RAG-mediated DNA breaks attenuate PU.1 activity in early B cells through activation of a SPIC-BCLAF1 complex

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Graphical Abstract

Highlights
- RAG DNA breaks upregulate SPIC, which induces genome-wide changes in PU.1 activity
- SPIC binds to gene-regulatory elements, resulting in loss of PU.1 at these regions
- SPIC complexes with BCLAF1 to suppress transcription in response to RAG DNA breaks
- SPIC/BCLAF1 inhibits SYK and promotes transition from large to small pre-B cells

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In Brief
ETS-family transcription factors are key regulators of early B cell development. Soodgupta et al. show that RAG-induced DNA breaks generated during antigen receptor gene recombination activate a SPIC/BCLAF1 transcription factor complex that counters PU.1 activity and regulates gene expression changes to promote transition from large to small pre-B cells.

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RAG-Mediated DNA Breaks Attenuate PU.1 Activity in Early B Cells through Activation of a SPIC-BCLAF1 Complex

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SUMMARY

Early B cell development is regulated by stage-specific transcription factors. PU.1, an ETS-family transcription factor, is essential for coordination of early B cell maturation and immunoglobulin gene rearrangement. Here we show that RAG DNA double-strand breaks (DSBs) generated during Ig light chain gene (IgI) rearrangement in pre-B cells induce broad transcription changes in PU.1 chromatin binding. RAG DSBs activate a SPIC/BCLAF1 transcription factor complex that displaces PU.1 throughout the genome and regulates broad transcriptional changes. SPIC recruits BCLAF1 to gene-regulatory elements that control expression of key B cell developmental genes. The SPIC/BCLAF1 complex suppresses expression of the SYK tyrosine kinase and enforces the transition from large to small pre-B cells. These studies reveal that RAG DSBs direct genome-wide changes in ETS transcription factor activity to promote early B cell development.

INTRODUCTION

B cell development requires the sequential assembly and expression of genes encoding the immunoglobulin heavy (IgH) and immunoglobulin light (IgL) chains to generate a mature B cell receptor (BCR) (Rajewsky, 1996). Ig genes are assembled through the process of V(D)J recombination, which joins distant variable (V), joining (J), and diversity (D) segments (Fugmann et al., 2000). The DNA double-strand breaks (DSBs) necessary for V(D)J recombination are generated by the RAG endonuclease, which is composed of the RAG1 and RAG2 proteins (Fugmann et al., 2000). RAG-mediated DNA breaks are generated in the G1 phase of the cell cycle and activate the DNA damage response (DDR) kinase ATM, which facilitates repair of the broken DNA ends through nonhomologous end joining (Helmink and Sleckman, 2012). In response to RAG DSBs, ATM also activates a broad transcriptional program that regulates genes involved in diverse B cell functions, including migration, cell-cycle arrest, survival, and differentiation (Bednarski et al., 2012, 2016; Bredemeyer et al., 2008; Helmink and Sleckman, 2012; Steinel et al., 2013). This genetic program is mediated by ATM-dependent activation of several transcription factors, including NF-kB1, NF-kB2, and SPIC (Bednarski et al., 2012, 2016; Bredemeyer et al., 2008).

The IgH gene is assembled first in pro-B cells and productive rearrangement results in its surface expression with surrogate light chains (µ5 and VpreB) to generate the pre-BCR, which signals transition to the large pre-B cell stage (Clark et al., 2014; Herzog et al., 2009; Rajewsky, 1996). Pre-BCR oligomerization signals through the SYK tyrosine kinase to promote proliferation and clonal expansion of large pre-B cells (Clark et al., 2014; Herzog et al., 2009). Activation of SYK also triggers Igk (IgK) gene recombination (Clark et al., 2014). RAG expression is suppressed in proliferating cells, and as such, Igk gene assembly requires induction of cell-cycle arrest and transition to the small, non-proliferating pre-B cell stage (Clark et al., 2014; Desiderio et al., 1996; Johnson et al., 2008; Ochiai et al., 2012). RAG DSBs activate ATM-dependent DDR signaling pathways that enforce cell-cycle arrest and promote survival to prevent proliferation of cells with unrepaired DSBs and permit time for proper assembly of Igk genes (Bednarski et al., 2012, 2016; DeMicco et al., 2016).

B cell development and assembly of Ig genes are carefully orchestrated by developmental stage-specific transcription factors, including E2A, EBF, Pax5, PU.1 and SPIB (Pang et al., 2014). The ETS-family transcription factor PU.1 is required for B cell lineage commitment and is constitutively expressed throughout B cell development (Polli et al., 2005; Schweitzer and DeKoter, 2004; Scott et al., 1994, 1997). PU.1 has critical functions during B cell maturation. In pre-B cells, PU.1 regulates expression of a diverse genetic program, including genes involved in B cell proliferation, differentiation, and Ig gene rearrangement (Batista et al., 2017; Heinz et al., 2010; Solomon et al., 2015). Expression of SYK and germline transcription of Igk, which are required for pre-BCR signaling and initiating V(D)J recombination, respectively, depend on PU.1 activity (Batista et al., 2017; Herzog et al., 2009; Schwarzenbach et al., 1995; Schweitzer and DeKoter, 2004). Interestingly, loss of PU.1 in B

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Figure 1. RAG DSB Signals Induce Genome-wide Changes in PU.1 Binding

(A) qPCR analysis of Igk genomic DNA from Rag1<sup>−/−</sup>:Bcl2 (red) and Art<sup>−/−</sup>:Bcl2 (blue) abl pre-B cells treated with imatinib for 48 h. Schematic shows germline (GL) Igk locus and unrepaired Jκ1 coding end with location of PCR primers. PCR is normalized to Rag1<sup>−/−</sup>:Bcl2 abl pre-B cells, which do not generate RAG DSBs and have only intact germline Igk DNA. Data are representative of three independent experiments.
cell progenitors results in only a mild defect in B cell development because of compensatory function of another ETS-family transcription factor, SPIB (Pollil et al., 2005; Sokalski et al., 2011; Ye et al., 2005). PU.1 and SPIB associate with nearly identical regions of the genome in B cells and regulate transcription of a similar cohort of genes (Solomon et al., 2015). Combined loss of PU.1 and SPIB impairs B cell maturation in the bone marrow and predisposes to the development of B cell leukemia (Sokalski et al., 2011).

We previously demonstrated that SPIC, an ETS-family transcriptional repressor with homology to PU.1 and SPIB, also functions in pre-B cells (Bednarski et al., 2016; Bemark et al., 1999; Hashimoto et al., 1999). Unlike PU.1 and SPIB, SPIC is not constitutively expressed in early B cells but, rather, is induced by signals from RAG DSBs (Bednarski et al., 2016). SPIC operates primarily as a transcriptional repressor and counters the activating functions of PU.1 and SPIB (Li et al., 2015; Zhu et al., 2008). In pre-B cells, SPIC suppresses expression of Syk and Blnk, which inhibits pre-BCR signaling and enforces cell-cycle arrest in pre-B cells with RAG DSBs (Bednarski et al., 2016). SPIC also inhibits transcription of Igk to prevent generation of additional RAG DSBs (Bednarski et al., 2016). Binding of SPIC to gene-regulatory elements for Syk, Blnk, and Igk is associated with loss of PU.1 at these genomic regions. Thus, expression of SPIC antagonizes PU.1 as these identified genes to suppress transcription and coordinate pre-B cell development.

Whether SPIC has broader functions in gene regulation and its mechanism of action in B cells have been defined. SPIC may oppose PU.1 at limited gene targets or, alternatively, may modulate PU.1 activity throughout the genome. In this regard, attenuation of PU.1 activity by SPIC could suppress pre-B cell genetic programs to promote continued B cell maturation. SPIC may function simply by displacing PU.1 through competition for DNA binding sites or may complex with other transcriptional regulators to repress transcription. We show here that, in response to RAG DSBs, SPIC binds throughout the genome of pre-B cells and elicits global changes in PU.1 chromatin association. SPIC associates with the transcriptional repressor BCLAF1 (Bcl2-associated factor 1) to regulate a distinct subset of RAG DSB-dependent gene expression changes and to enforce transition from large to small pre-B cells. These experiments provide insight into the regulation of ETS transcription factors in early B cells and the impact of DDR signaling on B cell development.

RESULTS

RAG DSB Signals Induce Genome-Wide Changes in PU.1 Binding

To determine the effects of DNA damage signaling on PU.1 activity in early B cells, we used Abelson-kinase transformed pre-B cells (abl pre-B cells) deficient in Rag1 or the Artemis endonuclease that express the Bcl2 transgene (Rag1^{-/-}/Bcl2 and Art^{-/-}/Bcl2, respectively) (Bredemeyer et al., 2008). Expression of the Abl kinase promotes pre-B cell proliferation and suppresses expression of Rag1 and Rag2. Treatment with the Abl kinase inhibitor imatinib triggers cell-cycle arrest, induction of RAG expression, and recombination of Igk (Bredemeyer et al., 2008). The Bcl2 transgene supports survival of imatinib-treated cells. Following treatment with imatinib, Rag1^{-/-}/Bcl2 abl pre-B cells do not generate RAG DSBs. In contrast, Art^{-/-}/Bcl2 abl pre-B cells generate RAG DSBs at Igk, but these DSBs are not repaired as Artemis is required to open hairpin-sealed coding DNA ends (Figure 1A) (Bredemeyer et al., 2008; Helmink and Sleckman, 2012). The RAG DSBs in Art^{-/-}/Bcl2 abl pre-B cells activate ATM-dependent DDRs (Bednarski et al., 2012, 2016; Bredemeyer et al., 2008).

Chromatin immunoprecipitation followed by next-generation DNA sequencing (ChIP-seq) reveals global changes in PU.1 binding in pre-B cells with RAG DSBs (Art^{-/-}/Bcl2) compared with pre-B cells without RAG DSBs (Rag1^{-/-}/Bcl2), despite no differences in PU.1 expression (Figures 1B, 1C, and S1A). Induction of RAG DSBs results in gain of few new binding sites but loss of approximately 20% of the PU.1 binding sites identified in Rag1^{-/-}/Bcl2 abl pre-B cells (Figure 1B). Gene Ontology analysis demonstrates that genes within 12 kb of lost PU.1 binding sites are involved in immune cell activation and differentiation (Figure S1B). In contrast, PU.1 binding sites that are conserved between Rag1^{-/-}/Bcl2 and Art^{-/-}/Bcl2 abl pre-B cells are proximal to genes involved in cell homeostasis and maintenance (i.e., signaling, nuclear transport, apoptosis). Novel RAG DSB-induced PU.1 binding occurred near genes involved in cell adhesion and developmental processes. Induction of RAG DSBs did not alter PU.1 binding across genomic regulatory elements as equal binding to promoters, genes, or intergenic regions (i.e., enhancers) is observed in both Rag1^{-/-}/Bcl2 and Art^{-/-}/Bcl2 abl pre-B cells (Figure S1C). Thus, in response to RAG DSBs, pre-B cells have a genome-wide reduction in PU.1 chromatin binding, which is expected to result in changes in gene expression that affect important cellular functions.

Expression of SPIC Alters PU.1 Binding in Pre-B Cells

RAG DSBs trigger ATM-dependent induction of SPIC (Figure 2A). Expression of SPIC, in turn, results in loss of PU.1 binding at genes required for pre-BCR signaling (Bednarski et al., 2016). To determine if expression of SPIC is responsible for the global changes in PU.1 binding observed in response to RAG DSBs, we stably transduced Rag1^{-/-}/Bcl2 abl pre-B cells with a lentiviral vector encoding a tetracycline-inducible FLAG-HA-tagged SPIC (Rag1^{-/-}/Bcl2:Spic^{tet}). Treatment with doxycycline induced equivalent SPIC mRNA expression as triggered by RAG DSBs (Figures 2A and 2B). We performed ChIP-seq for PU.1 in Rag1^{-/-}/Bcl2:Spic^{tet} abl pre-B cells treated with imatinib alone or in combination with doxycycline to induce expression of SPIC (Figure 2B). Expression of SPIC does not alter PU.1 expression

(B) Dot plot and heatmap of fold changes and signal intensity for PU.1 peaks identified by ChIP-seq in Rag1^{-/-}/Bcl2 and Art^{-/-}/Bcl2 abl pre-B cells treated with imatinib for 48 h. Data are from common peaks identified in two replicates for each cell.
(C) Representative tracks at indicated regions for PU.1 ChIP-seq from (B). ChIP-qPCR validation for PU.1 binding at each locus is also shown. Data are mean and SE for three independent experiments. **p ≤ 0.01 and ****p ≤ 0.0001; ns, not significant. See also Figure S1.
but results in significant changes in PU.1 chromatin binding (Figures 2C and 2D). Moreover, expression of SPIC results in changes in PU.1 binding that are similar to changes induced by RAG DSBs (compare Figures 1B and 2D). These findings demonstrate that changes in PU.1 binding in response to RAG DSBs are, in large part, due to RAG DSB-mediated induction of SPIC.
SPIC and PU.1 Bind to Identical Genomic Regions

SPIC and PU.1 have homologous DNA binding domains and have been previously shown in vitro to bind to the same DNA sequence (Bemark et al., 1999; Hashimoto et al., 1999). Current commercial antibodies against endogenous SPIC do not work for ChIP. Thus, to determine if SPIC and PU.1 binding to chromatin is similarly distributed throughout the genome, we performed ChIP-seq with anti-HA antibodies to precipitate FLAG-HA-SPIC in performed ChIP-seq with anti-HA antibodies to precipitate FLAG-HA-SPIC in Rag1−/−;Bcl2;Spicabl pre-B cells treated with doxycycline (to induce SPIC). Results were compared with findings from ChIP-seq for PU.1 in Rag1−/−;Bcl2 abl pre-B cells without SPIC expression. Peaks with ≥1 bp of overlap between the two ChIP-seq datasets were considered as enriched for binding to both transcription factors. We find that SPIC and PU.1 bind to similar locations throughout the genome (Figure 3A). Additionally, PU.1 binding is lost at sites where SPIC is bound (Figures 3B, 3C, and S2).

The ChIP peaks for SPIC and PU.1 in regions where both transcription factors bind (common peaks in Figure 3A) have significant nucleotide overlap (Figures 3D). Indeed, the majority of these shared binding sites overlap by >70%, and the greatest number of ChIP peaks have >90% overlap. Furthermore, SPIC and PU.1 bind to similar regions throughout the genome (Figure 3E). Collectively, these findings demonstrate that SPIC and PU.1 bind to similar regulatory elements in pre-B cells and that SPIC binding results in displacement of PU.1 from these regions.

SPIC Recruits BCLAF1 to Chromatin

PU.1 forms heterodimeric complexes with IRF4 or IRF8 to regulate transcription initiation (Brass et al., 1996; Heinz et al., 2010). SPIC does not complex with IRF4 or IRF8 but binds to similar DNA sequences as PU.1 (Carlsson et al., 2003). These findings raise the question of whether SPIC complexes with distinct protein partners to regulate gene expression. To identify SPIC interacting partners, we generated Art−/−;Bcl2 abl pre-B cells expressing either a tetracycline-inducible FLAG-HA-tagged SPIC (Art−/−;Bcl2;Spicabl) or a tetracycline-inducible FLAG-HA-tagged PU.1 (Art−/−;Bcl2;Pu1abl). Cells were treated with imatinib to induce RAG DSBs and with doxycycline to induce comparable expression of the FLAG-tagged transcription factors (Figure S3). SPIC and PU.1 were immunoprecipitated using anti-FLAG antibodies, and associated proteins were identified by tandem mass spectrometry. Unique peptides were compared with SPIC- and BCLAF1-dependent genes are enriched for immune processes in B cells (Figure S4B). Gene Ontology analysis revealed that BCLAF1- and SPIC-dependent genes are enriched for immune processes in B cells (Figure S4B). Importantly, loss of BCLAF1 does not alter SPIC binding to the Syk promoter, suggesting that recruitment of BCLAF1 is needed for SPIC-mediated transcriptional changes but not for SPIC binding to chromatin (Figure S5D).

We then determined the contribution of SPIC/BCLAF1 to the genetic program regulated by RAG DSBs in pre-B cells. Gene profiling revealed that BCLAF1 regulates a significant portion of RAG DSB-mediated genes (540 of 717 genes, ≥2-fold change, adjusted p < 0.05; Figure 5E; Table S3). Comparison of RAG DSB-dependent (Art−/−;Bcl2 versus Rag1−−;Bcl2; Figure 5E), SPIC-dependent (Rag1−/−;Bcl2;Spicabl expressing SPIC versus Rag1−−;Bcl2; Figure 5A), and BCLAF1-dependent (Art−/−;Bcl2 expressing shBCLAF1 versus Art−/−;Bcl2; Figure 5E) gene expression changes identified 141 genes whose expression is modulated by all three variables (Figures 5F, 5G, and S5A; Table S4). Approximately 25% of these genes have concordant changes in expression (repressed by RAG DSBs, repressed by SPIC, and repressed by loss of BCLAF1; Figure S5A). Pathway analyses are enriched for diverse B cell functions, including proliferation, cell adhesion, and cell death (Figure S5B). These findings demonstrate that the SPIC/BCLAF1 complex regulates a distinct genetic program in pre-B cells with RAG DSBs.
Figure 3. SPIC and PU.1 Bind to Identical Genomic Regions

(A) Dot plot and heatmap of fold changes and signal intensity for PU.1 and SPIC (by anti-HA ChIP) peaks identified by ChIP-seq in Rag1^−/−:Bcl2:Spic^WT abl pre-B cells treated with imatinib for 48 h in the absence (for PU.1 ChIP) or presence (for SPIC ChIP) of 2 μM doxycycline (Dox). Data are from common peaks identified in two replicates of each cell line.

(B) Representative ChIP-seq binding of PU.1 and SPIC at indicated regions. PU.1 ChIP-seq was performed in Rag1^−/−:Bcl2:Spic^WT abl pre-B cells treated with imatinib alone (−Dox, no SPIC) or with imatinib and doxycycline to induce expression of SPIC (+Dox, +SPIC) for 48 h. ChIP-seq for SPIC was performed as in A in Rag1^−/−:Bcl2:Spic^WT abl pre-B cells treated with imatinib and doxycycline for 48 h.

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BCLAF1 Regulates Pre-BCR Signaling in Primary Pre-B Cells

To determine if BCLAF1 is required for regulation of SPIC function in primary pre-B cells, we expanded pre-B cells from Rag<sup>1<sup>−/−</sup></sup>:Bcl2<sup>Spictet</sup> and Art<sup>−/−</sup>:Bcl2<sup>Spictet</sup>:Pu.1<sup>tet</sup> abl pre-B cells, respectively, after treatment with imatinib and 2 μM doxycycline for 48 h. Scatterplot shows number of total peptides per protein identified by mass spectrometry analysis of co-immunoprecipitation of SPIC (y axis) versus PU.1 (x axis).

(D) Nucleotide overlap between PU.1 and SPIC peaks identified in (A). Peaks were grouped in bins on the basis of percentage of overlap as shown.

(E) Enrichment of PU.1 and SPIC binding across genomic regions on the basis of ChIP-seq data in (A).

See also Figure S2.

(C) ChIP-qPCR validation for PU.1 and SPIC binding at each locus shown in (B). Data are mean and SE for three independent experiments. **p < 0.01, ***p < 0.001, and ****p < 0.0001.

(D) ChIP-qPCR of BCLAF1 binding at the Syk promoter in Rag<sup>1<sup>−/−</sup></sup>:Bcl2<sup>Spictet</sup> abl pre-B cells treated with imatinib for 48 h in the absence (–) or presence (+) of 2 μM doxycycline (Dox) to induce SPIC expression.

(F) Re-ChIP for BCLAF1 after primary ChIP for SPIC or PU.1 (using anti-HA antibodies) in Rag<sup>1<sup>−/−</sup></sup>:Bcl2<sup>Spictet</sup> or Rag<sup>1<sup>−/−</sup></sup>:Bcl2<sup>Spictet</sup>:Pu.1<sup>tet</sup> abl pre-B cells, respectively, treated with imatinib and 2 μM doxycycline for 48 h.

(G) ChIP-qPCR of BCLAF1 binding at the Syk promoter in Rag<sup>1<sup>−/−</sup></sup>:Bcl2<sup>Spictet</sup> abl pre-B cells treated with imatinib for 48 h. Data in (A–C) are representative of three independent experiments. Data in (E–G) are mean and SE for three independent experiments. **p < 0.01, ***p < 0.001, and ****p < 0.0001.

See also Figure S3 and Table S1.
Figure 5. SPIC and BCLAF1 Regulate Gene Expression in Pre-B Cells in Response to RAG DSBs

(A) Volcano plot of gene expression changes (fold change ≥ 2, p ≤ 0.05) between \(\text{Rag}^{1/-}:\text{Bcl}2:\text{Spic}^{\text{mt}}\) abl pre-B cells with and without SPIC induction. RNA-seq was performed on \(\text{Rag}^{1/-}:\text{Bcl}2:\text{Spic}^{\text{mt}}\) abl pre-B cells treated with imatinib alone (-SPIC) or with imatinib and 2 μM doxycycline (+SPIC) for 48 h. Data are from two independent cultures for each treatment.

(legend continued on next page)
Loss of BCLAF1 does not alter induction of Spic but does lead to increased expression of Syk in Art−/−:Igh:Bcl2 small pre-B cells (Figures 6A and 6B). Consistent with the rescue of Syk mRNA levels, Syk protein is increased in Art−/−:Igh:Bcl2 pre-B cells lacking BCLAF1 to levels equivalent to those observed in Rag1−/−:Igh:Bcl2 pre-B cells (Figure 6C). On the basis of these results, we conclude that BCLAF1 is necessary for repression of Syk in response to RAG DSBs in primary small pre-B cells.

To assess BCLAF1 binding to the Syk promoter during wild-type pre-B cell development in vivo, we used Spicgyfp/gfp mice, which contain an IRES-EGFP targeted to the 3′ non-coding exon of Spic (Haldar et al., 2014). Approximately 2% of small pre-B cells from Spicgyfp/gfp mice are EGFP positive, indicative of Spic expression (Figures 6D and 6E). EGFP-expressing small pre-B cells are not observed in Atm−−:Spicgyfp/gfp, indicating that induction of Spic (and EGFP) depends on DNA damage signaling (Figure 6E). Spic-expressing Spicgyfp/gfp small pre-B cells (EGFP positive) have reduced PU.1 binding and increased BCLAF1 binding to the Syk promoter as well as decreased Syk expression (Figures 6F–6H) (Bednarski et al., 2016). These results suggest that Spic/BCLAF1 complex is induced by DNA damage signals from transient RAG DSBs generated during IgI rearrangement in wild-type small pre-B cells.

**Loss of BCLAF1 Alters Large to Small Pre-B Cell Transition**

Activation of SYK downstream of the pre-BCR can promote pre-B cell proliferation in the absence of IL-7 signaling (Clark et al., 2014; Herzog et al., 2009; Ochiai et al., 2012; Rolink et al., 2000; Wossning et al., 2006). Given that loss of BCLAF1 prevents Spic-mediated repression of SYK, we hypothesized that loss of BCLAF1 may alter pre-B cell proliferation and the transition from large to small pre-B cells during early B cell development. To test this, we generated Bclaf1f/f:Mb1-cre mice, which have selective loss of BCLAF1 in B cells (Figure 7A) (Hobeika et al., 2006). Pre-B cells from Bclaf1f/f:Mb1-cre and Bclaf1f/f mice were expanded in the presence of IL-7. Following IL-7 withdrawal, Bclaf1-deficient pre-B cells from Bclaf1f/f mice have increased S-phase progression and increased Syk expression compared with pre-B cells from Bclaf1f/f mice (Figures 7B–7D). These findings support a role for BCLAF1 in the regulation of pre-B cell proliferation possibly through modulation of SYK activity downstream of pre-BCR signaling.

We next assessed B cell populations in vivo. In our breeding, Mb1-cre mice have normal numbers of pro-B cells but reduced pre-B cells relative to littermate wild-type Bclaf1f/f mice (Figures 7E and 7F). In contrast, Bclaf1f/f:Mb1-cre mice have increased numbers of pre-B cells compared with Mb1-cre mice and are similar to Bclaf1f/f mice (Figure 7E). Interestingly, the increase in pre-B cells in Bclaf1f/f:Mb1-cre mice is due primarily to larger numbers of large pre-B cells (Figure 7E). Loss of Bclaf1 does not alter numbers of pro-B cells or small pre-B cells. Consistent with findings in cultured cells, in vivo large, proliferating pre-B cells from Bclaf1f/f:Mb1-cre mice have increased Syk mRNA levels (Figure 7F). Syk expression is not altered in small pre-B cells (Figure 7F). We propose that BCLAF1 functions in response to RAG DSBs in pre-B cells to suppress Syk and enforce transition from the large to small pre-B cell developmental stage.

**DISCUSSION**

Here we show that RAG DSBs induce genome-wide changes in PU.1 localization and function, which coordinates a distinct genetic program in B cells undergoing Ig gene rearrangement. This modulation of PU.1 activity is mediated by RAG DSB activation of a Spic/BCLAF1 transcriptional repressor complex. Spic displaces PU.1 at gene regulatory sites but requires association with BCLAF1 to suppress transcription. This antagonistic function of Spic/BCLAF1 coordinates a broad genetic program and enforces transition from large to small pre-B cells in response to RAG DSBs.

PU.1 is a key regulator of cell fate decisions during early hematopoiesis and is essential for generating B cells from hematopoietic progenitors (Dakic et al., 2007; DeKoter et al., 2002; Pang et al., 2018; Scott et al., 1994, 1997). PU.1 expression is high in myeloid cells, in which it is required to promote lineage specific gene expression (Heinz et al., 2010). In contrast, PU.1 expression is reduced during B cell differentiation and remains low in established B cells (Back et al., 2005; Nutt et al., 2005). This differential activity of PU.1 is critical for directing appropriate lineage commitment. Dysregulation of PU.1 expression leads to aberrant differentiation and can result in leukemic transformation (Anderson et al., 2002; Pang et al., 2016; Rosenbauer et al., 2004, 2006; Sokalski et al., 2011). PU.1 activity is also regulated through interaction with other transcription factors, which modulate its DNA binding properties or its transcriptional function (Maia and Atchison, 2000; Nerlov et al., 2000; Rogers et al., 2016). For example, in early lymphoid precursors, E2A
Figure 6. BCLAF1 Regulates SYK Expression in Primary Pre-B Cells

(A–C) Art^{-/-}:Igh:Bcl2 pre-B cells were transduced with a retrovirus expressing a scrambled shRNA (-) or shBclaf1 (+) and then subsequently withdrawn from IL-7.

(A and B) Spic and Syk mRNA expression assessed in indicated small pre-B cells 2 days after IL-7 withdrawal. Data are mean and SE for three independent experiments.

(C) Western blot of SYK and BCLAF1 in indicated small pre-B cells 2 days after IL-7 withdrawal. Data are representative of three independent experiments.

(D) Flow cytometric analysis showing EGFP (y axis) and FSC (x axis) in bone marrow pre-B cells (B220^loCD43^IgM^-) from wild-type and Spic^{gfp/gfp} mice. Data are representative of five independent experiments.

(E) Percentage of EGFP-positive small pre-B cells in Spic^{gfp/gfp} (circles) and Atm^{-/-}:Spic^{gfp/gfp} (squares) mice was quantified by flow cytometry as in (D). Data are mean and SE from three independent mice of each genotype.

(F–H) Syk mRNA expression (F), ChIP-PCR of PU.1 at Syk promoter (G), and ChIP-PCR of BCLAF1 at Syk promoter (H) in EGFP-negative (-) and EGFP-expressing (+) small pre-B cells sorted from Spic^{gfp/gfp} mice. Data in (F) are the mean and SE from three independent experiments. Data in (G) and (H) are representative of two independent experiments.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001; ns, not significant.
association with PU.1 inhibits PU.1-induced transcription of myeloid genes and promotes B lymphoid differentiation (Rogers et al., 2016). We find that PU.1 activity is regulated at the pre-B cell developmental stage through RAG DSB-mediated induction of SPIC, which binds chromatin and displaces PU.1. This transcription factor exchange results in changes in expression of genes involved in pre-BCR signaling, B cell proliferation, and B cell differentiation.

SPIC and PU.1 have homologous DNA binding domains (Bemark et al., 1999; Hashimoto et al., 1999). As such, SPIC can compete for DNA binding sites occupied by PU.1, and binding of SPIC results in displacement of PU.1 from these sites. Interestingly, SPIC associates with >90% of the PU.1 sites, but PU.1 binding is lost at only approximately 20% of the regions it binds in the absence of SPIC expression (Figures 2D and 3A). It is conceivable, then, that SPIC and PU.1 may simultaneously bind specific regions of the genome, and SPIC binding may not always fully displace PU.1. Rather, binding of SPIC nearby PU.1 may alter PU.1 transcriptional activity or other transcriptional machinery at these sites. Alternatively, in an individual cell, each ETS site may be occupied by either SPIC or PU.1, but ChIP analysis on a bulk population is not

Figure 7. BCLAF1 Regulates Large to Small Pre-B Cell Transition

(A) Western blot of BCLAF1 in sorted CD19− (non-B cell) and CD19+ B cell populations from bone marrow of 5-week-old Bclaf1ff:Mb1-cre mice. Data are representative of three independent mice.

(B) Flow cytometric analysis of BrdU incorporation (y axis) and DNA content (7AAD, x axis) performed 24 h after IL-7 withdrawal. Data are mean and SE for at least three independent experiments.

(C) Quantitation of flow cytometric analysis of pro-B cells (F4/80+CD43+) and pre-B cells (B220+B220−CD43+) in bone marrow of 5-week-old Bclaf1ff (black bars, n = 12), Mb1-cre (gray bars, n = 10), and Bclaf1ff:Mb1-cre (white bars, n = 10) mice. Data are mean and SE for three independent experiments.

(D) Syk mRNA expression 24 h after IL-7 withdrawal. Data are mean and SE for three independent experiments.

(E) Syk mRNA expression in small and large pre-B cells sorted from 5-week-old Bclaf1ff (black bars, n = 4), Mb1-cre (gray bars, n = 4), and Bclaf1ff:Mb1-cre (white bars, n = 5) mice. Data in (E) and (F) are mean and SE for indicated numbers of mice. *p < 0.05, **p < 0.01, and ***p < 0.001; ns, not significant.

See also Figure S6.
sensitive enough to discriminate between these two different states.

In early B cells, PU.1 and SPIB are constitutively expressed and have complementary functions (Schweitzer and DeKoter, 2004; Scott et al., 1994, 1997; Sokalski et al., 2011; Solomon et al., 2015). As such, conditional deletion of either PU.1 or SPIB alone mildly alters B cell development, but loss of both transcription factors results in a block in B cell differentiation at the pro-B cell stage (Polli et al., 2005; Sokalski et al., 2011, Su et al., 1997; Ye et al., 2005). PU.1 and SPIB bind to similar regions throughout the genome of pro-B cells and regulate transcription of key developmental genes, including Syx and Blnk, which are necessary for pre-BCR signaling and induction of proliferation of large pre-B cells (Solomon et al., 2015). We find that SPIC also binds to the same genomic sites as PU.1. Given that SPIB and PU.1 bind identical regions and have complementary functions in early B cells, SPIC is also expected to counter SPIB similar to our observed results for PU.1. In contrast to PU.1 and SPIB, SPIC is inducibly expressed in pre-B cells in response to RAG DSBs and functions primarily as a transcriptional repressor. Expression of SPIC opposes PU.1 and SPIB activity resulting in suppression of pre-BCR and BCR signaling in early B cells and mature B cells, respectively, leading to a block in B cell maturation or function (Bednarski et al., 2016; Zhu et al., 2008). Importantly, complete or permanent inhibition of PU.1 and SPIB could be detrimental to B cell development, as combined loss of these transcription factors results in leukemic transformation (Sokalski et al., 2011). In this regard, induced expression of SPIC by RAG DSBs permits for stage-specific and transient inhibition of PU.1 (and SPIB). SPIC expression is expected to be lost after RAG DSBs are repaired and associated DDR signaling is terminated. The reduction in SPIC would allow PU.1 (and SPIB) to rebind to chromatin and resume transcriptional activities necessary for mature B cell function. Thus, RAG DSBs regulate a temporary suppression of PU.1 to promote transition from large to small pre-B cells and then permit continued transition to antibody-producing mature B cells.

PU.1 forms heterodimeric complexes with IRF4 or IRF8 to promote transcription (Brass et al., 1996; Heinz et al., 2010; Pongubala et al., 1992). As such, combined loss of IRF4 and IRF8 results in similar abnormalities in B cell development as loss of PU.1 (Lu et al., 2003; Ma et al., 2006). SPIC binds the same DNA sequence as PU.1 but has a distinct protein-interaction domain and does not bind IRF4 or IRF8 (Carlsson et al., 2003). Thus, SPIC could mediate suppression of transcription simply through displacement of PU.1 and loss of associated transcription activation machinery (i.e., IRF4). Displacement of the PU.1/IRF4 complex alone, though, may be insufficient to repress transcription as this is not expected to result in rapid changes in histone modifications or RNA polymerase activity, which drive gene expression. Alternatively, in a manner similar to PU.1, SPIC may effect transcriptional inhibition by recruiting additional proteins to gene-regulatory elements. In this regard, we find that SPIC, but not PU.1, binds BCLAF1. BCLAF1 is not necessary for SPIC binding to chromatin but is required for transcriptional repression. On the basis of these findings, we propose that antagonism of PU.1 activity is mediated by a SPIC-BCLAF1 complex that binds to chromatin and suppresses key PU.1-regulated genes. Further studies are needed to determine the mechanism by which the SPIC-BCLAF1 complex regulates transcription (i.e., activity on histone epigenetics, RNA polymerase activity, and locus accessibility).

BCLAF1 was first identified as a transcriptional repressor but also functions as an activator to promote expression of p53 and cytokines in response to DNA damage (Kasof et al., 1999; Liu et al., 2007; Shao et al., 2016). BCLAF1 also has been identified as a component of the RNA splicing complex (Savage et al., 2014; Vohodina et al., 2017). We find that in early B cells, BCLAF1 complexes with SPIC to repress gene expression in response to RAG-mediated DSBs. BCLAF1 chromatin binding nearly completely overlaps with SPIC-bound genomic regions. SPIC and BCLAF1 could bind DNA independently and then cooperatively suppress transcription. In this regard, in vitro studies have shown that BCLAF1 binds the interferon-stimulated response element (ISRE) (Qin et al., 2019). The sequence for binding of the PU.1/IRF4 heterodimer contains a portion of the ISRE site in series with an ETS motif. BCLAF1 and SPIC could bind this same sequence, or, alternatively, BCLAF1 may be recruited to gene regulatory regions through protein-protein interactions with SPIC, which binds ETS DNA sequences. The domains that govern SPIC and BCLAF1 protein interactions and DNA binding are currently being investigated.

We find that loss of BCLAF1 prevents RAG DSB- and SPIC-mediated repression of Syx mRNA expression. SYK is a key signaling molecule downstream of the pre-BCR and is required for the pre-BCR to promote proliferation of large pre-B cells (Clark et al., 2014; Herzog et al., 2009). We previously showed that in response to RAG DSBs, induction of SPIC suppresses pre-BCR signaling to enforce cell-cycle arrest in small pre-B cells (Bednarski et al., 2016). Thus, loss of BCLAF1 is expected to mitigate RAG DSB-induced inhibition of proliferation. Indeed, Bclaf1-deficient pre-B cells have increased cell cycle entry, and mice with B cell-specific deletion of BCLAF1 have increased numbers of proliferating, large pre-B cells, consistent with increased SYK activity. Loss of BCLAF1 does not result in a complete block in B cell development, which may reflect that additional mechanisms, such as p53, exist to regulate G1 arrest in small pre-B cells undergoing Ig gene rearrangement.

In summary, we find that SPIC/BCLAF1 functions to modulate PU.1 activity in pre-B cells. High activity of PU.1 promotes proliferation and expansion of large pre-B cells. As cells transition to small pre-B cell stage and initiate Ig gene assembly, RAG DSBs induce expression of SPIC, which partners with BCLAF1, to oppose PU.1 activity resulting in gene expression changes, including suppression of Syk, that promote transition from large to small pre-B cells. After rearrangement of Ig, the block to BCR signaling is lifted, and the immature B cell stage. We propose that RAG DSB-dependent activation of SPIC/BCLAF1 functions as rheostat to titrate PU.1 activity during early B cell development.
STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2019.09.026.

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AUTHOR CONTRIBUTIONS

D.S., L.S.W., W.Y., J.M.A., and R.J. performed experiments and data analyses. N.M. and J.E.P. provided expertise and assisted with data analyses. M.K. and K.M.M. provided mice and expertise for the studies. J.J.B. supervised the project, interpreted experiments, wrote the manuscript, and secured funding.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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### STAR METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| APC human CD25 Clone BC96 | BioLegend | Cat# 302610; RRID: AB_314280 |
| biotin conjugated anti-hCD2 | BD Biosciences | Cat# 555325; RRID: AB_395732 |
| PE- anti-hCD25 Clone BC96 | BioLegend | Cat# 302605; RRID: AB_314275 |
| FITC-conjugated anti-CD45R/B220 (clone RA3-6B2) | BD Biosciences | Cat# 553088; RRID: AB_394618 |
| PE-conjugated anti-CD43 (clone S7) | BD Biosciences | Cat# 553271; RRID: AB_394748 |
| FITC-conjugated anti-CD43 (clone S7) | BD Biosciences | Cat# 553270; RRID: AB_394747 |
| PE-Cy7-conjugated anti-CD45/B220 (clone RA3-6B2) | BD Biosciences | Cat# 552772; RRID: AB_394458 |
| allophycocyanin (APC)-conjugated anti-IgM (clone II/41) | BD Biosciences | Cat# 60642; RRID: AB_1727443 |
| PE-conjugated anti-hCD2 | BD Biosciences | Cat# 555327; RRID: AB_398464 |
| APC-conjugated anti-hCD2 | BD Biosciences | Cat# 560642; RRID: AB_1727443 |
| Anti-SYK (clone D115Q) | Cell Signaling Technology | Cat# 12358; RRID: AB_2687923 |
| Anti-BCLAF1 antibody (A300-608A) | Bethyl Laboratories | Cat# A300-608A; RRID: AB_513581 |
| PU.1 (PA5-17505) | Thermo Fisher | Cat# 17505; RRID: AB_10989141 |
| Anti-GAPDH | Cell Signaling | Cat# 5174; RRID: AB_10622025 |
| anti-FLAG (clone M2) | Sigma | Cat# 1804; RRID: AB_262044 |
| HRP-conjugated anti-mouse IgG | Cell Signaling | Cat# 7074; RRID: AB_2099233 |
| HRP-conjugated anti-rabbit IgG | Cell Signaling | Cat# 7065; RRID: AB_10890862 |
| Anti-HA | Abcam | Cat# ab9110; RRID: AB_307019 |
| control rabbit IgG | Millipore | Cat# 06-371; RRID: AB_390146 |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| SuperScriptII | Life Technologies | 18064-014 |
| Brilliant II SYBR Green | Agilent | 600828 |
| Interleukin-7 (IL-7) | Miltenyi Biotec | 130-098-222 |
| Imatinib | Novartis | 00078-0438-15 |
| Lipofectamine 2000 | Life Technologies | 11668-019 |
| PEG-8000 | Fisher | P156-500 |
| Sequabrene | Sigma | S 2667 |
| ATM inhibitor KU55933 | Tocris | 3544 |
| Micrococcal nuclease | New England Biolabs | M0247S |
| **Critical Commercial Assays** |        |            |
| RNasey | QIAGEN | 74104 |
| Protein A Dynabeads | Thermo Fisher Scientific | 10002D |
| QIAquick PCR purification kit | QIAGEN | 28106 |
| Anti-biotin magnetic beads | Miltenyi Biotec | 130-090-485 |
| Anti-hCD25 magnetic beads | Miltenyi Biotec | 130-092-983 |
| Anti-hCD2 magnetic beads | Miltenyi Biotec | 130-091-114 |
| High sensitivity DNA ChiPs | Agilent Technologies | 5067-4626 |
| Magnetic Separation columns | Miltenyi Biotec | 130-042-201 |
| ECL | Pierce | 32209 |
| ECL Prime | GE Healthcare | RPN2232 |
| FITC BrdU Flow Kit | BD Biosciences | 559619 |
| **Deposited Data** |        |            |
| Chip-seq | This paper | NCBI GEO #: GSE129130 (subseries: GSE129124) |
| RNA-seq | This paper | NCBI GEO #: GSE129130 (subseries: GSE129129) |

(Continued on next page)
**LEAD CONTACT AND MATERIALS AVAILABILITY**

Further information and request for resources and reagents should be directed to the Lead Contact, Jeff Bednarski (bednarski_j@wustl.edu). All unique/stable reagents, including plasmids and mouse lines, are available from the Lead Contact with a completed Materials Transfer Agreement.

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Experimental Models: Cell Lines** |
| Art<sup>–/–</sup>:Bcl2 abl pre-B cell | Barry Sleckman | N/A |
| Rag<sup>–/–</sup>:Bcl2 abl pre-B cell | Barry Sleckman | N/A |
| PlatE | Cell Biolabs, Inc | RV-101 |
| **Experimental Models: Organisms/Strains** |
| Mb1-cre (Cd79a<sup>tm1(Cre)Reth</sup>) mice | Jackson Laboratory | 20505 |
| Bclaf1<sup>fl/fl</sup> mice | KOMP Repository | Bclaf1<sup>fl/fl</sup> |
| Rag1<sup>–/–</sup>:<sup>μ</sup>lgh:Bcl2 | Barry Sleckman | N/A |
| Art<sup>–/–</sup>:<sup>μ</sup>lgh:Bcl2 | Barry Sleckman | N/A |
| B6.Cg-Tg(ACTFLPe)9205Dym/J | Jackson Laboratory | 5703 |
| Spic<sup>gefp/igfp</sup> | Kenneth Murphy | N/A |
| **Oligonucleotides** |
| Primers are listed in Table S5 | This paper | N/A |
| **Recombinant DNA** |
| MSCV-hCD2-mir30 vector | Mark Schlissel | N/A |
| pFLRU-TRE-FLAG-HA-PU.1-Ubc-rTA-ires-Thy1.2 | This paper | N/A |
| pFLRU-TRE-FLAG-HA-SPIC-Ubc-rTA-ires-Thy1.2 | This paper | N/A |
| pCMV-VSV-G | Stewart et al., 2003 | Addgene #8454 |
| pCMV-d8.2R dvpr | Stewart et al., 2003 | Addgene #8455 |
| **Software and Algorithms** |
| Bowtie v. 1.1.2 | Langmead and Salzberg, 2012 | http://bowtie-bio.sourceforge.net/bowtie2/index.shtml RRID:SCR_005476 |
| MACS version 2 | Zhang et al., 2008 | http://liulab.dfci.harvard.edu/MACS/ RRID:SCR_013291 |
| Galaxy V18.09 | Afgan et al., 2016 | https://usegalaxy.org RRID:SCR_006281 |
| GenomicRanges | Lawrence et al., 2013 | https://www.bioconductor.org/packages//2.10/bioc/html/GenomicRanges.html RRID:SCR_000025 |
| Bedtools V2.25.0 | Quinlan and Hall, 2010 | https://github.com/arq5x/bedtools2. RRID:SCR_006646 |
| EdgeR- TMM | Robinson et al., 2010 | http://bioconductor.org/biocLite.R RRID:SCR_012802 |
| Limma-Voom | Law et al., 2014 | https://omicstools.com/limma-fool RRID:SCR_010943 |
| TreeView Version 1.1.6r4 | Saldanha, 2004 | https://bitbucket.org/TreeView3Dev/treeview3/ RRID:SCR_016916 |
| Heatmap3 | Zhao et al., 2014 | https://bioconductor.org/packages/release/bioc/html/heatmaps.html |
| DAVID (v.6.8) | Huang et al., 2009 | https://david.ncifcrf.gov RRID:SCR_001881 |
| EaSeq (v1.111) | Lerdrup et al., 2016 | http://easeq.net |
| MAnorm | Shao et al., 2012 | http://bcb.dfci.harvard.edu/~gcyan/MAnorm/ MAnorm.htm RRID:SCR_010869 |
| Prism 8 (v8.0.2) | GraphPad Software | https://www.graphpad.com |
**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Mice:** All mice were bred and maintained under specific pathogen-free conditions at the Washington University School of Medicine and were handled in accordance to the guidelines set forth by the Division of Comparative Medicine of Washington University. Mb1-cre (Cd79atm1(cre)Reth) mice were purchased from The Jackson Laboratory. Bclaf1f/f mice were generated by the trans-NIH Knock-Out Mouse Project (KOMP) and obtained from the KOMP Repository (www.komp.org). Rag1−/−;mIgh:Bcl2 and Art−/−;mIgh:Bcl2 were generated as previously described (Bednarski et al., 2012, 2016). Spicf/f(Rspictm2.1Kmm) were kindly provided by K. M. Murphy (Haldar et al., 2014). Spicf/f(Rspictm2.1Kmm), Mb1-cre, Bclaf1f/f and Bclaf1f/f;Mb1-cre mice are on a B6 background. All other mice are on a mixed genetic background. Both sexes were used equivalently in all experiments. In vivo studies were conducted on 4-5 week old mice.

**Cell Lines and Primary Cultures**

Rag1−/−;Bcl2 and Art−/−;Bcl2 abl pre-B cells were a gift from Barry Sleckman. Cell lines were authenticated by genotyping. To induce cell cycle arrest and induction of RAG DSBs, cell lines were treated with 3 µM imatinib for indicated times (Bredemeyer et al., 2008). Primary pre-B cell cultures were generated by culturing bone marrow from 4-6 week old mice at 2 × 10^6 cells/mL in media containing 5 ng/mL of IL-7 (Miltenyi Biotec) for 7-10 days (Bednarski et al., 2012, 2016). Both sexes were used equivalently in all experiments. For IL-7 withdrawal experiments, cells were resuspended in media without IL-7 and maintained at 2 × 10^6 cells/mL for the indicated times. ATM inhibitor KU55933 (15 µM; Tocris) was added to cultures at time of addition of imatinib or IL-7 withdrawal.

**METHOD DETAILS**

**cDNA Expression and shRNA-Mediated Knock-down**

cDNAs for SPIC and PU.1 with 5′ FLAG-HA tag were individually cloned into the pFLRU-TRE-Ubc-rtTA-IRES-Thy1.2 lentiviral vector. shRNA targeting Bclaf1 (sequence: 5′-CCCTCATAGTCTTCACCTATT-3′) was cloned into the MSCV-hCD2-mir30 vector (Bednarski et al., 2012). Retrovirus was produced in platE cells by transfection of the retroviral plasmid with Lipofectamine 2000 (Life Technologies) according to the manufacturer’s protocol. Lentivirus was produced in 293T cells by transfection of the lentiviral plasmid along with pCMV-VSV-G and pCMV-d8.2R plasmids with Lipofectamine 2000 (Stewart et al., 2003). Viral supernatant was collected and pooled from 24-72 hours after transfection. Viral supernatant was used immediately to transduce cells or was concentrated prior to transduction. To concentrate viral particles, PEG-8000 (Sigma; final concentration 8%) was added to viral supernatant followed by centrifugation at 2500 RPM for 20 minutes. Precipitated virus was resuspended at 300x concentration in sterile PBS. Pre-B cells were transduced with unconcatenated virus (10 × 10^6 cells in 1 mL viral supernatant) or with concentrated virus (40 × 10^6 in 1 mL with 10x viral particles) in media with polybrene (6 µg/mL; Sigma) by centrifugation for 90 min at 1300 RPM at room temperature. Four hours later fresh media was added and the cells were incubated overnight. Virus-containing media was removed and cells were cultured in fresh media (2 × 10^6/mL). Cells expressing the retrovirus construct were identified by flow cytometric assessment of hCD25 or hCD2 expression using a FACSCalibur (BD Biosciences). Transduced cells were sorted using biotin conjugated anti-hCD2 or anti-hCD25 (BD Biosciences) and anti-biotin magnetic beads (Miltenyi Biotec) on MS columns (Miltenyi Biotec) according to the manufacturer’s protocol.

**Flow Cytometric Analyses and Cell Sorting**

Flow cytometric analyses were performed on a FACSCalibur or BD LSRFortessa (BD Biosciences). Sorting was conducted on a Sony Sy3200 through the Siteman Cancer Center Flow Cytometry Core Facility. Fluorescein isothiocyanate (FITC)-conjugated anti-CD45R/B220 (clone RA3-6B2), phycoerythrin (PE)-conjugated anti-CD43 (clone S7), FITC-conjugated anti-CD43 (clone S7), PE-Cy7-conjugated anti-CD45/B220 (clone RA3-6B2), allophycocyanin (APC)-conjugated anti-IgM (clone II/41), APC-conjugated anti-hCD2, and PE-conjugated anti-hCD2 were purchased from BD Biosciences. PE-conjugated anti-hCD25 (clone BC96) and APC-conjugated anti-hCD25 (clone BC96) were purchased from BioLegend.

**Cell Cycle Analysis**

To assess pre-BCR driven proliferation, pre-B cells were resuspended in media without IL-7 and maintained at 2 × 10^6 cells/mL. Twenty-four hours after removal from IL-7 cells were pulsed BrdU for two hours using the BrdU-FITC kit (BD Biosciences) per the manufacturer’s instructions. DNA content was assayed by 7AAD (BD Biosciences).

**Western Blot**

Western blots were done on whole cell lysates (Bednarski et al., 2016). Anti-SYK (clone D11SQ) and anti-GAPDH (clone D16H11) antibodies were from Cell Signaling Technology. Anti-BCLAF1 antibody (A300-608A) was from Bethyl Laboratories. Anti-PU.1 (PA5-17505) was from Thermo Fisher Scientific. Anti-FLAG (clone M2) was from Sigma. Secondary reagents were horseradish peroxidase (HRP)-conjugated anti-mouse IgG (Cell Signaling; catalog # 7076) or anti-rabbit IgG (Cell Signaling; catalog # 7074). Westerns were developed with ECL (Pierce) and ECL Prime (GE Healthcare).
RT-PCR
For genomic DNA isolation, cells were lysed in lysis buffer (100 mM TRIS pH8.5, 5 mM EDTA, 200 mM NaCl and 0.2% SDS) and DNA was precipitated by addition of isopropanol, washed with 70% ethanol and then resuspended in Tris-EDTA buffer (Bredemeyer et al., 2009). RNA was isolated using RNeasy (QIAGEN) and reversed transcribed using a polyT primer with SuperScriptIII (Life Technologies) according to the manufacturers’ protocol. RT-PCR was performed using Brilliant II SYBR Green (Agilent) and acquired on an Mx3000P (Stratagene). Each reaction was run in triplicate. Values were normalized to housekeeping genes as indicated, and fold change was determined by the ΔΔ cycle threshold method. Primer sequences are listed in Table S5.

Chromatin Immunoprecipitation (ChIP) and ChIP-Seq
ChIP was performed using anti-PU.1 (PAS-17505, Thermo Fisher Scientific), anti-FLAG (clone M2, Sigma), anti-HA (ab9110, Abcam), anti-BCLAF1 (A300-608A, Bethyl Laboratories), control rabbit IgG (Millipore) and control mouse IgG antibodies (clone P3.6.2.8.1, eBioscience) as previously described (Bednarski et al., 2016). Briefly, DNA was cross-linked with 2% formaldehyde for 10 min at room temp (1 × 10⁶ cells/ml). Reaction was stopped with 125 μM Glycine. Cells were lysed with NP-40 and nuclei were frozen in liquid nitrogen then lysed with SDS. DNA was fragmented by sonicating with 30 s pulses for 60 cycles using a Bioruptor (Diagenode). DNA fragmentation was in the range of 200-500 bp and was monitored by agarose gel electrophoresis. Immunoprecipitation was performed with anti-PU.1 (1:100), anti-HA (1 μg), anti-BCLAF1 (2 μg), or control rabbit IgG and Protein A Dynabeads (Life Technologies). DNA was eluted, reverse cross-linked and then purified with QiAquick PCR purification kit (QIAGEN). For ChIP-PCR analysis, PCR was performed using Brilliant II SYBR Green (Agilent) and acquired on an Mx3000P (Stratagene). Primers are listed in Table S5. For ChIP-seq analysis, fragmented DNA was quantified using 2100 Bioanalyzer (Agilent Technologies) and DNA libraries were prepared using Illumina TrueSeq. Sequencing was performed using an Illumina HiSeq 3000 by the Washington University Genome Technology Access Center. Input controls were used for all samples. FASTQ files were aligned to mm9 using Map with Bowtie for Illumina v. 1.1.2 to the reference genome (NCBI37/mm9) (Langmead and Salzberg, 2012). MACS version 2 was used to call peaks with a tag size set to 45, band width of 300 and a p value of 1 × 10⁻⁵ (Zhang et al., 2008). Input. bed files of total reads for MM-ChIP were generated using Convert from BAM to BED tool v0.1.0 in Galaxy V18.09 (Afgan et al., 2016). Promoter regions were defined as regions extending 12 kb upstream of transcription start site. R package (GenomicRanges) and Bedtools V2.25.0 were used to determine overlapping ChIP peaks (Lawrence et al., 2013; Quinlan and Hall, 2010). MANorm using parameters –w 300–s1 50–s2 50 was used to calculate normalized fold changes for each ChIP-seq comparison (Shao et al., 2012). A 1.5 fold change magnitude was used to separate enriched and unbiased peaks for each comparison. EaSeq v1.111 was used to generate ratiometric heatmaps from RPM-normalized ChIP-seq signal (Lerdrup et al., 2016). Data will be deposited in NCBI’s Gene Expression Omnibus.

Ultra-Low-Input Native ChIP
EGFP-negative (-) and EGFP-expressing (+) small pre-B cells were sorted from SPI/Cigfp/igfp mice. ULI-NChIP was performed as previously described (Brind’Amour et al., 2015). Briefly, chromatin was fragmented using micrococcal nuclease (New England Biolabs) at 37 °C for 5 mins and diluted in complete immunoprecipitation buffer (20 mM Tris-HCl pH 8.0, 2 mM EDTA, 15 mM NaCl, 0.1% Triton X-100, protease and phosphatase inhibitors). Fragmented chromatin was precleared with Protein A Dynabeads (Life Technologies). Immunoprecipitation was performed with anti-PU.1 (1:100), anti-BCLAF1 (10 μg), or control rabbit IgG and Protein A Dynabeads (Life Technologies). The antibody-beads complex was washed with low salt (20 mM Tris-HCl, pH 8.0, 0.1% SDS, 1% Triton X-100, 0.1% deoxycholate, 2 mM EDTA and 150 mM NaCl) and high salt (20 mM Tris-HCl, pH 8.0, 0.1% SDS, 1% Triton X-100, 0.1% deoxycholate, 2 mM EDTA and 300 mM NaCl) buffer. DNA was eluted in high salt buffer. DNA was purified and ChIP-PCR was performed as above.

RNA-Seq Analysis
RNA was extracted using RNeasy Kit (QIAGEN). Libraries were prepared using Illumina TrueSeq Adapters and paired-end sequencing was performed using an Illumina HiSeq 3000 by the Washington University Genome Technology Access Center according to the manufacturer’s protocols. Sequencing data were analyzed as previously described (Andley et al., 2018). Briefly, RNA-seq reads were aligned to mm9 assembly with STAR version 2.0.4b1. Gene counts were derived from uniquely aligned unambiguous reads by Subread-featureCount version 1.4.5. Gene-level counts were imported into the R/Bioconductor package EdgeR and TMM normalization size factors were calculated to adjust for differences in library size (Robinson et al., 2010). Differential expression analysis was then performed to analyze for differences between conditions using the R/Bioconductor package limma-voom (Law et al., 2014). Results were filtered for only those genes with Benjamini-Hochberg false-discovery rate adjusted p values less than or equal to 0.05. DAVID (Database for Annotation, Visualization and Integrated Discovery, v6.8) was used to test if differentially expressed genes resulted in perturbations in known Gene Ontology (GO) terms and KEGG pathways (Huang et al., 2009). Volcano plots were generated using R (ggplot2). Java TreeView Version 1.1.6r4 and R/Bioconductor package heatmap3 were used to display heatmaps (Saldana, 2004; Zhao et al., 2014). DAVID was used to display annotated KEGG graphs across groups of samples for each GO term or KEGG pathway with a Benjamini-Hochberg false-discovery rate adjusted p value ≤ 0.05.

Tandem Affinity Purification and MS Analysis
FLAG-HA-tagged SPIC and PU.1 were immunoprecipitated using anti-FLAG antibody as previously described with the following modifications (Mosammaparast et al., 2013; Nakatani and Ogryzko, 2003). Cells were lysed lysis of cells (1 × 10⁶ cells/1.5 ml) in
TAP buffer (50 mM Tris, pH 7.9, 150 mM NaCl, 1% NP-40, and protease and phosphatase inhibitor cocktails (Sigma). The lysate was cleared by centrifugation and incubated with anti-FLAG beads (40 μl/10^9 cells; clone M2; Sigma-Aldrich) for 4 hours. After extensive washing in the same buffer, bound material was eluted with FLAG peptide (Sigma-Aldrich) and analyzed by western blotting. Coomassie-stained bands were cut from SDS-PAGE and sent to Taplin Biological Mass Spectrometry Facility at Harvard Medical School (taplin.med.harvard.edu). In-gel trypsin digestion was performed and the detection of complexed proteins was done using Orbitrap ion-trap mass spectrometers (ThermoFisher Scientific). Interacting proteins were identified by matching protein database with acquired fragmentation pattern by using Sequest (ThermoFisher Scientific) (Eng et al., 1994).

QUANTIFICATION AND STATISTICAL ANALYSIS

RNA-seq and ChIP-seq were analyzed for statistical significance using the software packages described above. For all other analyses, statistics and figures were generated using Prism 8 (v8.0.2). P values were generated via Student’s t test (unpaired, two-tailed). Error bars are SE. *p value ≤ 0.05, **p value ≤ 0.01, ***p value ≤ 0.001, ****p value ≤ 0.0001.

DATA AND CODE AVAILABILITY

The ChIP-seq and RNA-seq data generated during this study are available at NCBI Gene Expression Omnibus under accession number GEO: GSE129130.
Supplemental Information

RAG-Mediated DNA Breaks Attenuate PU.1 Activity in Early B Cells through Activation of a SPIC-BCLAF1 Complex

Deepti Soodgupta, Lynn S. White, Wei Yang, Rachel Johnston, Jared M. Andrews, Masako Kohyama, Kenneth M. Murphy, Nima Mosammaparast, Jacqueline E. Payton, and Jeffrey J. Bednarski
Figure S1. RAG DSBs do not alter PU.1 expression or distribution among genomic regions. Related to Figure 1. (A) Western blot of PU.1 levels in Rag1−/-:Bcl2 and Art−/-:Bcl2 abl pre-B cells treated with imatinib for 48 hours. Data are representative of three independent experiments. (B) Gene ontology analysis identifying the biological processes related to genes neighboring PU.1 ChIP-seq peaks from Figure 1B. (C) Graph depicting enrichment of PU.1 binding across genomic regions based on PU.1 ChIP-seq in Figure 1B.
Figure S2

Figure S2. SPIC and PU.1 bind similar genomic regions. Related to Figure 3. Representative tracks for PU.1 and SPIC ChIP-seq at indicated positions from Figure 3B. ChIP-seq was performed as described in Figure 3B.
Figure S3. Inducible expression of SPIC and PU.1 in pre-B cells. Related to Figure 4. Western blot shows expression of FLAG-HA-PU.1 and FLAG-HA-SPIC in Art<sup>+/−</sup>:Bcl2:Pu.1<sup>tet</sup> and Art<sup>+/−</sup>:Bcl2:Spic<sup>tet</sup>, respectively, after treatment with imatinib and 2 mM doxycycline for 48 hours (as in Figure 4A). * indicates non-specific band. Data are representative of three independent experiments.
Figure S4

A.

![Graphs showing gene expression changes](image)

B. GO analysis of genes repressed by SPIC and rescued by BCLAF1 knockdown

| Enriched Pathways                    | p-value (adj) |
|--------------------------------------|--------------|
| Immune response                      | 0.00105      |
| Response to cytokine stimulus        | 5.60 e-8     |
| B cell mediated immunity             | 0.00105      |
| Negative regulation of migration     | 0.00398      |

Figure S4. SPIC and BCLAF1 regulate a gene expression in pre-B cells. Related to Figure 5. (A) RT-PCR validation of representative gene expression changes from Figure 5C. Data are mean and standard error for three independent experiments. **p-value ≤ 0.01, ***p-value ≤ 0.001, ****p-value ≤ 0.0001. (B) Gene ontology analysis depicting immune and B cell function pathways related to genes from Figure 5C that are repressed upon SPIC induction in Rag1⁻/⁻:Bcl2:Spictet cells and rescued following BCLAF1 knockdown.
Figure S5

A. Heatmap of the 141 genes identified in Figure 5F. Cells were treated as in Figure 5C and 5E. (B) Gene ontology analysis depicting immune and B cell function pathways related to the 141 genes identified in the Figure 5F. Representative genes in each category are listed. All pathways have p-value ≤ 9.77 x 10^{-5}.
Figure S6. Analysis of bone marrow B cell populations. Related to Figure 7. Whole bone marrow was stained with B220, IgM and CD43 to quantify B cell populations as indicated. Data are representative of three independent experiments.
Table S4. RAG DSBs, SPIC and BCLAF1 cooperatively regulate expression of a cohort of genes in pre-B cells. Related to Figure 5. Comparison of RAG DSB-dependent, SPIC-dependent, and BCLAF1-dependent gene expression changes from Table S2 and S3.

| Gene1 | Gene2 | Gene3 | Gene4 | Gene5 | Gene6 |
|-------|-------|-------|-------|-------|-------|
| A2ml1 | Fmd4b | Ppm1j | Ceacam9 | Lhfpl4 | Sfmbt2 |
| AA467197 | Gh | Ptafr | Cfap70 | Lrrc15 | Slc15a2 |
| Ackr2 | Gm14326 | Pygl | Clmn | Lrrc36 | Slc4a5 |
| Acot7 | GnaI1 | Rab11fip5 | Cntd1 | Lm4 | Slc5a11 |
| Aes | Gprasp2 | Rab44 | Col28a1 | Lyplal1 | Slc5a5 |
| Ahnak2 | H2-Aa | Rasgrp3 | Cpeb1 | Mcoln3 | Sncb |
| Angptl4 | H2-DMa | Rbp2 | Crhbp | Mfsd2a | Sntb1 |
| Ankrd37 | Hapln1 | Rgs1 | Crisp1 | Mmp14 | Spaca9 |
| Arrdc5 | Hck | Rgs11 | Ctl4a | Ms4a1 | Stac |
| Art5 | Hhatl | Rgs13 | Ctnnd2 | Myc | Steap4 |
| Atf3 | Hist1h2ae | Rimkla | Dll1 | Myh7 | Taf9b |
| Bambi | Hpn | Rpl3 | Dmpk | Nid2 | Tbxas1 |
| BC049352 | Hspg2 | Rps3a1 | Dnajc22 | Nkd1 | Tcaim |
| Bcat1 | Iftim3 | Rshp1 | Dmn1 | Nox1 | Tdrd9 |
| C4bp | Igf2 | Rxta | Dpt | Npr2 | Tigit |
| Cabp1 | Il17f | Ryr3 | Dse | Nrxn3 | Tmprss4 |
| Cacna1e | Il1m | S100a4 | Dtx1 | Nxph4 | Tnfrsf9 |
| Cacna2d1 | Il2rb | Scd4 | Eda2r | Olfm1 | Tox |
| Capn11 | Iqgap2 | Scn11a | Etv5 | Otog | Trp73 |
| Cend2 | Itih3 | Sell | Fam129a | Paqr4 | Tsnaxip1 |
| Ccne1 | Kcnn4 | Serpina3f | Fcer2a | Pcdh19 | Uaca |
| Cd86 | Kcd14 | Serpina3g | Fdg4 | Pinlyp | |
| Fos | Pitpm1 | Wfdc5 | Flrt1 | Uchl1 | |
| Zan | Plekg3 | Wnt10b | Pipox | Vldlr | |
### Table S5. Primer sequences. Related to STAR methods.

List of primers used for genotyping, RT-PCR and ChIP-PCR.

| Primer Name | Application                  | Sequence                                      |
|------------|-----------------------------|-----------------------------------------------|
| Jk1_F      | qRT PCR for DNA breaks      | GCTACCCACTGCTCTGFTCC                         |
| Jk1_R      | qRT PCR for DNA breaks      | CTTGGGAGAGTGCCAGAATC                         |
| Syk_F      | qRT PCR                     | TCTCTGCTAGCTCTGAGCC                         |
| Syk_R      | qRT PCR                     | CAGAGGCCTCAGAGCCTCC                         |
| Spic_F     | qRT PCR                     | TCTCTGCTAGCTCTGAGCC                         |
| Spic_R     | qRT PCR                     | CAGAGGCCTCAGAGCCTCC                         |
| B-Actin_F  | qRT PCR                     | AAACATTTCAAGAGCCCATGGAC                     |
| B-Actin_R  | qRT PCR                     | CTCTGACGTTGAGATAAGGTC                      |
| Xcrr3_F    | qRT PCR                     | ATCCCTAGTGCCCTCCTT                          |
| Xcrr3_R    | qRT PCR                     | TTTGCTGAGACCTGCTTC                          |
| Zap70_F    | qRT PCR                     | TGGTACCTTGTGCGAGAATG                        |
| Zap70_R    | qRT PCR                     | ATCTCCGAGATCTTCATGCAAT                     |
| Cd86_F     | qRT PCR                     | GGGGTCTTGGACTGCTGCTG                       |
| Cd86_R     | qRT PCR                     | GCCCTGGCTGATGAGGCTGTG                      |
| Syk_CHIP_F | ChIP qRT PCR                | GGGGAACTGAGCCCTAAAG                        |
| Syk_CHIP_R | ChIP qRT PCR                | TATAGGGCAGTGGTGAGAG                        |
| Dapk1_CHIP_F | ChIP qRT PCR            | TCATAGCTACCAGCAGACTG                       |
| Dapk1_CHIP_R | ChIP qRT PCR          | AACAGATGTGCTAAGGTC                         |
| Cd200_CHIP_F | ChIP qRT PCR           | GCTAGGATCAAAGAATCCAGTC                     |
| Cd200_CHIP_R | ChIP qRT PCR           | GCTAGGATCAAAGAATCCAGTC                     |
| Sos2_CHIP_F | ChIP qRT PCR              | TGAAGTTAGAGGGCTGTCG                        |
| Sos2_CHIP_R | ChIP qRT PCR              | AGGAAAGGACGGCTTCGAC                       |
| Ragl_1     | Genotyping                 | AGAAGGAGAGGATTCTCAAGAGGG                    |
| Ragl_2     | Genotyping                 | TTGGAGAGCTGCTGAGCTG                        |
| Ragl_3     | Genotyping                 | ACCGCCATCACAGATACTGGG                      |
| IgH_F      | Genotyping                 | CAGGGTTATTTGCTACTAGGC                      |
| IgH_R      | Genotyping                 | CATTCCTACCTCCACCTGCTCCCT                   |
| Bcl2_F     | Genotyping                 | CGAGATGTGCTGACAGCTGACCTG                   |
| Bcl2_R     | Genotyping                 | TCACCTTGAGCGCCAGATAGCGACCACCA              |
| Art_1      | Genotyping                 | CAAGAGGATCATCGTGATAGGGTCC                   |
| Art_2      | Genotyping                 | CCCGTAACACAGCAGATGACAGGACCCGG              |
| Art_3      | Genotyping                 | ACCCCAGGCTATCCTGACCACC                     |
| Mb1 Cre_F  | Genotyping                 | CATTTCGAGGAGACCTCA                         |
| Mb1 Cre_R  | Genotyping                 | ACTGAGGCAGAGATTGG                          |
| Bcalf1_1   | Genotyping                 | AGACCTTGAGTACACATACCTGTGAATCCC             |
| Bcalf1_2   | Genotyping                 | ACTAACACATCACAATGACAGCTCC                  |
| Bcalf1_3   | Genotyping                 | AGCAGAAACTTACGCGAACAGTGGTC                |
| FlpE_F     | Genotyping                 | ACTCCGTTAGGGCCCTTAC                       |
| FlpE_R     | Genotyping                 | GCCCTTATTCAATCCTT                           |