## Exposure to Mites Sensitizes Intestinal Stem Cell Maintenance, Splenic Marginal Zone B Cell Homeostasis, And Heart Development to Notch Dosage and Cooperativity.

Francis M. Kobia<sup>1†\*</sup>, Kristina Preusse<sup>1†</sup>, Quanhui Dai<sup>1,2</sup>, Nicholas Weaver<sup>3</sup>, Praneet Chaturvedi<sup>1</sup>, Sarah J. Stein<sup>4</sup>, Warren S. Pear<sup>4</sup>, Zhenyu Yuan<sup>5</sup>, Rhett A. Kovall<sup>5</sup>, Yi Kuang<sup>1</sup>, Natanel Eafergen<sup>6</sup>, David Sprinzak<sup>6</sup>, Brian Gebelein<sup>1</sup>, Eric Brunskill<sup>1</sup>, Raphael Kopan<sup>1,‡</sup>

- <sup>1</sup> Division of Developmental Biology, Department of Pediatrics, University of Cincinnati College of Medicine, Cincinnati, OH, USA.
- <sup>2</sup> State Key Laboratory of Genetic Engineering, School of Life Sciences, Fudan University, Shanghai, China
- <sup>3</sup> Immunology Graduate Program, University of Cincinnati College of Medicine, Cincinnati, OH, USA.
- Department of Pathology and Laboratory Medicine, Abramson Family Cancer Research Institute, Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA 19104, USA
- Department of Molecular Genetics, Biochemistry and Microbiology, University of Cincinnati College of Medicine, Cincinnati, OH, USA
- <sup>6</sup> School of Neurobiology, Biochemistry, and Biophysics, The George S. Wise Faculty of Life Sciences Tel Aviv University, Israel
- † The authors made equal contribution
- ‡ Author for correspondence: Raphael.Kopan@CCHMC.ORG
- \* Present Address: Mount Kenya University, Department of Research and Innovation,
- P. O. Box 342-01000, Thika Kenya.

**Key words**: Notch, heart, intestine, stem cells, MZB, Lymphoma, SMZL, ectoparasite

#### Abstract

Cooperative DNA binding is a key feature of transcriptional regulation. Here we examined the role of cooperativity in Notch signaling by CRISPR-mediated engineering of mice in which neither Notch1 nor Notch2 can homo- or heterodimerize, essential for cooperative binding to sequence paired sites (SPS) located near many Notch-regulated genes. While most known Notch-dependent phenotypes were unaffected in Notch1/2 dimer-deficient mice, a subset of tissues proved highly sensitive to loss of cooperativity. These phenotypes include heart development, compromising viability in combination with low gene dose, and the gut, developing ulcerative colitis in response to 1% DSS. The most striking phenotypes – gender imbalance and splenic marginal zone B cell lymphoma – emerged in combination with dose reduction or when challenged by chronic fur mite infestation. This study highlights the role of the environment in malignancy and colitis, and is consistent with Notch-dependent anti-parasite immune responses being compromised in the dimer deficient animals.

## **Highlights**

- Notch dimerization has an in vivo role in contributing to intestinal homeostasis
- Loss of cooperativity can manifest as Notch gain or loss of function phenotypes
- Mite infestation exacerbates all phenotypes, triggers MZB hyperproliferation in mutant animals
- Mite-infested mutant mice develop SMZL with age

### Introduction:

The evolutionarily conserved Notch receptors and ligands influence metazoan development and adult tissue homeostasis by directly translating an *inter*-cellular interaction into *intra*-cellular transcriptional outputs that control cell fate, proliferation, differentiation, and apoptosis (Artavanis-Tsakonas et al., 1999; Bray, 2006; Kopan and Ilagan, 2009). Mammals possess four Notch receptors (N1 to N4) and five Delta/Jagged ligands; all of which are Type I transmembrane proteins. The Notch pathway stands out relative to other signaling pathways in lacking signal amplification: canonical Notch signaling is initiated when a ligand on one cell engages a Notch receptor on a neighboring cell. This interaction unfolds the receptor's juxtamembrane region enabling cleavage by the metalloprotease ADAM10. The truncated, cell membrane bound polypeptide is then cleaved by the γ-secretase complex freeing the Notch Intracellular Domain (NICD), which subsequently translocates into the nucleus (Kopan and Ilagan, 2009; Kovall et al., 2017). NICD associates with the DNA-binding protein CSL (CBF1/Suppressor of Hairless/LAG-1, also known as RBPJ in vertebrates) and recruits the coactivator Mastermind; thereby assembling a Notch Transcription Complex (NTC) that activates Notch target gene expression (Bray, 2016; Gordon et al., 2008; Kovall et al., 2017).

The Notch pathway plays complex and context-dependent roles during development and adult tissue homeostasis. Perturbations in the Notch pathway are associated with developmental syndromes (Masek and Andersson, 2017) and cancers (Aster et al., 2017; Siebel and Lendahl, 2017). For example, *Notch1* promotes T-cell development (Pui et al., 1999; Radtke et al., 2010; Radtke et al., 1999), while *Notch2* is indispensable for Marginal Zone B-cell (MZB) development (Gibb et al., 2010; Hozumi et al., 2004; Liu et al., 2015; Maillard et al., 2004; Moran et al., 2007; Saito et al., 2003; Tanigaki et al., 2002; Witt et al., 2003). Accordingly, elevated Notch1 signaling is oncogenic in T-cells driving Acute Lymphoblastic Leukemia (T-ALL) (Ellisen et al., 1991; Weng et al., 2004) while increased Notch2 signaling is associated with splenic marginal zone B-cell transformation (Arcaini et al., 2016; Piris et al., 2017; Santos et al., 2017). Inversely, where Notch signals promote differentiation, the pathway can have a tumor suppressor function with diminished Notch signaling being associated with cancer (Demehri et al., 2009; Koch and Radtke, 2007).

How can the Notch pathway control multiple, dissimilar outcomes? Because each Notch receptor is consumed as it generates a signal and cannot be reused, signal strength has emerged as a key in controlling outcome. Indeed, some Notch-dependent processes are exquisitely sensitive to dosage and manifest both haploinsufficient and triplomutant effects in tissues such as the fly wing (Bray, 2006; Guruharsha et al., 2012). In mammalian tissues, signal

strength, defined as the sum of NICD released from all ligand-bound Notch receptors on the cell surface, is a far more important determinant of Notch signaling outcome than NICD composition (i.e. N1ICD vs N2ICD) (Liu et al., 2015; Liu et al., 2013; Ong et al., 2006). Several mechanisms have been found to modulate signal strength including receptor glycosylation (Haltiwanger, 2008; Kakuda and Haltiwanger, 2017), force-generating ligand endocytosis (Gordon et al., 2015; Langridge and Struhl, 2017), the contact interface between cells (Shaya et al., 2017), and the ratio of receptor/ligand constitution within the cell (Sprinzak et al., 2010). More recently, ligand-dependent signal dynamics were demonstrated to be another key determinant of signaling outcome (Nandagopal et al., 2018).

Interestingly, NICD can assemble into dimeric, cooperative NTCs (Arnett et al., 2010) on Sequence Paired Sites (SPSs), first described in the regulatory regions of the *Drosophila* E(spl) (enhancer of split) locus (Bailey and Posakony, 1995). SPSs consist of two DNA binding sites orientated in a head to head manner (Bailey and Posakony, 1995; Nellesen et al., 1999) separated by 15-17 nucleotides (Nam et al., 2007). NICD dimerization is facilitated via a conserved interface in NICD's ANK (Ankyrin repeat) domain. In human N1ICD, this interface consists of Arg<sup>1985</sup>, Lys<sup>1946</sup> and Glu<sup>1950</sup>. Dimerization in Notch1 is effectively abolished by mutating Arg<sup>1985</sup> into Ala<sup>1985</sup> (R<sup>1985</sup>A), Lys<sup>1946</sup> into Glu<sup>1946</sup> (K<sup>1946</sup>E) or Glu<sup>1950</sup> into Lys<sup>1950</sup> (E<sup>1950</sup>K), and loss of dimerization results in reduced activation of dimer-dependent targets (Arnett et al., 2010; Liu et al., 2010a; Nam et al., 2007).

Given the conservation of the dimer interface in most Notch receptors (*C. elegans* being the exception), the conservation of SPSs near known Notch targets (Hass et al., 2015; Liu and Posakony, 2012, 2014), and the ability of synthetic SPSs to regulate NICD levels (Kuang et al., 2019), we hypothesized that the precise mixture of agnostic and dimer-sensitive targets in a given cell will couple with Notch signal strength and dynamics to shape the responses to Notch signals and contribute to the context-specificity of Notch-related pathologies. To test this hypothesis in a mammalian species, we evaluated mice homozygous or hemizygous for dimerization-deficient alleles of *Notch1* (*N1*<sup>RA</sup>) and *Notch2* (*N2*<sup>RA</sup>) and found that dimerization contributes to context-specific Notch activity. We report that in the hemizygous state, dimer deficient Notch molecules are haploinsufficient in the heart and intestine, with lethal consequences when modified by the presence of the ectoparasite, *Demodex musculi* (fur mites). Mice homozygous for a dimer deficient *N1*<sup>RA/RA</sup> allele displayed a female-biased, mild cardiac phenotype consistent with Notch loss-of-function. Conversely, *N2*<sup>RA/RA</sup> mice displayed a striking over proliferation of marginal zone B cells in parasite-infested mice, but not in mite-free mice. In time, the cell-type specific gain of function activity of *N2*<sup>RA/RA</sup> produced a splenic

marginal zone lymphoma-like phenotype in mite-exposed mice. Mechanistically, these effects reflect a shift in target amplitude favoring monomer-driven targets, and reduced negative feedback.

## **Results**

### Generation of N1 and N2 dimerization deficient mice

To interrogate the physiological role of dimerization, we first quantified the effective binding cooperativity of NTC to SPS and CSL sites using purified RBPj, the N1 RAM-ANK domain, and a MAML protein in electrophoretic mobility shift assays (EMSA; see supplemental information). We compared the mobility of NTC complexes containing either wild type N1ICD or a mutant version with the substitution of a single critical Arginine residue (Arg<sup>1974</sup>) located in the mouse N1 ANK domain to alanine (Figure 1A-B, Supplemental Figure S1A-D). Quantifying the intensity levels of the bands in the image and fitting the data to a binding model that takes into account cooperative binding (see supplemental methods), we measured the cooperativity contributed by NTC binding to either an SPS probe (1xSPS) or a CSL probe containing two sites (2xCSL). We find that both wild type and mutant NTC had no cooperative binding to the 2xCSL probe. When the 1st site was already bound, the wild type NTC displayed a 5-fold stronger binding to the 2nd site in the 1xSPS probe compared to the binding to the 1st site. Importantly, NTC containing N1<sup>RA</sup>ICD had no cooperative binding to the 2nd site in the 1xSPS probe (Fig. 1A, Supplemental Figure S1A-D).

To ask if loss of cooperativity impacts function *in vivo*, we used CRISPR-Cas9 to introduce amino acid substitutions at Arg<sup>1974</sup> and Arg<sup>1934</sup> in N1 and N2, respectively. Following the generation of a double strand break in Exon 32 of N1 or N2, a short oligonucleotide harboring two linked mutations was homologously recombined. Nucleotides coding for Arg<sup>1974</sup> in N1 (N1R<sup>1974</sup>) and Arg<sup>1934</sup> in N2 (N2R<sup>1934</sup>) were changed to code for Ala, creating the N1<sup>R1974A</sup> (N1<sup>RA</sup>) and N2<sup>R1934A</sup> (N2<sup>RA</sup>) mutations, respectively (Figure 1B, E). To facilitate genotyping of N<sup>RA</sup> animals silent mutations were introduced to abolish a BgIII restriction site while creating an Xbal restriction site in N1<sup>RA</sup> (Figure 1B), or to generate a BgIII restriction site in N2<sup>RA</sup> (Figure 1E). Animals that carry one (N1<sup>+/RA</sup> or N2<sup>+/RA</sup>) or two (N1<sup>RA/RA</sup> or N2<sup>RA/RA</sup>) mutated chromosomes were first identified by digesting PCR products of the targeted chromosomal regions in Exon 32 with Xbal and/or BgIII. Xbal cuts the N1<sup>RA</sup> chromosome but not the wild type PCR product; BgIII cuts the *N2<sup>RA</sup>* but not the N2 wild type PCR products (Fig1 D, G). The presence of each respective RA mutation in *N1<sup>RA</sup>* and *N2<sup>RA</sup>* founders was subsequently confirmed by direct DNA sequencing

(Fig1 C, F). Animals deficient for N1 dimerization ( $N1^{RA/RA}$ ), N2 dimerization ( $N2^{RA/RA}$ ) or both N1 and N2 dimerization ( $N1^{RA/RA}$ ;  $N2^{RA/RA}$ ) were generated by crossing founders. All were viable, fertile and without any overt phenotype at P360 in mixed or C57BL6 background (unless otherwise stated, all subsequent results were generated in the mixed background). Because Notch1 signaling plays a central role in T-cell development (Radtke et al. 1999, 2010), we expected that loss of Notch1 dimerization would negatively impact the T-cell compartment. However, an analysis of the T-cell compartments in the thymus and spleen as well as thymic T-cell sub-compartments (DN, SP and DP), revealed a normal T-cell compartment in N1<sup>RA/RA</sup> mice relative to wild type controls (Supplemental Figure S2). This finding and the normal lifespan of  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  animals, was surprising since we anticipated many dimerization-dependent genes (e.g., Nrarp, Hes1, Hes5 and Myc; (Arnett et al., 2010; Hass et al., 2015) would be negatively impacted as seen when constitutively active wild type and dimer mutant Notch proteins lacking the extracellular domain ( $N\Delta E$  and  $N^{RA}\Delta E$ ) are overexpressed in cell culture (Supplemental Figure S1E).

# Homologous $N1^{RA/RA}$ ; $N2^{RA/RA}$ mice display barrier defects in the colon and reduced proliferation in the crypt when challenged with 1% DSS

Notch1 and Notch2 act redundantly during the development and maintenance of the gut to block Klf4-induced niche exit of intestinal stem cells (Pellegrinet et al., 2011; Riccio et al., 2008), and Math1-induced differentiation of secretory cells (Noah and Shroyer, 2013; Shroyer et al., 2007). However, no pathology was revealed in histological examination of N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> intestines (see below). Several studies have shown that a chronic decrease in Notch signaling compromised the intestinal barrier and exacerbated colitis in different mouse models (Mathern et al., 2014; Obata et al., 2012; Okamoto et al., 2009). Under the assumption that a compromised barrier may increase sensitivity to dextran sulfate sodium (DSS)-induced colitis. we asked if intestinal homeostasis and intestinal barrier were robust enough in N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> by exposing mice to DSS. When given 2.5% DSS in their drinking water, wild type mice developed colitis as evident by a mild weight loss whereas co-housed N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> mice exhibited severe weight loss necessitating euthanizing within one week (Figure 2A). To ask if N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> were predisposed to develop colitis we exposed them to 1% DSS, which does not affect the weight of control littermates even after repeated exposures (Figure 2B, C). By contrast, co-housed  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  mice experience significant weight loss, some had blood in stool at end of the third and fourth DSS cycle, and one had to be removed during the first cycle of DSS following severe weight loss and blood in stool. However, all remaining mice recovered

well when DSS was removed from the drinking water, even after multiple cycles of treatment (Figure 2B). Histological analysis of the colons revealed injury in wild type mice and a complete disruption of colonic epithelium of N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> mice treated with 2.5% DSS (Figure 2D, top). After four periods of 1% DSS exposure N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> mice had a more severe injury than controls or 2.5% DSS exposed wild type mice (Figure 2D). Relative to baseline (Figure 2E), proliferation in wild type crypt epithelia (Krt8/18 positive) was elevated in 1% or 2.5% DSS treated controls (Figure 2F). By contrast, we noticed a surprising decrease in proliferation competence (less Ki67) (and mitosis (Phospho-H3 positive cells)) in colonic epithelium following the fourth 1% DSS treatment in crypts of N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> mice (Figure 2G) relative to wild type or untreated crypts, suggesting a role for Notch cooperativity within colonic stem cells recovering from injury. Finally, RA-specific while changes in barrier markers were not detected, a significant increase in IL4. Th17A and IL1ß was observed in DSS-treated N1<sup>RA/RA</sup>: N2<sup>RA/RA</sup> relative to wild type (Figure 2H) consistent with enhanced inflammatory response driving colitis in these animals. Thus, these results are consistent with N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> alleles being hypomorphic loss-of-function alleles in maintaining intestinal homeostasis and immune response.

## N1<sup>RA</sup>; N2<sup>RA</sup> hemizygotes display impaired intestinal cell proliferation and fate allocation in the crypt and cause ventricular septum defects in the heart.

The wild type and R-A mutant monomeric NICD protein bind CSL sites similarly (Figure 1A, Supplemental Figure S1), supporting the hypothesis that the signal generated in  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  mice activated sufficient non-SPS gene targets to a level needed to support Notch-dependent decisions. If the Notch RA alleles are hypomorphic, challenging R-A mutant mice by lowering protein levels might reveal additional phenotypes. To test this idea, we generated an allelic series in which one copy of N1 or N2 was deleted in the  $N1^{RA/RA}$ ;  $N2^{+/+}$  or  $N1^{+/-}$ ;  $N2^{RA/RA}$  backgrounds, respectively, as well as deleting one copy of N1 and one copy of N2 deleted in the  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  background. Single N1 or N2 RA hemizygotes resembled R-A mutants; however, crossing  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  with  $N1^{+/-}$ ;  $N2^{+/-}$  animals generated significantly fewer  $N1^{RA/-}$ ;  $N2^{RA/RA}$  pups at birth than expected (p<3x10<sup>-9</sup>, Chi-squared distribution analysis, Supp Table S1).

*N1*<sup>+/-</sup>; *N2*<sup>+/-</sup> double hemizygous animals have a normal lifespan and no overt phenotype under normal housing conditions, but have been reported to develop mild cardiac phenotypes (Garg et al., 2005; Nigam and Srivastava, 2009). Given the critical role the circulatory system plays during gestation, we suspected that the poor representation of *N1*<sup>RA/-</sup>; *N2*<sup>RA/-</sup> pups at birth was due to defects in cardiac development. Analysis of E16.5 embryos born to mite-free dams

revealed highly penetrant and severe Ventricular Septal Defects (VSD) in the hearts of  $N1^{RA/-}$ ;  $N2^{RA/-}$  embryos (Figure 3), compromising viability. Milder VSD was also observed with lower penetrance in N1<sup>RA/-</sup>; N2<sup>+/RA</sup> embryos (Figure 3). Around this time, the colony became infected with fur mites, but the infestation was not immediately detected on the sentinels. Strikingly, when we retroactively segregated the fecundity data in our colony management software, we observed a significant (p=0.02,  $\chi^2$ ) skewing of the gender ratio in C57BL6  $N1^{RA/RA}$  carriers, with male:female ratio nearing 2:1 in fur mite infested mice, as opposed to the normal 1:1 ratio in mite-free mice (Supp Table S2).

The longest-lived N1<sup>RA/-</sup>; N2<sup>RA/-</sup> pup born to mite infested, mixed background dams was much smaller than its litter mates and was sacrificed at P30 when it became moribund. Necropsy revealed a striking and complete loss of the intestinal crypts and villi (Carulli et al., 2015; Riccio et al., 2008; van Es et al., 2005; Vooijs et al., 2007). Since complete failure to form an intestine would have not permitted survival outside the womb, we assumed this phenotype reflected loss of intestinal stem cells (Figure 4A, A') consistent with the loss of Ki67 in DSSchallenged N1<sup>RA/RA</sup>: N2<sup>RA/RA</sup> mice (Figure 2G). To investigate this phenotype further, we assessed proliferation and fate allocation in the developing intestine of E18-P1 animals with various allele combinations. To examine fate allocation, we stained sections with alcian blue and analyzed the number of secretory Goblet cells, which are known to expand in Notch hypomorphic background (Noah and Shroyer, 2013; Vooijs et al., 2007). The small intestines of control or N1<sup>+/-</sup>; N2<sup>+/-</sup> P0 pups were indistinguishable (Figure 4B, C). By contrast, alcian blue staining revealed a significant expansion in the goblet cell compartment at P0 in N1<sup>RA-</sup>; N2<sup>RA-</sup> pups born to fur mites infested dams (Figure 4D). In mice free of mites, the overall morphology appeared normal with few regions displaying excess goblets. To assess proliferation, we stained adjacent intestinal sections with antibodies against Ki67 and Phospho-H3. Strikingly, whereas trans-hemizygote  $N1^{+/-}$ :  $N2^{+/-}$  intestines resembled wild type (Figure 4E-F), the  $N1^{RA-}$ : N2<sup>RA/-</sup> intestine from mite-infested dames contained very few Ki67 positive cells (Figure 4G-H). We also noted a reduction in the number of cells undergoing DNA replication in particular, and competence to self-renew in general, in N1+/-; N2RA/RA intestines (Supplemental Figure S3C-C', G,G'). However, upon eradication of Demodex musculi with permethrin, all E18.5 and all but 1/8 surviving P0 N1<sup>+/-</sup>; N2<sup>+/-</sup> pups had at least some regions within the small intestine in which Ki67positive tissue could be observed (Figure 4J, the percentage of Ki67+ crypts; Supplemental Figure S3I-J': note reduced proliferation in Ki67-positive crypts of a P1 pup post-mite eradication). Overall, these data reflect an impaired ability to maintain intestinal stem cells in N1<sup>RA/-</sup>; N2<sup>RA/-</sup> mice, which was exacerbated by Demodex musculi infestation. Because these phenotypes can reflect complex interaction between multiple tissues in the  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  and  $N1^{RA/-}$ ;  $N2^{RA/-}$  mice, we deferred the mechanistic analysis of this gene-environment interaction in ISC maintenance to future investigation.

## The N2<sup>RA</sup> allele drives expansion of MZ B-cells in *Demodex musculi* infested mice.

Marginal zone B-cells (MZB) are exquisitely sensitive to N2 dosage (Gibb et al., 2010; Hozumi et al., 2004; Liu et al., 2015; Maillard et al., 2004; Saito et al., 2003; Tanigaki et al., 2002; Witt et al., 2003). Even a mild reduction in N2ICD can generate a noticeable decline in MZB cell numbers (Liu et al., 2015). We used FACS to analyze the splenic B cell population and asked if  $N2^{RA}$  displayed haploinsufficient phenotypes in the dose-sensitive MZ compartment. Intriguingly, we observed a significant increase in the MZB compartment in  $N2^{RA/RA}$  spleens in a mixed-background mouse exposed to *Demodex musculi* (Figure 5A). The size of other splenic B-cell subsets, including the immediate MZB precursors (MZP), Follicular B-cells (FoB) and transitional type-2 cells (T<sub>2</sub>) did not change significantly in either  $N2^{RA/RA}$  (Figure 5A),  $N2^{+/-}$  or  $N2^{RA/-}$  (Figure 5B), inconsistent with a fate switch.

MZB cells reside in the splenic marginal zone where they surveil for blood-borne pathogens. Upon encountering antigens, they rapidly differentiate into IgM producing plasmablasts that secrete vast amounts of IqM before undergoing apoptosis (Cerutti et al., 2013). To test IqM levels in vivo, we performed ELISA on serum collected from unstimulated N2<sup>RA/RA</sup> and littermate controls and detected no significant difference in the levels of circulating IgM (Figure 5C). To test for the possibility that loss of N2 dimerization negatively impacted differentiation into IgM producing plasmablasts upon stimulation, we FACS isolated MZB and FoB cells from mite-exposed N2<sup>RA/RA</sup> and wild type spleens. We then stimulated these cells in vitro with LPS for 5 days followed by imaging to assess for proliferation, and by ELISA to quantify the IgM levels secreted into the medium. Relative to wild type MZB cells, there was no appreciable difference in the ability of N2RA/RA MZB cells to respond to LPS stimulation (Supplemental Figure S4A), and ELISA analysis detected no significant difference in the amount of IgM secreted by N2<sup>RA/RA</sup> MZB cells relative to wild type controls (Figure 5C). Finally, we tested if loss of dimerization affected the development of B1 B cells, which predominantly reside in the peritoneal cavity (Witt et al., 2003). FACS analysis revealed no significant changes in the levels of either the B1a or B1b subsets of B1 B-cells between N2<sup>RA/RA</sup> and wild type animals (Figure 5D). Together, these data suggest that MZB cell numbers did not increase due to a fate switch or a failure to differentiate properly, and are consistent with a proliferative phenotype triggered by the N<sup>RA</sup> mutation.

In the course of these experiments, the Demodex musculi infestation was eradicated with permethrin. Within a few months, as we saw with other phenotypes, mite-free N2RA/RA animals no longer showed MZB expansion (Figure 5E, post-permethrin, PP). Since permethrin has been reported to suppress the immune system in mice (Blaylock et al., 1995; Punareewattana et al., 2001), we assumed permethrin transiently suppressed MZB proliferation. Multiple subsequent analyses over many months, however, failed to detect a difference in MZB numbers between N2<sup>RA/RA</sup> and control mice post eradication and permethrin treatment, suggesting that this allele was indistinguishable from wild type in untreated and pathogen free animals. Accordingly, we compared MZB numbers between  $N2^{+/-}$  spleens, which have decreased MZB numbers due to haploinsufficiency, with N2RA/- spleens and found no further reduction in MZB numbers (Liu et al., 2015; Saito et al., 2003). These data suggest that the  $N2^{RA}$  has neither a haploin sufficient character for the  $N2^{RA}$  allele in the MZB linage (Figure 5B). nor a proliferative phenotype triggered cell-autonomously by the N2RA protein. To ask if environmental stimuli interacted with N2<sup>RA</sup> within the spleen, mice were exposed to bacterial lipopolysaccharide (LPS) or to dermatitis, produced by twice weekly exposure of the ear epidermis to dust mite extract (HDM, Dermatophagoides farinae) for 6 weeks (see methods). MZB cell numbers were analyzed 8 weeks after the first LPS injection or HDM extract application. LPS induced transient IgM production but did not change MZB census in the spleen at 8 weeks (Figure 5E, LPS). Importantly, HDM-induced dermatitis elevated MZB numbers only in the N2<sup>RA/RA</sup> mice but not in controls (Figure 5E, HDM). Collectively, these experiments establish that specific gene-environment interactions produce the MZB phenotype in N2RA/RA mice.

To assess the rate of splenic B-cell proliferation in mite-infested mice, we stained C57BL/6J  $N2^{RA/RA}$  and WT spleen sections with antibodies against Ki67, B220 (to mark the B-Cells) and Moma-1 (to mark the marginal zone). We detected defuse B220 staining with elevated Ki67 in marginal zone cells in the  $N2^{RA/RA}$  spleen (Figure 5F), as well as in germinal centers (rich in FoB-cells and strongly B220 positive, Supplemental Figure S4), not seen in WT spleen of co-housed animals (Supplemental Figure S4B-D). Caspase 3 activation was not observed in  $N2^{RA/RA}$  sections (not shown). Increased marginal zone proliferation in  $N2^{RA/RA}$  relative to wild type control was also detected in spleens after six weeks of HDM treatment (Figure 5Fd, d'). Combined, these data suggest that mite infestation or prolonged HDM exposure leads to enhanced MZB proliferation.

## Aged N2<sup>RA/RA</sup> mice have enlarged spleens but only *Demodex musculi* infested mice progress to a Splenic Marginal Zone B-cell Lymphoma (SMZL)-like state

As noted above, we observed no change in lifespan in N2<sup>RA/RA</sup> or N1<sup>RA/RA</sup>; N2<sup>RA/RA</sup> animals. However, as old animals were culled, we noticed enlarged spleens in animals carrying the N2<sup>RA/RA</sup> allele post mite eradication (Supplemental Figure S5A). Spleens from mice infested by Demodex musculi were much larger, the most severe splenomegaly exhibited by N2<sup>RA/RA</sup>; N1+/R4 (Figure 6A-E), perhaps because they contained a higher contribution of the inflammationprone FVB strain (Quigley et al., 2011; Quigley et al., 2009). Histologically, spleens from miteinfested N2<sup>RA/RA</sup> carriers lost their typical architectures with N2<sup>RA/RA</sup>; N1<sup>+/RA</sup> spleens showing the greatest disruption. Splenic morphology showed an expansion of the white pulp at the expense of the red pulp. In the most severe cases, the spleens appeared to consist of white pulp only (Figure 6F-K). In addition, two  $N2^{RA/RA}$ :  $N1^{+/RA}$  animals had multiple enlarged abdominal lymph nodes. H/E staining revealed a striking morphological similarity to the enlarged spleens from the same animals, suggesting infiltration by a splenic population, or lymphoproliferation within the LN (Figure 6L-L"). To further characterize the nodes, we stained the enlarged spleens and lymph nodes for the B-cell markers B220 and Pax5. Both were positive for the B-cell markers (Figure 6M-N'). These features are reminiscent of human SMZL (Splenic Marginal Zone B-cell -Lymphoma), a clonal B-cell neoplasm that manifests itself in elderly patients and consists of small lymphocytes that infiltrate the lymph nodes and other organs (Arcaini et al., 2016; Piris et al., 2017; Santos et al., 2017). As would be expected, the enlarged lymph nodes and spleens were highly proliferative as indicated by Ki67 and phospho-H3 staining (Supplemental Figure S5B,D). Paradoxically, human SMZL are associated with hyperactive NOTCH2 signaling due to an N2ICD-stabilizing truncation upstream to the WSSSP sequence in the PEST domain of NOTCH2 (Kiel et al., 2012; Peveling-Oberhag et al., 2015; Rossi et al., 2012), whereas the N2<sup>RA</sup> alleles are hypomorphic in expression assays and in the context of the heart, crypt, and gut barrier. To confirm that this line of CRISPR-modified mice did not inherit a truncated Notch2 allele, we sequenced Exon 34 DNA isolated form lymph node filled with B-cells and found a perfect match to the wild type *Notch2* allele (ENSMUSG00000027878).

## Loss of Notch dimerization in *Demodex musculi* infested mice did not stabilize N2ICD in MZB cells but promoted their proliferation of MZB cells via the Myb-FoxM1 axis.

Recently, we identified an unanticipated consequence of increasing the number of SPS sites in the *Drosophila* genome, namely, accelerated degradation of phospho-NICD (Kuang et al., 2019). Further, we demonstrated that this enhanced degradation affects some, but not all,

Notch dependent decisions in *Drosophila*. Bristle precursor cells requiring a pulse of Notch were refractory to the destabilizing effect, whereas wing margin cells reliant on prolonged Notch signals were sensitive to NICD (Kuang et al., 2019). To test if the similarity of our phenotype to SMZL might in part be reflective of N2ICD stabilization, we assessed N2ICD levels in nuclear extracts from wild type and  $N2^{RA/RA}$  MZB cells. We did not detect a significant accumulation of nuclear N2ICD in sorted  $N2^{RA/RA}$  MZBs (Supplemental Figure S6), suggesting that the gain-of-function phenotypes observed in  $N2^{RA/RA}$  MZB cells is not due to enhanced N2ICD stabilization. Thus, unlike in many human SMZL patients, increased N2ICD stability is unlikely to be the cause of the SMZL phenotype in the N2<sup>RA/RA</sup> mice.

In N2<sup>RA/RA</sup> mice, Notch pathway activity in T-cells and skin is intact as it relies on Notch1. Assuming the MZB effect is cell autonomous we addressed the mechanism involved in enabling a haploinsufficient allele to drive MZB expansion via transcriptome and chromatin analyses. We sorted and performed RNAseg on MZB cells from spleens of three N2<sup>RA/RA</sup> and three wild type littermates infested with Demodex musculi. The data were mapped to the mouse genome (mm10), and PCA analysis indicated that one of the triplicates in each group was an outlier. To stabilize the variance across replicates a two-way normalization strategy was used (see methods). Differentially expressed transcripts were identified using EdgeR. 1207 differentially expressed transcripts (>1 LogTPM, >1 log fold changes, 0.1 FDR, <0.05 pVal) were clustered into upregulated (270) and downregulated (937) N2<sup>RA/RA</sup> genes (Figure 7A). Down regulated genes included the validated Notch MZB targets Hes1, Hes5, and Dtx1 and upregulated genes included FoxM1, E2F1, and Myb (Figure 7B). These three genes are known to promote cell cycle, DNA replication and DNA repair in B-Cells (Figure 7C, (Lefebvre et al., 2010; Lei et al., 2005)). Indeed, all the GO term for biological process for the upregulated genes were associated with the cell cycle (Figure 7B'; Table S3), with clear enrichment for E2F1 targets (Table S3). The downregulated genes did not enrich for any significant GO terms.

Given that Myb can act as an inducer of FoxM1 and E2F1 (Lefebvre et al., 2010), we confirmed its transcript levels were elevated by RT-qPCR (Figure 7D). Fos and Pax5 were also found to be increased within N2<sup>RA/RA</sup> MZB. Myb could be activated by altered accessibility or by increased burst frequency and/or size, affecting amplitude. To assess accessibility, we performed ATACseq on MZB cells isolated from wild type and N2<sup>RA/RA</sup> mice (4 biological replicates each). We mapped reads under the peaks and used EdgeR to identify differential accessibility across the genome between samples (see methods for detail). 89059 peaks were mapped, of which 87575 were present in both N2<sup>RA/RA</sup> and wild type MZB cells with all eight samples being highly related (R>92%; Spearman Correlation; Figure 7G). Hence, genome

accessibility changed minimally between these MZB cells with only 984 peaks (1.1%) enriched in N2<sup>RA/RA</sup> and 500 (0.56%) in wild type. We next used GREAT to assign genes to all peaks. Based on these assignments, 14698 peaks mapped near differentially expressed (DE) genes, and of these only 108 were unique to wild type and 122 were unique to  $N2^{RA/RA}$ . Importantly, none of the differently enriched peaks were present at the Myb, FoxM1, Dtx1, Hes1 or Hes5 loci (Supplemental Figure S7). Next, we analyzed DNAse hypersensitive peak upstream of the Myb locus in the ENCODE datasets, and identified an enhancer accessible in several cell types, including the kidney. The genome of a kidney-derived cell line used to overexpress constitutively active N∆E and a dimer deficient version, N∆ERA (Supplemental Figure S1) was examined by SplitDamID (Hass et al., 2015). We detected strong binding to the Myb enhancer using Notch/RBP complementing pairs, but not by Notch dimers. This data suggested that the Myb enhancer responds to monomeric NTC but not to dimeric NTC. Collectively, these data suggest that the changes in MZB gene expression in the N2<sup>RA/RA</sup> animals are not due to dramatic changes in chromatin accessibility but are rather due to changes in gene expression caused by the loss of Notch dimer NTC complexes at some other loci, most likely the SPSdependent negative regulators Hes1, Hes5, and Nrarp (Hass et al., 2015; Severson et al., 2017).

### **Discussion**

The possibility that DNA binding site architecture contributes to Notch signaling outcomes has been considered in past studies (Liu and Posakony, 2012, 2014). NTCs can bind to enhancers with CSL binding sites as monomers that function independently of one another. In such a model, the probability of target activation will depend upon the relative abundance of nuclear NICD and the number of CSL sites at a given enhancer (Gomez-Lamarca et al., 2018; Kueh et al., 2016; Schroeter et al., 1998) and/or burst size (Falo-Sanjuan et al., 2019; Lee et al., 2019). SPSs are found in enhancer regions of up to 30% of mammalian Notch targets including Nrarp/NRARP, Hes1/HES1, Hes5, and Myc (Arnett et al. 2010, Cave et al. 2005, Liu et al. 2010, Ong et al. 2006, Severson et al. 2017, Yashiro-Ohtani et al. 2014). We have previously documented that ~2500 SPSs were bound by N1ICD dimers in the mouse genome, and that dimerization-deficient Notch molecules bind poorly to SPSs in vivo, even at high NICD concentrations (Hass et al., 2015). In the murine kidney cell line, mK4, ≃15% of Notch targets required NICD dimerization. SPSs and cooperativity also proved critical for the oncogenic activity of N1ICD in murine T-cells, where NICD dimerization at a distant enhancer was required for Myc activation (Liu et al., 2010b). Thus, Notch targets are either agnostic to dimerization or sensitive to its presence, leading to the hypothesis that cooperativity may modulate Notch responses at physiological NICD concentrations. Recently, evidence that SPS contributed to increased probability of activation and increased duration during Drosophila embryonic development were reported (Falo-Sanjuan et al., 2019).

While our study concludes that embryonic development in the mouse proceeds normally in the presence of Notch receptors lacking cooperativity, it nonetheless uncovered important contributions of cooperativity to mammalian development and homeostasis under environmental stress. Notch<sup>RA</sup> phenotypes are greatly enhanced by reduced gene dosage in Notch1 (causing ventricular septal defects (Donovan et al. 2002, MacGrogan et al. 2010, Sakata et al. 2002)) or in both Notch1 and Notch2 (gut). The developmental phenotype uncovered when N1<sup>RA</sup> dose is reduced (in *N1<sup>RA/-</sup>*) resembles other Notch pathway deficiencies known to cause a wide range of heart defects including ventricular septal defects (VSD) (Donovan et al. 2002, MacGrogan et al. 2010, Sakata et al. 2002), and are milder than a weak hypomorphic Notch1/2 allele on the B6 background (Liu et al. 2015). Interestingly, dosage alone did not impact the ability of the *N2<sup>RA</sup>* allele to control development and homeostasis of the dosage-sensitive marginal zone B cells in parasite-free mice, even in the hemizygote state (*N2<sup>RA/-</sup>*).

Strikingly, loss of cooperativity-dependent contributions to homeostasis becomes acute when mutant animals are burdened with exoparasites, namely fur mites. Interestingly, when the

environment includes unmanaged exoparasite infestation, dimerization-deficient cooperativity mutant receptors can behave as hypomorphic (loss-of-function) alleles in some contexts (intestinal barrier formation, ventricular septum formation, intestinal stem cell self-renewal, female survival), remain indistinguishable from the wild type allele in many other tissues, or can drive the same disease as a gain-of-function allele (splenic marginal zone lymphoma, SMZL) in a cell type that otherwise seemed to be unaffected by the mutation. Note that in  $N2^{RA/RA}$  mice, Notch pathway activity in T-cells and skin, where Notch1 is present, are identical to the wild type. This provided the rationale for the molecular analysis – the defect is more likely to reside in the B-cell linage than elsewhere, though this will remain to be examined more fully in the future. We propose these behaviors can be explained by assuming that the balance between targets agnostic to dimerization (Myb) and those dependent on cooperativity on SPS sites (Hes1, Hes5, Nrarp) varies by cell type (Figure 8). Moreover, these differences in dependence upon SPS vs non-SPS target gene regulation between tissues can be further exposed by additional stressors such as environmental insults (i.e. exoparasites) or changes in gene dose. We have proposed before that accessible SPS can act as a "sink", holding on to NICD that otherwise will be available to regulate monomer-dependent enhancers. If true, more N2RAICD may available to regulate Myb. Importantly, this insight may translate to other transcription factors where variants of uncertain significance are associated with developmental syndromes or neoplastic disease. Such mutations may control molecular behaviors integrating the environment with specific cooperative interactions in the affected tissues.

The diverse cell types responding to a cutaneous parasite must reflect a systemic change driven by the parasite. Since we could replicate the impact on MZB proliferation in short exposures to dust mite extract, live parasite is not needed and the response is likely mediated by an immune cell (or cytokine) intermediate. Dermatitis is typically associated with both skin-produced cytokines (e.g., IL-33, -17C and TSLP), and a skewing of CD4+ T-cells into T Helper 2 (Th2) and their associated cytokines (e.g., IL-4,-5,-13, (Huang et al., 2003)). Similarly, fur mite infestation has been shown to increase the level of different cytokines (e.g. IL-6 and IL-12; (Johnston et al., 2009)), including Il-4 and -5 (Moats et al., 2016; Pochanke et al., 2006). While the exact axis linking anti-parasite immune responses and splenic marginal zone lymphoma, intestinal homeostasis, female/male ratios and heart development are yet to be identified, this study highlights the role of the environment in chronic disease, including ulcerative colitis and a smoldering malignancy such as SMZL. Combined with the significantly better outcomes in SMZL patients with activated *Notch2* alleles (Rossi et al., 2012), one has to wonder about the

role of the immune environment in onset and progression of human SMZL patients with *Notch2* mutations.

#### Methods

### **Animals:**

Animals were housed at CCHMC's animal facility in accordance with IACUC's animal welfare regulations. The use of animals for the experiments described in this work was approved through protocol number IACUC2018-0105. To generate the RA alleles, three gRNAs (g25, g26 and g27) that target sites surrounding the R<sup>1934</sup> codon in N2 were tested for their genome editing activities. Briefly, plasmids carrying the guide sequence under a U6 promoter and Cas9 were transfected into MK4 cells. Cells were harvested two days after transfection and PCR amplification, with primers flanking the target sites, was performed on genomic DNA isolated from the transfected cells. PCR products were then subjected to T7 endonuclease I (T7E1) assay. Cleavage products generated by T7E1 are indicative of indels present in the PCR products as a result of genomic DNA breakage caused by Cas9-gRNA and the subsequent repair by NHEJ. A gRNA (g3) for Tet2, which was known to have good editing activity, was included as positive control. Sixteen mice were obtained from injection of g27 (GGCATCCAGATCGGTTACA), Cas9 mRNA and a donor oligo (shown in Figure 1, g27 underlined in blue) into one-cell embryos. Digestion of PCR products with BgIII showed that eight contained products with the BallI site, four of which were homozygotes. Sequencing of PCR products confirmed that all had the correct R<sup>1934</sup>A mutations. Since we mated the homozygote mice to each other, we identified potential second site mutations with the CCTop tool ((Stemmer et al., 2015); https://crispr.cos.uni-heidelberg.de). g27 matches to Notch2 and to a few loci with four mismatches (Lemd1, Ccm2, Papolg, Tubb2b and Tubb6) falling within an Exon and next to a PAM sequence (no loci had fewer than four mismatches to this gRNA). PCR products from potential off-target sites were sequenced and no mutations were identified in DNA isolated from any of the founders. For Notch1, a similar process led to the selection of a gRNA (CATTCGGGCATCCAGATCTG), which was injected with Cas9 mRNA and a donor oligo carrying mutations that change R<sup>1974</sup> to alanine into one cell embryos. PCR products from 24 animals were digested with BgIII and XbaI, respectively. Only one heterozygote founder carried the R<sup>1974</sup>A mutation, confirmed by sequencing. This founder was outcrossed into our mixed colony, having a single possible 4-mismatch target in the pseudogene Vmn1r-ps6. In the C57BL6line, the same qRNAs were used. A single heterozygote founder pair was bred within the C57BL6line.

#### **DSS** treatment/Colitis induction:

For colitis induction mice were given 1% to 2.5% Dextran sulfate sodium (DSS) in autoclaved drinking water for up to 14 days. Weight was measured every day and stool was checked for changes to monitor disease progression; in case of severe weight loss (>15%) or blood in stool mice were euthanized. For recovery mice were given normal drinking water for 14 days before starting the next cycle of DSS treatment, up to 4 cycles of injury and recovery. At the end of treatment mice were euthanized and tissue was collected for analysis.

## LPS injection/dermatitis induction:

To stimulate the immune system with bacterial antigens we injected adult mice intraperitoneally twice with LPS (0.5ug/kg bodyweight, once a month) over an 8-week period. Blood was collected from the submandibular vein once per week to analyze IgM production by ELISA. The mice were euthanized after the eighth week and their spleen was harvested for histological analyses.

Mild dermatitis was induced by application of house dust mite (HDM, *Dermatophagoides farinae*) extract on the ears as described (Jin et al., 2016). In brief, the ear lobes of adult mice were gently stripped five times with surgical tape and 20 ug of HDM dissolved in autoclaved PBS was administered to each ear twice weekly for 6 weeks. Mice were euthanized and spleen harvested for histology and MZB numbers determination.

### **Antibodies:**

A full list of antibodies is provided in the Key Resource table. The following antibodies were used for FACS analysis: Anti-mouse/human CD45R/B220, anti-CD23, anti-CD21/CD35 (CR2/CR1), anti-CD93 [AA4.1], anti-mouse IgM, anti-IgD, anti-CD45.1, anti-CD45.2, anti-CD3ɛ, anti-Ly-6G/Ly-6C(Gr-1), anti-CD11b, anti-TCRb and anti-CD5 were all purchased from Biolegend. Anti-Mouse CD19 was purchased from BD Biosciences. Rabbit Anti-Ki67 (Novocastra), rabbit anti-cleaved caspase 3 (Cell signaling), rabbit anti-Phospho-Histone H3 (Ser10) (Cell signaling), rat anti-Pax5 (Biolegend), Guinea pig anti-Cytokeratin 8+18 antibody and FITC rat anti-mouse/human CD45R/B220 (BioLegend) were used for fluorescence immuno-

histochemistry. Anti-rabbit Alexa Fluor Cy3 (Jackson ImmunoResearch), anti-rabbit Alexa Fluor 647 (Jackson Immunoresearch), anti-rat Cy3 (Jackson Immunoresearch) and Alexa fluor anti-guinea pig 647 secondary antibodies were used for in fluorescence immuno-histochemistry. Notch2 (D76A6) XP (Cell signaling) and HRP-linked rabbit IgG (GE health care) secondary antibody were used for western blot analysis.

## Protein purification and electrophoretic mobility shift assay (EMSA).

The mouse RBPj (aa 53-474), mouse N1ICD (aa 1744-2113), mouse N1<sup>RA</sup>ICD (aa 1744-2113) and human SMT3-MAML1 (aa 1-280) proteins were expressed and purified from bacteria using affinity (Ni-NTA or Glutathione), ion exchange, and/or size exclusion chromatography as previously described (Friedmann et al., 2008). The purity of proteins was confirmed by SDS-PAGE with Coomassie blue staining and concentrations were determined by absorbance measurements at UV280 with calculated extinction coefficients. EMSAs were performed using native polyacrylamide gel electrophoresis as previously described (Uhl et al., 2010; Uhl et al., 2016). Fluorescent labeled probes (1.75 nM/each probe) were mixed with increasing concentrations of purified RBPj protein in 4-fold steps (from 2.5 to 640 nM) with or without the indicated purified N1ICD and MAML proteins at 2μM. Acrylamide gels were imaged using the LICOR Odyssey CLx scanner.

### **EMSA** quantitation

To quantitatively analyze the EMSA experiments we first extracted the gray scale intensity of each band in each lane of the EMSA gels (Supplementary Fig. S1 A-B). We then fitted the data to a 2-site binding model that takes into account cooperative binding to the 2nd site. The model calculates the binding probability assuming equilibrium binding kinetics (Michaelis-Menten) to two sites. Cooperativity is taken into account by assuming that the binding dissociation constant of a complex to the second site is divided by a cooperativity factor C, namely that  $K_{d,2nd} = \frac{1}{C}K_{d,1st}$ . The probabilities to find the probe bound by 0,1 or 2 complexes are given by:

$$P_0 = \frac{1}{1 + 2\alpha + C\alpha^2}$$
 ,  $P_1 = \frac{2\alpha}{1 + 2\alpha + C\alpha^2}$  ,  $P_2 = \frac{C\alpha^2}{1 + 2\alpha + C\alpha^2}$ 

where  $\alpha = \frac{[NTC]}{K_d}$  is the statistical weight associated with binding of the NTC complex to a CSL site.  $K_d$  is the dissociation constant to a single site. If the cooperativity factor, C, is equal to 1, then the binding to the two sites is non-cooperative. C > 1 corresponds to positive cooperativity (2<sup>nd</sup> binding is enhanced). C < 1 corresponds to negative cooperativity (2<sup>nd</sup> binding is suppressed).

Since we observe that even at high concentrations of RBPj the 1-site state is never depleted (e.g. see RBPj on CSL) we assumed that there is only some fraction of the probes, f < 1, that binds two complexes and another fraction, (1 - f), that can only bind one complex. In this case the probability to find the probe is modified to:

$$P_0 = f \frac{1}{1 + 2\alpha + C\alpha^2} + (1 - f) \frac{1}{1 + 2\alpha} \ , \quad P_1 = f \frac{2\alpha}{1 + 2\alpha + C\alpha^2} + (1 - f) \frac{2\alpha}{1 + 2\alpha} \ , \quad P_2 = f \frac{C\alpha^2}{1 + 2\alpha + C\alpha^2}$$

We then fit the experimental data (normalized band intensities) to these expressions. The fitting parameters are  $K_d$ , C, and f. The parameters are extracted for each experiment separately.

To get an estimation of the confidence interval on the fitting parameters, we use a bootstrap method to randomly generate 5000 data sets with the same statistical properties as the experimental data sets (the same mean and standard deviation of band intensities). We then perform the same fitting procedure on all bootstrapped data to obtain the 95% confidence interval on the fitting parameters.

#### Flow cytometry:

MZB and FoB cells were sorted by FACS as described in (Chen et al. 2016). Single cell splenocytes were prepared by placing the spleen in a petri dish and mincing it with a razor blade. The disrupted spleen tissue was then transferred into a 50ml falcon tube and pipetted up and down in 5 ml of ice cold 1% BSA (Fisher Scientific) in PBS and briefly vortexed to ensure thorough disruption. Single-cell suspensions were obtained by passing the disrupted spleen tissue through a 70 um strainer (Falcon). The strainer was then rinsed with 5 ml ice cold 1% BSA/PBS and the single cell preparation pelleted by centrifuging at 2000 rpm, for 5 min followed by red blood cell lysis (BioLegend), after which the cells were pelleted at 2000 rpm, for 5 min, and resuspended in ice cold 3% BSA in PBS before cell counting. Single color antibody stained as well as unstained controls were prepared with 3 million splenocytes each, 1.5ul (0.75ug) of each antibody was added into the corresponding single-color control tube. For MZB detection,

approximately 10 million cells (in 1ml 3% BSA) were distributed into Eppendorf tubes and 7.5ul of the antibody mix (2.5ul each of FITC anti-B220, PE/Cy7 anti-CD23 and PerCP/Cy5.5 anti-CD21) added into each tube. They were then incubated at 4°C for 60 minutes before sorting for B220+CD21hiCD23lo MZB cells and B220+CD21htCD23hi FoB cells. To isolate MZB (CD93-B220<sup>+</sup>CD19<sup>+</sup>IgM<sup>+</sup>IgD<sup>+</sup>CD23<sup>lo</sup>), FoB (CD93<sup>-</sup>B220<sup>+</sup>CD19<sup>+</sup>IgM<sup>+</sup>CD21<sup>+</sup>), MZP (CD93<sup>-</sup> B220<sup>+</sup>CD19<sup>+</sup>CD21<sup>hi</sup>IgM<sup>hi</sup>IgD<sup>hi</sup>CD23<sup>hi</sup>) and T2 B-cells (CD93<sup>+</sup>B220<sup>+</sup>IgM<sup>+</sup>CD23<sup>+</sup>), single cell splenocytes were incubated with the APC/Cy7 anti-B220, PE/Cy7 anti-CD93, Hoechst blue anti-CD19, FITC anti-IgM, AmCyan anti-IgD, Percp/Cy5.5 anti-CD21 and Pacific blue CD23 and FACS sorted. To analyze peritoneal B1 B-cell populations, peritoneal cells were collected as described in (Ray and Dittel, 2010) and incubated with APC/Cy7 anti-B220, PE anti-TCRb, FITC anti-CD11b, PE/Cy7 anti-CD23 and APC anti-CD5. They were then gated into B2 B-cells (B220<sup>+</sup>CD23<sup>+</sup>) and B1 B-cells (B220<sup>+</sup>CD23<sup>-</sup>). B1 B-cells were then gated into B1a (CD11b<sup>+</sup>CD5<sup>+</sup>) and B1b (CD11b<sup>+</sup>CD5<sup>-</sup>) B-cells.

For T cell analysis, thymus and spleen were harvested and disrupted by gentle grinding of tissue with a pestle (CellTreat) on a 40um cell strainer (Falcon). Red blood cells in spleen samples were lysed using ACK lysis buffer (Lonza) for 5 minutes at 4 degrees. Cells were counted and stained at 20x106 cells/ml. Cells were resuspended in PBS with Zombie Violet viability dye (Biolegend) and incubated at 4°C for 20 minutes. Cells were washed, and then resuspended in an antibody cocktail in PBS supplemented with 2% FBS (Hyclone) in the presence of 5% 2.4G2 Fc blocking antibody (in house), and incubated for 30 minutes at 4 degrees. Cells were washed, resuspended in PBS with 2% FBS, and analyzed using an LSRII (BD). Data were analyzed using FlowJo software v9.7 and average population percentages and absolute numbers were graphed using GraphPad Prism (GraphPad Software, Inc.).

#### MZB and FOB culture and ELISA:

MZB and FoB cells were cultured in RPMI-1640 (Sigma-Aldrich) containing 10% FBS (Sigma-Aldrich), 10μg/ml gentamicin (Gibco B.R.L) and 50μM β-mercaptoethanol (Sigma-Aldrich). 10<sup>5</sup> MZB and FoB cells were seeded into U-bottom 96-well plates (Thermo Scientific) in 200μL of media. Differentiation was stimulated by adding lipopolysaccharides from *Salmonella enterica* serotype typhimurium -LPS (Sigma-Aldrich) into each well at a final concentration of 2μg/ml. The LPS-stimulated cells were cultured at 37°C, 5% CO<sub>2</sub>, for 5 days. To measure IgM levels by ELISA, MaxiSorp flat-bottom 96-well plates (Nunc) were coated with 200μL of 5μg/ml goat anti-

mouse IgM (Southern Biotech) in 0.05% PBST (PBS-tween20) at 4°C, overnight. The plates were then washed thrice with 0.05% PBST and blocked with 3% BSA (Fisher Scientific) in PBS for 30 minutes at room temperature. Purified mouse IgM (Southern Biotech) was used to prepare a control standard. 150µl of supernatant collected from the stimulated B-cells at the end of 5-day incubation was diluted in 0.05% PBST (PBS containing 0.05% Triton-100), and along with the standard, was loaded on to the MaxiSorp flat bottom (Thermo Fisher Scientific) ELISA plates for 1hr incubation at room temperature followed by three washes with PBST. HRP goat anti-mouse IgM (Southern Biotech), diluted at 1:5000 in PBST, was added and incubated for 1hr at room temperature followed by three washes with PBST. TMB substrate solution (Thermo Fisher Scientific) was then added and developed for 10 minutes. Development was stopped with 0.16M sulfuric acid before colorimetric absorption analysis at 450nm.

## **Histology**

Mouse tissues were fixed overnight in 4% PFA and embedded in paraffin. Adult intestinal tissue was divided into sections corresponding to the jejunum, duodenum, ileum and colon. Each section was then folded into a swiss-roll (Moolenbeek and Ruitenberg, 1981) and then fixed, mounted and sectioned for histological analysis. For P0 animals, whole intestinal tissue was collected and then fixed and mounted whole before sectioning for histological or immunofluorescence analysis. 5µm thick sections were deposited on Superfrost plus microscope slides (Fisher Scientific), deparaffinized with xylene (Fisher Scientific), and rehydrated. Sections were stained with hematoxylin for 2.5 minutes and Eosin for 1 minute before dehydration, clearing and mounting. For alcian blue staining, sections were rinsed in 3% acetic acid (Fisher Scientific) for 3 minutes and incubated at room temperature for 30 minutes in 1% Alcian Blue 8GX (Sigma-Aldrich) in 3% acetic acid solution. The sections were then rinsed for 3 minutes in 3% acetic acid followed by 10 minutes in running water. After counterstaining with Nuclear Fast Red (Vector) for 5 minutes, slides were rinsed for 10-minutes in running water prior to dehydration, clearing and mounting. For each sample, images of at least 10 random fields of view were acquired on a Nikon 90i Upright widefield microscope. Where indicated, quantitative analysis of images was done using Nikon's NIS-Elements Advanced Research software (NIS-AR). In order to automatically identify and quantify Alcian Blue positive goblet cells, an inhouse NIS-AR plugin was applied to identify and mask the discrete, large, dark blue Alcian Blue spots scattered on the pink-white Nuclear Fast Red counterstain (Figure 3). We then manually traced the villi in the region of interest (ROI) to exclude the Alcian Blue stained

mucus. The plugin then counted the Alcian Blue positive foci within the ROI. Goblet cell numbers were determined for all acquired fields of view for each animal analyzed. The average number of goblet cells per field of view was obtained for each analyzed animal. Where multiple animals of the same genotype were analyzed, the average goblet cell number per field of view was obtained by averaging the goblet cell numbers in all fields of view from all the animals of that genotype.

## Fluorescent Immunohistochemistry

Paraffin embedded sections were deparaffinized with xylene, rehydrated, and subjected to Heat-Induced Epitope Retrieval (HIER) by boiling in Trilogy (Cell Marque) for 30 minutes. After cooling, the sections were rinsed with distilled water for five minutes followed by PBS with 0.3% Triton-X100 (Fisher Scientific) for five minutes. The sections were then incubated at room temperature for two hours in blocking solution (10% normal donkey serum (Jackson ImmunoResearch) in PBS-Triton), followed by overnight incubation with respective primary antibodies at 4°C. The sections were washed 3x10 minutes with PBS-Triton and incubated for two hours at room temperature with Fluorophore-labeled secondary antibodies in blocking solution. Unbound secondary antibody was washed three times for 10 minutes each in PBS-Triton and where indicated, counterstained with DAPI before mounting with Prolong Gold Antifade (Invitrogen). Confocal imaging was performed on a Nikon Ti-E Inverted Microscope.

### Cytoplasmic/nuclear separation and Western Blot

N2ICD stability was determined in FACS-isolated MZB and FoB cells. Sorted cells were pelleted for 5 min with 2000 rpm at 4°C and resuspended in 500 ul hypotonic buffer (HB; 20mM Tris-HCl, pH7.5, 10mM NaCl, 3 mM MgCl<sub>2</sub>) with added protease inhibitor (Roche). After 30min incubation on ice, 25 ul of 10 NP-40 was added and samples vortexed for 10 seconds. Nuclei were pelleted at 3000 rpm/10min at 4°C and the cytoplasmic fraction transferred into a fresh tube. The nuclei pellets were washed twice in 500ul HB and then lysed in 20ul 2X Laemmli sample buffer (120mM Tris-HCl, pH6.8, 20% glycerol, 4% SDS). The cytoplasmic fractions were concentrated using AmiconUltra – 0.5mL – 30K centrifugal filters (Merck Millipore) and mixed

1:1 with 2X Laemmli sample buffer. The extracts were separated on 6% polyacrylamide (Biorad) gel and transferred onto nitrocellulose membranes in Tris/glycine transfer buffer. Membranes were blocked with blocking solution (5% milk in 0.1% PBS-Tween20 (Fisher Scientific) for 1 hour, room temperature, and incubated overnight at 4°C with primary antibody in blocking solution. Membranes were then washed three times for five minutes with 0.1% PBS-Tween20 and incubated for 1 hour at room temperature with anti-rabbit HRP secondary antibody (GE healthcare) in blocking solution. Membranes were developed with a Supersignal West femto chemiluminescent substrate kit (Fisher scientific) and developed using Chemidoc (Biorad) detection system. Signal intensities were quantified using Image Lab (Biorad) software.

## **qPCR**

RNA from tissue and cells was extracted using the PureLink<sup>TM</sup> RNA Mini kit (Invitrogen) and cDNA was synthesized with SuperScript® II reverse transcriptase (Invitrogen) following the manufacturers instruction. Quantitative PCRs were performed using iTaq Universal SYBR® Green Supermix (BioRad) on the StepOnePlus<sup>TM</sup> RT PCR system. Data were analyzed using the Delta-Delta-CT methods. A full list of oligos is provided in In Key Resources Table.

### **RNAseq**

Total RNA was isolated from FACS-sorted MZB cells using a Qiagen MicroRNAeasy kit according to manufacturer's instructions. RNA-SEQ libraries were generated using the Nugen Ovation RNA-Seq System V2 and 75 b paired-end reads were obtained on an Illumina Hi-Seq 2500 machine with read depths of ~38 million following to Illumina protocols. Quality checked RNA-seq reads were mapped to mouse genome (mm10) using RSEM (v1.2.19) and quantified. Raw counts and transcripts per million (TPM) units were reported for all genes. Principal component analysis (PCA) was performed using R package prcomp. ~60% variance was accounted for by PC 1 and 2. Based on the PCA, analysis, we excluded biological replicate 3 in both WT MZB and RA MZB from further analysis (See Table S3). Differential Expression was performed using R package RUVSeq, which normalizes and contains EdgeR. A two-way normalization strategy was used to stabilize the variance across replicates for each transcript: an upper quantile normalization followed by an empirical gene normalization using ~5000 least differentially expressed transcripts. Two-way normalized counts were then used to perform

differential expression with EdgeR. Upregulated and downregulated genes were filtered (see Table S3) and loaded to the Toppgene suite (<a href="https://toppgene.cchmc.org/enrichment.jsp">https://toppgene.cchmc.org/enrichment.jsp</a>) to identify significant enrichment.

## **ATACseq**

For sample library preparation we followed the Omni-ATAC method outlined by (Buenrostro et al., 2015; Corces et al., 2017) and purified Tn5 was generated as described (Picelli et al., 2014). Briefly, 50,000 nuclei from FACS-sorted MZB cells were processed for Tn5 transposasemediated tagmentation and adaptor incorporation at sites of accessible chromatin. FACS isolated MZB cells were pelleted and washed with ice-cold PBS. The pellet was resuspended in ATAC-Resuspension Buffer (10mM Tris-HCl pH 7.4, 10mM NaCl, 3mM MgCl2 0.1% NP40, 0.1% Tween-20, 0.01% Digitonin) and incubated on ice for 3 minutes. The lysed cells were washed in ATAC-Wash Buffer (10mM Tris-HCl pH 7.4, 10mM NaCl, 3mM MgCl<sub>2</sub>, 0.1% Tween-20), inverted three times and the nuclei pelleted. The nuclei were resuspended in ATAC transposition mix (10mM Tris-HCl pH 7.6, 10mM MgCl<sub>2</sub>, 20% Formamide, 100nM Tn5 transposase) and incubated at 37°C for 30 minutes in a thermomixer at 1000 RPM. Following tagmentation, the DNA fragments were purified using a Zymo DNA Clean and Concentrator Kit and library amplification was performed using customized Nextera PCR primer Ad1 in combination with any of Ad2.1 through Ad2.12 barcoded primers as described (Buenrostro et al., 2015). The quality of the purified DNA library was performed utilizing an Agilent Bioanalyzer 2100 using High Sensitivity DNA Chips (Agilent Technologies Inc., Santa Clara, CA). The samples were pooled at a concentration of 5nM and run on an Illumina HI-SEQ 2500 sequencer (Illumina, Inc. San Diego, CA) to obtain paired-end reads of 75 bases (PE75). sequencing was carried out on two conditions MZB wild-type and RA. For each condition, 4 biological replicates were sequenced, in single end fashion. Read depth for replicates varied from ~ 50 to 98 million reads thereby giving us average read depth of ~67 million reads. Quality check, mapping and peak calling was performed using CSBB-v3.0 [Process-ChIP-ATAC\_SingleEnd]. CSBB uses fastqc, bowtie2 and macs2 [parameters: --nomodel --shift 37 -extsize 73] respectively. Duplicate mapped reads were removed before peak calling. Mapped reads in bam format was converted to bigwigs using deeptools (deeptools.ie-freiburg.mpg.de) for visualization purposes. For assessing open nucleosome region differences from MZB WT to RA MZB, we performed differential peak analysis. Within each sample type only peaks with 75% replication and at least 50% overlap among biological replicates were used for differential peak

analysis. Further, peak sets passing above defined criteria were merged (at least 1bp overlap) using bedtools merge and then number of reads mapping under each peak for each replicate was inferred using FeatureCounts program. Finally, differential peak analysis was performed using EdgeR. No statistically significant changes in peaks (ATAC regions) between WT MZB and RA MZB were identified.

## **Key Resources**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Goat Anti-Mouse IgM-UNLB	Southern Biotech	1021-01
Goat Anti-Mouse IgM-HRP	Southern Biotech	1021-05
Lyophilized Rabbit Polyclonal Antibody Ki67		
Antigen	Novocastra	NCL-Ki67p
Cleaved Caspase-3 (Asp175) (5A1E) Rabbit mAb	Cell signaling	9664P
Phospho-Histone H3 (Ser10) Antibody	Cell signaling	9701S
Purified anti-Pax-5 Antibody	Biolegend	649702
	Jackson_	
Cy3-AffiniPure Donkey Anti-Rabbit IgG (H+L)	ImmunoResearch	711-165-152
Cy3 AffiniPure Donkey Anti-Rat IgG (H+L)	Jackson	740 405 450
, , ,	ImmunoResearch	712-165-150
Alexa Fluor® 594 anti-mouse/human CD45R/B220	Diologond	103254
[RA3-6B2]	Biolegend Abcam	ab51814
Anti-Metallophilic Macrophages antibody [MOMA-1] Alexa Fluor® 488-AffiniPure F(ab')2 Fragment	Jackson	au31014
Donkey Anti-Rabbit IgG (H+L)	ImmunoResearch	711-546-152
Alexa Fluor® 647 anti-mouse IFN-γ [XMG1.2]	Biolegend	505816
Biotin anti-mouse IL-5 [TRFK4]	Biolegend	504401
Alexa Fluor® 488 anti-mouse IL-17A [TC11-	Biologona	001101
18H10.1]	Biolegend	506909
Alexa Fluor® 647 anti-mouse CD3ε Antibody	Biolegend	152319
Anti-Cytokeratin 8+18 antibody (ab194130)	Abcam	ab194130
Donkey anti-Guinea Pig IgG, Alexa -2788875	Millipore Sigma	AP193SA6
Mouse NKp46/NCR1 Antibody	R&D Systems	AF2225-SP
Alexa Fluor® 647 AffiniPure F(ab') <sub>2</sub> Fragment	Jackson	
Donkey Anti-Goat IgG (H+L)	ImmunoResearch	705-606-147
anti-mouse/human CD45R/B220 [RA3-6B2]	Biolegend	103206
anti-mouse CD23 [B3B4]	Biolegend	103222
anti-mouse CD21/CD35 (CR2/CR1) [7E9]	Biolegend	123416
anti-mouse CD93 (AA4.1)	Biolegend	136505
anti-mouse IgM [RMM-1] 50 μg	Biolegend	406505
anti-mouse IgD [11-26c.2a]	Biolegend	405723
anti-mouse CD45.1 [A20]	Biolegend	110706
anti-mouse CD45.2 [104]	Biolegend	109814
Anti-Mouse CD3e Clone 145-2C11	BD Biosciences	553063
anti-mouse Ly-6G [1A8]	Biolegend	127615
anti-mouse/human CD11b [M1/70]	Biolegend	101216
anti-mouse/human CD45R/B220	Biolegend	103227
anti-mouse TCR β chain [H57-597]	Biolegend	109208
APC/Cy7 anti-mouse CD45R/B220	Biolegend	103224
anti-mouse CD5 [53-7.3]	Biolegend	100626
Rat Anti-Mouse CD19	Biolegend	bdb563557
CD4 Monoclonal Antibody (RM4-5), PerCP-Cyanine5.5	eBiosciences	45-0042-82
CD8a Monoclonal Antibody (53-6.7), FITC	eBiosciences	11-0081-82

CD8a Monoclonal Antibody (53-6.7), PE-Cyanine7	eBiosciences	25-0081-82
TCR beta Monoclonal Antibody (H57-597), APC-		47-5961-82
eFluor 780	0210001011000	0001 02
APC anti-mouse CD25 Antibody (clone PC61)	Biolegend	102011
CD4 Monoclonal Antibody (GK1.5), PE	eBiosciences	12-0041-82
PE/Cy7 anti-mouse CD117 (c-Kit) Antibody	Biolegend	105814
CD3e Monoclonal Antibody (145-2C11), FITC	eBiosciences	11-0031-82
CD19 Monoclonal Antibody (eBio1D3 (1D3)), PE	eBiosciences	12-0193-82
Notch2 (D76A6) XP USE	Cell signaling	5732
monoclonal anti-b-actin (AC-15)	Sigma-Aldrich	A5441
ECL Rabbit IgG, HRP-linked whole AB (from		NA934
Donkey)		
ECL Mouse IgG, HRP-linked whole AB (from	GE Healthcare	NA931
Donkey)		
Bacterial and Virus Strains		
BL21 (DE3) Tuner Competent cells	Novagen	70623
Biological Samples		
Chemicals, Peptides, and Recombinant Proteins		
Xbal	NEB	R0145
BgIII	NEB	R0144
Mouse IgM-UNLB	Southern Biotech	0101-01
DyLight649(Cy5)-conjugated Streptavidin	Jackson	016-490-084
by Lights 10 (6) of sorting action of the pravious	Immunoresearch	010 100 001
Dextran sulfate sodium salt (DSS, M.W. ~36000 -		0216011050
50000, Colitis grade)		
Red blood cell lysis buffer (10x)	BioLegend	420301
BSA	Fisher Scientific	Bp1600-100
FBS	HyClone	AE24573269
FBS	Sigma-Aldrich	F2442
Nunc MaxiSorp <sup>™</sup> flat bottom	Thermo Fisher	44-2404-21
·	Scientific	
TMB substrate solution	Thermo Fisher	N301
RPMI-1640	Sigma-Aldrich	R0883
Gentamicin	Gibco B.R.L	15750-060
β-mercaptoethanol	Sigma-Aldrich	M7522
U-bottom 96-well	Thermo Scientific	174925
Superfrost Plus microscope slides	Fisher Scientific	12-550-15
Xylene	Fisher Scientific	X3P
Alcian Blue 8GX	Sigma-Aldrich	A3157
Nuclear Fast Red	Vector/Sigma-	H-3403/N3020
	Aldrich	
Trilogy	Cell Marque	920P-10
Normal donkey serum	Jackson	017-000-121
	ImmunoResearch	
Prolong Gold Antifade Reagent	Invitrogen	P36930
Complete Protease Inhibitor Cocktail	Roche	11697498001
Red Blood cell lysis buffer	BioLegend	420301
ACK lysis buffer	Lonza	10-548E

Cell strainer (40 μm)	Falcon	08-771-1
Cell strainer (70 µm)	Falcon	08-771-2
Critical Commercial Assays		
Supersignal West Femto Chemiluminescent substrate kit	Thermo Scientific/Fisher Scientific	P134095
PureLink <sup>™</sup> RNA Mini kit	Life Technologies (Invitrogen)	12183025
Superscript®II reverse transcriptase	Invitrogen	18064014
iTaq Universal SYBR®Green Supermix	BioRad	1725121
Zombie VioletTM Fixable Viability kit	BioLegend	423113
Micro RNAeasy kit	Qiagen	74106
Zymo DNA Clean and Concentrator kit	Zymoresearch	D4029 ?
Deposited Data		
Raw and analysed RNA seq data	This paper	GEO?
Raw and analysed ATAC seq data	This paper	GEO?
SplitDam data	(Hass et al., 2015)	GEO: GSE70402
ATACseq DAT	This paper	GEO?
Experimental Models: Cell Lines		
mK4 cells	(Valerius et al., 2002)	N/A
Experimental Models: Organisms/Strains		
N1RA	This paper	N/A
N2RA	This paper	N/A
N1 deletion	(Conlon et al., 1995)	Notch1 <sup>tm1Con</sup>
N2 lacZ	(Skarnes et al., 1995)	N/A
C57BL/6	Charles River	027
FVB/N	Harlan	N/A
B6D2F2	Harlan	N/A
Oligonucleotides		
N1-deletion fwd: ACGATATCGTGGTGCATACGCTCCTGTGCG	(Conlon et al., 1995)	N/A
N1-deletion rev: GTCAGTTTCATAGCCTGAAGAACG	(Conlon et al., 1995)	N/A
N1RA fwd: GCGGGATTCCTAGCCTGGTTACTC	This paper	N/A
N1RA rev: GTCCTTGTTGGCTCCGTTCTTCAG	This paper	N/A
N2-LacZ fwd: GTTGCAGTGCACGGCAGATACACTTGCTGA	(Skarnes et al., 1995)	N/A
N2-lacZ rev:	(Skarnes et al.,	N/A
GCCACTGGTGTGGGCCATAATTCAATTCGC	1995)	
N2RA fwd: ACGGCTCATCCCTGACATGAG	This paper	N/A
N2RA rev: TCCTCTCAGAAGGGTAGCAAGTC	This paper	N/A
Actb qPCR fwd: GGCTGTATTCCCCTCCATCG	Harvard PrimerBank	6671509a1
Actb qPCR rev: CCAGTTGGTAACAATGCCATGT	Harvard PrimerBank	6671509a1
Fos qPCR fwd: CGGGTTTCAACGCCGACTA	Harvard	6753894a1
	PrimerBank	

Fos qPCR rev: TTGGCACTAGAGACGGACAGA	Harvard	6753894a1
FOS GEORIEV. TIGGCACTAGAGACGGACAGA		073309 <del>4</del> a1
0    000 (	PrimerBank	0070007 4
Gapdh qPCR fwd:	Harvard	6679937a1
AGGTCGGTGTGAACGGATTTG	PrimerBank	
Gapdh qPCR rev:	Harvard	6679937a1
TGTAGACCATGTAGTTGAGGTCA	PrimerBank	
Myb qPCR fwd: AGACCCCGACACAGCATCTA	Harvard PrimerBank	19526459a1
Myb qPCR rev: CAGCAGCCCATCGTAGTCAT	Harvard PrimerBank	19526459a1
Notch1 qPCR fwd:	Harvard PrimerBank	13177625a1
GATGGCCTCAATGGGTACAAG		
Notch1 qPCR rev:	Harvard PrimerBank	13177625a1
TCGTTGTTGATGTCACAGT		
Notch2 qPCR fwd:	Harvard PrimerBank	33859592a1
ATGTGGACGAGTGTCTGTTGC		
Notch2 qPCR rev:	Harvard PrimerBank	33859592a1
GGAAGCATAGGCACAGTCATC		
Pax5 qPCR fwd: CCATCAGGACAGGACATGGAG	Harvard PrimerBank	6679213a1
Pax5 qPCR rev: GGCAAGTTCCACTATCCTTTG	Harvard PrimerBank	6679213a1
IL1β-qPCR-fwd:	Cao et al., 2012	N/A
CAACCAACAAGTGATATTCTCCATG		
IL1β-qPCR-rev: GATCCACACTCTCCAGCTGCA	Cao et al., 2012	N/A
IL4-qPCR-fwd:	Stellato et al., 2011	N/A
AGATGGATGTGCCAAACGTCCTCA		
IL4-qPCR-rev:	Stellato et al., 2011	N/A
AATATGCGAAGCACCTTGGAAGCC		
Claudin4-qPCR-fwd:	Harvard PrimerBank	6753440a1
GTCCTGGGAATCTCCTTGGC		2==2442
Claudin4-qPCR-rev:	Harvard PrimerBank	6753440a1
TCTGTGCCGTGACGATGTTG	Obile 1 - 1 - 0040	N1/A
ZO1-qPCR-fwd: GCACCATGCCTAAAGCTGTC	Chihara et al., 2013	N/A
ZO1-qPCR-rev: ACTCAACACACCACCATTGC	Chihara et al., 2013	N/A
TNFα-qPCR-fwd:	Harvard PrimerBank	7305585a1
CCCTCACACTCAGATCATCTTCT	Llaw card Drive and and	700550504
TNFα-qPCR-fwd: GCTACGACGTGGGCTACAG	Harvard PrimerBank	7305585a1
IL17A-qPCR-fwd:	This paper	N/A
GACGCGCAAACATGAGTCC	This man an	N1/A
IL17A-qPCR-rev: TTTGAGGGATGATCGCTGCT	This paper	N/A
N1RA sgRNA fwd:	This paper	N/A
caccgCATTCGGGCATCCAGATCTG	This paper	N/A
N1RA sgRNA rev: aaacCAGATCTGGATGCCCGAATGc	This paper	IN/A
N2RA sgRNA fwd:	This paper	N/A
caccGGCATCCAGATCGGTTACA	Tillo papel	IN/A
N2RA sgRNA rev:	This paper	N/A
aaacTGTAACCGATCTGGATGCC	ττιιο ραροί	14//
44401017170007110100/11000		

N1RA donor oligo: g*c*t*tgcatttagatcaccctgcctgaacccatccctgccttccagat cctgctccggaacgcagctactgatctagatgcccgaatgcatgatg gcacaactccactgatcctggctgcgcgcctggccgtg*g*a*g	This paper	N/A
N2RA donor oligo:  a*g*c*caggcgggcagccaggatcaggggggtagtaccatcgttc  attctggcatccagatctgttaccgcgttgcggatcagaatctagaag  agaagcagagaagtgtcccttagataaggaaaaaga*a*t*g	This paper	N/A
2xCSL(EMSA): cgaaCGTGGGAAacctaggctagaggcacCGTGGGAA actagtgcgggcgtggct	This paper	N/A
1xSPS(EMSA): gctaCGTGGGAAaggagcaaactgcgtTTCCCACGttcgt agtgcgggcgtggct	This paper	N/A
5'IRDye-700_complementary_oligo(EMSA): agccacgcccgcact	This paper	N/A
5'IRDye-800_complementary_oligo(EMSA): agccacgcccgcact	This paper	N/A
Recombinant DNA	Γ	
Plasmid: N1∆E	(Ong et al., 2006)	N/A
Plasmid: N1RA∆E	This paper	N/A
pSpCas9(BB)	Ran et al., 2013	48139
pTXB1-Tn5	Picelli et al., 2014	N/A
Software and Algorithms		
Image lab	Bio-Rad Laboratories	https://www.bio-rad.com/en-us/product/image-lab-software?ID=KRE6P5E8Z
Prism 8	Graph Pad	www.graphpad.co m
CCTop tool	Stemmer et al., 2015	https://crispr.cos.u ni-heidelberg.de
NIS-Elements Advanced Research software	Nikon	N/A
GeneSpring	Agilent	N/A
FlowJo software v9.7	Becton Dickinson	https://www.flowjo. com
Altanalyze	(Emig et al., 2010)	http://www.altanaly ze.org/
RSEM (v1.2.19)	(Li and Dewey, 2011)	http://deweylab.bio stat.wisc.edu/rsem
RUVSeq	(Risso et al., 2014)	http://www.biocond uctor.org/packages /devel/bioc/html/R UVSeq.html

EdgeR	(Robinson et al., 2010)	http://bioconductor. org/packages/relea se/bioc/html/edge R.html
deeptools	Ramirez et al., 2016	deeptools.ie- freiburg.mpg.de
CSBB-v3.0 software	CSBB - Computational Suite For Bioinformaticians and Biologists	https://omictools.c om/csbb-tool
ToppGene Suite	Cincinnati Children's Hospital	http://toppgene.cch mc.org
Other		
Lipopolysaccharides from Salmonella enterica (LPS)	Sigma-Aldrich	L6143
House dust mite extract (HDM, Dermatophagoisdes farina)	Greer Laboratories Inc./Fisher Scientific	NC0277827

## **Acknowledgments**

The authors wish to thank Dr. Matt Kofron with help with imaging, Dr. Yueh-Chiang Hu and the genome editing core for generating the NRA alleles, and members of the Kopan lab for comments and encouragement during the study. Dr. Stephan Waggoner, Kelli VanDussen and members of the DB and Notch community for valuable advice. Many thanks to Dr Sai Tammula and the veterinarians at CCHMC. Cell sorting was supported by NIH S100D023410 and 01DK106225 to the CCHMC Research Flow Cytometry Core. BG and DS were supported at a grant from the National Science Foundation and the Binational Science Foundation (NSF/BSF - 1715822). SJS was supported by NIH T32CA009140 and the American Cancer Society PF-15-065-01-TBG. WSP was supported by NIH R01 CA215518 (WSP). RK, FMK, KP, NW, and EB were supported by MIH R01 CA163353 (RK). RK and QD were supported by the William K. Schubert Endowed Chair at Cincinnati Children's Hospital Medical Center. QD receives a small stipend from the Chinese government.

### **Authors contributions**

FMK generated most of the data in mite infested mice and contributed to writing of the manuscript; KP contributed most of the data in the mite-free colony and supervised QD during the DSS experiments. NW contributed the initial establishment of the MZB phenotype. RAK/ZY purified the proteins, YK, NE, DS and BG analyzed cooperativity, with YK/BG performing EMSA experiments and NE/DS calculating the cooperativity factor. SJS and WSP analyzed *N1*<sup>RA</sup> T-cells. EB performed the RNA and chromatin analyses and supervised NW. BA performed the RNAseq clustering and enrichment analysis. EB/RK conceptualized the study and analyzed the data with input from KP, FMK, and QD. RK wrote the manuscript. All authors read and edited the final version.

## Figure Legends:

**Figure 1. Generation of Notch dimerization deficient mice.** (A) Electrophoresis mobility shift assay for purified proteins binding to CSL (magenta) or SPS (green) probes. Balls mark occupancy of one or two sites. Note that cooperative binding by wild type NTC, but not the RA mutant NTC, specifically depletes the SPS probe, but not the CSL probe (compare lanes 8 and 13). See text and Supplemental Figure S1 for detail. (B-C) CRISPR-Cas9-mediated double-strand break was used to mediate homologous recombination of a short oligonucleotide into Exon 32 substituting Arg  $N1^{R1974}$  (B-B") and  $N2^{R1934}$  into Ala (C-C"), generating  $N1^{RA/RA}$  and  $N2^{RA/RA}$  animals. To facilitate genotyping, two silent mutations (in red) were included in the oligo, abolishing the BgIII restriction site while generating an Xbal site in *Notch1* (B'). In Notch2, a silent mutation (in green) was included to create a BgIII site (C'). Sequencing PCR products containing these regions confirmed the presence of the Arg to Ala substitution in founders; digestion of these PCR products with Xbal (for N1, B") and BgIII (for N2, C") confirmed the presence of one (N1) or two (N2) mutant alleles.

Figure 2. Notch dimerization deficient mice are sensitized to DSS-induced colitis. (A) All  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  mice exposed to 2.5% DSS treatment had to be euthanized due to severe weight loss before day seven. (B) Daily weight measurements of wild type (blue) or mutant (red) mice treated with alternate cycles of 1%DSS (gray sections) or no DSS (white sections). (C) Survival curve of 1% DSS treated mice. Three of four mutant mice had to be euthanized due to sever weight loss by the fourth cycle. (D) Hematoxylin-eosin staining of colonic tissue from DSS treated mice. Dashed black boxes are enlarged below. Yellow dashed box showed a section with crypts in an otherwise injured colon in 1% DSS treated  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  mice. (E) Ki67 staining of untreated mice with the indicated genotypes, crypt regions (Ki67+) are boxed. (F) Increased proliferation competence in colon of DSS treated wild type mice (Krt8/18+, Ki67+ staining outside the box). (G) Decreased proliferation competence in colon of  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  mice exposed to four rounds of 1% DSS. (H) qPCR on RNA extracted from distal colon of  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  and WT mice treated for 11 days with 1% DSS; n=3. (\*, p<0.05).

Figure 3. Notch1<sup>R1974A</sup> substitution compromised ventricular septum development. (A) Normal heart with complete septum. (B)  $N1^{RA/-}$ ;  $N2^{RA/-}$  heart with a severe, highly penetrant VSD. The penetrance by age shown below the image. (C)  $N1^{RA/-}$ ;  $N2^{+/RA}$  heart with a milder, less penetrant VSD.

Figure 4. Notch dimerization deficient mice are hypomorphic for Notch activity in the gut. (A-C) Alcian blue staining of  $N1^{RA/RA}$ ;  $N2^{RA/RA}$  and  $N1^{+/-}$ ;  $N2^{+/-}$  adult intestines. (D) Deletion of one N1 and one N2 allele in the RA background ( $N1^{RA/-}$ ;  $N2^{RA/-}$ ) caused lethality; loss of the intestine (presumably due to stem cell exhaustion) was evident in the longest surviving pup at P30. (E-G) Alcian blue analysis of P0 intestines detected an increase in goblet cell numbers in  $N1^{RA/-}$ ;  $N2^{RA/-}$  (G) relative to wild type or  $N1^{+/-}$ ;  $N2^{+/-}$  intestines (E-F), quantified in (H). (I-J) Ki67 staining in P0 intestine from control  $N1^{+/-}$ ;  $N2^{+/-}$  (I) or  $N1^{RA/-}$ ;  $N2^{RA/-}$  (J) newborn. (K) quantification of Ki67-positive area at the indicated age. Among surviving  $N1^{RA/-}$ ;  $N2^{RA/-}$  pups during (n=2) and after (n=12) fur mite infestation.

Figure 5. Loss of Notch2 dimerization expands the splenic MZ B-cell compartment in the presence of mites. (A) FACS analysis of  $N2^{RA/RA}$  fur mite infested mouse spleens revealed an expansion the MZ B-cell compartment relative to wild type (+/+). Values for MZ P, F0, and T2 splenic B-cell populations were not significantly different; n=11. (B) The MZB-cell compartment in heterozygotes. (C) LPS-induced differentiation and IgM production *in vitro* and serum levels of IgM *in vivo*; n=4. (D). Peritoneal B1a B-cell compartment in mite infested wild type and  $N2^{RA/RA}$  mice. (ns=not significant, n=2). (E). The MZB-cell compartment in  $N2^{RA/RA}$  mice after treatment with permethrin, an immunosuppressant used to manage fur mite infestation (PP), after LPS, or after 6 weeks of treatment with house dust mites (HDM) extract. (\*, p<0.05; ns=not significant). In (A-E) P-values generated by two-tailed t-test. Error bars represent s.d. (F). Proliferation in the splenic marginal zone of mite infested mice: (Fa) The spleen from a mite infested control animal or (Fb) from  $N2^{RA/RA}$  mice. Boxed region magnified in Fc and again, in Fc', asterisk and dot provided for orientation. Magnification noted. (Fd, d') 60x view of marginal zone from control and HDM treated  $N2^{RA/RA}$  mice.

Figure 6. Aged N2<sup>RA/RA</sup> and N2<sup>RA/RA; N1+/RA</sup> mice develop severe splenomegaly and tumors reminiscent of Splenic Marginal Zone B-cell Lymphoma (SMZL). (A-E) Images of spleens from mice P600-P700 with indicated genotypes. (F-K) Severely enlarged spleens are associated with disorganized morphology and the appearance of cells with large nuclear:cytoplasmic ratio (genotypes indicated, see low magnification image in Supplemental Figure S6E). (L) large enteric lymph nodes detected in the individual shown in (J), densely populated with cells with large nuclear:cytoplasmic ratio (L'-L"). Staining the spleen (M) and lymph node (N) with Ki67 and the B-cell markers B220 and Pax5 identifies infiltrating cells as spleen-derived B cells. Magnification: 10x.

Figure 7. Mite infested N2<sup>RA/RA</sup> activate a proliferation module in MZB. (A) RNAseq analysis identifies enrichment for genes trending down or up in N2<sup>RA/RA</sup> MZB realative to wild type controls. (B) Down trending genes include classical dimer-dependent Notch targets in MZB, Hes1 and Hes5, and Dtx. Up regulated genes include FoxM1, Myb, and E2F1. (B') clusters 6 and 7 are highly enriched for GO terms associated with replication. (C) A schematic of the replication module controlled by Myb. (D) QPCR validation for Fos, Myb, and Pax5. (F) The Myb locus analyzed for chromatin accessibility (Encode DHS and ATAC-seq data generated in this study), transcription (RNA), and DNA methylation by DAM methyl transferase complementation (SplitDAM). Control D/AM haves methylation patterns are compared to methylation by Notch/RBPj pairs (both dimers and monomers) and Notch/Notch pairs (only dimers). See text and (Hass et al., 2015) for additional detail. (G) Spearman correlation table measures ATACseq peaks similarity matrix between all replicates. All are >92% similar.

Figure 8. A schematic summary of the findings with a hypothetical mechanism. *Notch* integrates environmental cues (mites, cytokines) to drive proliferation in MZB (*Notch2*) and block differentiation in intestinal stem cells (ISC. *Notch1* & 2). Activation of SPS-dependent *Hes* repressors creates a negative feedback tuning of transcription amplitude in MZB, and driving the response magnitude in ISC. In *Notch*<sup>RA/RA</sup> mutants, SPS-dependent *Hes* gene expression is dampened, and increased availability of NICD<sup>RA</sup> to monomer driven target due to reduce trapping on accessible SPS may also contribute. Repeated DSS treatment, or hemizygosity in *Notch*<sup>RA/RA</sup> mutants, exposed insufficient blockade of pro-differentiation signals in ISC, most likely delivered via immune cells or cytokines (IL1ß, 4, or 17A) in DSS treated mice. Fur mite infestation creates an environment for runaway proliferation in MZB, and enhanced dosage effects in *Notch*<sup>RA/RA</sup> mutants leading to complete loss of ISC postnatally even without DSS challenge.

#### References

Arcaini, L., Rossi, D., and Paulli, M. (2016). Splenic marginal zone lymphoma: from genetics to management. Blood *127*, 2072-2081.

Arnett, K.L., Hass, M., McArthur, D.G., Ilagan, M.X., Aster, J.C., Kopan, R., and Blacklow, S.C. (2010). Structural and mechanistic insights into cooperative assembly of dimeric Notch transcription complexes. Nat Struct Mol Biol *17*, 1312-1317.

Artavanis-Tsakonas, S., Rand, M.D., and Lake, R.J. (1999). Notch signaling: cell fate control and signal integration in development. Science *284*, 770-776.

Aster, J.C., Pear, W.S., and Blacklow, S.C. (2017). The Varied Roles of Notch in Cancer. Annu Rev Pathol 12, 245-275.

Bailey, A.M., and Posakony, J.W. (1995). Suppressor of Hairless directly activates transcription of Enhancer of split complex genes in response to Notch receptor activity. Genes and Development 9, 2609-2622.

Blaylock, B.L., Abdel-Nasser, M., McCarty, S.M., Knesel, J.A., Tolson, K.M., Ferguson, P.W., and Mehendale, H.M. (1995). Suppression of cellular immune responses in BALB/c mice following oral exposure to permethrin. Bull Environ Contam Toxicol *54*, 768-774.

Bray, S.J. (2006). Notch signalling: a simple pathway becomes complex. Nat Rev Mol Cell Biol 7, 678-689.

Bray, S.J. (2016). Notch signalling in context. Nat Rev Mol Cell Biol 17, 722-735.

Buenrostro, J.D., Wu, B., Chang, H.Y., and Greenleaf, W.J. (2015). ATAC-seq: A Method for Assaying Chromatin Accessibility Genome-Wide. Current protocols in molecular biology / edited by Frederick M Ausubel [et al] 109, 21 29 21-21 29 29.

Carulli, A.J., Keeley, T.M., Demitrack, E.S., Chung, J., Maillard, I., and Samuelson, L.C. (2015). Notch receptor regulation of intestinal stem cell homeostasis and crypt regeneration. Dev Biol 402, 98-108.

Cerutti, A., Cols, M., and Puga, I. (2013). Marginal zone B cells: virtues of innate-like antibody-producing lymphocytes. Nat Rev Immunol *13*, 118-132.

Corces, M.R., Trevino, A.E., Hamilton, E.G., Greenside, P.G., Sinnott-Armstrong, N.A., Vesuna, S., Satpathy, A.T., Rubin, A.J., Montine, K.S., Wu, B., et al. (2017). An improved ATAC-seq

protocol reduces background and enables interrogation of frozen tissues. Nat Methods 14, 959-962.

Demehri, S., Turkoz, A., and Kopan, R. (2009). Epidermal Notch1 Loss Promotes Skin Tumorigenesis by Impacting the Stromal Microenvironment. Cancer Cell *16*, 55-66.

Ellisen, L.W., Bird, J., West, D.C., Soreng, A.L., Reynolds, T.C., Smith, S.D., and Sklar, J. (1991). TAN-1, the human homolog of the Drosophila notch gene, is broken by chromosomal translocations in T lymphoblastic neoplasms. Cell *66*, 649-661.

Falo-Sanjuan, J., Lammers, N.C., Garcia, H.G., and Bray, S.J. (2019). Enhancer Priming Enables Fast and Sustained Transcriptional Responses to Notch Signaling. Developmental Cell *50*, 1-15.

Garg, V., Muth, A.N., Ransom, J.F., Schluterman, M.K., Barnes, R., King, I.N., Grossfeld, P.D., and Srivastava, D. (2005). Mutations in NOTCH1 cause aortic valve disease. Nature *21*, 180-184.

Gibb, D.R., El Shikh, M., Kang, D.J., Rowe, W.J., El Sayed, R., Cichy, J., Yagita, H., Tew, J.G., Dempsey, P.J., Crawford, H.C., *et al.* (2010). ADAM10 is essential for Notch2-dependent marginal zone B cell development and CD23 cleavage in vivo. J Exp Med *207*, 623-635.

Gomez-Lamarca, M., Falo-Sanjuan, J., Stojnic, R., Abdul Rehman, S., Muresan, L., Jones, M.L., Pillidge, Z., Cerda-Moya, G., Yuan, Z., Baloul, S., *et al.* (2018). Activation of the Notch Signaling Pathway In Vivo Elicits Changes in CSL Nuclear Dynamics. Dev Cell *44*, 611-623 e617.

Gordon, W.R., Arnett, K.L., and Blacklow, S.C. (2008). The molecular logic of Notch signaling--a structural and biochemical perspective. J Cell Sci *121*, 3109-3119.

Gordon, W.R., Zimmerman, B., He, L., Miles, L.J., Huang, J., Tiyanont, K., McArthur, D.G., Aster, J.C., Perrimon, N., Loparo, J.J., *et al.* (2015). Mechanical Allostery: Evidence for a Force Requirement in the Proteolytic Activation of Notch. Dev Cell *33*, 729-736.

Guruharsha, K.G., Kankel, M.W., and Artavanis-Tsakonas, S. (2012). The Notch signalling system: recent insights into the complexity of a conserved pathway. Nat Rev Genet *13*, 654-666.

Haltiwanger, R.S. (2008). Unsweetened Notch leads to myeloproliferation. Blood *112*, 214-215.

Hass, M.R., Liow, H.H., Chen, X., Sharma, A., Inoue, Y.U., Inoue, T., Reeb, A., Martens, A., Fulbright, M., Raju, S., *et al.* (2015). SpDamID: Marking DNA Bound by Protein Complexes Identifies Notch-Dimer Responsive Enhancers. Mol Cell *59*, 685-697.

Hozumi, K., Negishi, N., Suzuki, D., Abe, N., Sotomaru, Y., Tamaoki, N., Mailhos, C., Ish-Horowicz, D., Habu, S., and Owen, M.J. (2004). Delta-like 1 is necessary for the generation of marginal zone B cells but not T cells in vivo. Nat Immunol.

Huang, C.H., Kuo, I.C., Xu, H., Lee, Y.S., and Chua, K.Y. (2003). Mite allergen induces allergic dermatitis with concomitant neurogenic inflammation in mouse. J Invest Dermatol *121*, 289-293.

Jin, M., Choi, J.K., Choi, Y.A., Kim, Y.Y., Baek, M.C., Lee, B.H., Jang, Y.H., Lee, W.J., Lee, S.J., Kim do, W., *et al.* (2016). 1,2,4,5-Tetramethoxybenzene Suppresses House Dust Mite-Induced Allergic Inflammation in BALB/c Mice. Int Arch Allergy Immunol *170*, 35-45.

Johnston, N.A., Trammell, R.A., Ball-Kell, S., Verhulst, S., and Toth, L.A. (2009). Assessment of immune activation in mice before and after eradication of mite infestation. J Am Assoc Lab Anim Sci 48, 371-377.

Kakuda, S., and Haltiwanger, R.S. (2017). Deciphering the Fringe-Mediated Notch Code: Identification of Activating and Inhibiting Sites Allowing Discrimination between Ligands. Dev Cell *40*, 193-201.

Kiel, M.J., Velusamy, T., Betz, B.L., Zhao, L., Weigelin, H.G., Chiang, M.Y., Huebner-Chan, D.R., Bailey, N.G., Yang, D.T., Bhagat, G., *et al.* (2012). Whole-genome sequencing identifies recurrent somatic NOTCH2 mutations in splenic marginal zone lymphoma. J Exp Med *209*, 1553-1565.

Koch, U., and Radtke, F. (2007). Notch and cancer: a double-edged sword. Cell Mol Life Sci *64*, 2746-2762.

Kopan, R., and Ilagan, M.X. (2009). The Canonical Notch Signaling Pathway: Unfolding the Activation Mechanism. Cell *137*, 216-233.

Kovall, R.A., Gebelein, B., Sprinzak, D., and Kopan, R. (2017). The Canonical Notch Signaling Pathway: Structural and Biochemical Insights into Shape, Sugar, and Force. Dev Cell *41*, 228-241.

Kuang, Y., Golan, O., Preusse, K., Cain, B., Salomone, J., Campbell, I., Okwubido-Williams, F.V., Hass, M.R., Eafergan, N., Moberg, K.H., *et al.* (2019). Enhancer architecture sensitizes cell specific responses to Notch gene dose via a bind and discard mechanism. bioRxiv.

Kueh, H.Y., Yui, M.A., Ng, K.K., Pease, S.S., Zhang, J.A., Damle, S.S., Freedman, G., Siu, S., Bernstein, I.D., Elowitz, M.B., *et al.* (2016). Asynchronous combinatorial action of four regulatory factors activates Bcl11b for T cell commitment. Nat Immunol *17*, 956-965.

Langridge, P.D., and Struhl, G. (2017). Epsin-Dependent Ligand Endocytosis Activates Notch by Force. Cell *171*, 1383-1396 e1312.

Lee, C., Shin, H., and Kimble, J. (2019). Dynamics of Notch-Dependent Transcriptional Bursting in Its Native Context. Dev Cell *50*, 426-435 e424.

Lefebvre, C., Rajbhandari, P., Alvarez, M.J., Bandaru, P., Lim, W.K., Sato, M., Wang, K., Sumazin, P., Kustagi, M., Bisikirska, B.C., *et al.* (2010). A human B-cell interactome identifies MYB and FOXM1 as master regulators of proliferation in germinal centers. Mol Syst Biol *6*, 377.

Lei, W., Liu, F., and Ness, S.A. (2005). Positive and negative regulation of c-Myb by Cyclin D1, cyclin-dependent kinases and p27 Kip1. Blood.

Liu, F., and Posakony, J.W. (2012). Role of architecture in the function and specificity of two Notch-regulated transcriptional enhancer modules. PLoS Genet 8, e1002796.

Liu, F., and Posakony, J.W. (2014). An enhancer composed of interlocking submodules controls transcriptional autoregulation of suppressor of hairless. Dev Cell 29, 88-101.

Liu, H., Chi, A.W., Arnett, K.L., Chiang, M.Y., Xu, L., Shestova, O., Wang, H., Li, Y.M., Bhandoola, A., Aster, J.C., *et al.* (2010a). Notch dimerization is required for leukemogenesis and T-cell development. Genes Dev *24*, 2395-2407.

Liu, H., Chi, A.W., Arnett, K.L., Chiang, M.Y., Xu, L., Shestova, O., Wang, H., Li, Y.M., Bhandoola, A., Aster, J.C., *et al.* (2010b). Notch dimerization is required for leukemogenesis and T-cell development. Genes Dev *24*, 2395-2407.

Liu, Z., Brunskill, E., Varnum-Finney, B., Zhang, C., Zhang, A., Jay, Y.P., Bernstein, I., Morimoto, M., and Kopan, R. (2015). The intracellular Domains of Notch1 and 2 Are Functionally Equivalent During Development and Carcinogenesis. Development *142*, 2452-2463.

Liu, Z., Chen, S., Boyle, S., Ilagan, M.X., Zhu, Y., Zhang, A., and Kopan, R. (2013). The Extracellular Domain of Notch2 Increases its Cell surface Abundance and Ligand Responsiveness During Kidney. Dev Cell *25*, 585-598.

Maillard, I., Weng, A.P., Carpenter, A.C., Rodriguez, C.G., Sai, H., Xu, L., Allman, D., Aster, J.C., and Pear, W.S. (2004). Mastermind critically regulates Notch-mediated lymphoid cell fate decisions. Blood *104*, 1696-1702.

Masek, J., and Andersson, E.R. (2017). The developmental biology of genetic Notch disorders. Development *144*, 1743-1763.

Mathern, D.R., Laitman, L.E., Hovhannisyan, Z., Dunkin, D., Farsio, S., Malik, T.J., Roda, G., Chitre, A., Iuga, A.C., Yeretssian, G., et al. (2014). Mouse and human Notch-1 regulate mucosal immune responses. Mucosal Immunol 7, 995-1005.

Moats, C.R., Baxter, V.K., Pate, N.M., and Watson, J. (2016). Ectoparasite Burden, Clinical Disease, and Immune Responses throughout Fur Mite (Myocoptes musculinus) Infestation in C57BL/6 and Rag1(-/-) Mice. Comp Med *66*, 197-207.

Moolenbeek, C., and Ruitenberg, E.J. (1981). The "Swiss roll": a simple technique for histological studies of the rodent intestine. Lab Anim 15, 57-59.

Moran, S.T., Cariappa, A., Liu, H., Muir, B., Sgroi, D., Boboila, C., and Pillai, S. (2007). Synergism between NF-kappa B1/p50 and Notch2 during the development of marginal zone B lymphocytes. J Immunol *179*, 195-200.

Nam, Y., Sliz, P., Pear, W.S., Aster, J.C., and Blacklow, S.C. (2007). Cooperative assembly of higher-order Notch complexes functions as a switch to induce transcription. Proc Natl Acad Sci U S A *104*, 2103-2108.

Nandagopal, N., Santat, L.A., LeBon, L., Sprinzak, D., Bronner, M.E., and Elowitz, M.B. (2018). Dynamic Ligand Discrimination in the Notch Signaling Pathway. Cell *172*, 869-880 e819.

Nellesen, D.T., Lai, E.C., and Posakony, J.W. (1999). Discrete enhancer elements mediate selective responsiveness of enhancer of split complex genes to common transcriptional activators. Developmental Biology *213*, 33-53.

Nigam, V., and Srivastava, D. (2009). Notch1 represses osteogenic pathways in aortic valve cells. J Mol Cell Cardiol *47*, 828-834.

Noah, T.K., and Shroyer, N.F. (2013). Notch in the intestine: regulation of homeostasis and pathogenesis. Annu Rev Physiol *75*, 263-288.

Obata, Y., Takahashi, D., Ebisawa, M., Kakiguchi, K., Yonemura, S., Jinnohara, T., Kanaya, T., Fujimura, Y., Ohmae, M., Hase, K., *et al.* (2012). Epithelial cell-intrinsic Notch signaling plays an essential role in the maintenance of gut immune homeostasis. J Immunol *188*, 2427-2436.

Okamoto, R., Tsuchiya, K., Nemoto, Y., Akiyama, J., Nakamura, T., Kanai, T., and Watanabe, M. (2009). Requirement of Notch activation during regeneration of the intestinal epithelia. Am J Physiol Gastrointest Liver Physiol *296*, G23-35.

Ong, C., Cheng, H., Chang, L.W., Ohtsuka, T., Kageyama, R., Stormo, D.G., and Kopan, R. (2006). Target selectivity of vertebrate Notch proteins: collaboration between discrete domains and CSL binding site architecture determine activation probability. J Biol Chem *281*, 5106-5119.

Pellegrinet, L., Rodilla, V., Liu, Z., Chen, S., Koch, U., Espinosa, L., Kaestner, K.H., Kopan, R., Lewis, J., and Radtke, F. (2011). Dll1- and Dll4-mediated Notch signaling is required for homeostasis of intestinal stem cells. Gastroenterology *140*, 1230-1240.

Peveling-Oberhag, J., Wolters, F., Doring, C., Walter, D., Sellmann, L., Scholtysik, R., Lucioni, M., Schubach, M., Paulli, M., Biskup, S., *et al.* (2015). Whole exome sequencing of microdissected splenic marginal zone lymphoma: a study to discover novel tumor-specific mutations. BMC Cancer *15*, 773.

Piris, M.A., Onaindia, A., and Mollejo, M. (2017). Splenic marginal zone lymphoma. Best Pract Res Clin Haematol *30*, 56-64.

Pochanke, V., Hatak, S., Hengartner, H., Zinkernagel, R.M., and McCoy, K.D. (2006). Induction of IgE and allergic-type responses in fur mite-infested mice. Eur J Immunol *36*, 2434-2445.

Pui, J.C., Allman, D., Xu, L., DeRocco, S., Karnell, F.G., Bakkour, S., Lee, J.Y., Kadesch, T., Hardy, R.R., Aster, J.C., *et al.* (1999). Notch1 expression in early lymphopoiesis influences B versus T lineage determination. Immunity *11*, 299-308.

Punareewattana, K., Smith, B.J., Blaylock, B.L., Longstreth, J., Snodgrass, H.L., Gogal, R.M., Jr., Prater, R.M., and Holladay, S.D. (2001). Topical permethrin exposure inhibits antibody production and macrophage function in C57Bl/6N mice. Food Chem Toxicol *39*, 133-139.

Quigley, D.A., To, M.D., Kim, I.J., Lin, K.K., Albertson, D.G., Sjolund, J., Perez-Losada, J., and Balmain, A. (2011). Network analysis of skin tumor progression identifies a rewired genetic architecture affecting inflammation and tumor susceptibility. Genome Biol *12*, R5.

Quigley, D.A., To, M.D., Perez-Losada, J., Pelorosso, F.G., Mao, J.H., Nagase, H., Ginzinger, D.G., and Balmain, A. (2009). Genetic architecture of mouse skin inflammation and tumour susceptibility. Nature *458*, 505-508.

Radtke, F., Fasnacht, N., and Macdonald, H.R. (2010). Notch signaling in the immune system. Immunity 32, 14-27.

Radtke, F., Wilson, A., Stark, G., Bauer, M., van Meerwijk, J., MacDonald, H.R., and Aguet, M. (1999). Deficient T cell fate specification in mice with an induced inactivation of Notch1. Immunity 10, 547-558.

Ray, A., and Dittel, B.N. (2010). Isolation of mouse peritoneal cavity cells. Journal of visualized experiments: JoVE.

Riccio, O., van Gijn, M.E., Bezdek, A.C., Pellegrinet, L., van Es, J.H., Zimber-Strobl, U., Strobl, L.J., Honjo, T., Clevers, H., and Radtke, F. (2008). Loss of intestinal crypt progenitor cells owing

to inactivation of both Notch1 and Notch2 is accompanied by derepression of CDK inhibitors p27(Kip1) and p57(Kip2). EMBO Rep 9, 377-383.

Rossi, D., Trifonov, V., Fangazio, M., Bruscaggin, A., Rasi, S., Spina, V., Monti, S., Vaisitti, T., Arruga, F., Fama, R., et al. (2012). The coding genome of splenic marginal zone lymphoma: activation of NOTCH2 and other pathways regulating marginal zone development. J Exp Med 209, 1537-1551.

Saito, T., Chiba, S., Ichikawa, M., Kunisato, A., Asai, T., Shimizu, K., Yamaguchi, T., Yamamoto, G., Seo, S., Kumano, K., *et al.* (2003). Notch2 is preferentially expressed in mature B cells and indispensable for marginal zone B lineage development. Immunity *18*, 675-685.

Santos, T.S.D., Tavares, R.S., and Farias, D.L.C. (2017). Splenic marginal zone lymphoma: a literature review of diagnostic and therapeutic challenges. Rev Bras Hematol Hemoter *39*, 146-154.

Schroeter, E.H., Kisslinger, J.A., and Kopan, R. (1998). Notch-1 signalling requires ligand-induced proteolytic release of intracellular domain. Nature 393, 382-386.

Severson, E., Arnett, K.L., Wang, H., Zang, C., Taing, L., Liu, H., Pear, W.S., Shirley Liu, X., Blacklow, S.C., and Aster, J.C. (2017). Genome-wide identification and characterization of Notch transcription complex-binding sequence-paired sites in leukemia cells. Sci Signal *10*.

Shaya, O., Binshtok, U., Hersch, M., Rivkin, D., Weinreb, S., Amir-Zilberstein, L., Khamaisi, B., Oppenheim, O., Desai, R.A., Goodyear, R.J., *et al.* (2017). Cell-Cell Contact Area Affects Notch Signaling and Notch-Dependent Patterning. Dev Cell *40*, 505-511 e506.

Shroyer, N.F., Helmrath, M.A., Wang, V.Y., Antalffy, B., Henning, S.J., and Zoghbi, H.Y. (2007). Intestine-specific ablation of mouse atonal homolog 1 (Math1) reveals a role in cellular homeostasis. Gastroenterology *132*, 2478-2488.

Siebel, C., and Lendahl, U. (2017). Notch Signaling in Development, Tissue Homeostasis, and Disease. Physiol Rev *97*, 1235-1294.

Sprinzak, D., Lakhanpal, A., LeBon, L., Santat, L.A., Fontes, M.E., Anderson, G.A., Garcia-Ojalvo, J., and Elowitz, M.E. (2010). Cis Interactions beween Notch and Delta generate mutally exclusive signaling states. Nature *465*, 86-90.

Stemmer, M., Thumberger, T., Del Sol Keyer, M., Wittbrodt, J., and Mateo, J.L. (2015). CCTop: An Intuitive, Flexible and Reliable CRISPR/Cas9 Target Prediction Tool. PLoS One *10*, e0124633.

Tanigaki, K., Han, H., Yamamoto, N., Tashiro, K., Ikegawa, M., Kuroda, K., Suzuki, A., Nakano, T., and Honjo, T. (2002). Notch-RBP-J signaling is involved in cell fate determination of marginal zone B cells. Nat Immunol *3*, 443-450.

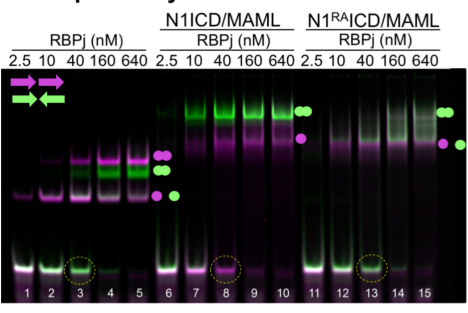
van Es, J.H., van Gijn, M.E., Riccio, O., van den Born, M., Vooijs, M., Begthel, H., Cozijnsen, M., Robine, S., Winton, D.J., Radtke, F., *et al.* (2005). Notch/gamma-secretase inhibition turns proliferative cells in intestinal crypts and adenomas into goblet cells. Nature *435*, 959-963.

Vooijs, M., Ong, C.T., Hadland, B., Huppert, S., Liu, Z., Korving, J., van den Born, M., Stappenbeck, T., Wu, Y., Clevers, H., *et al.* (2007). Mapping the consequence of Notch1 proteolysis in vivo with NIP-CRE. Development *134*, 535-544.

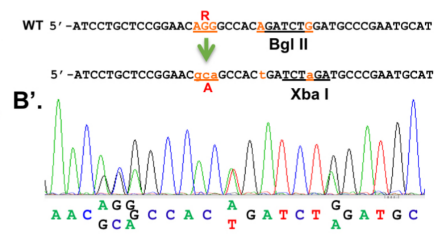
Weng, A.P., Ferrando, A.A., Lee, W., Morris, J.P., IV, Silverman, L.B., Sanchez-Irizarry, C., Blacklow, S.C., Look, A.T., and Aster, J.C. (2004). Activating Mutations of NOTCH1 in Human T Cell Acute Lymphoblastic Leukemia. Science *306*, 269-271.

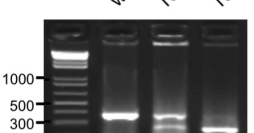
Witt, C.M., Won, W.J., Hurez, V., and Klug, C.A. (2003). Notch2 haploinsufficiency results in diminished B1 B cells and a severe reduction in marginal zone B cells. J Immunol *171*, 2783-2788.

## A. Cooperativity



## B. Notch1 → Notch1+/RA



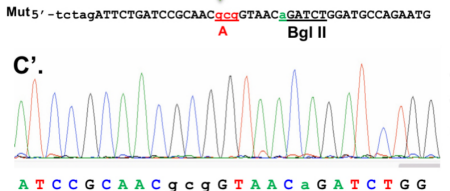


В".

100-

## C. Notch2 → Notch2<sup>RA/RA</sup>





· "RA

