A B-cell acute lymphoblastic leukemia regulatory network defines novel therapeutic targets in *IGH-CRLF2* patients

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ABSTRACT:

CRLF2 overexpression in B-ALL patients with an *IGH-CRLF2* translocation activates JAK-STAT, PI3K and ERK/MAPK signaling pathways. Although inhibitors of these pathways are available, investigating alternate targets could reduce treatment-associated toxicities. Comparing RNA-seq from *IGH-CRLF2* and non-translocated patients we defined a translocation gene signature. Next, we assembled a B-ALL cancer-specific regulatory network using 529 B-ALL patient samples from the NCI TARGET database coupled with priors generated from ATAC-seq peak TF-motif analysis. The network was used to infer differential changes in TF activities predicted to control *IGH-CRLF2* deregulated genes, thereby enabling identification of translocation-associated pathways and potential new therapeutic targets.

INTRODUCTION:

Acute lymphoblastic leukemia (ALL) is the most common cancer in children¹. Historically, clinical criteria have driven risk stratification for these patients, however over time many genetic alterations have been identified as prognostic predictors. Several groups have described varied genetic signatures associated with pediatric ALL with the aim of elucidating their contributions to leukemogenesis^{2,3}. Improved risk stratification based on genetic signature has altered treatment and led to significant improvements in overall survival¹. However, about 20% of patients fail current treatment strategies or die following relapse. Moreover, adults with ALL have a worse prognosis and an average overall survival of 35-50%¹. Therefore, it is important to gain a better understanding of the mechanisms by which genetic alterations drive leukemogenesis to refine therapies that target the disease-essential pathways involved⁴.

Genetic alterations that lead to overexpression of the cytokine receptor-like factor 2 (*CRLF2*) gene have been associated with a high-risk subset of pediatric patients with B-cell acute lymphoblastic leukemia (B-ALL). The *CRLF2* gene encodes the thymic stromal lymphopoietin receptor (TSLPR) which forms a heterodimer with IL-7 receptor alpha (IL7RA) to bind TSLP⁵. Binding of TSLP to the IL7RA-TSLPR complex signals the phosphorylation of Janus kinase 1 (JAK1) and Janus kinase 2 (JAK2), leading to the activation of the JAK-STAT signaling pathway⁶. Studies have shown that stimulation of TSLP in B-ALL not only induces activation of the JAK-STAT pathway, but also activates the PI3K/mTOR⁷ and ERK/MAPK signaling pathways¹.

CRLF2 overexpression occurs in 5-15% of patients with B-ALL and in 50-60% of pediatric B-ALL patients with Down syndrome^{6,8}. CRLF2 overexpression in B-ALL can occur either from a chromosomal translocation between CRLF2 and the immunoglobulin heavy chain locus (IGH) on chromosome 14 (IGH-CRLF2) or from an interstitial deletion of the pseudoautosomal region of the X/Y chromosomes resulting in the fusion of CRLF2 to the P2RY8 gene (P2RY8-CRLF2)⁹. CRLF2 deregulation very rarely occurs via activating mutations

of *CRLF2*. The *IGH-CRLF2* translocation occurs in precursor cells and is thought to result from aberrant rejoining during V(D)J-recombination^{10,11}. This translocation is most commonly found in adolescents and adult patients and is typically associated with a poor prognoses, while the *P2RY8-CRLF2* fusion is found in younger patients¹.

CRLF2 chromosomal alterations are often accompanied by mutations in *JAK1*, *JAK2* and *Ikaros* (*IKZF1*) genes. Several groups have suggested that aberrant *CRLF2* signaling cooperates with mutant JAK and IKZF1 activity to promote the development of leukemia^{9,12,13}. As a result, the focus has shifted towards the use of signal transduction inhibitors (STIs) to target JAK-STAT, PI3K and MAPK signaling pathways¹. Although, STIs have shown promise in early clinical trials¹⁴⁻¹⁶, broad application of signal transduction inhibitors is challenging due to the interconnected roles of their targets in biological processes (i.e. JAK kinases) including immunity and hematopoiesis¹⁷. Moreover, it has been shown that mutated JAK2 is required for the initiation of leukemia, but it is not necessary for its maintenance ¹⁸. Therefore, there is a need to more closely investigate the genome-wide impact of the *IGH-CRLF2* alteration in B-ALL to aid in the identification of novel therapeutic targets.

To do so, we first sought to identify transcriptional regulators that control differentially expressed genes associated with the *IGH-CRLF2* translocation through a comparative analysis of translocated versus non-translocated (Non-T) *IGH-CRLF2* B-ALL samples in both patients and cell lines. While robust genome-wide changes in gene expression were separately observed in patients and cell lines, only a small subset of changes were consistent amongst the two groups. Thus, we chose to define the *IGH-CRLF2* associated gene set using only patient samples. We constructed a B-ALL transcriptional network to define the interactions between transcription factors (TFs) and the genes they regulate 19-21 using 529 B-ALL patient samples from the NCI TARGET database. We inferred the targets of TFs linked to the gene set associated with the *IGH-CRLF2* cohort using the Inferelator algorithm 22,23, along with RNA-seq, ATAC-seq, and TF-motif analysis. The network was then used to predict differential transcription

factor activity (TFA) in *IGH-CRLF2* versus non-translocated (Non-T) samples. This approach enabled the identification of ten potential regulators of differentially expressed genes (including DNMT1, EGR1, FOXP1, ZBTB7A). The differentially expressed gene targets of these TFs are enriched in anticipated and novel *CRLF2*-associated pathways. Transcription of several of these gene targets (*IL1RAP*, *LEF1*, *PPP1R13B*, etc.) change across all patients and cell line samples. It is of note that these genes have been implicated in both Acute myeloid leukemia (AML)^{24,25} and ALL^{26,27}. Thus, it is plausible that the genes along with their regulators could contribute to the maintenance of leukemia and be potential candidates for therapeutic targeting in *IGH-CRLF2* B-ALL. The network-based approach we applied has been similarly implemented in several other systems to infer regulatory interactions that have been experimentally validated²⁸⁻³¹.

RESULTS:

Genome-wide transcriptional changes in primary IGH-CRLF2 patient samples

Traditional chemotherapy is non-specific and targets rapidly dividing cells. As a result, toxicity results in injury to healthy cells, causing further morbidity and at times, can be dose-limiting ³². To reduce overall toxicity and improve prognosis, most research has been directed towards understanding the underlying molecular pathology of the leukemia. In this study, we focused on identifying pathways associated with the *IGH-CRLF2* translocation (**Fig. 1a**) with the goal of finding new potential therapeutic targets in this subset of B-ALL patients.

Gene expression profiles associated with the *IGH-CRLF2* translocation were identified by comparing RNA-sequencing from 17 primary B-ALL patient samples, with the translocation (n=13) with non-translocated (Non-T, n=4) patients. Differential analysis performed using DESeq2³³ uncovered a total of 1,179 de-regulated genes with an adjusted p-value of less than 0.01 and |log₂FoldChange| >1 (**Fig. 1b**). Of these 507 genes (~43%) were up-regulated and 669 (~57%) down-regulated in the *IGH-CRLF2* translocated patients. To determine whether the

gene expression changes were localized to the translocated chromosomes we analyzed the percentage of differentially expressed genes per chromosome. The results clearly demonstrate that expression changes are distributed across all chromosomes and there is no enrichment of deregulated genes on chromosome 14 or the pseudoautosomal region of X/Y (Supplementary Fig. 1a).

To further analyze the changes in gene expression between the *IGH-CRLF2* and the Non-T patient samples we performed a principal component analysis. This revealed a clear separation between the two groups of patients (**Supplementary Fig. 1b**). In addition, hierarchical clustering of the 1,179 differentially expressed genes not only separates the *IGH-CRLF2* and Non-T samples, but also clearly divides the *IGH-CRLF2* patients into two groups, referred to as Group 1 and Group 2 (**Fig. 1c**). These two groups could not be distinguished by the presence or absence of activating mutations in *JAK* and *IKZF1* which were found in most of the translocated cohort: 12/13 and 10/13 *IGH-CRLF2* patients, respectively (**Fig. 1c**). As expected the *IGH-CRLF2* translocation clearly results in the up-regulation of *CRLF2* (log₂FoldChange= 5.53) in the translocated versus Non-T control patient samples as shown by RNA-seq tracks on IGV (**Fig. 1d**), suggesting activation of JAK-STAT signaling. Furthermore, *SOCS6*, a suppressor gene of cytokine signaling³⁴, is statistically significantly down-regulated (log₂FoldChange= -1.68) in *IGH-CRLF2* samples (**Supplementary Fig. 1c**), further supporting the observation that JAK-STAT^{1,4} signaling is activated in these samples.

Using the log2 fold changes of the 1,179 differentially expressed genes, ingenuity pathway analysis (IPA) was performed. Positive and negative z-scores indicate pathways that are activated or repressed respectively in the *IGH-CRLF2* samples. The IPA analysis (**Supplementary Fig. 1d**) identified 20 significant pathways, the majority of which (19/20) had negative z-scores, indicating that the majority of the pathways are repressed. The PI3K and ERK pathways were included in the repressed cohort which was unexpected as activation of these pathways is normally linked to *CRLF2* overexpression⁷, Additionally, the JAK-STAT

signaling pathway did not emerge as being statistically enriched according to the gene set we defined, despite the fact that translocated patients have activating mutations in *JAK* genes. One possible explanation for this outcome is that heterogeneity between patient samples leads to noise in gene expression changes which blurs the analysis. In summary, analysis of RNA-seq data in primary patient samples identifies many significant transcriptional changes associated with the translocated *IGH-CRLF2* chromosomal alteration. Furthermore, hierarchical clustering of the patient samples using the differentially expressed gene list, not only clearly distinguishes *IGH-CRLF2* patients from Non-T, but also defines two distinct *IGH-CRLF2* group of patients.

Differential analysis between the two distinct Group 1 and Group 2 patients results in 456 differentially expressed genes. Although pathway analysis results in no significant pathway enrichment we identified six differentially expressed genes involved in p53 signaling, including the proto-oncogene *MDM2* and p53 regulator *MDM4* that are down-regulated in Group 1 (**Fig. 1e**). We postulated there maybe a link between p53 signaling and relapse but were not able to confirm whether p53 signaling in one subtype provides any advantage in this context, as there were 25% (1 of 4) Group 1 patients that relapsed, compared to 33% (3 of 9) of patients in Group 2.

IGH-CRLF2 patient samples have limited changes in chromatin accessibility

To further investigate the impact of the *IGH-CRLF2* translocation, we hypothesized *IGH-CRLF2* transcriptional changes could be accompanied by changes in chromatin accessibility. To test this hypothesis, we performed ATAC-sequencing and compared chromatin accessibility in *IGH-CRLF2* versus Non-T patient samples (18 *IGH-CRLF2* and 6 Non-T). These patient numbers are different to the numbers used for RNA-seq analysis as some of the patient samples for RNA-seq did not pass quality control. Differential analysis of all the ATAC-seq peaks identified 162 regions that were more accessible and 126 regions with reduced accessibility in the *IGH-CRLF2* translocated condition (**Fig. 2a**). The majority of ATAC-seq

peaks were not significantly differential, indicating accessibility is for the most part stable between *IGH-CRLF2* and Non-T patient samples. Significant accessibility changes were evenly distributed across promoters, gene bodies, and intergenic regions (**Supplementary Fig. 2b**).

Differential peaks were assigned to promoters (within 3kb of TSS), UTRs, exons and introns of genes (Fig. 2b). About 82.6% (238) of differential ATAC-seq peaks were associated with genes (n=224), either on promoters or gene bodies. Of the genes that had at least one differential ATAC-seq peak in the promoter or gene body, 14.2% (32) were significantly differentially expressed in IGH-CRLF2 versus Non-T patient samples. The majority of the differential ATAC-seq peaks that overlap differentially expressed genes were regulated in the same direction, with the exception of 7 ATAC-gene pairs (Fig. 2b). In the example shown in Fig. 2c, three significantly more accessible peaks were associated with overexpressed DPP4 in IGH-CRLF2 patients. Conversely, the down-regulated PKIA gene was linked to a significant reduction in accessibility (Supplementary Fig. 2c). We also observed several instances where a significant change in accessibility was linked to a gene whose expression was not affected by the IGH-CRLF2 translocation (Supplementary Fig. 2d). Importantly, the majority of transcriptional changes were not associated with significant changes in chromatin accessibility near the promoter or within the gene bodies (Fig. 2d), as demonstrated for the RRAS gene and two other examples shown in Fig. 2e, Supplementary Fig. 2e-f. Thus, we conclude that overall, the IGH-CRLF2 translocation leads to limited genome-wide chromatin accessibility changes and accessibility remains stable even at promoter regions of differentially expressed genes. However, we postulated that differences in transcription factor (TF) binding at stable ATAC-seg peaks could influence gene expression changes and turned to patient-derived cell lines as a model to address this question.

Transcriptional changes in patient *IGH-CRLF2* derived cell lines are not recapitulated in patient samples

The patient-derived cell lines used in our analysis were MHH-CALL-4³⁵ (CALL) and MUTZ5³⁶ (MUTZ) that harbor *IGH-CRLF2* translocations, and a control non-T pre-B leukemic cell line, SMS-SB³⁷ (SMS). As seen in the RNA-seq tracks of **Fig. 3a**, *CRLF2* is clearly overexpressed in the *IGH-CRLF2* translocated CALL and MUTZ cell lines, compared to the Non-T SMS control. Principal component analysis (PCA) of cell line RNA-seq samples (**Supplementary Fig. 3a**) indicates that the majority of the variance (71%) lies between *IGH-CRLF2* cell lines and Non-T cell lines, while about 26% of variance separates the two translocated cell lines. Thus the PCA analysis indicates that the *IGH-CRLF2* translocation is the major cause of differences between the two conditions, consistent with what was observed in patient samples.

Differential gene expression analysis of CALL versus SMS (3,045 DE genes) and MUTZ versus SMS (2,702 DE genes) identified thousands of differentially expressed genes (**Fig. 3b**), with roughly equivalent numbers of up and down-regulated genes in each case. The number of overlapping differentially expressed genes (ie those likely to be related to the translocation event) is shown in **Fig. 3c**. As with the patient samples, the gene expression changes were distributed across all chromosomes (**Supplementary Fig. 3b**).

As shown in **Fig. 3d**, the log₂ fold change of (CALL/SMS) and (MUTZ/SMS) demonstrated that about 97% of the genes were in convergent orientation (960 up-regulated, 772 down-regulated). However, of the common 1,732 convergent differentially expressed genes in the cell lines, only 221 overlapped with the differentially expressed genes found in translocated versus non-T patient samples (**Fig. 3e**). Furthermore, only 181 out of the 221 genes were in convergent orientation (**Fig. 3f**), indicating that the profile of gene expression changes in cell lines may not be the best representation of what is occurring in patients. Furthermore, no significantly enriched pathways were identified from the overlapping gene set.

To identify the gene list that best separates the translocated class from the non-translocated class, we performed PCA using the differentially expressed genes from cell lines

(Set 1) and patient samples (Set 2). The PCA results associated with the 1,792 differentially expressed genes from Set 1, demonstrate a clear separation between translocated cell lines and non-translocated cell lines. In contrast, the patient samples cluster together and do not separate according to the PC scores calculated using Set 1 genes (Fig. 3g – upper panel). In contrast, when the PCA associated with Set 2 was performed (1,176 differentially expressed genes in patients), we found a clear separation between cell lines and patients on the first principal component that separates the *IGH-CRLF2* from Non-T condition on the second principal component, indicating that Set 2 genes are more representative of the effect of the translocation in both primary patient samples and cell line samples (Fig. 3g – lower panel). In summary, we found many significant transcriptional changes in cell lines but only a subset of these were recapitulated in patients. On the other hand, the top 500 patient-associated transcriptional changes can be used to not only distinguish between patients and cell lines (PC1), but also clearly separate the *IGH-CRLF2* samples from the Non-T samples (PC2).

Chromatin accessibility changes in *IGH-CRLF2* translocated cell lines are more numerous than those in patient samples

Our analyses reveal little compatibility in expression changes between *IGH-CRLF2* and Non-T in cell lines and patient samples. To investigate this further we analyzed chromatin accessibility using ATAC-sequencing. Significant ATAC-seq peaks were called and a reference peakome of 41,111 peaks was created to include all possible ATAC-seq peaks across the three cell lines, CALL, MUTZ, and SMS. DESeq2 analysis on the cell line peakome identified 2,550 altered ATAC-seq peaks between CALL and the SMS control, and 2,718 between MUTZ and the SMS control (Fig. 4a). The overlap between the two comparisons identified 1,396 differentially accessible ATAC-seq peaks (Fig. 4b), of which 99.7 % (1392/1396) were in convergent orientation (Fig. 4c). This number is incompatible with the total number of differentially accessible ATAC-seq peaks (288) identified in patient samples. However, we

postulated that regions that were changing in patients should change similarly in cell lines. To determine this, we ranked all differential ATAC-seq peaks in patient samples (**Fig. 2a**) according to fold change and adjusted p-value, and took the 162 most accessible peaks and 126 least accessible peaks in *IGH-CRLF2* patient samples and calculated the ATAC signal of the cell lines at these regions. The limited alterations in accessibility in patients were weakly recapitulated in cell lines as shown in **Fig. 4d** However, compared to the stability of ATAC-seq peaks in patients, we observed more alterations in accessibility in cell lines. This finding further supports the conclusion that using cell lines as a model to study the *IGH-CRLF2* translocation is not ideal. Therefore, all the downstream analysis focused solely on the transcriptional changes identified in patient samples.

Construction of a B-ALL regulatory network using ATAC-motif derived priors

CRLF2 overexpression activates a signaling cascade that involves many TF regulators and gene targets. Although, we identified a subset of gene expression changes that could be important for the pathogenesis of the leukemia, it is not clear how all of these genes are regulated and connected. Here, our aim was to first identify which TFs could potentially be regulating the differentially expressed genes we identified, and second to infer the relationship between these TFs and their target genes. To address this, we performed TF-motif analysis, using FIMO³⁸, at ATAC-seq peaks that fall within promoter regions (20kb upstream of TSS) of all genes genome-wide. We used a hyper-geometric test to test for enrichment of TF-motifs at differentially expressed genes, which resulted in 102 unique significant TF motifs enriched at significantly up- and down-regulated genes with four motifs DMRTC2, DUXA, PITX1, and PITX3 were found in both sets (Fig. 5a). Thus, motif enrichment analysis resulted in a list of 102 unique potential TF regulators that could be important for the regulation of the *IGH-CRLF2* associated gene signature.

To better understand the relationship between the candidate TF regulators identified in Fig. 5a and the genes they potentially regulate, we sought to infer a transcriptional regulatory network using the Inferelator algorithm^{22,23}. The main limitation of network inference is the sample size, to model transcriptional interactions in a B-ALL specific context requires a large dataset that included hundreds of B-ALL patient samples, not limited to samples with an *IGH-CRLF2* rearrangement. Making use of the TARGET initiative we analyzed 529 B-ALL RNA-seq samples and obtained a normalized gene expression matrix that could be used for the construction of a B-ALL specific regulatory network.

Recent studies have incorporated prior information of TF target genes from different data types, like ChIP-seq, ATAC-seq, and TF-motif analysis to considerably improve network inference ^{28-30,39}. Here, we focused on combining chromatin accessibility data together with TF-motif analysis of the *IGH-CRLF2* cohort to generate priors and infer the B-ALL network (**Fig. 5b**). We selected only regulatory interactions of TFs that were significantly enriched at promoters of differentially expressed genes between *IGH-CRLF2* and Non-T patient samples (102 TFs – **Fig. 5a**). Transcription factor activities estimated using the ATAC-seq motif derived priors and the gene expression matrix obtained from the TARGET database were used for the Inferelator algorithm (**Fig. 5b**). Finally, a B-ALL specific regulatory network involving 102 TFs and 37,086 interactions with combined confidences > 0.5 was inferred. The number of gene targets inferred for each individual TF is shown in **Supplementary Fig. 4a**, and the number of differentially expressed gene targets in the *IGH-CRLF2* cohort are shown in **Supplementary Fig. 4b**.

Defining a sub-regulatory network with TFs affected by CRLF2 alteration

Transcriptional regulatory networks shed light on the relationship between transcription factors (TFs) and their gene targets. Here, we constructed a B-ALL specific regulatory network involving 102 TFs that may be important regulators of the *IGH-CRLF2* gene signature. To

narrow down the list, we created a sub-network and identified TF regulators in the IGH-CRLF2 cohort that were altered either at the mRNA level or the protein activity level. First, we analyzed the expression levels of all genes encoding the 102 TF regulators and identified five TF regulators that were significantly differentially expressed between IGH-CRLF2 and Non-T patients (Supplementary Fig. 4c). Next, we determined TFs that had differences in their activity in the IGH-CRLF2 cohort. Transcription factor activity (TFA) is commonly estimated from mRNA levels, however since posttranslational modifications can influence TF activity, use of mRNA as a proxy for TFA is not the best approach. If prior knowledge of interactions involving TFs and their target genes is available it can considerably improve TFA estimation and network inference^{28,30}. Thus, to estimate the activities of the 102 TFs identified from motifs of significantly up- and down-regulated genes in the IGH-CRLF2 versus Non-T patients, a normalized gene expression matrix was used. This was combined with priors of TF-gene interactions identified from the inferred B-ALL regulatory network (Fig. 5b). Sixteen TFs had significant differences in the mean estimated TFA in the IGH-CRLF2 cohort (Supplementary Fig. 4b) and five had significantly altered levels of expression giving a total of twenty significant TF regulators (ZNF713 was identified in both analyses (Fig. 5c). TFs with no significant differences in mean TFA are shown in Supplementary Fig. 5a. We note that many TFs with significantly altered activity are implicated in cancer. For instance, FOXP1, a member of the forkhead family of transcription factors is known to play important roles in B-cell development and lymphoid malignancies⁴⁰ and could therefore potentially contribute to tumorigenesis in B-ALL patients with the IGH-CRLF2 alteration.

We previously suggested that heterogeneity between patients could lead to noise in the *IGH-CRLF2* gene signature. To filter gene expression changes we focused on inferred target genes of TFs that are affected by *CRLF2* overexpression, either at the transcriptional level or TF activity level. The 6,545 regulatory interactions in **Fig. 5c** describe the relationship between significantly differential TFs and their gene targets. Of these, 387 interactions are regulatory

interactions involving differentially expressed genes (329 out of a total of 1177 unique genes - 28%) in the *IGH-CRLF2* versus Non-T patient cohort (**Fig. 6a,b**). We define the filtered list of 329 differentially expressed genes and their twenty TF regulators as the *CRLF2* specific subnetwork (**Fig. 6c**). The number of differentially expressed gene targets for each TF regulator is shown in **Supplementary Fig. 6a.** As seen in **Supplementary Fig. 6b**, several differentially expressed gene targets are regulated by more than one TF.

Differentially expressed gene targets of TFs with predicted altered activity are enriched in expected and novel *CRLF2*-associated pathways

We performed pathway analysis focusing on the differentially expressed gene targets in the *CRLF2*-altered sub-network (**Fig. 7a**) and identified 21 enriched pathways including the anticipated activation of canonical JAK-STAT, PI3K, and ERK pathways. Thus, in contrast to our initial analysis which focused on all differentially expressed genes in the *IGH-CRLF2* translocated versus Non-T patient samples, we were able to identify known *CRLF2*-associated pathways including the JAK-STAT signaling pathway. These findings validate the approach taken in order to identify a more *CRLF2*-relevant differentially expressed gene list. In **Fig. 7a**, differentially regulated genes (33) are labeled and connected with the appropriate pathway. For example, PI3K up-regulation of signaling is linked to the differential expression of *SOCS6*, *RALB*, *RRAS*, and *PTPN6*. Additionally, the FLT3 signaling ^{41,42} pathway, which is known to be associated with *CRLF2* overexpression, is identified as being upregulated.

The other pathways identified by our analyses could also be playing an important role in *CRLF2*-overexpressing B-ALL. As such, we wanted to define a handful of interesting genes that could have the highest potential as therapeutic targets. For this, we compared all 33 differentially expressed genes in the patient samples and in each pairwise cell line comparison (CALL versus SMS, and MUTZ versus SMS). Although, we previously found that patient-derived transcriptional changes and cell line-derived transcriptional changes do not strongly overlap,

there are some patient-derived transcriptional changes that are recapitulated in the cell lines. Thus, we compared log2 fold changes and significance of the 33 differentially expressed genes across all patient and cell line comparisons (**Fig. 7b**) and identified six common genes. Upregulation of *IL1RAP*, *LEF1*, *CD79B*, *CTGF*, and repression of *PPM1H*, *PPP1R13B* was robust across all *IGH-CRLF2* patient and cell lines. Previous studies^{24-27,43,44} implicate the majority of these genes in hematologic malignancies but their affect on *IGH-CRLF2* B-ALL is not known. Therefore, analysis of the changes in TF activity and gene target mRNA level in the *CRLF2* affected sub-network identified a list of genes that are strong candidates for targeted therapies in *IGH-CRLF2* translocated leukemia.

Discussion

Patients with B-ALL who carry the IGH-CRLF2 translocation are at increased risk for refractory disease and relapse. This alteration leads to *CRLF2* overexpression and activation of JAK-STAT and other associated pathways¹. Available treatments include non-specific cytotoxic chemotherapies which are often effective but responsible for many of the toxicities seen in these patients. The addition of Tyrosine-Kinase Inhibitors and JAK inhibitors in the treatment of leukemias has allowed for targeted treatments with fewer toxicities, however, alternate pathway-specific therapies are needed to further tailor treatment. To better study the impact of the *IGH-CRLF2* translocation, we analyzed RNA-seq from rearranged *IGH-CRLF2* and Non-T B-ALL patient samples and identified hundreds of differentially expressed genes, but were unable to link these changes to the expected pathway alterations associated with the *IGH-CRLF2* translocation. We subsequently analyzed ATAC-seq data to evaluate the effect of this genetic alternation on DNA accessibility. We found accessibility to be mostly stable near the promoters of genes with altered gene expression and conclude that chromatin accessibility is not driving changes in the transcriptional landscape. Instead, we inferred binding of specific proteins at accessible sites at differentially expressed gene promoters that influence transcription.

Leveraging hundreds of available B-ALL RNA-seq samples from the TARGET database along with prior knowledge of TF-motifs at accessible promoters, we inferred a global B-ALL network. Deeper investigation of the TF regulators and their targets in this network identified a candidate list of potential therapeutic targets.

IL1RAP, one of the significantly overexpressed candidates we identified in *IGH-CRLF2* rearranged patients, encodes a component of the interleukin complex. Upregulation of this gene is known to occur in Acute myeloid leukemia (AML), another lymphoid malignancy associated with poor prognosis²⁵. Using antibodies targeting the IL1RAP receptor expressed on the surface of immature AML cells, Agerstam et al. demonstrated clear antileukemic effects in xenograft models.. Targeting IL1RAP also blocks IL-1 signaling and inhibits proliferation of human AML cells²⁵. Using antibodies, RNA interference, and deletion of *IL1RAP*, a more recent study demonstrated it was possible to inhibit pathogenesis *in vivo* and *in vitro* without disrupting normal hematopoietic function²⁴. *IL1RAP* plays an important role in potentiating AML cells, and there is strong evidence for its therapeutic effects in AML. However, its function in B-ALL is not known.

Other candidates, *LEF1* and *PPP1R13B* have also been implicated in cancer. Lymphoid enhancer-binding factor 1, *LEF1*, is a transcription factor that acts downstream of the Wnt/β-catenin signaling pathway. This TF can independently regulate gene expression and is necessary for stem cell maintenance and organ development²⁶. Disrupted *LEF1* is associated with cancer progression and proliferation of cells in ALL, chronic lymphocytic leukemia (CLL), Burkitt lymphoma (BL), and colorectal cancer (CRC)²⁶. In particular, one study clearly shows high expression of *LEF1* associated with shorter relapse-free survival (RFS) in B-ALL²⁷. Overall, this gene is considered a biomarker for patient prognosis in many hematological malignancies. Though, we now provide evidence suggesting that it has a specific role in leukemogeneis in translocated *IGH-CRLF2* patients. Additionally, *PPP1R13B*, another candidate target, encodes a member of the apoptosis stimulating p53 family of proteins (ASPP), this gene is known to be

repressed in ALL and its reduced expression is due to hypermethylation of the *PPP1R13B* gene promoter^{43,44}.

DNA methylation is associated with transcriptional repression and is maintained by the DNMT1 protein. The DNMT1 motif was one of the top regulators identified via motif enrichment analysis at differentially expressed gene promoters. The role of DNMT1 as a transcription factor is not clear, however, it still may have regulatory potential through its function in maintaining DNA methylation and in particular for patients with the *IGH-CRLF2* translocation. For instance, Loudin et al. studied transcription and methylation profiles in overexpressing *CRLF2* ALL patients with Down syndrome. Their methylation profiles indicate high methylation levels correlate with reduced gene expression including genes involved in cytokine-receptor interactions⁴⁵. We hypothesize that methylation could be playing a role in this context.

Overall, we have investigated the transcriptional impact of *CRLF2* overexpression in a high-risk subset of B-ALL patients. Using an integrative approach, we derived regulatory interactions that have identified strong candidates for targeted therapies in B-ALL patients with an *IGH-CRLF2* translocation or other leukemias. *IL1RAP*, *LEF1*, and *PPP1R13B* are amongst the most interesting candidates as there is strong evidence for their role in hematological malignancies. The strategy we have used here can further be adapted to elucidate important regulators responsible for additional subsets of B-ALL and other challenging pediatric malignancies.

Methods

All methods are available in supplemental note section of supplemental information

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Author contributions

B.C, S.B, R.B and J.S designed and managed the project. B.C and P.L collected samples and

performed experiments. S.B performed data analyses. D.C, R.R, and A.W aided in analysis

design. B.C, R.B, and J.S secured funding. S.B and J.S wrote the manuscript. B.C, R.B and J.S

revised the manuscript.

Declaration of Interests

The authors declare no competing interests.

Figure Legends

Fig. 1. Genome-wide transcriptional changes in primary IGH-CRLF2 patient samples. a,

Schematic of IGH-CRLF2 translocation between chromosome 14 and chromosome X/Y. b,

Volcano plot of the differentially expressed genes (FDR=5%, |log₂ fold change| >1) between

IGH-CRLF2 patients and Non-T patients. Red points and blue points correspond to up- and

down regulated genes (n=507 and 669, respectively) in the IGH-CRLF2 patient samples. c,

Heatmap of 1,176 differential expressed genes from the 17 patient samples, with known

information corresponding to B-ALL associated variants and other metadata indicated at the top

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of the heatmap. **d**, Screenshot of RNA-seq tracks for Non-T, *IGH-CRLF2* Group 1, and *IGH-CRLF2* Group 2 at *CRLF2* locus. **e**, Volcano plot of the differentially expressed genes between *IGH-CRLF2* Group 1 Group 2 patients. Red and blue points correspond to up- and down-regulated (n=77 and 379, respectively).

Fig. 2. IGH-CRLF2 patient samples have limited changes in chromatin accessibility. a,

Heatmap of 162 more accessible ATAC-seq peaks in IGH-CRLF2 patients (purple bar) and 126

less accessible ATAC-seq peaks in IGH-CRLF2 patients. b, Differentially accessible peaks

(n=36) linked to differentially expressed genes (n=32) and the log2 fold change of gene

expression (x-axis) plotted against the log₂ fold change of ATAC-seq reads (y-axis). Linked

genes are labeled and colored according to genomic annotation. c, IGV screenshot of ATAC-

seq tracks at up-regulated DPP4 gene across all patients. The highlighted region indicates

significantly more accessible ATAC-seq peaks. d, Average ATAC-seq signal of Non-T (blue)

and IGH-CRLF2 (red) patients at 631 ATAC-seq peaks at up-regulated gene promoters in IGH-

CRLF2 patients (upper panel), and 706 ATAC-seq peaks at down-regulated gene promoters in

IGH-CRLF2 patients (lower panel). e, IGV screenshot of ATAC-seq tracks at up-regulated

RRAS gene across all patients, orange highlighted region indicates ATAC-seq region with no

significant difference.

Fig. 3. Transcriptional changes in patient IGH-CRLF2 derived cell lines are not

recapitulated in patient samples. a, IGV screenshot of RNA-seq tracks at CRLF2 gene on

chromosome Y in CALL, MUTZ, and SMS cell lines. CALL and MUTZ samples harbor the IGH-

CRLF2 translocation whereas the control cell line SMS does not. b, Number of differential

expressed genes between IGH-CRLF2 cell line samples (CALL and MUTZ) compared to SMS

control, red and blue bars indicate up- and down-regulated genes in CALL and MUTZ. c.

Overlap of differentially expressed gene sets between CALL and MUTZ compared to SMS

control (1792 common differentially expressed genes) **d**, Scatter plot of the 1,792 common genes plotting \log_2 fold change of CALL and MUTZ (y and y-axis, respectively) and \log_2 fold change of over SMS control. **e**, Overlap of differentially expressed genes (221) between cell lines (Set 1) and patient samples (Set 2). **f**, Scatterplots of \log_2 fold change of RNA-seq counts of translocated cell lines (CALL and MUTZ) over SMS control (x-axis) plotted against \log_2 fold change of *IGH-CRLF2* patient over Non-T patient RNA-seq counts (y-axis). Red and blue points indicate common convergent up- and down-regulated genes (n=120 and 61, respectively) in cell line and patient samples. **g**, Principal component analysis of all patient and cell line samples computed according to the differentially expressed genes in cell lines (Set 1) (upper panel) and patients (Set 2). Cell lines and patient samples are clearly separated on the first principal component, while translocated and non-translocated samples cluster according to condition on the second principal component (highlighted by red ellipses).

Fig. 4. Chromatin accessibility changes are more numerous in cell lines than patient samples. a, Number of differential ATAC-seq peaks between *IGH-CRLF2* cell line samples (CALL and MUTZ) compared to SMS control, red and blue bars indicate increased and decreased peaks in CALL and MUTZ. b, Overlap of ATAC-seq peaks between CALL and MUTZ indicate 1,396 -seq peaks commonly change in both CALL and MUTZ compared to SMS control. c, Scatter plot of the 1,396 ATAC-seq peaks showing log₂ fold change of CALL and MUTZ (y- and x-axis, respectively) over SMS control. d, Average ATAC-seq signal of Non-T patients (blue), *IGH-CRLF2* patients (red), SMS cell line samples (orange), CALL cell line samples (green), MUTZ cell line samples (purple) at 162 and 126 significantly increased and decreased peaks (upper and lower panels, respectively). Heatmaps show average signal 1kb on either side of the midpoint of each ATAC-seq peak, .

Fig. 5. Inferred B-ALL network identifies significant TFs in the *IGH-CRLF2* patient cohort.

a, Significantly enriched TF motifs (102) at accessible regions 20 kb upstream of the promoters of differentially up and down expressed genes. Plot indicates $-\log_{10}(p\text{-value})$ on x-axis obtained from the hyper-geometric test performed to evaluate enrichment (p-value <0.05) of a particular TF motif. Blue and red bars indicate motif enrichment at down- and up-regulated genes, respectively. b, Workflow for construction of B-ALL regulatory network. Inferelator algorithm was used to infer a regulatory network with ATAC-motif derived priors and 529 RNA-seq B-ALL patient samples from the TARGET database. Transcription Factor Activity (TFA) was calculated using the prior matrix and TARGET gene expression matrix for enriched TF motifs at differentially expressed genes in the *IGH-CRLF2* cohort. The resulting network consists of 102 TFs and 37,086 significant regulatory interactions with combined confidences > 0.5. c, Filtered B-ALL network involving regulatory interactions between significant TFs and their gene targets. TFs are the source nodes labeled in black and the gene targets are the target nodes in blue. The size of the source nodes reflects the number of targets each TF has. TF activation and

Fig. 6. *CRLF2* specific sub-network depicting significant TFs and their differentially regulated target genes. **a.** Pie chart representing the percentage of differentially expressed genes controlled by TFs affected by *CRLF2* overexpression, either at the transcriptional level or TF activity level (387). **b.** Pie chart representing the percentage of differentially expressed genes (329) in the *IGH-CRLF2* sub-network controlled by TFs with significantly altered expression or activity. **c.** *CRLF2* specific sub-network depicting significant TFs and their differentially regulated target genes. TFs are labeled in black and their respective gene targets are connected to them by red and blue edges indicating the type of regulation (activation/repression).

repression of a gene are represented by red and blue sedge, respectively.

Fig. 7. Differentially expressed gene targets of predicted significant TFs are enriched in expected and novel *CRLF2*-associated pathways. a, Significant pathways (–log₁₀(B-H p-value > 0.5 & |z-score| > 1) identified from analysis of differentially expressed gene targets in the *CRLF2* specific sub-network are shown at the top of the figure, ranked by the – log₁₀(Benjamini-Hochberg p-value). Z-score in the bar plot indicates sign of the pathway with red and blue representing up- and down-regulation of the pathway. Differentially expressed genes enriched in these pathways are indicated by purple bins while green bins indicate TF-gene interactions. b, Positive and negative log₂ fold changes of all the differentially expressed genes in IGH-CRLF2 vs Non-T patients and cell lines (CALL vs SMS, MUTZ vs SMS) are indicated as red (positive) and blue (negative) bins with significantly deregulated genes labeled with white stars.

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Fig. 1

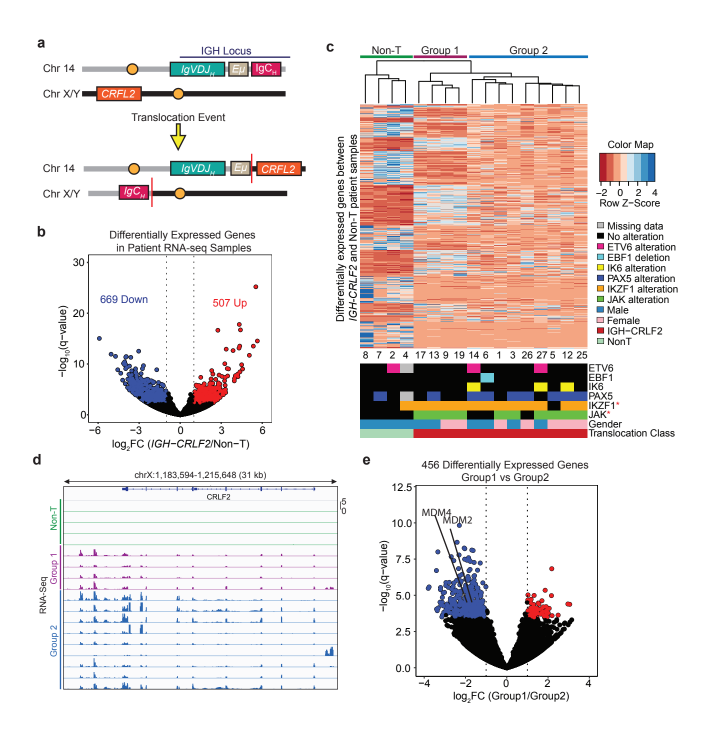


Fig.2

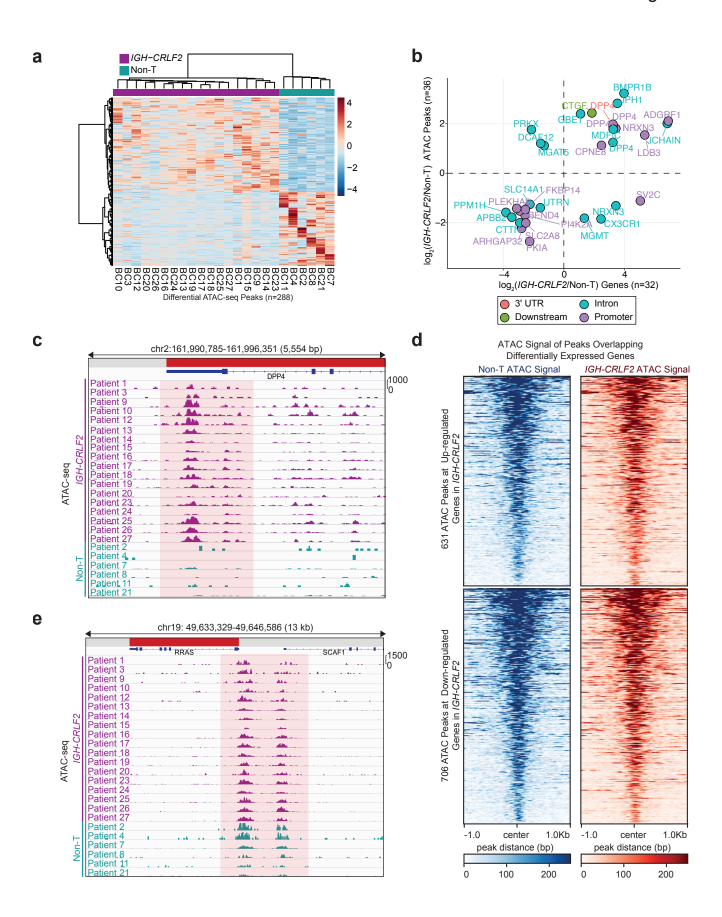


Fig. 3

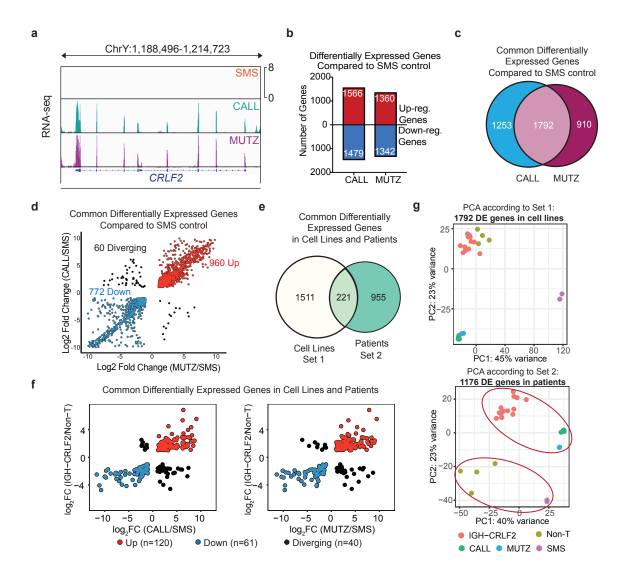


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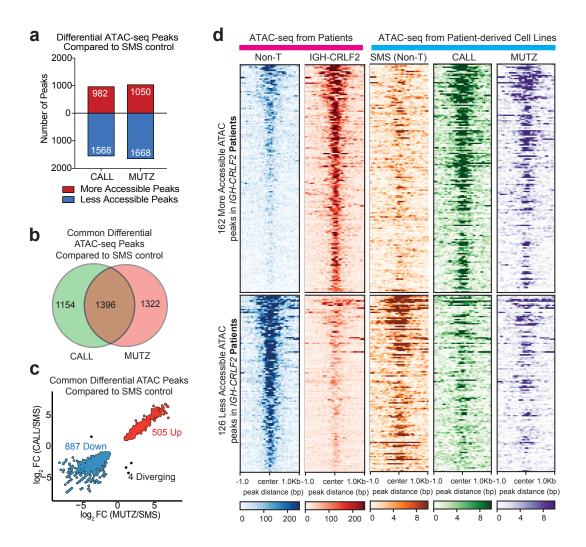
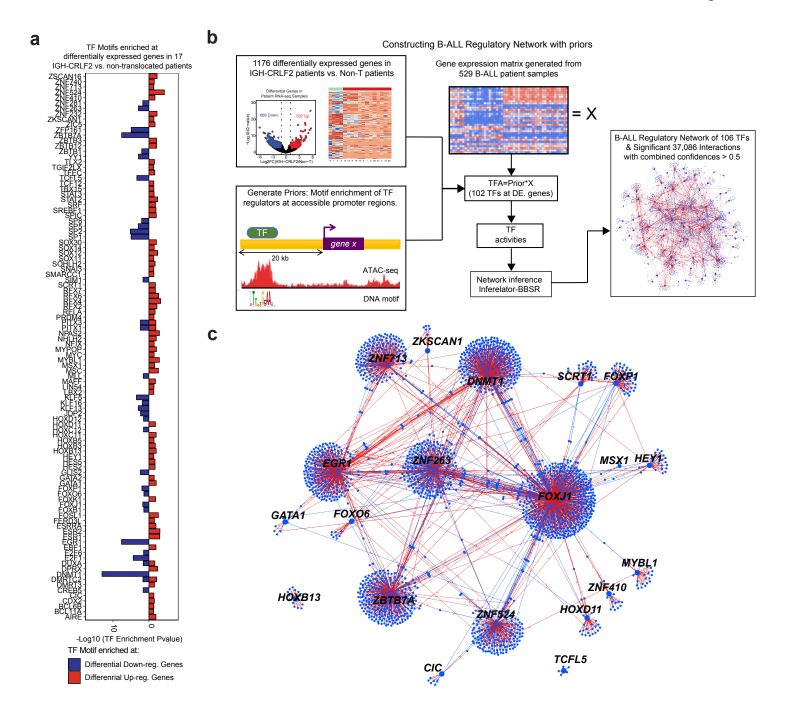


Fig. 5



ZNF410

MSX1

Fig. 6

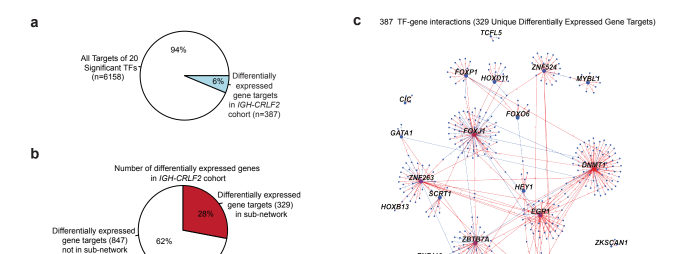


Fig. 7

