessed for CRLF2
rearrangements, transcript and protein, dominant-
negative IKZF1 isoforms, and phosphorylation of surro-
gate markers of Jak2, ABL, and Ras signaling pathways.

Concurrence of CRLF2 abnormalities with B-ALL
gene fusions
The co-occurrence of B-ALL recurrent gene fusions with
CRLF2 rearrangements and overexpression is an uncom-
mon finding and only 20 patients have been previously
reported (see supplementary material, Table S1). Most of
these cases have been detected by screening CRLF2
overexpression and/or rearrangements in B-ALL children
or adults. BCR-ABL1 is the most frequent fusion found
concurrent with CRLF2 abnormalities; this condition
confers an aggressive disease progression with clones
resistant to treatment with ABL inhibitors [18,19].

CRLF2 rearrangements and overexpression are com-
monly determined in the BCR-ABL1-like B-ALL subtype,
but not in B-ALL patients with recurrent gene fusions,
who are not screened for CRLF2 abnormalities [10,12–
19]. This exclusion criterion could underestimate the fre-
quency of patients with the coexistence of recurrent gene
fusions, and CRLF2 rearrangements and overexpression.

Table 2. Analysis of surrogate markers of Jak2/Stat5, ABL, and Ras pathways in patients with B-ALL and recurrent gene fusions.

| Patients | Primary abnormality | Pathway activation |
|----------|---------------------|--------------------|
|          |                     | Stat5 (pY694)      |
|          |                     | CrkL (pY207)       |
|          |                     | Erk 1/2 (pY204)    |
| L1       | BCR-ABL1            | Low                |
| L2       | Low                 |
| L5       | Med                 |
| L6       | Med                 |
| L7       | Low                 |
| L8       | High                |
| L9       | Low                 |
| L10      | Med                 |
| L11      | Med                 |
| L12      | Low                 |
| L13      | Med                 |
| L14*     | Low                 |
| L15      | Neg                 |
| L21†     | ETV6-RUNX1          | ND                 |
| L22      | Neg                 |

Cells in gray represent positive results.
Low, 1-30%; Med, 31-60%; High, 61-100%; ND, not determined; Neg, negative.
*Inhibited with ruxolitinib.
†Inhibited with imatinib and dasatinib.
A

Jaks/Stat5 Pathway

BCR-ABL1

CD19

ABL Pathway

CD19

Ras Pathway

CD19

TCF3-PBX1

Sta5 (pY694)

C-KIT (pY559)

Erk (pT202/pY204)

ETV6-RUNX1

Sta5 (pY694)

C-KIT (pY559)

Erk (pT202/pY204)

B

Blasts

BCR-ABL1

CD19

Dp 22.0

C-KIT (pY559)

Dp 13.6

TCF3-PBX1

Sta5 (pY694)

Dp 72.8

Blasts CRLF2−

Blasts CRLF2+

Sta5 (pY694)

C-KIT (pY559)

Dp 22.5

Dp 64.1

Sta5 (pY694)

C-KIT (pY559)

Dp 68.3

Dp 68.3

Figure 2. Legend on next page.
The incidence of B-ALL among populations, the frequencies of genetic variants for B-ALL susceptibility, and recurrent gene fusion rates are influenced by ethnic differences [33–35]. According to our literature review, the coexistence of BCR-ABL1 fusion with CRLF2 rearrangements and overexpression could be more common in Mexican American patients (see supplementary material, Table S1). Therefore, it is essential to consider that ethnicity could influence these findings [13,18,19]. Another related fact is that the incidence of childhood B-ALL in Mexicans (79.8/million) is much higher compared to other populations [36,37], and Hispanic children have worse outcomes and the lowest survival rates [33,38,39]. In addition, overrepresentation of CRLF2 rearrangements and overexpression has been observed in the B-ALL Hispanic population [7]. Besides the natural history of each genetic subtype of B-ALL, the Mexican population’s genetic background may impact patients’ recurrence with the coexistence of CRLF2 abnormalities and recurrent gene fusions [7,18,19]. Further studies in a higher number of Mexican B-ALL patients and in non-Hispanic B-ALL populations must be performed to prove this hypothesis.

Surrogate markers to evaluate signaling pathway activation in patients with CRLF2 abnormalities and recurrent gene fusions

It has been observed that in vitro TSLP stimulation of blasts carrying CRLF2 rearrangements or overexpression activates multiple signaling molecules, including Stat5, Erk, and Src [40]. In the context of B-ALL recurrent gene fusions, coexisting with CRLF2 abnormalities, we found phosphorylation of Stat5, CrkL and Erk.

Three BCR-ABL1-positive patients were positive for CrkL and Stat5 phosphorylation in the same group of blasts (Figure 2B). These findings could be associated with CRLF2 overexpression through the activation of CRLF2-IL7Ra by the TSLP ligand, which induces the Jak2/Stat5 pathway increasing blast proliferation and survival. Another plausible option is that BCR-ABL1 p190 protein, the most frequent isoform in ALL patients, contributes to the direct activation of CrkL, but also of Stat5 through Jak2 [41]. One BCR-ABL1 patient (L6) was negative for CrkL activation, and a possible explanation is that both CRLF2 overexpression and JAK2 mutation are required for phosphorylation of Stat5; unfortunately, the mutational status of JAK2 was not determined [40]. Our results suggest crosstalk and a synergistic effect between BCR-ABL1 and CRLF2 to activate the Jak2/Stat5 pathway [42,43], which could be prone to induce chemoresistance [18].

One ETV6-RUNX1 patient was positive for P2RY8-CRLF2 fusion, but not for CRLF2 overexpression (L19). This fusion might be present in a minor clone, not being enough to allow the detection of CRLF2 overexpression; unfortunately, a sample for FISH analysis was not available to determine the size of clones [44]. As in BCR-ABL1-positive patient L6, patient L22 presented intracellular CRLF2 protein in a high percentage of the blasts; however, it was negative for Stat5 phosphorylation. This is probably related to the absence of a JAK2 mutation [40]. Patient L21 was positive for CrkL phosphorylation. This finding is not common in the ETV6-RUNX1 context. Nevertheless, we cannot discard a genetic abnormality in a member of the ABL pathway. An ETV6-RUNX1-positive patient with a BCR-ABL1 fusion as a secondary abnormality has already been reported [45]. Although this fusion is not present in the L21 patient, other ABL-type mutations may cause the pathway activation.

Among the eight TCF3-PBX1 patients with CRLF2 protein, four presented P2RY8-CRLF2 fusion and Stat5 phosphorylation; this finding suggests that Jak2-Stat5 activation occurs through the CRLF2-IL7Ra-TSLP pathway. Besides, seven of eight patients with CRLF2 expression showed CrkL phosphorylation. In a conditional E2a-Pbx1 mice model, it has been reported that this fusion is responsible for Src protein-tyrosine kinase family overexpression. These kinases are downstream targets of functional pre-BCR (B-cell receptor), a distinctive feature of human TCF3-PBX1 leukemia, and two of its main targets after pre-BCR stimulation are CrkL and PLCg2 [46,47]. Therefore, the two patients negative for CrkL phosphorylation might be pre-BCR negative [48,49]. Besides the pathway activation findings, these patients could present other mutations as part of their leukemia’s natural history. Mutations in several members of the Jak/Stat pathway (Jak1, Jak3, Ptpn11, and Il7r genes) and the Ras pathway (Kras and Nras) have been reported in the E2a-Pbx1 mouse

Figure 2. (A) Surrogate markers of the Jak2/Stat5, ABL, and Ras signaling pathways analyzed by phosphoflow assay in the blast populations of patients with coexistence of CRLF2 rearrangements and B-ALL gene fusions. (B) Coexistence of phosphorylated markers of the Jak2/Stat5 and ABL (double positive, DP) signaling pathways through the phosphoflow assay in the blast populations of BCR-ABL1- and TCF3-PBX1-positive patients.
model [1]. Interestingly, patients L10 and L11 showed phosphorylation for all the analyzed markers, including Erk; this characteristic is related to aggressiveness as, in the mouse model, Ras mutations conferred a shorter latency to develop leukemia [1].

It is impossible to ascertain if these results will be statistically significant, given the limited number and short time of patients’ follow-up; however, our findings could be relevant for treatment selection. To investigate this possibility, we studied a relapse patient positive for TCF3-PBX1. Our results are in line with other reports regarding resistance to treatment in patients with B-ALL and the coexistence of BCR-ABL1 and CRLF2 abnormalities [18]. A similar situation could be present in this relapsed patient and the other TCF3-PBX1-positive cases we examined (Tables 1 and 2).

In T-cell acute lymphoblastic leukemia (T-ALL) patients, Stat5 phosphorylation was induced by TSLP, although CRLF2 protein was localized intracellularly [32]. It is unknown if there is activation through the intracellular protein, but it is possible that the surface receptor, even in low quantity, could be capable of inducing the signaling pathway through a low amount of ligand entering into the cell [32,50]. Considering that CRLF2 surface protein was present in a low fraction of blasts, proteomic characterization of intracellular CRLF2 is required to determine if a mutation prevents the protein from reaching the cell surface or if it is a soluble isoform [51–53]. The transcript was detected at high or low expression levels in almost all the patients with the cell surface or intracellular CRLF2 protein. The cases with low expression could be explained by the presence of mutations or single-nucleotide variants modifying the CRLF2 transcript translation rate or half-life [54,55].

A group of patients showed high expression of CRLF2 but were negative for P2RY8-CRLF2 fusion. The CRLF2 overexpression may have been caused by a nondetermined IGH-CRLF2 rearrangement, CRLF2 or JAK2 mutations, or Ck2 kinase overactivity [46].

Figure 3. Three-step model of B-ALL pathogenesis. This model postulates (1) an initiating genetic lesion, in this case TCF3-PBX1 or BCR-ABL1, that confers self-renewal properties to hematopoietic stem cell or lymphoid progenitors; (2) a second lesion, such as IKZF1 deletion or P2RY8-CRLF2 fusion, causing differentiation block at progenitor B-cell level; and (3) requirement for a third class cooperating mutation to fully transform leukemia cells, affecting pathways such as cytokine receptors (CRLF2 mutation) and/or Ras signaling activation.
it has been reported that expression of the dominant-negative IKZF1 isoforms, Ik6 and Ik8, can cause overexpression of CRLF2 as IKZF1 acts as a negative transcriptional regulator of this gene. The expression of CRLF2 is negatively regulated through epigenetic changes produced by IKZF1; thus, the loss of function of IKZF1 is in part responsible for the high expression of CRLF2. In our patients, the dominant-negative IKZF1 isoforms were particularly frequent (54%) [20,47].

Here, we report a new group of B-ALL Mexican children with recurrent gene fusions in coexistence with CRLF2 rearrangements and/or overexpression, and with phosphorylation of surrogate markers of Jak2, ABL, or Ras signaling pathways. In contrast to previously reported cases, most of our patients are TCF3-PBX1 positive. In BCR-ABL1 and TCF3-PBX1 patients, a signature of signaling pathway marker phosphorylation was identified, where gene fusions are the initial genetic alteration and confer self-renewal properties to lymphoid progenitors [1,48]. We observed second lesions affecting B-cell development’s essential transcription factors, such as the dominant-negative IKZF1 isoforms or P2RY8-CRLF2 fusion. Finally, CRLF2 gene alterations or Ras pathway activation were detected as a third class of cooperating lesions to fully transform leukemia cells and affecting functions such as cytokine receptors and associated kinases [1,27,56]. Thus, we propose for our patients this multistep model of B-ALL pathogenesis following the findings previously reported in cell lines and mice models [1,40] (Figure 3).

As stated in the TCF3-PBX1-positive patient at relapse, the analyses of CRLF2 rearrangements and expression, in addition to phosphorylation of surrogate markers of diverse signaling pathways, contribute relevant information to identify cell clones that could be resistant to treatment and are prone to be treated with specific inhibitors of Src, Ras, Ck2, and/or Jak2. These inhibitors could be combined with conventional chemotherapy to reduce leukemic cells’ viability [57] or could be used to sensitize the cells to commonly used therapeutic agents [58,59]. To obtain useful information for B-ALL patients with recurrent gene fusions, we suggest assessing them at diagnosis for CRLF2 gene abnormalities, activation of signaling pathways, and in vitro inhibition.

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Author contributions statement

DML conceived the study; designed, performed, and interpreted the flow cytometry experiments; drafted the manuscript; and designed the figures. MdRJV conceived the study; designed, performed, and interpreted CRLF2 molecular experiments; drafted the manuscript; and designed the figures. ARL designed, performed, and interpreted molecular experiments (IKZF1). DMA designed and performed molecular (CRLF2 and IKZF1) and FISH experiments; partial results of this study are part of his Master’s degree dissertation (Posgrado en Ciencias Biológicas, Universidad Nacional Autónoma de México). AHM performed CRLF2 molecular experiments. CSL analyzed the data and worked on the manuscript. MdPNM supported results interpretation and worked on the manuscript. MZT and LJV diagnosed the patients and obtained clinical data. BJR performed FISH and cytogenetic analysis; partial results of this study are part of her Specialist in Human Cytogenetics Thesis (Instituto Nacional de Pediatría, Secretaría de Salud). RCC and RPA recruited patients. MHA acquired clinical data and developed a database. PPV conceived the study, interpreted results, wrote the manuscript, and designed the tables.

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SUPPLEMENTARY MATERIAL ONLINE

Figure S1. Phosphoflow analysis of the in vitro inhibition assay of Jak2/Stat5 and ABL pathways in blast populations from TCF3-PBX1 and ETV6-RUNX1 patients

Figure S2. Patient with TCF3-PBX1 at relapse

Table S1. B-ALL patients with coexistence of CRLF2 abnormalities and gene fusions reported in the literature and in this study