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HIGHLIGHTS
Setd2/H3K36me3 is essential in maintaining a normal hematopoiesis
Loss of Setd2/H3K36me3 impairs lymphogenesis and V(D)J recombination
Loss of Setd2/H3K36me3 and ATM kinase activity leads to mis-repaired recombination
Setd2/H3K36me3 prevents apoptosis of post-mitotic neuronal cells

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Loss of H3K36 Methyltransferase SETD2 Impairs V(D)J Recombination during Lymphoid Development

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SUMMARY

Repair of DNA double-stranded breaks (DSBs) during lymphocyte development is essential for V(D)J recombination and forms the basis of immunoglobulin variable region diversity. Understanding of this process in lymphogenesis has historically been centered on the study of RAG1/2 recombinases and a set of classical non-homologous end-joining factors. Much less has been reported regarding the role of chromatin modifications on this process. Here, we show a role for the non-redundant histone H3 lysine methyltransferase, Setd2, and its modification of lysine-36 trimethylation (H3K36me3), in the processing and joining of DNA ends during V(D)J recombination. Loss leads to mis-repair of Rag-induced DNA DSBs, especially when combined with loss of Atm kinase activity. Furthermore, loss reduces immune repertoire and a severe block in lymphogenesis as well as causes post-mitotic neuronal apoptosis. Together, these studies are suggestive of an important role of Setd2/H3K36me3 in these two mammalian developmental processes that are influenced by double-stranded break repair.

INTRODUCTION

In early normal lymphocyte development, gene segments that will eventually encode the immunoglobulin (Ig) and T cell receptor (TCR) variable regions are recombined from Variable (V), Diversity (D), and Joining (J) gene segments in a process known as V(D)J recombination (Alt et al., 2013). During the DNA recognition and cleavage stage, recombination signal sequences (RSSs) that flank the individual V, D, and J gene segments are targets of RAG1/2 endonucleases and result in the generation of hairpinned coding ends (CEs) and blunt-ended signal ends (SEs) (Alt et al., 2013; Schatz and Swanson, 2001). In the second phase of V(D)J recombination, and-processing and end-ligation of CEs and SEs are mediated by classical non-homologous end-joining (C-NHEJ) factors and produce an imprecisely repaired coding joint (CJ) consisting of V(D)J exons and a precisely repaired but discarded circular signal joint (SJ) (Alt et al., 2013; Schatz and Swanson, 2001). A set of core C-NHEJ factors (KU70, KU80, XRCC4, and LIG4) is absolutely essential for end-joining and is evolutionarily conserved (Alt et al., 2013; Kumar et al., 2014). Loss or defects of C-NHEJ factors can impair end-processing (DNA-PKcs, ARTEMIS) or end-joining (KU proteins, XRCC4, XLf, LIG4) and results in severe immunodeficiencies in both mouse models and human disease (Alt et al., 2013; Kumar et al., 2014; Bassing et al., 2002). The DNA damage protein, ataxia telangiectasia mutated (ATM), its target, histone H2AX; and DNA damage response adaptor protein, MRE11, are all also involved in the end-ligation process (Bredemeyer et al., 2006; Yin et al., 2009; Hung et al., 2018). The single loss of any of these factors, or C-NHEJ factor XLf, has only modest effects on lymphogenesis and V(D)J recombination (Bredemeyer et al., 2006; Yin et al., 2009; Hung et al., 2018; Li et al., 2008). In addition to loss of the core C-NHEJ factors, combined deficiencies of proteins non-essential for the end-joining reaction can severely impair C-NHEJ to a similar extent, as in the case of combined loss of XIfl and Atm or XIfl and Mre11 (Kumar et al., 2016; Hung et al., 2018; Li et al., 2008; Lescale et al., 2016; Zha et al., 2011).

Another mammalian developmental process that utilizes C-NHEJ for repair of double-strand breaks (DSBs) is embryonic neurogenesis (Frappart and McKinnon, 2008). Neural progenitors that have exited the cell cycle and are migrating out of the embryonic ventricular zones as they differentiate are thought to rely on NHEJ-mediated repair of DSBs (Frappart and McKinnon, 2008). In mice, loss of core C-NHEJ factors leads to apoptosis of post-mitotic neurons and embryonic lethality (Gao et al., 1998; Frank et al., 2000; Gu et al., 2001). This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
Histone H3 lysine-36 tri-methylation (H3K36me3) is a histone modification that is catalyzed by the non-redundant histone methyltransferase, SETD2 (Wagner and Carpenter, 2012). H3K36me3 is associated with actively transcribed genes, and SETD2 plays important roles in the control of gene expression (Wagner and Carpenter, 2012). Loss-of-function mutations in SETD2 or dominant negative “onco-histone” mutations in the H3K36 residue itself have been described in a broad array of malignancies, particularly in hematopoietic and central nervous system (CNS) tumors (Parker et al., 2016; Zhang et al., 2012; McKinney et al., 2017; Moffitt et al., 2017; Zhu et al., 2014; Lu et al., 2016). In mammalian cells, SETD2 regulates specific steps of the DNA damage response during mismatch repair (MMR) and homologous recombination (HR) (Li et al., 2013; Pfister et al., 2014; Aymard et al., 2014). More recently, a role for Setd2 in normal thymocyte development and V(D)J recombination was described (Ji et al., 2019). Although a role for H3K36 methylation in NHEJ had been previously suggested in yeast (Fnu et al., 2011), insights into the mechanism for how this post-translation histone modification in mammalian cells may impact this mode of repair remains unknown. Thus, to determine the role, if any, of Setd2 and H3K36me3 in this mode of DNA repair in mammals, we studied its loss in two developmental pathways that utilize NHEJ. Here, we specifically show that whereas loss of Setd2/H3K36me3 does not completely abrogate repair of DSBs, loss leads to mis-repair in B-cell lymphoid development of Rag-induced DNA DSBs, especially when combined with loss of ATM kinase activity. Loss of Setd2 leads to the increased formation of aberrant hybrid joints and additionally leads to reductions in overall B cell repertoire. Finally, loss of Setd2 also leads to post-mitotic neuronal apoptosis.

RESULTS

Loss of Setd2 Disrupts Normal Hematopoiesis, Particularly Lymphopoiesis

The complete loss of Setd2 is embryonic lethal at embryonic day 10.5 (E10.5)–E11.5 (Hu et al., 2010). Therefore, to study the role of Setd2 in normal and malignant hematopoiesis, we previously generated a conditional mouse model expressing Cre-recombinase under inducible (Mx1) or constitutive (Vav1) hematopoietic lineage-restricted promoters (Mar et al., 2017). The loss of Setd2 ablated H3K36me3 in hematopoietic tissues through excision of exon 3 of Setd2 (Figure 1A). Heterozygous mice had no overt hematopoietic phenotype (Figures S1A, S1B and S2D), whereas homozygous loss of Setd2 resulted in a significant perturbation of normal hematopoiesis, including decreased overall bone marrow cellularity (Figure 1B), significant loss of mature lymphoid cells (B220+ B cells and CD3+ T cells) in the bone marrow, and expansion of erythroid (Ter119+) cells (Figures 1C and 1D). The significant reduction in T cells in the bone marrow observed upon complete Setd2 loss was also mirrored by a severe diminution of thymic size (Figure 1E), which was concomitant with significant splenomegaly (Figure 1F). Strikingly, the splenomegaly was due to the aberrant expansion of erythroid cells and significant ablation of B-lymphoid (B220+) populations (Figure 1G). In addition, loss of Setd2 induced qualitative and quantitative defects in hematopoietic stem cells, as well as abnormal erythroid progenitor expansion in the bone marrow (Figures 1D, 1G, 1H, S1C–S1F, and S2A–S2C). These hematopoietic phenotypes are consistent with other reports on Setd2 knockout mice (Zhou et al., 2018; Zhang et al., 2018; Ji et al., 2019). Altogether, these data indicate that loss of Setd2 disrupts normal hematopoiesis and severely impacts lymphoid development.

Setd2/H3K36me3 Is Important in Normal Lymphocyte Development

We and others (Zhou et al., 2018; Zhang et al., 2018; Ji et al., 2019) observed that loss of Setd2 early in hematopoiesis resulted in significant deletions of the lymphoid populations in the bone marrow, spleen, and thymus (Figures 1C, 1D, 1G and S1). To rule out that these early developmental deficiencies were not solely the result of reduced numbers of early lymphoid progenitors (Figure S1C), we crossed our knockout mice with multiple B lymphoid lineage-restricted Cre-recombinase-expressing lines and found that whereas an early and profound proB block in development was only observed upon early deletion of Setd2 (Mx1, Vav1cre) (Figure 2A), the deletion of Setd2 in later stages of B cell development (with hCD2, Mb1, and Cd19cre) resulted in abnormal lymphocytic output that was more apparent in more mature B cell populations (Figures 2B and 2C). Specifically, loss of Setd2 induced at later stages of B cell development significantly reduced detectable mature B cells (IgM+IgD+) in the bone marrow (Figures 2B and 2C) and resulted in the significant depletion of B cell lineage cells in the spleen (Figure 2C). These data
suggest that Setd2/H3K36me3 is important in B lymphopoiesis at different stages, but that the severe block at the proB cell stage was only apparent with early loss in hematopoiesis.

**Setd2/H3K36me3 Is Crucial for Normal Immunoglobulin Rearrangement in Early Lymphocyte Development Recombination**

The severe block in early B cell development was striking and warranted further examination. Early loss of Setd2/H3K36me3 blocked B cell development at the proB cell stage with similar total numbers of Fraction A (FrA)-defined pre-proB cells when compared with littermate controls (Figures 3A and S3A–S3E). This block at the
proB cell stage was concomitant with a near-complete ablation of immature IgM+ B cells in the bone marrow and spleen (Figures 3A and S3B). Similarly, Setd2Δ/Δ mice exhibited a block in early T cell development at the double-negative stage (DN: CD4−/CD8−), with an accumulation at the DN3 stage (Figure 3B). This arrest at the DN3 stage was similarly observed in Mx1-cre-driven exon 6-7 deletion Setd2 knockout mouse (Ji, et al., 2019). Thus, lymphopoiesis in Setd2Δ/Δ mice appeared to be arrested at stages wherein V(D)J recombination occurs and is reminiscent of the lymphopenia observed with deficiency of factors necessary for V(D)J recombination (Alt et al., 2013; Kumar et al., 2014; Bassing et al., 2002). In all early B/T cell progenitor populations of Setd2Δ/Δ mice, developmental blocks also coincided with increased levels of apoptosis and phospho-γH2ax (Figures 3C, S3F, and S3G).

Figure 2. Setd2/H3K36me3 Important for B Cell Development at Different Stages
(A) Schematic of ontological expression of various B-lineage-restricted cre-recombinase mouse lines crossed to Setd2f/f mice with Igh locus rearrangement status indicated, and representative flow cytometry of B220+ early B cells progenitors (proB and preB cells) of control, Vav/Mx1, hCD2, Cd19, and Mb1cre Setd2Δ/Δ mice.
(B) (i) Representative flow cytometry of bone marrow stained for early B cell progenitors and mature and immature B cell markers (IgM and IgD). (ii) Total bone marrow cellularity. (iii) Fraction composition of B220+, proB (B220+ Cd43− IgM−), preB (B220+ Cd43+ IgM−), immature B (B220+ Cd43+ IgM−), and mature B cell (B220+ IgM+ IgD+) populations of hCD2 (n = 7), Mb1 (n = 14), Cd19 (n = 9) Setd2Δ/Δ, and sex- and age-matched littermate controls.
(C) Spleen (i) weight (n = 14 for all groups), (ii) total cellularity (n = 3 for all groups), and (iii) percentage composition of different B cell populations in bone marrow and spleen (n = 6 for all groups).
Significance indicated as comparison with controls. *p < 0.05, **p < 0.01, ***p < 0.001, error bars represent SD.
despite similar in vivo proliferation rates and cell cycle status (Figures 3D and S3H). Furthermore, Setd2Δ/Δ proB cells did not display any significant differences in the expression of factors related to V(D)J recombination at the gene or protein level (Figures S4A–S4C). The arrest in B cell, but not in T cell, development in Setd2Δ/Δ mice could be partially rescued by crossing knockout mice with a transgenic mouse expressing a fully rearranged Ig heavy chain complex (IgHelMD4) specific for hen egg lysozyme (HEL) (n = 4 for Setd2f/f, n = 5 for Setd2Δ/Δ MD4, and n = 3 for Setd2Δ/Δ).

*** p<0.001, error bars represent SD. See also Figures S3 and S4.
Figure 4. Loss of Setd2/H3K36me3 Does Not Alter Chromatin Architecture or Accessibility of the Early proB Igh Locus and Causes Abnormal V(D)J Recombination

(A) H3K36me3 ChIP sequencing of the Igh locus of Cd19−/− proB (B220+Cd43+IgM−) cells from bone marrow of Vav Setd2−/− and controls. (n = 3 for all groups). Magnification of region of Igh with focal H3K36me3, loss of representative H3K36me3 tracks overlaid with assay for transposase-accessible chromatin (ATAC)-seq of same region from control proB cells for reference. Annotation of critical regulatory sites as indicated.

(B) Representative ATAC-seq tracks of regulatory region of the Igh locus described in (A) of sorted Cd19− proB cells from two matched control and Setd2 knockout sorted proB cells.

(C) Representative ChIP-PCR of regulatory region for histone H3 marks K36me3, K9ac, and K4me3. Data representative of n = 3 independent experiments.

(D) Representative ChIP-PCR of Rag1 and Hmgb2 of proB cells from Setd2−/− and controls at the same regulatory region with standard deviations as indicated, n = 3 independent experiments.

(E) Relative quantitation of sterile transcription of DH genes, Cμ, and enhancer RNAs of proB cells from Setd2−/− and controls. Data represented as an average of three independent ChIP-PCRs from three independently sorted proB cell populations.
and productive immunoglobulin heavy chain (Igh) locus (Figure 3E), indicating a role for Setd2/H3K36me3 in enforcing normal V(D)J recombination.

**Loss of Setd2/H3K36me3 Does Not Abrogate Chromatin Architecture or Accessibility of the Early proB Igh Locus and Causes Aberrant V(D)J Recombination**

To determine the impact of loss of Setd2/H3K36me3 on the Igh locus at the proB stage, we conducted chromatin immunoprecipitation (ChIP) sequencing and found both a global loss of H3K36me3 across the genome and a focal loss on the Igh locus where a well-studied critical regulatory region near the Eμ enhancer resides (Chowdhury and Sen, 2001) (Figures 4A and S5A). As accessibility of this region is critical for B cell development (Chowdhury and Sen, 2001; Chakraborty et al., 2009), we wanted to ascertain if the loss of H3K36me3 affected local chromatin architecture or accessibility. In proB cells, ablation of H3K36me3 neither affected chromatin accessibility (Figures 4B and S5B) nor disrupted the local levels of H3K4me3 and H3K9ac (Figure 4C), two histone modifications essential for maintaining an open and actively transcribed chromatin structure at this regulatory region (Chowdhury and Sen, 2001; Chakraborty et al., 2009) and for H3K4me3, the recruitment and activation of the Rag2 protein itself (Shimazaki and Lieber, 2014; Johnson et al., 2010; Ji et al., 2010; Matheson and Corcoran, 2012; Bettridge et al., 2017). Loss of H3K36me3 did not significantly affect the methylation states of mono-, di-, or tri-methyl lysine-27 or mono- and di-methyl lysine-36 residues in this region (Figure S5C). We detected equivalent recruitment of Rag1 and Hmg2 to this same region on the Igh locus (Shimazaki and Lieber, 2014; Johnson et al., 2010; Ji et al., 2010; Matheson and Corcoran, 2012) (Figure 4D), suggesting that the initiation phase of the V(D)J recombination reaction was intact. We were also not able to detect in sorted FrA proB cells any evidence of Rag1 recruitment or H3K36me3 at variable gene families on the Igh locus in either Setd2-deficient cells or controls (Figure S5D).

In addition, the level of sterile transcription of Igh genes was only mildly reduced (Figure 4E), particularly when compared with deletion of the Eμ enhancer, which causes significant transcriptional dysregulation (Chakraborty et al., 2009). Upon closer examination of recovered V(D)J recombination products from proB cells, we observed that Setd2 deficiency resulted in aberrant recombination and, in some cases, lack of expected rearrangement products altogether (Figures 4F and S5E). Combined, these data are suggestive that the V(D)J recombination defect is not due to decreased expression or regulation of the Igh locus, but due to defects in the repair phase of the reaction.

**Aberrant End-Joining of Rag-Induced DSB with Setd2/H3K36me3 Deficiency**

To ascertain if defects in Setd2-deficient lymphogenesis could be a consequence of impaired Rag-induced DSB repair during V(D)J recombination, we generated murine Setd2-deficient (Setd2−/−), Ku80−/− (Xrcc5−/−), and Lig4−/− late-proB v-Abelson (v-Abl)-transformed lines expressing a Bcl2 transgene by CRISPR/Cas9-mediated inactivation (Figures S6A–S6D) (Hung et al., 2018; Jacobsen et al., 2006). Loss of Setd2 in v-Abl cells neither perturbed the expression of factors involved in V(D)J recombination nor affected the cell cycle distribution of these cells (Figures S7A–S7C) in vitro. Treatment with Abl kinase inhibitor imatinib (STI) induces G1 cell-cycle arrest and Rag1/2 expression leading to light chain rearrangement (Hung et al., 2018; Jacobsen et al., 2006) (Figure S8A). We additionally introduced a chromosomally integrated inversion recombination substrate (pMG-INV), which can be used to assess the efficiency of V(D)J recombination by measuring GFP expression in cells and/or visualization of the repair products and intermediates by Southern blotting and PCR-based strategies (Hung et al., 2018) (Figure 5A). Inactivation of ATM kinase activity with an inhibitor (ATMi) is sufficient to induce the formation of hybrid joints (HJs, joining of CEs to SEs) (Bredemeyer et al., 2006) and could additionally be used to observe aberrant V(D)J recombination.

Comparative rearrangement efficiency of the pMG-INV substrate was observed in both wild-type (WT) and Setd2−/− v-Abl cells treated with imatinib, as indicated by GFP expression (Figure 5B). As expected, treatment of WT v-Abl cells with ATMi resulted in a modest (25%) decrement in GFP expression (Figure 5B). Strikingly, treatment of Setd2−/− cell lines with ATMi resulted in a ~60% reduction in GFP expression (Figure 5C). This finding is reminiscent of the severe defect in GFP expression found in Xlf-deficient v-Abl cells...
Figure 5. Aberrant End-Joining of Rag-Induced DSB with Setd2/H3K36me3 Deficiency
(A) Schematic of recombination substrate pMG-INV. Unrearranged (UR) and SE and CE intermediates and resulting SJs and CJs. Long-terminal repeats (LTRs), XbaI and NheI restriction digestion sites, recombination signal sequences (RSSs), GFP, Thy1.2 cDNA, and corresponding probes shown.

(B) Representative flow cytometric analysis of GFP expression in control and Setd2−/− pMG-INV v-Abl cells treated with Abl kinase inhibitor imatinib (STI-571) and ATM kinase inhibitor (ATMi, KU55933) for 48 and 96 h.

(C) GFP expression of pMG-INV harboring v-Abl cell lines treated for 72 h with imatinib + ATMi assessed by flow cytometry. Control, Lig4−/−, and at least four independently derived Xrcc5+/− and Setd2+/− v-Abl clones were treated in n = 4 independent induction experiments. Significance to controls to STI-571 treatment alone condition was calculated. ***p < 0.001, error bars represent SD.

(D) Southern blot analysis of genomic DNA from induced Setd2−/− and control lines that were digested with (i and ii) XbaI and (iii) NheI hybridized with (i and iii) Thy1 or (ii) GFP probe. Hybrid Joins (HJ) indicated as well (joint of CEs and SEs).

(E) Schematic of PCR method to detect the formation of a coding joint and hybrid joint recombination product of the pMG-INV retroviral recombination substrate, and PCR result of pMG-INV coding and hybrid joints from indicated v-Abl cell clones treated for 72 h with ABLki with or without ATMki (KU55933). Il-2 gene PCR was used as a loading control. Blue arrow indicates coding joint product, and red arrow indicates hybrid joint product.

(F) PCR strategy to detect endogenous Vκ6-23 to Jκ1 coding joints (CJ) and hybrid joints (HJ) in (i) control and Setd2-deficient v-Abl lines treated for 72 h with STI-571 ± ATMi and (ii) Setd2fl and Setd2Δ/Δ splenocytes. Il-2 gene PCR was used as a loading control and analyzed and quantified by high-sensitivity TapeStation (D1000). Blue arrow indicates CJ product, and red arrow indicates HJ product.

See also Figures S6–S9.
Figure 6. Loss of Setd2/H3K36me3 Reduces CDR3 Repertoire and Variable Gene Usage in proB Cells

(A) PCR and next-generation sequencing (NGS) strategy of V(D)J recombination products of VH families to JH4; 500-bp product (blue box) was extracted and submitted for NGS amplicon sequencing. Sequencing was analyzed with MiXCR T/B cell repertoire software (Bolotin et al., 2017).

(B) (i) Number of unique CDR3 clones, (ii) fraction and total clone counts by amino acid length, (iii) number of N-nucleotide additions, (iv) total nucleotide deletions of CDR3s recovered, and (v) total sequencing reads of Setd2Δ/Δ (KO) and controls for each VH family. proB cells were sorted from n = 4 for each genotype and subjected to NGS analysis. Significance measured by Wilcoxon rank-sum test.

RNA sequencing analysis of four independently sorted control and four Setd2Δ/Δ proB cell compartments was also conducted, and de novo assembly of CDR3 sequences was conducted using the TRUST algorithm (Li et al., 2017; Hu et al., 2018, 2019). (C) Unique CDR3 counts for heavy and light chain Ig loci from TRUST analysis. (D) Estimated B cell fraction of reads and B cell diversity recovered from analysis of RNA transcripts from TRUST analysis. Estimate B cell fraction was calculated by taking the fraction of number of reads mapped to BCR(IGV/IGJ/IGC) region to
treated with ATMi (Zha et al., 2011; Lescale et al., 2016) and suggests a functional redundancy between Setd2/H3K36me3 and Atm kinase activity in ensuring proper repair. To investigate potential effects on repair, we conducted Southern blotting, which revealed pMG-INV SJ and CJ formation without obvious accumulation of free unrepaired SEs and CEs in STI-treated WT and Setd2−/− v-Abl B cells (Figures 5D and S8B). Thus, like Xif, Mri, and Atm, Setd2 is not essential for C-NHEJ during V(D)J recombination (Bredemeyer et al., 2006; Zha et al., 2011; Hung et al., 2018). The non-essentiality of Setd2/H3K36me3 for end-joining was further intimated by the inability of dual loss of Setd2/H3K36me3 and p53 to generate translocations leading to the development of proB-cell lymphomas (Figures S9A and S9B), unlike loss of core C-NHEJ factors in a p53-null background (Difilippantonio et al., 2000; Gao et al., 2000; Frank et al., 2000).

Despite not being required for end-joining, ATMi-treated Setd2−/− v-Abl cells exhibited significantly increased mis-repaired recombination products that corresponded to either repaired SJs but unrepaired CEs (SJ + CE) (Figures 5A, 5D(i-ii), and S8B) or the formation of hybrid joins (Figures 5A, 5D(i), and S8B), products consistent with the observed loss of GFP signal. The enhanced formation of HJs with loss of Atm kinase activity and Setd2 was further corroborated by the increased detection of aberrant HJ products of the pMG-INV substrate in ATMi-treated Setd2−/− cells (Figure 5E). Although we could detect evidence of HJ formation by PCR of the recombination substrate in Setd2−/− lines without ATMi (Figure 5E), these products were below detection by Southern blotting (Figures 5D(i) and S8B), but consistent with the modest decrease in GFP signal with STI treatment alone (Figures 5B and S5). Furthermore, we could detect HJ formation not only from the endogenous k light chain locus of v-Abl Setd2 knockout cells but also in Setd2Δ/Δ splenocytes (Figure 5F). Together, these data indicate a novel role for Setd2/H3K36me3, especially in combination with Atm kinase activity, in the repair phase of V(D)J recombination to ensure proper joining.

**Loss of Setd2/H3K36me3 Reduces Overall B Cell Repertoire**

Even without loss of Atm kinase activity, however, there was abundant evidence of aberrant and abnormal rearrangement of the endogenous Igh locus in primary Setd2Δ/Δ proB cells (Figures 4F and S5E). We sequenced a similarly sized recombination product, present in both control and Setd2Δ/Δ proB cells, for three different heavy chain Variable gene families joined to the Jα4 fragment and found additional abnormalities (Figure 6A). Loss of Setd2/H3K36me3 not only reduced the overall number of unique productive Igh rearrangements (assessed by the number of hypervariable complementarity defining region-3 (CDR3)) but also resulted in shortening of CDR3 length and reductions in N-nucleotide additions (Figures 6A and 6B) (Bolotin et al., 2017). Similarly, when we looked at RNA-based transcripts of productive rearrangements of the Igh locus in proB cells using the hyper-variable region calling algorithm TRUST (Li et al., 2017; Hu et al., 2018, 2019), Setd2Δ/Δ proB cells exhibited significant decreases in detectable unique CDR3 sequences and an overall decrease in overall B cell repertoire diversity (Figures 6C and 6D). Furthermore, global variable and joining gene usage was reduced (Figure 6E) in proB cells from Setd2Δ/Δ mice and coincided with evidence of increased clonality in variable gene usage, which did not appear to be dependent on the proximity to the Diversity and Joining gene segment regions on the Igh locus (Figure 6F). These observations indicate that, in addition to its role with Atm during end-joining to ensure appropriate repair, Setd2/H3K36me3 is also involved in other mechanisms that prevent mis-processing and mis-repair of broken DNA ends to ensure the fidelity of V(D)J recombination and is critical in maintaining a normal immune repertoire.

**Setd2/H3K36me3 Prevents Post-mitotic Neuronal Apoptosis and Perinatal Lethality**

As Setd2/H3K36me3 seemed to be important for V(D)J recombination, we next wondered if it could play a role in Rag-independent C-NHEJ repair. To first test this, we assessed the sensitivity of asynchronous WT and Setd2−/− v-Abl cells to ionizing radiation and found that Setd2−/− v-Abl lines were more sensitive to DSBs induced by ionizing radiation (Figure 7A(i)). This increased sensitivity to ionizing radiation of
Figure 7. Post-mitotic Neuronal Apoptosis and Perinatal Lethality with Loss of Setd2/H3K36me3

(A) (i) Asynchronous and (ii) STI-571 G1-arrested WT, Xrcc5, and two different Setd2−/− v-Abl cells subjected to different doses of ionizing radiation and serially diluted five times in triplicate to assess survival. Values plotted as a mean of each dose as a percent of non-irradiated controls; bars represent standard deviation. Data are representative for three different independent treatment experiments. Viability was assessed at 72 h. No viable cells were detected beyond the 0.5 Gy dose for G1-arrested Xrcc5 v-Abl clones.

(B) Expected and observed genotype distributions of matings between Nestin-cre Setd2D+/+ mice to Setd2f/+ mice (n = 15 litters) and Nestin-cre Setd2D+/+ mice to Setd2f/f mice (n = 16 litters) 21 days post-partum (dpp).

(C) Embryo genotypes at different embryonic stages (n = 5 litters for each stage).

(D) Litter sizes of E14.5, E16.5, E18.5, and 0.25 dpp from Nestin-cre Setd2D+/+ mice to Setd2f/f breedings. Significance to 0.25 dpp was measured (n = 5 litters for each stage except 0.25 dpp, n = 11).
Setd2-deficient cells, at levels even greater than that of core NHEJ factors Xrcc5 (Ku80) and Lig4 (Figures 7A(i) and Figure S10A), could be in part due to previously described roles in HR-mediated DSB repair ascribed to Setd2/H3K36me3 (Li et al., 2013; Pfister et al., 2014; Aymard et al., 2014). To exclude the effects that may be driven by cells that may be cycling, we chose to study the sensitivity of G1-arrested v-Abl cells by inhibition with STI-571 or the Cdk4/6 inhibitor PD0332991 for 48 h before exposure to ionizing radiation. In both cell-cycle-arresting conditions, G1-arrested Setd2−/− v-Abl cells exhibited increased sensitivity to DSBs induced by ionizing radiation compared with WT controls (Figures 7A(ii), S10B, and S10C). Together, these data suggest that Setd2/H3K36me3 is important in the DSB repair in the G1 phase of cell cycle.

In addition to lymphogenesis, core C-NHEJ factors are critical for enforcing normal neurogenesis, where differentiating post-mitotic neurons migrating out of the ventricular zone do not have HR available for repair of DNA DSBs and instead rely on C-NHEJ (Frappart and McKinnon, 2008). Given our findings for a role for Setd2/H3K36me3 in the repair phase of V(D)J recombination, we tested its potential functions in neurogenesis. Consistent with the defective CNS development and embryonic lethality observed in knockout mouse models of core C-NHEJ factors (Gao et al., 1998; Frank et al., 2000; Gu et al., 2000), we observed post-mitotic neuronal apoptosis and perinatal lethality in mice with homozygous deletion of Setd2 in neural progenitor-restricted Nestin-cre-expressing mice, where deletion has been demonstrated to occur before post-mitotic neurogenesis (Liang et al., 2012). The severity of this phenotype is demonstrated by the complete absence of Nestin-cre Setd2−/−/Δ pups at 21 days post-partum (dpp) (Figures 7B and 7C) with perinatal lethality by 0.25 dpp (Figure 7D). Histopathological analysis of embryos revealed evidence of apoptosis of post-mitotic neurons as early as E14.5 (Figures S11A and S11B), as well as more widespread apoptosis in the developing brain in E18.5 embryos and in 2-h post-partum pups (Figures 7E, 7F, S11C, and S11D) suggesting a role for Setd2/H3K36me3 in embryonic neurogenesis that is reminiscent of deficiency for C-NHEJ factors.

**DISCUSSION**

The preservation of a genome is predicated on proper repair of DNA DSB and a balance between HR, which is a highly accurate and also slower form of repair that is restricted to the availability of a sister chromatid strand, and NHEJ, which is highly efficient but intrinsically error prone (Takata et al., 1998). Although there have been some studies highlighting the importance of chromatin accessibility, nucleosome positioning, DNA looping, and recognition of histone modifications (e.g., H3K4me3 by RAG2) (Shimazaki and Lieber, 2014; Johnson et al., 2010; Bettridge et al., 2017; Matthews et al., 2007; Matheson and Corcoran, 2012) in the regulation of the V(D)J recombination process and phosphorylation of γH2AX in repair (Yin et al., 2009; Celeste et al., 2003), little has been described in terms of the impact of other chromatin modifications, especially in the end-joining phase of the V(D)J recombination reaction.

Here, we discovered a strong dependency of normal hematopoiesis and, in particular, lymphogenesis on Setd2/H3K36me3 phenotypes consistent with previously published reports of three distinctly engineered knockout mouse models (Zhou et al., 2018; Zhang et al., 2018; Ji et al., 2019). Early loss of Setd2/H3K36me3 leads to severely impaired B and T cell development that could be partially rescued, in the case of B lymphogenesis, with the expression of a fully rearranged Igh locus, pointing to a role for Setd2/H3K36me3 in V(D)J recombination. This role of Setd2 in V(D)J recombination is further corroborated by the partial rescue of T lymphopoiesis with the over-expression of a rearranged TCR in Lck-cre Setd2Δ/Δ mice (Ji et al., 2019). Ji et al. also report decreased Rag1 occupancy across the Igh locus in B220+Cd19+ cells from Cd19cre Setd2Δ/Δ mice (Ji et al., 2019), including at variable region family genes. In sorted proB cells (B220+Cd43+IgM+) where we induced loss of Setd2/H3K36me3 early in hematopoiesis, however, we could not detect a significant effect on the recruitment of Rag1 to the critical regulatory region near where the Eμ enhancer resides and where Rag1 has previously been shown to bind in vivo at the proB cell stage of development (Ji et al., 2010). The observed Rag1 occupancy differences observed by Ji et al. (2019) in populations of lymphoid cells are likely impacted by their inclusion of
more mature B cell populations (B220⁺Cd19⁺ includes Cd43⁺ late proB and more mature Cd43⁻preB cells) that are depleted in Setd2⁺/Δ mice, as well as by incomplete and leaky deletion of Setd2 via the use cre-expressing mouse lines (Cd19cre) that delete after the proB cell developmental stage of interest (Rickert et al., 1997; Hobeka et al., 2006; Kraus et al., 2004; Siegemund et al., 2015).

We wanted to understand the nature of the defect in repair with early loss of Setd2 in proB cells, especially with respect to end-igation, as we could not detect significant differences in the initiation phase of V(D)J recombination and as we could detect recombination of the Igh locus, albeit abnormal recombination. We therefore turned to an inducible system to measure the repair defect and uncovered new roles for Setd2/H3K36me3 in the fidelity of the V(D)J recombination reaction, especially in combination with ATM kinase activity. We determined that Setd2/H3K36me3 is not absolutely required for end-igation by the lack of detectable unrepaired CEs and SEs of an ectopically integrated recombination substrate, pMG-INV. This non-essentiality for repair is consistent with our ability to detect recombination products in proB cells of Setd2⁺/Δ mice, albeit abnormal ones, and the lack of proB lymphomas as a consequence of translocation events in dual Setd2⁺/Δ p53⁺/Δ mice. It is noteworthy that unlike dual loss of ATM and XLF, loss of Setd2/H3K36me3 and Atm kinase activity did not result in complete abolishment of proper end-joining, but did lead to increased abnormal joining (e.g., SJ + CE joins and HJs), suggesting that Setd2/H3K36me3’s role in end-igation is functionally distinct from C-NHEJ XRCC4 paralogs XLF and PAXX (Lescale et al., 2016; Kumar et al., 2016). Although we could detect robust and efficient conversion of fully and properly recombined SJ + CJ products of the pMG-INV recombination substrate in induced Setd2-deficient v-Abl cells, we could also detect, with combined loss of Atm kinase activity, an aberrant SJ + CE recombination product. This raises the possibility of a CE-specific hairpin opening/repair defect, which would certainly require further study as CEs are thought to be efficiently processed and repaired at a much higher rate than SEs and CE-only defects have not previously been reported (Ramsden and Gellert, 1995; Schlissel et al., 1993; Roth, et al., 1992; Canela et al., 2016; Meek et al., 2016).

Paradoxically, whereas the end-igation defect of Setd2/H3K36me3 loss, as determined by the assessment of a recombination substrate in transformed v-Abl cells, may appear subtle, in mice, its loss severely arrests normal B/T lymphocyte development, similar to what is observed in mice with loss of C-NHEJ factors (Alt et al., 2013; Kumar et al., 2014) and in contrast to loss of factors involved in end-igation, such as Atm, Xlf, Paxx, or Mri individually (Bredemeyer et al., 2006; Kumar et al., 2016; Hung et al., 2018; Li et al., 2008; Lescale et al., 2016; Zha et al., 2011). This difference suggests that there are other determinants that contribute to the severe block in B (and T) lymphoid development in Setd2 knockout mice. For example, certain factors present at the Igh locus in proB cells may not be fully recapitulated in the assessment of rearrangement of an ectopic recombination substrate in transformed v-Abl cells, potentially affecting both the efficiency and accuracy of repair.

Likewise, it is also possible that there are other roles of ATM, or ATM targets, and SETD2 in end-joining that are not directly related and are mediated by distinct processes that, when combined, exacerbate mis-repair overall. Specifically, the loss of Setd2/H3K36me3 could influence the recruitment of factors associated with NHEJ in DNA DSB that recognize H3K36me3 (e.g., PHRF1 and PHF1) that could impact end-joining efficiency (Chang et al., 2015; Hong et al., 2008; Musselman et al., 2012), rather than playing a direct role in the stabilization of broken chromosomal ends, a function that has been ascribed to ATM (Bredemeyer et al., 2006). Additionally, we did not find any overt gene expression differences in alternative-NHEJ (A-NHEJ) factors and were not able to detect higher rates of deletions or translocations thought to accompany some forms of this A-NHEJ (Corneo et al., 2007) in either our CDR3 repertoire analysis, products of recombination of the recombination substrate in v-Abl cells, or in mice with dual loss of Setd2 and p53. These correlative lines of evidence however, do not fully preclude the possibility of rare forms of A-NHEJ contributing to the observed phenotype, warranting further investigation.

When we examined the abnormal recombination in proB cells of Setd2⁺/Δ mice more closely, we additionally noted significant loss of B cell repertoire and reduction of variable gene usage, characterized by less diversity in re-arranged, productive CDR3 sequences, as well as other abnormalities. Shortening of CDR3 length, without concomitant reduction of immune repertoire, has been previously connected to end-igation factor, XLF (Uspepet et al., 2016). In addition, shortening of CDR3 sequences of Ig heavy chain rearrangements has previously been observed in Pol X family knockouts (pol λ, μ), which with terminal deoxyribonucleotidyl transferase participates in nucleotide end-processing of heavy chain junctions during V(D)J
recombination (Bertocci et al., 2006). Thus, it is possible that Setd2/H3K36me3 may play additional roles in the end-processing of junctions, in addition to preventing the mis-repair in end-ligation we observed.

We also found that loss of Setd2/H3K36me3 increases sensitivity of cells to DSBs induced by ionizing radiation in both asynchronous and G1-arrested v-Abl cells, suggesting that Setd2/H3K36me3 may play a role in DSB repair activities in cells where HR would not be available, due to the lack of a sister chromatid. This result seemed to be supported by our discovery that neural-specific deletion of Setd2/H3K36me3 resulted in post-mitotic neuronal apoptosis. Although it is possible that other factors may be contributing to the neurogenesis defect, the temporal and spatial localization of apoptotic neuronal cells are highly suggestive of a role for Setd2/H3K36me3 in post-mitotic neurogenesis. The perinatal lethality we observed in Nestin-cre Setd2Δ/D mice is in stark contrast to Nestin-cre-driven conditional loss of core C-NHEJ factors, Xrcc4 or Lig4, where mice survive to adulthood (Frappart et al., 2009; Yan et al., 2006). Instead, it parallels the severity of the developmental arrest we observed in lymphogenesis, despite Setd2/H3K36me3 not being essential for C-NHEJ repair.

Previous studies have indicated a role for H3K36me3 in not only the recruitment of components on DNA damage repair in both MMR (MSH2/6) (Li et al., 2013) and HR (CtIP) (Pfister et al., 2014; Aymard et al., 2014) but also in a myriad of other cellular processes (Wagner and Carpenter, 2012). Although we have not exhaustively ruled out every potential role of Setd2/H3K36me3 in DSB repair, transcription, and splicing, we have attempted to study the role of Setd2/H3K36me3 in two systems that utilize NHEJ-mediated DSB repair by taking advantage of the non-redundancy of Setd2 in two independent mammalian developmental processes. Given the high frequency of SETD2 mutations in B and T cell lymphomas (Parker et al., 2016; Zhang et al., 2012; McKinney et al., 2017; Moffitt et al., 2017), primary human immunodeficiency (Ji et al., 2019), and neurological/developmental disorders (e.g., autism, intellectual disability, high-grade pediatric glioma) (Fontebasso et al., 2013; D’Gama et al., 2015; Tlemsani et al., 2016; Lelieveld et al., 2016), our findings demonstrating a role of Setd2/H3K36me3 in normal lymphogenesis and neurogenesis are especially noteworthy.

Limitations of Studies
Limitations of our studies include our inability to assess localization of critical NHEJ factors (e.g., Rag2, Ku70, Ku80, Xrcc4, Lig4, and others) due to the lack of readily available antibodies to perform ChIP of these proteins. In addition, although the use of v-Abl-transformed late-proB cell lines with an ectopically expressed recombination substrate can determine the essentiality of factors for end-joining in NHEJ, the use of such a system fails to recapitulate the factors or genomic structure present at the endogenous immunoglobulin loci, which may also influence the V(D)J recombination process during early lymphocyte development. Thus, it is possible that the severity of the block in normal lymphocyte development observed in vivo could be influenced by factors and mechanisms beyond the role of Setd2/H3K36me3 in end-joining during the repair phase of V(D)J recombination. Also, we were unable to assess if there was a non-enzymatic role for the Setd2 protein itself in any of the NHEJ processes in lymphocyte development, as expression of a full-length Setd2 (~2,500 amino acids, ~8-kb nucleotide coding sequence) in primary murine cells was unable to be achieved (by us and also not reported by other groups with genetically engineered Setd2 knockout mouse models, including but limited to Zhou et al., 2018; Zhang et al., 2018; Ji et al., 2019). We also did not fully and exhaustively investigate the role of A-NHEJ factors or the potential contribution of other roles that Setd2 and H3K36me3 play in normal cellular processes in both V(D)J recombination or post-mitotic neurogenesis settings.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

DATA AND CODE AVAILABILITY
RNA-seq, ATAC-seq, ChIP-seq, TRUST, and MIXCR data generated and analyzed during the current study have been deposited into the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO). The accession numbers for the RNA-seq, ATAC-seq, ChIP-seq, TRUST, and MIXCR data reported in this paper are GSE130904, GSE131588, GSE131608. These are unified under SuperSeries GSE131690.
SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.100941.

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AUTHOR CONTRIBUTIONS

S.H.C. and S.A.A designed experiments and wrote the paper. S.H.C., J.R.C., C.N.M., J.C.M., Y.X., and B.-R.C. performed experiments. J.Z. performed MiXCR and TRUST analyses, and R.P.K. performed bioinformatics analyses. V.K., E.C., P.J.H., Z.F., X.S.L., J.C., A.N., and B.P.S. provided experimental support and reagents for these studies as well as critical review of this paper.

DECLARATION OF INTERESTS

S.A.A. has a consultant and/or shareholder for Epizyme Inc, Imago Biosciences, Cyteir Therapeutics, C4 Therapeutics, Syros Pharmaceuticals, OxStem Oncology, Accent Therapeutics, and Mana Therapeutics. S.A.A. has been a consultant and/or shareholder for Epizyme Inc, Imago Biosciences, Cyteir Therapeutics, C4 Therapeutics, Syros Pharmaceuticals, OxStem Oncology, Accent Therapeutics, and Mana Therapeutics. S.A.A. has received research support from Janssen, Novartis, and AstraZeneca. S.H.C. is currently an employee at Beam Therapeutics. The authors have no additional financial interests.

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Supplemental Information

Loss of H3K36 Methyltransferase

SETD2 Impairs V(D)J Recombination
during Lymphoid Development

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Figure S1. Loss of Setd2/H3K36me3 disrupts normal hematopoiesis, Related to Figure 1
(A) CBC and differentials from Setd2Δ/Δ, Setd2Δ/+ and controls (Setd2f/f) (n=3 for all groups) *, p<0.05 **, p<0.01 ***, p<0.001 (B) (i) total cellularity of spleen, thymus, whole bone marrow and lineage negative BM, total cellularity of cells positive for differentiated markers B220, Cd3, Mac1, Gr1, or Ter119 in (ii) bone marrow or (iii) spleen and (iv) spleen and thymus weights for heterozygous Setd2 mice and controls (n= 3 for all groups). All values were non-significant. (C) Representative flow cytometry plot of HSC fractions, multipotent progenitors (MPP1-4) and percent composition of lineage negative cells of HSC stem cell and progenitor populations (n= 10 for all groups). (D) Representative flow cytometry plots for early myeloid and erythroid progenitor populations in bone marrow. MkP: KithSca1-Cd41+Cd150+, GMP: KithSca1-Cd41+Cd150-FcyRII/III+, pre-GM: KithSca1-Cd41+FcyRII/III+Cd150*Endoglin+, pre-CFU-E: KithSca1-Cd41+FcyRII/III+Cd150*Endoglin+ and proportion of HSC and erythroid progenitor populations of control and Setd2Δ/Δ (n = 7 for all groups). (E) Representative flow cytometry of erythroid progenitor populations. proE, 1-Basophilic erythroblasts (Baso-E), 2-late Baso-E and chromatophilic erythroblasts (Poly-E), and 3-orthochromatophilic erythroblasts (Ortho-E) in control and Setd2Δ/Δ bone marrow. (F) Percent composition of different erythroid progenitor populations in bone marrow and spleen of control and Setd2Δ/Δ mice (n = 3). *, p<0.05 **, p<0.01 ***, p<0.001, error bars represent SD.
Figure S2. Setd2Δ/Δ HSCs are defective in competitive and non-competitive bone marrow (BM) reconstitution assays, Related to Figure 1

(A) (i) Peripheral blood engraftment, as measured by Cd45.2+ of competitive reconstitution of lethally irradiated (9cGy) recipient mice with 1x10^6 WBM Mx1 Setd2Δ/Δ and control bone marrow with 1x10^6 WBM from syngeneic Cd45.1+ mice after induction of Mx1-cre with polyI:polyC (plpC). (ii) Total BM cellularity of mice 16 weeks after induction with plpC (iii) percent Cd45.2 involvement in BM, spleen and peripheral blood, (iv) total cell counts of Cd45.2+ donor HSC stem and progenitor populations. (B) Non-competitive reconstitution assay as conducted in (A) without 1x10^6 Cd45.1+ wildtype competitor WBM cells. Arrows indicate when plpC treatment was initiated. (ii-iv) graphs were generated from mice 28 days post plpC induction as Setd2Δ/Δ mice were moribund due to complete BM failure. (C) Competitive and non-competitive reconstitution of lethally irradiated (9cGy) recipient mice with 1x10^6 WBM Vav-cre Setd2Δ/Δ and control BM with 1x10^6 WBM from syngeneic Cd45.1+ mice. Overall peripheral blood Cd45.2+ chimerism in (i) competitive and (ii) non-competitive reconstitution assays. (iii) total cellularity of Cd45.2+ whole and lineage negative BM (iv) Cd45.2+ chimerism (v) percentage and total numbers of Cd45.2+ donor derived HSC of competitive and non-competitive transplants 16 weeks post-transplant (D) Competitive and non-competitive reconstitution of lethally irradiated recipient mice with 1x10^6 WBM Vav-cre Setd2Δ/Δ and Vav-cre Setd2Δ/Δ control BM with 1x10^6 WBM from syngeneic Cd45.1+ mice. (i) Overall peripheral blood Cd45.2+ chimerism and (ii) BM Cd45.2+ chimerism in competitive and non-competitive reconstitution assays 16 weeks post-transplant. All data is representative of at least 3 independent transplant experiments. n = 5 for each transplant group. *, p<0.05 **, p<0.01 ***, p<0.001, error bars represent SD.
Figure S3. Loss of Setd2/H3K36me3 arrests development at a proB cell stage, Related to Figure 3.  
(A) Representative flow cytometry plots with gating strategy for different B cell progenitor populations for controls and Setd2Δ/Δ B220+ BM cells. Fraction A (FrA): B220+IgM-Cd43+Cd19-Cd24loCd25; pre-proB: B220+IgM+Cd43+Cd19+Cd24loCd25; preB: B220+IgM+Cd43+Cd19+Cd24hiCd25; FrC: B220+IgM-Cd43+Cd19+Cd24loCd25; proB: B220+IgM+Cd43; immature B: B220+Cd43+Cd19+. (B) Percent composition of B220+ WBM of different B cell population in the bone marrow (n = 6). (C) B cell progenitor proportions in B220+ BM of Setd2Δ/Δ and controls (n= 14 for all groups). (D) Total cellularity of B cell progenitor proportions in B220+Cd43+IgM BM of Setd2Δ/Δ and controls. n= 9 for all groups. (E) Ratio of preB to proB compartment and DN4 to DN3 compartment total cellularity from BM of Setd2Δ/Δ and controls (n= 8 for all groups, from Fig. 3A-B) (F) Quantification of phospho-γH2ax foci by immunofluorescence from sorted FrA proB cells from Setd2Δ/Δ (n=8 mice) and controls (n=7 mice). (i) Fraction of total cells containing foci (n=366 for Setd2Δ/Δ, n= 337 for controls). (ii) Foci/cell in cells containing phospho-γH2ax foci (n=260 for Setd2Δ/Δ, n= 191 for controls). (iii) Frequency of foci/cell for all cells containing phospho-γH2ax foci for Setd2Δ/Δ and controls. (G) percent of different B cell populations positive for phospho-γH2ax by flow cytometry (n=15 for Setd2Δ/Δ, n=13 for controls). All groups were significant (p<0.01) except for preB population. (H) Representative flow cytometry plot of cell cycle status indicated by co-staining with DAPI and BrdU incorporation of sorted FrA proB cells from Setd2Δ/Δ (n=3) and controls (n= 2) and summary graph of each cell cycle stage. All values for cell cycle status were non-significant. ** p<0.01 *** p<0.001, error bars represent SD.
**Figure S4.** Loss of Setd2/H3K36me3 does not alter expression of C-NHEJ and A-NHEJ proteins, Related to Figure 3.

(A) Volcano plot of fold expression of a FrA proB cell compartment sorted from n=3 independent Setd2Δ/Δ and littermate control mice and subjected to RNA-Seq. Genes highlighted include: Rag1, Rag2, Xrcc4, Lig4, Xrcc5, Xrcc6, Xrcc7, Prkdc, Hmgb2, Mre11a, Dclre1c, Atm, Parp1, Polq, Brca2, Ctbp1, Rad52, Mrnip. (B) Real-time PCR of C-NHEJ and A-NHEJ genes of FrA proB cells sorted from n=2 Setd2Δ/Δ mice and a littermate control. Expression was normalized to Gapdh expression and calculated relative to the wildtype control by a ΔΔCt method. (C) (i-v) Immunoblotting of FrA proB cells sorted from n=3 Setd2Δ/Δ mice and n=3 littermate controls (Setd2ff) for C-NHEJ proteins Ku70, Ku80, Xrcc4, Rag1, Rag2, Hmgb2 with loading controls Vinculin and Gapdh. Mb – low molecular weight ladder, Ms – high molecular weight ladder.
Figure S5. Loss of H3K36me3 does alter local chromatin architecture or accessibility of the Igh locus but leads to aberrant V(D)J recombination, Related to Figure 4.

(A) Composite Heatmap and volcano plot of H3K36me3 ChIP-Seq signal of Cd19⁻ and Cd19⁺ control (WT) and Setd2Δ/Δ (KO) proB cells (B220⁺Cd43⁺Cd19-IgM⁻) (n=3 for each genotype). (B) ATAC-sequencing tracks from Cd19⁻ and Cd19⁺ control (WT) and Setd2Δ/Δ (KO) proB cells (B220⁺Cd43⁺Cd19-IgM⁻) (n=2 for each genotype) across the Igh locus. (C) Representative ChIP-PCR of relative abundance of H3K36me1-3 and H3K27me1-3 at regulatory region of Igh locus. Regions R1-R5 were regions with H3K36me signal in between the Eμ enhancer binding site and Cμ gene region with HoxA9, Cy3 and actin as controls. Data is representative of n=2 independent experiments. (D) (i-iii) H3K36me3, H3K4me3, and H3K9ac ChIP-PCR of variable heavy chain gene families and from n=4 independent ChIP experiments and (iv) Rag1 occupancy by ChIP-PCR variable gene family genes with primers from (Ji et al 2019 and (Ji et al., 2010; Hauser et al., 2014; Subrahmanyam et al., 2012; Chakraborty et al., 2009; Hesslein, et al, 2003). (n=1 ChIP). (E) (i-iii) replicates of PCR assay to detect V(D)J recombination products of rearrangement shown in Figure 2E of the Igh locus of different V₅₉ families from independently sorted proB cells from Setd2Δ/Δ (KO) and controls (WT). No product meant germline non-rearrangement.
Figure S6. Generation of v-Abl transformed Setd2, Xrcc5 and Lig4 deficient cell lines, Related to Figure 5.

(A) H3K36me3 levels of several individual v-Abl clones (used in Figure 5) that were either Setd2 wildtype (wt, indicated with asterisk) or Setd2 deficient lines (Setd2−/− or KO). Total H3 and Gapdh were used as loading controls.

(B) (i-ii) Immunoblots of Setd2 wildtype or Setd2−/− v-Abl lines for Kmt3a (Setd2) and Vinculin loading controls.

(C) Immunoblot of v-Abl lines for Xrcc5 (Ku80) null lines. Gapdh was used as a loading control.

(D) Representative flow cytometric analysis of GFP expression in Lig4 and Xrcc5−/− pMG-INV v-Abl cells treated with Abl kinase inhibitor imatinib (STI-571) and ATM kinase inhibitor (ATMi, KU55933) for 48 and 96 hours. Representative of 4 independent induction experiments.
Figure S7. Deletion of Setd2 in v-Abl cells does not alter expression of C-NHEJ and A-NHEJ proteins or cell cycle status, Related to Figure 5.

(A) (i-iii) Immunoblotting of WT and Setd2−/− v-Abl clones for C-NHEJ proteins Ku70, Ku80, Xrcc4, Rag1, Rag2, Hmgb2 with loading controls Vinculin and Gapdh. Mb – low molecular weight ladder, Ms – high molecular weight ladder. (B) Real-time PCR of C-NHEJ and A-NHEJ genes of Setd2 deficient and WT v-Abl transformed cells. Expression was normalized to Gapdh expression and calculated relative to the wildtype control by a ΔΔCt method. (C) Representative flow cytometry plot of cell cycle status indicated by co-staining with DAPI and BrdU incorporation of Setd2 deficient and WT v-Abl cells. All data representative of n=2 independent experiments. Error bars represent SD.
Figure S8. Dual loss of Setd2/H3K36me3 and Atm activity leads to abnormal recombination and end-joining repair, Related to Figure 5.

(A) (i-iii) Immunoblotting for NHEJ proteins and phospho- and total γH2ax of WT (Setd2+/+) and Setd2 deficient (Setd2−/−) v-Abl cells treated for 48 hours with STI or STI+ATMi. Data representative of 3 independent induction experiments. Mb – low molecular weight ladder, Ms – high molecular weight ladder. (B) Replicate Southern blot analysis of genomic DNA from Setd2−/− clone (D10) and control line that were digested with NheI (left) and XbaI (middle and right panels) and hybridized with Thy1 probe (middle, right panels) or GFP probe (left panel). Bands indicated are for URs, SJs, CJs, SEs, CJ+SEs or SJ+CEs. Hybrid Joins (HJ) indicated as well. Southern blots were generated from an independent induction experiment of v-Abl clones with imatinib and ATMi shown in Figure 5.
Figure S9. Dual loss of Setd2/H3K36me3 and p53 does not lead to the development of pro-B lymphomas, Related to Figure 5. (A) Kaplan-Meier survival of double knockout Vav (n= 1), Mx1 (n=7), Cd2 (n=4), Mb1cre (n=9) Setd2Δ/Δ p53Δ/Δ mice and controls (n = 11). All mice developed thymic lymphomas. (B) Weight of thymus in mouse thymic lymphomas from double knockout mice and controls. ***, p<0.001 calculated to control (Setd2ff p53+/-). Error bars represent SD.
Figure S10. Asynchronous and synchronous Setd2 deficient v-Abl cells exhibit increased sensitivity to ionizing radiation, Related to Figure 7.

(A) Asynchronous and STI G1-arrested WT, Lig4−/− and 2 different Setd2−/− v-Abl cells subjected to different doses of ionizing radiation and serially diluted 5 times in triplicate to assess survival. Values plotted as a mean of each dose as a percent of non-irradiated controls, bars represent standard deviation. Data is representative for 3 different independent treatment experiments. Viability was assessed at 72 hours. 

(B) Representative flow cytometric analysis of cell cycle status as determined by DNA content with DAPI staining 48 hours after treatment with 1mM Cdk4/6 inhibitor PD0332991. Data representative of n=3 independent experiments.

(C) PD0332991 G1-arrested WT Setd2−/− v-Abl cells subjected to different doses of ionizing radiation and serially diluted 5 times in triplicate to assess survival. Values plotted as a mean of each dose as a percent of non-irradiated controls, bars represent standard deviation. Data is representative for 3 different independent treatment experiments. Viability was assessed at 72 hours.
Figure S11. Post-mitotic neuronal apoptosis with loss of Setd2/H3K36me3, Related to Figure 7.  
(A,B) TUNEL assay of two independently matched control and Nestin-cre Setd2Δ/Δ E14.5 coronal sections of the lateral ventricle displaying the ganglionic eminence (GE), Lateral Ventricle (V), Ventricular zone and intermediate zone (IZ) of the telencephalon. Distances as indicated in microns. (C) Higher magnification of cleaved Caspase-3 immunohistochemistry of E18.5 sagittal regions of lateral ventricle of control and Nestin-cre Setd2Δ/Δ embryos from Figure 7D. Distances as indicated in microns. (D) Cleaved caspase-3 immunohistochemistry of coronal sections of the lateral ventricle of 2 hour post-partum control and Nestin-cre Setd2Δ/Δ pups from same samples as shown in Figure 7E. Magnifications indicated by corresponding red and green boxes. Arrows indicate cleaved caspase-3 staining and pyknotic nuclei. Distances as indicated in microns.
Transparent Methods
Mice and Isolation of primary B cells and v-Abl B cells

Strategy for generating Setd2Δ/Δ mice was previously described (Mar et al., 2018). Setd2ff mice were bred to Mx1, Vav1, Cd19, Mb1, hCD2-cre mice to homozygosity. At 6-8 weeks of age, mice were sacrificed and bone marrow from femurs, tibias, hips, and spines were isolated. BM was then RBC lysed and stained with B220-biotin (BD Pharmingen) and subsequently stained with Anti-biotin microbeads (Miltenyi) and applied on a LS column (Miltenyi). Subsequent staining for cellular antigens was conducted. For V(D)J recombination product assessment, RNA-Seq, ChIP-Seq, ChIP-PCR, and qPCR were stained and sorted by FACS (BD-Aria) for B220+IgM+IgD-CD25-CD19-proB cells and B220+IgM-CD25-CD19-preB. IgHeIMD4 (Goodnow et al., 1988) and Mb1cre mice were generously provided by Jayanta Chaudhuri. All mouse experiments were approved by the Institutional Animal Care and Use Committees at Dana-Farber Cancer Institute and Memorial Sloan Kettering Cancer Center.

WT and Lig4−/− v-Abl B cell lines were generously provided by Andre Nussenzweig and Barry Sleckman (Bredemeyer et al., 2006). Xrcc5 and Setd2 knockout v-Abl cells were generated as before (Bredemeyer et al., 2006; Jacobsen et al., 2006). Briefly, CMV expression vectors containing Cas9 and a BFP tagged expression vector (Hung et al., 2018) with sgRNAs targeting gene of interest were nucleofected with Amaxa nucleofection Kit P4 as per manufacturer’s instructions (Lonza). Less than 24 hours later, BFP+ cells were single cell sorted into 96 well plates and expanded and western blotting was conducted to confirm protein knockout. Single cell clones with confirmed knockout of protein were transduced with retrovirus containing pMG-INV (Hung et al., 2018) vector and subsequently sorted for expression of cell surface marker Thy1.2. The recombination substrate vector pMG-INV was generously provided by Barry Sleckman. To induce G1 arrest, cells were treated with 3 μM imatinib (STI571) with or without 15 μM ATMi (KU55933) for up to 96 hours and assayed for GFP expression by flow cytometry and genomic DNA isolation for downstream PCR analysis. sgRNAs for CRISPR/Cas9 used were as follows: Xrcc5: sgXrcc5-1 (5'-GAATGATATCAGTTCCGTAG-3'), sgXrcc5-2 (5'-GAGCTTGGTAAGAAAAACG-3'), sgXrcc5-3 (5'-GTCTAAGCTATCGGACGA-3'), sgXrcc5-4 (5'-TGTCCTTGAAGCCGAGAC-3'), sgXrcc5-5 (5'-TGACACATCATCGGGGTCC-3'). Setd2: sgSetd2-1 (5'-GCAATTGCCCTTAATATCCCG-3'), sgSetd2-2 (5'-GGAGTTCCCTTACGAGGCTC-3'), sgSetd2-3 (5'-TTCGAGCATCTGACGACG-3'), sgSetd2-4 (5'-ATAATAGGGAGCCGAGACG-3').

RNA-sequencing of LSK and proB cells

B cells were isolated as indicated above. LSKs were obtained by lineage depletion of WBM and conducted as per manufacturer’s protocols with biotin-labeled antibodies for CD3, Gr1, Ter119, and B220 (BD Pharmingen) and subsequently subjected to magnetic depletion with anti-biotin microbeads and depletion on a LD column (Miltenyi). Lineage depleted cells were then stained with cKit and Sca1 and sorted on a FACS cell sorter (BD-BR-Pari). Qiagen RNA kits were used as per manufacturer’s protocol for RNA isolation and purity was confirmed with RNA Tapestation (Illumina). RNA-seq libraries
were prepared with NEBNext UltraKits and for proB cells. For LSKs, RNA was amplified with SMARTer Ultra Low Input RNA Kit for Illumina Sequencing.

**Hematopoietic Reconstitution assays**
For competitive/non-competitive reconstitution assays, 1x10⁶ total, unfractionated whole BM cells of control and fully excised *Setd2* mice were injected into lethally irradiated mice (9 cGy) and bled every 4 weeks up to 16 weeks before these mice were sacrificed and hematopoietic reconstitution was assessed in the bone marrow. Non-excised *Mx1cre Setd2Δ/Δ* mice were treated with pl:pC after 8 weeks of engraftment in reconstitution assays.

**BrdU incorporation assays**
For *in vivo* assessment of BrdU incorporation, BrdU was injected intraperitoneally 4 hours before mice sacrificed and BrdU incorporation was assessed as per manufacturer’s protocols (BD Biosciences). Recommended manufacturer’s protocols were also followed for *in vitro* labelling of v-Abl cells. Cells were labelled for 2 hours before fixed and permeabilized.

**Phospho-γH2ax Immunofluorescence**
Cytospins of 20 000 sorted FrA proB cells were prepared at 500 rpm for 5 minutes, permeabilized and fixed in 4% paraformaldehyde/PBS at 4°C for 10 minutes. Phospho-γH2AX (Abcam) primary antibody was used at 1:500 and incubated at 4°C overnight. Secondary antibody Anti-rabbit Alexa647 (CST) was applied for 30 minutes at room temperature. DAPI (1 µg/ml) counterstaining was conducted for 5 minutes at room temperature and covered with Prolong Gold Antifade Reagent (ThermoFisher). Slides were analyzed by confocal microscopy (Leica TCS SP5) (Leica) and foci were quantified using Image J software (NIH).

**Ionizing Radiation Sensitivity Assay**
Ionizing radiation sensitivity assays were performed for v-Abl lines that were either asynchronous or G1 arrested with STI-571 (Selleckchem) or PD-0332991 (Sigma-Aldrich) before irradiation. Cells were treated with 0.1, 0.5, 1, or 2.5 Gy ionizing radiation with percent survival measured relative to a non-irradiated and non-treated control for each cell line assayed. Cells that were G1 arrested were treated 48 hours prior to irradiation before washing off medium containing 3µM STI571 or 1µM PD-0332991 and re-plated with fresh media. 50,000 cells were initially plated in 96 non-tissue culture treated plates and 1:1 serially diluted 5 times with 3 replicates for each condition and cultured at for 3 days. Viability was measured by staining with DAPI and measured by flow cytometry on BD Fortessa in HTS mode.

**Genomic DNA isolation**
Up to 5x10⁶ proB cells were harvested and genomic DNA extracted using PureLink Genomic DNA kit (Invitrogen) as per manufacturer’s protocols.

**PCR analyses**
PCR of retroviral substrate coding joints (CJs) and hybrid joints (HJs): pMG-INV was generously provided by Barry Sleckman (Hung et al., 2018). Oligonucleotides CJ_F and CJ_HJ_R were used to amplify CJs in pMG-INV. Oligonucleotides HJ_F and CJ_HJ_R were used to amplify HJs. CJ_F: (5’-TCAGCCAGAAATTCACTGGCA-3’); HJ_F: (5’-TTGTACACCATACGCTCCG-3’); CJ_HJ_R: (5’-GCTTATCGATACCCTGACCT-3’). All PCRs were done on genomic DNA from cells that had been treated with STI571 for 96 hours. The Il2 gene was amplified using the IMR42 and IMR43 oligonucleotides (Bredemeyer et al., 2006). PCRs for the Il2 gene, and all retroviral HJs and CJs, were carried out in 50μL with cycling conditions of 95°C for 2 minutes followed by 30 cycles 94°C 30s, 55°C 30s, 72°C 60s (Bredemeyer et al., 2006). Murine Il-2: forward (5’-CTAGGCCACAAGGAGATCTGG-3’); reverse (5’-GTAGGTGAAATTCACTGGATGC-3’).

PCR analyses of endogenous receptor gene rearrangements: PCR of Vκ6-23 HJs and CJs was carried out by amplifying 0.5μg of genomic DNA from B220 enriched splenocytes from Setd2Δ/Δ and Setd2f/f controls in 50μl with primers for amplification as follows: pkJa and pk6a for HJ and pkJa and pk6d for CJ. PCR conditions were 95°C for 5 minutes followed by 17 cycles 94°C 30s, 64°C 30s, 72°C 30s. Products from this reaction were amplified in 50μl using the above conditions with primers pkJa and pk6b for HJ and pkJa and pk6c for CJ and 25 amplification cycles oligonucleotides (Bredemeyer et al., 2006). pk6a: (5’-TGATGTCAGAGGGCACAACTG-3’); pk6d: (5’-GAAATACATCAGACCAGCATGG-3’); pk6b: (5’-CTACCAAAACTTTGCAACACAGGC-3’); pk6c: (5’-ACATGGTCTGGTTGCTGGTGTTG-3’); pkJa: (5’-GGAGATGGCAGAAATTCACTGG-3’).

PCR results were analyzed with high sensitivity D1000 TapeStation reagents (Agilent Technologies).

V(D)J recombination PCR: Assay was conducted as previously described (ten Boekel et al., 1995; Ehlich et al., 1994; Corcoran et al., 1998). Briefly, two rounds of PCR were conducted on 100-300ng of gDNA using forward primers amplifying VH558, VHQ52, VH7183 family genes with nested primers located in the JH4 gene segment. Products were visualized on a 1% agarose gel and the V-(D)JH4 recombination product was gel extracted (Qiagen) and submitted for NGS sequencing. Vh7183_F1: (5’-CTCGCCACTGGACCTTCGGGTCAGTTGG-3’); Vh7183_F2: (5’-CAGCTGGTGAGTCTGGGGAGGC-3’); Vh558_F1: (5’-ACCATGGGATGGAGCTGKATCWTBC-3’); Vh558_F2: (5’-GTGARCCCTGGRCCCTCAGTGAAG-3’); VhQ52F: (5’-GCGAAGCTTGTGGAGTCTGGGGGAGCTTA-3’); DhQ52_F1: (5’-CACAGAGAATTCTCCATAGTTGATAGCTCAG-3’); DhQ52l_F2: (5’-GCCTCAGAATTCCTGTGGTGCTGACTGTTG-3’); Jh4_R1: (5’-AGGCTCTGAGATCCCTCAGACAG-3’); Jh4_R2: (5’-GGGTCTAGACTCTCAGCAGCCTGCTCCCTCAGG-3’); actin_738: forward (5’-GGTGTCATGAGTATGGGT-3’), reverse (5’-CGCAACAAATCTCACGTTCCAT-3’).
Chromatin Immunoprecipitation (ChIP) and ChIP-sequencing

Chromatin immunoprecipitation was coupled with high-throughput sequencing (ChIP-seq). 0.5-2x10^6 primary sorted proB cells were crosslinked with 1% formaldehyde for 10 minutes followed by 0.125 M glycine for 5 minutes. Fixed cells were washed twice with ice-cold phosphate-buffered saline and resuspended in ChIP lysis buffer and sheared using a Covaris E220 ultrasonicator (Covaris). Sheared chromatin was incubated overnight at 4°C with rabbit polyclonal anti-H3K36me3 (61101 pAb, Active Motif). Immune complexes were collected with protein A/G dynabeads (Invitrogen) and washed sequentially in low-salt wash buffer (20 mM Tris pH 8.0, 150 mM NaCl, 0.1% SDS, 1% Triton X-100, 2mM EDTA), high-salt wash buffer (20 mM Tris pH 8.0, 500 mM NaCl, 0.1% SDS, 1% Triton X-100, 2mM EDTA), LiCl wash buffer (10 mM Tris pH 8.0, 250 mM LiCl, 1% NP-40, 1% sodium deoxycholate, 1 mM EDTA), and TE. Chromatin was eluted buffer (1% SDS, 0.1 M NaHCO3), and then reverse cross-linked with 0.2 M NaCl at 65°C overnight. DNA was purified with a PCR purification kit (QIAGEN) and subjected to quantitative PCR or processed for ChIP-sequencing.

ChIP-PCR oligonucleotides used in this study were as follows (Ji et al., 2010; Hauser et al., 2014; Subrahmanyam et al., 2012; Chakraborty et al., 2009; Hesslein, et al, 2003):

- Vha: forward (5'-CCTTCGCCCCAATCCACC-3'), reverse (5'-CAAGTAACCCTCAAGAGAATGGAGACTC-3');
- Vh47: forward (5'-CAATCCTCACGACACGCTTA-3'), reverse (5'-TCAGGCTGTGATTACAACACTGTGT-3');
- Vh77: forward (5'-AAATCCTCCGACACAGCTTA-3'), reverse (5'-TAGACCGCAGAGTCCTCAGA-3');
- Dsp2: forward (5'-CAACAAAAACCCAGTATGCCAG-3'), reverse (5'-GTGCTTTCACCTGTCTGTGGG-3');
- Dfl-4.5: forward (5'-AGGCATCTCATCTCAGTCAGC-3'), reverse (5'-CTACAAACCCTCAAGAGC-3');
- Jh1: forward (5'-TGCTACTGGTACTTCGATGTCTG-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Jh2: forward (5'-CAGTCTCCTCACGGTAGTCT-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Jh3: forward (5'-GCCTGGTTGCTTACTGG-3'), reverse (5'-GACAAAGGTTGAATCT-3');
- Jh4: forward (5'-CAGTCTCCTCACGGTAGTCT-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Jh5: forward (5'-GCTAAACTGAGGTGATTACTCTT-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Jh6: forward (5'-CAATCTACAGAGGTTGAGTTG-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Jh7: forward (5'-GCCACCTCCAGGGAGACTC-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Jh8: forward (5'-GCCACCTCCAGGGAGACTC-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Sµ: forward (5'-GCCACCTCCAGGGAGACTC-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Cµ: forward (5'-GCCACCTCCAGGGAGACTC-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- Sγ: forward (5'-GCCACCTCCAGGGAGACTC-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- γ: forward (5'-GCCACCTCCAGGGAGACTC-3'), reverse (5'-GCCACCTCCAGGGAGACTC-3');
- IgHK36me3R1: forward (5'-TGAGCCCTCAGTCTGAGT-3'), reverse (5'-TGAGCCCTCAGTCTGAGT-3');
- IgHK36me3R2: forward (5'-TGAGCCCTCAGTCTGAGT-3'), reverse (5'-TGAGCCCTCAGTCTGAGT-3');
The following variable region ChIP-PCR primers were used for variable region were from (Ji et al 2019): V1-1: forward (5’-ACGTCACTGAGGTGTGAGCA-3’), reverse (5’-CTAGGCACATACGCTTATGC-3’); V1-7: forward (5’-TCATCAAGCTAGTTAGGTCTC-3’), reverse (5’-AGACAGTGTTGCAACCACAT-3’); V1-59: forward (5’-CATACTACACACCATCTTGGCT-3’), reverse (5’-AACCCTGGAGGAGTAGCAAACT-3’); V6-1: forward (5’-CCTCTACACAGCCATGGGTATGCT-3’), reverse (5’-GCAACATGCTCATGGGTATGCT-3’); D1-1: forward (5’-CTAGACTCAGTTTTGAGCTCAGCT-3’), reverse (5’-GAGGGTAGAGGACTCACCTGTA-3’); D2-8: forward (5’-CTGTGTAGTTACATAGTGGC-3’), reverse (5’-CTGACTGGCTAAACACTGTAG-3’); D5-4: (5’-CTGACTGGCTAAACACTGTAG-3’), reverse (5’-CACAAGGAGTGGATCTCTGATGT-3’); J2: forward (5’-GAGGTAGAGCTCAGCTCAGCT-3’), reverse: (5’-ACTTTAGGCTCAGCTCAGCT-3’); J3: forward (5’-CTGATAGCTGAGAGCTCAGCT-3’), reverse (5’-TGGTGGGACGAACATTTACA-3’).

For ChIP-Sequencing libraries were prepared using a ThruPLEX DNA-seq Kit (Rubicon Genomics) and validated using a TapeStation (Agilent Technologies) and Qubit 2.0 Flurometer (Thermo Fisher Scientific). Libraries were pooled and sequenced on a HiSeq2000 platform (Illumina).

**Quantitative PCR**

RNA isolated from sorted B cell populations and v-Abl cells were subjected to quantitative PCR with the following primers and normalized to Gapdh expression:

From (Chakraborty et al., 2009): Dfl-4.5: forward (5’-AGGCATCTCATCTCCTAAGC-3’), reverse (5’-CTGTGCCTCTTCTAAGC-3’); Dq52: forward (5’-TGGTGGCAGTTTTCTGACTAAGC-3’), reverse (5’-CTCTGGCCCTCACCAGAACAT-3’); Dsp2: forward (5’-TGTTACCTTACTTGGCAGG-3’), reverse (5’-TGGGTTTTGTGTGGTATATAC-3’); g-actin: forward (5’-GTGTGTCGGTGCTGATATGAT-3’), reverse (5’-GGTGTGGGAAGTTGCTGCTGAT-3’); Cμ: forward (5’-AGAGATCTCGCATGCTGATAC-3’), reverse (5’-TGTTCCGGAGAACACATTTACA-3’); Eμ (5’): forward (5’-CTGACATTACTAAAGGATACCTGAGCGG-3’), reverse (5’-CTCCAATCTACCTGGATCATT-3’); Eμ(3’): forward (5’-ATTCAGCCGAAAACTGGAGGTC-3’), reverse (5’-GGGGAAACTGAGAACTCTTGA-3’); From Zan et al 2017: Aicda: forward (5’-AGAAAGTCACGCTGGAGACC-3’), reverse (5’-CTCCTCCTCACCACGTGAC-3’); Rad52: forward (5’-
AGCCAGTATACAGCGGATGAA-3'), reverse (5'-GCCATGCGGCTGCTAATGTA-3'); PolI: forward (5'-
TGGCTATATGGGCAGCACCT-3'), reverse (5'-GCCATGCGGCTGCTAATGTA-3'); Ku70: forward
(5'-CACCAAGCGGTCTCTGACTT-3'), reverse (5'-AGAGAGGGCGCTAGGTAGTG-3'),
Ku80: forward (5'-AGGCCCAAGAAGCTCTATCA-3'), reverse (5'-
GCACTCTTGAGATCCCCACA-3'); Gapdh: forward (5'-TTACACCATTAGGAGAAGGC-3'),
reverse (5'-GGCATGGACTGTGGTCATGA-3')
Gapdh: forward (5'-CAAGCAGATGATGTTTCCTGC-3'), reverse (5'-
AGAACTAAGGGTGTTGGTAGC-3') (Wu et al., 2003).
Lig4: forward (5'-TCTGCCTTAAAGCGAAG mating-3'), reverse (5'-
GTTGAGAGGGCTCAGGTAGTG-3'); Xrcc4: forward (5'-
CACCAAGCGGTCTCTGACTT-3'), reverse (5'-TCATCGGTGCTTCCATCATA-3') (Okamura et al., 2016); Lig3: forward
(5'-CCTCTCCAAGCTCACCAAAG-3'), reverse (5'-TGCTCATTGTG-3');
Parp1: forward (5'-GCACTCTTGGATTCCCCACA-3'), reverse (5'-
CCTCTCCAAGCTCACCAAAG-3') (Meador et al, 2008); Rag1: forward (5'-
TGAGTAGAGCCTTCCTGTGG-3'); Rag2: forward (5'-CCTCTCCAAGCTCACCAAAG-3'), reverse (5'-
GCTTCAGGAAGACAGGGAGAG-3') (Bender et al. 2004); Rag2: forward (5'-
CAACATCCCTCAGCCTTCTCG-3'), reverse (5'-
ACATCTGCCTTCACGTCGAT-3');
Rad51: (5'-CTCATCGCTACAGGGAGAG-3') (Pandit et al., 2012).

ATAC-seq was performed as previously described (Buenrostro et al., 2013). For each
sample, cell nuclei were prepared from 5x10^4 cells and incubated with 2.5 µL
transposase (Illumina) in a 50 µL reaction for 30 minutes at 37°C. Following purification
of transposase-fragmented DNA, the library was amplified by PCR and subjected to
high-throughput sequencing on the HiSeq 2000 platform (Illumina).

NGS Data analysis and statistical methods
Reads from ChIP-seq and ATAC-seq libraries were trimmed for quality using
‘trim_galore’ and aligned to mouse genome assembly mm9 with bowtie2 using the
default parameters and duplicates removed with the Picard tool MarkDuplicates
(http://broadinstitute.github.io/picard/). Density profiles were created by extending each
read to the average library fragment size for ChIP and 0 bp for ATAC, then computing
density using the BEDTools suite (http://bedtools.readthedocs.io). Enriched regions
were discovered using MACS (v1.4) and scored against matched input libraries (fold
change > 2 and p-value < 1e-5). Genome browser tracks and read density tables were
normalized to a sequencing depth of ten million mapped reads.

ChIP and immunoblotting antibodies
Whole cell extracts for immunoblotting and Chromatin Immunoprecipitations were
conducted with H3K36me3 (61101 pAb, Active Motif), H3K36me3 (ab9050 Abcam),
H327me1 (61015 Active Motif), H3K27me2 XP (D18C8, CST), H3K9ac (ab4441,
Abcam), Hmgb2 (ab67282, Abcam), H3K27me3 (07-449, Millipore), H3K36me1
(Ab9048, Abcam), H3K36me2 (ab9049, Abcam), H3K4me3 (Ab8580, Abcam), Rag1
Southern Blot analyses
Southern blot analyses were conducted as described previously (Bredemeyer et al., 2006; Hung et al., 2018). Briefly, 10μg of genomic DNA from v-Abl B cells containing the pMG-INV recombination substrate were digested with XbaI or NheI and hybridized to a P32-labeled probe for Thy1 or GFP. Thy1 and GFP probes were made from 800bp and 700bp cDNA fragments respectively. After incubating with Thy1 probe, XbaI digested Southern blots were stripped and re-probed with the GFP probe.

Detection of Igh CDR3 sequences from NGS of proB genomic DNA
MiXCR v2.1.11 (Bolotin et al., 2017) was used to detect Igh CDR3 sequences from the next generation sequencing data. Only the best matched V/D/J genes were kept for each CDR3 sequences. The unproductive CDR3s, which are out of frame or contain stop codons, were excluded from the downstream analysis. The number of N-nucleotide additions and total deletions were evaluated based on the refPoints column in MiXCR output file. Wilcoxon rank sum test was used to compare the difference for the number of CDR3 sequences, the length of CDR3 amino acid sequences, the number of N-nucleotide additions and total deletions between knockout (KO) and WT group. All the statistical tests were implemented using R.

RNA-seq data processing for TRUST
RNA-seq fastq files were aligned to mouse reference genome mm10 using STAR2 (Dobin et al., 2013). TRUST v3.0.2 (Li et al., 2017; Hu et al., 2018; Hu et al., 2019) was used to infer both partial and complete BCR CDR3 sequences and gene coverage from the aligned RNA-seq BAM files. For each RNA-seq sample, the B cell percentage was estimated by the number of reads mapped to Igh gene region divided by the number of total sequencing reads and the BCR diversity was evaluated by the normalized unique Igh CDR3 calls (Hu et al., 2019).

Generation, analysis and histology of embryos for neurogenesis
C57Bl6 Setd2ff mice were bred to Nestin-cre mice to heterozygosity and timed matings between Setd2ff and Nestin Setd2Δ/+ were established to obtain E14.5, E16.5, E18.5 embryos and 2 hour post-partum pups. Tissues were fixed in 10% buffered formalin and embedded in paraffin and serially sectioned for sagittal and coronal sections (4 μm). TUNEL was performed on sections using TUNEL In Situ Cell Death Detection kit, POD (Roche) with Terminal Deoxynucleotidyl Transferase buffer (Takara Bio) and counterstained with DAPI. Immunostaining of cleaved caspase-3 was performed using anti-cleaved caspase-3 (Asp175) (5A1E) (CST 9664) with anti-rabbit conjugated to HRP secondary antibodies and visualized with DAB. Nuclei counterstained with hematoxylin. Whole tissues on slides were scanned by a digital slides scanner (3D Histech, MIDI) and viewed with Caseviewer (3D Histech).
**Statistical analyses**
Error bars in all data shown represent standard deviation. Unless otherwise indicated, determination of statistical significance and standard deviations were calculated using unpaired two-tailed Student’s t test (when comparing two conditions e.g. control vs knockout) or one way ANOVA (when comparing across multiple conditions concurrently) using Prism 7 software (GraphPad).
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