EndoU is a novel regulator of AICD during peripheral B cell selection

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Balanced transmembrane signals maintain a competent peripheral B cell pool limited in self-reactive B cells that may produce pathogenic autoantibodies. To identify molecules regulating peripheral B cell survival and tolerance to self-antigens (Ags), a gene modifier screen was performed with B cells from CD22−/−C57BL/6 (CD22−/−[B6]) mice that undergo activation-induced cell death (AICD) and fail to up-regulate c-Myc expression after B cell Ag receptor ligation. Likewise, lysozyme auto-Ag–specific B cells in Igκ hen egg lysozyme (HEL) transgenic mice inherit the spleen but undergo AICD after auto–Ag encounter. This gene modifier screen identified EndoU, a single-stranded RNA–binding protein of ancient origin, as a major regulator of B cell survival in both models. EndoU gene disruption prevents AICD and normalizes c-Myc expression. These findings reveal that EndoU is a critical regulator of an unexpected and novel RNA-dependent pathway controlling peripheral B cell survival and Ag responsiveness that may contribute to peripheral B cell tolerance.

Molecules that regulate lymphocyte homeostasis, proliferation, and survival operate in concert to enable robust adaptive immune responses to foreign antigens (Ags). For the B cell lineage, the optimal outcome of these processes is a diverse antibody (Ab) repertoire purged of pathological (self-reactive) B cells. The elimination of pathological B cells occurs either through clonal deletion or receptor editing during B lymphopoiesis in the bone marrow, or in the periphery through the induction of energy (Goodnow et al., 1988; Nemazee and Bürki, 1989; Gay et al., 1993; Tiegs et al., 1993). Anergic B cells primarily inhabit the spleen, are short-lived, and undergo activation-induced cell death (AICD) in response to B cell Ag receptor (BCR) stimulation (Goodnow et al., 1995; Shlomchik, 2008). BCR ligation by agonistic anti-IgM Abs induces 30–50% of spleen B cells from WT mice to blast and undergo proliferation ex vivo (DeFranco et al., 1982). However, the threshold for B cell AICD can be influenced by genetically altering the stimulatory and inhibitory pathways that regulate BCR–induced activation (Inaoki et al., 1997). The B cell–restricted surface protein CD22 is generally considered to negatively regulate BCR signaling by recruiting potent intracellular phosphatases after BCR ligation (Doody et al., 1995; O’Keefe et al., 1996; Otipoby et al., 1996; Sato et al., 1996; Nitschke et al., 1996; Tedder et al., 1997; Poe et al., 2000), and CD22−/− mice produce augmented levels of isotype-switched auto–Abs against DNA and some protein Ags (O’Keefe et al., 1999; Poe et al., 2011). Nevertheless, B cells from inbred CD22−/− mice with a B6/129 genetic background (CD22−/−[B6]) are phenotypically and functionally normal ex vivo (Poe et al., 2004). In contrast, spleen B cells from C57BL/6 (B6) mice genetically deficient in CD22 (CD22−/−[B6]) undergo AICD after BCR stimulation (Poe et al., 2004), which is likely to be a result of their inability to induce c-Myc transcription factor expression that balances B cell proliferation versus AICD (Donjerković and Scott, 2000; Poe et al., 2004). These striking phenotypic differences in B cells between mouse lines with a common deletion of Cd22 indicate that important B cell signaling events that promote AICD are influenced differently by the B6 and

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129 genetic backgrounds. These two CD22−/− mouse lines were therefore used to identify genetic and molecular factors regulating B cell AICD.

In these studies, a forward genetic screen was used to identify an evolutionarily conserved single-stranded RNA (ssRNA) binding protein, EndoU, as a novel regulator of AICD in CD22−/− B6 mice. EndoU was also overexpressed by anergic peripheral B cells from double-transgenic mice expressing BCRs specific for hen egg lysozyme (HEL) along with soluble HEL (sHEL) as the cognate auto–Ag (Ig15sHEL mice; Goodnow et al., 1989; Hippen et al., 2000; Shlomchik, 2008). EndoU deficiency in Ig15sHEL mice also reversed AICD ex vivo and led to augmented anti-HEL auto-Ab responses in vivo. Thus, EndoU defines a new posttranscriptional regulatory pathway that controls B cell AICD, particularly in response to auto–Ag.

RESULTS
A genetic modifier locus/loci regulates BCR-induced AICD and CD5 expression
Spleen B cells from an inbred B6/129 founder line (CD22−/−[nub]), their WT littermates (WT[nub]), and WT B6 (WT[B6]) mice developed into blasts at normal frequencies and proliferated similarly after ex vivo BCR ligation using agonistic anti-IgM Abs (Fig. 1, A and B). In contrast, B cells from CD22−/− mice that were extensively backcrossed onto the B6 genetic background (CD22−/−[B6]) underwent AICD after BCR ligation. CD22−/−[B6] B cells also expressed CD5 after BCR stimulation but failed to up-regulate c-Myc transcript expression, whereas B cells from CD22−/−[nub] had normal CD5 and c-Myc expression (Fig. 1, C and D). Similarly, B cells from Ig15sHEL mice with a B6 background underwent AICD, expressed CD5, and failed to up-regulate c-Myc after ex vivo BCR stimulation (Fig. 1, E–G). In contrast, B cells from Ig15 mice (lacking the sHEL auto–Ag) blasted robustly, expressed CD5 at normal levels, and expressed high c-Myc levels after BCR ligation. These striking phenotypic similarities between CD22−/−[B6] and Ig15sHEL mice suggested that shared genetic and signaling programs cell B AICD.

CD22−/−[nub] mice were maintained as syngeneic brother/sister pairings for >8 yr, with genetic loci homozygous for either B6 germline DNA (B6:B6) or 129 germline DNA (129:129). To identify the genetic basis for the CD22−/−[B6] B cell phenotype, CD22−/−[B6] and CD22−/−[nub] mice were first crossed to produce an F1 generation (CD22−/−[B6]F1, Fig. 2A) with either homozygous B6 (B6:B6) or heterozygous B6 and 129 (B6:129) genetic loci. CD22−/−[B6] F1 B cell blast development after BCR stimulation was intermediate between B cells from CD22−/−[B6] and CD22−/−[nub] mice (unpublished data), indicating that maximal AICD of CD22−/−[B6] B cells required B6:B6 homozygosity at a critical locus (or loci). CD22−/−[F1] and CD22−/−[B6] mice were subsequently crossed to generate genetically distinct CD22−/−[N1] littermates due to independent chromosomal segregation and crossovers between regions of heterozygous B6:129 genomic DNA during gametogenesis. Remarkably, B cells from CD22−/−[N1] mice underwent AICD (Fig. 2 B, top dot plots) or developed into blasts at substantial

**Figure 1.** CD22−/−[B6] B cells are phenotypically similar to anergic Ig15sHEL B cells. (A) Representative B cell blast assay in which WT or CD22−/−[B6] spleen B cells purified from B6 (WT[B6] CD22−/−[B6]) or inbred B6:129 (WT[nub] CD22−/−[nub]) mice were cultured in medium alone or containing F(ab′)2 anti-IgM Abs. After 48 h, the cells were analyzed for viability (7-amino-actinomycin D [7AAD] exclusion) and cell size (forward scatter [FSC]) by flow cytometry. Percentages indicate viable B cell blasts (7AAD−FSC<median> ) within the gate shown. Results are representative of >10 mice of each genotype producing similar results. (B) The proliferation of purified spleen B cells from CD22−/−[nub] or CD22−/−[B6] mice, or their WT littermates, was measured by [3H]-thymidine incorporation. Values represent mean ±SEM cpm for triplicate wells from one of three independent experiments producing similar results. *, P < 0.05, Student’s t test. (C) Representative CD5 immunofluorescence staining of freshly isolated B220+ splenocytes from ≥5 mice/group of the indicated genotypes. Background staining using an isotype-matched control mAb is shown (Neg Ctrl). (D) Spleen B cells from WT[B6] or CD22−/−[B6] mice were cultured with F(ab′)2 anti-IgM Abs for 18 h. [3H]-radiolabeled cDNA was generated from total RNA, followed by hybridization to GEArray signaling pathway gene arrays. Regions of the arrays containing the c-Myc gene (arrowheads) and other representative genes expressed in B cells, including the c-Myc binding partner Max, are shown. (E) Mean ±SEM frequencies of B cell blasts in untreated or F(ab′)2 anti-IgM Ab-stimulated cultures (48 h) from four each Ig15 and Ig15sHEL mice assessed in independent experiments. **, P < 0.01, Student’s t test. (F) Representative CD5 expression by spleen B cells from Ig15sHEL mice and Ig15 mice as assessed by flow cytometry. A nonreactive isotype-matched Ab was used as a negative control (Neg Ctrl). (G) Purified spleen B cells from Ig15 or Ig15sHEL mice were cultured alone or with F(ab′)2 anti-IgM Abs. After 18 h, the cells were analyzed for c-Myc expression by intracellular staining, with flow cytometry analysis. Percentages of c-Myc+ B cells (those above the dashed line) are indicated in each plot. Bar graphs represent the mean ±SEM percentage of c-Myc+ B cells in BCR-stimulated cultures compared with unstimulated cultures from three mice of each genotype assessed in independent experiments. **, P < 0.01, Student’s t test.
A genetic locus regulates AICD of B cells from CD22<sup>−/−</sup>[B6] mice

Segregation of the B cell survival versus AICD phenotypes in CD22<sup>−/−</sup>[N1] mice indicated that the genomes of the parental CD22<sup>−/−</sup>[B6] mouse lines were stable and amenable to genetic analyses. Therefore, a forward genetic linkage analysis was used to identify the gene locus/loci responsible for CD22<sup>−/−</sup>[B6] B cell AICD. Genome-wide genotyping of 250 informative single nucleotide polymorphisms (SNPs) between the B6 and 129 mouse strains used genomic DNA from 44 CD22<sup>−/−</sup>[N1] mice; 22 mice had viable B cells after BCR ligation, whereas 22 mice had B cells that underwent AICD (Fig. 2 C).

Quantitative trait loci (QTL) mapping regression analysis revealed a single locus with high logarithm of odds (LOD) scores (P < 10<sup>−6</sup>) on the distal end of chromosome (Chr) 15 (Fig. 2 D), with four consecutive SNPs spanning the region from 68–91 megabase pairs (Mb, Fig. 2 E). LOD scores from the first three SNPs (68, 71, and 78 Mbp) were 8.3, increasing to 12.6 for the distal SNP (91 Mbp). The increased LOD score at 91 Mbp resulted from two mice (#14 and 28) with apparently discordant AICD phenotype mapped to the 94–98 Mbp region for all 44 CD22<sup>−/−</sup>[N1] mice typed in the original 44 N1 mice. Using this method, the primary genetic element responsible for CD22<sup>−/−</sup>[B6] gene location (Chr 7) is indicated. (Fig. 2 D, green squares indicating regions of and 91 Mbp with LOD scores of 8.3–12.6. Columns represent individual CD22<sup>−/−</sup>[N1] mice shown in C–F. All 103 mice (55 with viable B cells and 48 with apoptotic B cells) were then assessed at homozygosity. (F) Refined PCR-based SNP mapping of the Chr 15 locus of CD22<sup>−/−</sup>[N1] mice were identified as viable or apoptotic (top box), or insufficient in blast development (bottom box). Shown for comparison are the parental mouse B cells assessed in each of the assays as controls: CD22<sup>−/−</sup>[B6] B cells (apo) and CD22<sup>−/−</sup>[N1] B cells (viable, gray circles). (D) QTL mapping results from the 44 CD22<sup>−/−</sup>[N1] experimental mice shown in C, revealing a Chr 15 locus with a LOD score peaking at >12 (P < 0.000001, χ<sup>2</sup> statistic, MapManager QTX permutation test). The Cd22 gene location (Chr 7) is indicated. (E) Detailed heat map display of the Chr 15 mapping data shown in D. The locus of significance was defined by four consecutive SNPs positioned at 68, 71, 78, and 91 Mbp with LOD scores of 8.3–12.6. Columns represent individual CD22<sup>−/−</sup>[N1] mice, with blue squares indicating homozygous regions for mice #14 and #28. Mice having apparently discordant results confirmed the existence of a genetic modifier locus/loci that regulates BCR-induced AICD and CD5 expression.

For Figure 2. A Chr 15 locus segregates relative to CD22<sup>−/−</sup> B cell survival after BCR ligation. (A) Crossing CD22<sup>−/−</sup>[B6] mice with CD22<sup>−/−</sup>[N1] mice generated genetically similar F1 progeny (CD22<sup>−/−</sup>[F1]). CD22<sup>−/−</sup>[F1] mice were then backcrossed to CD22<sup>−/−</sup>[B6] mice to generate genetically dissimilar N1 littermates (CD22<sup>−/−</sup>[N1]) that were used for QTL mapping. (B) Spleen B cells from CD22<sup>−/−</sup>[N1] mice were assessed for CD5 expression before BCR stimulation and for blast development after BCR stimulation as in Fig. 1. B cells from some CD22<sup>−/−</sup>[N1] mice were anergic (were CD5<sup>high</sup> and underwent AICD [7AAD<sup>high</sup>, FSC<sup>low</sup>] after BCR stimulation; top), whereas others had a normal phenotype (were CD5<sup>low</sup> and developed into blasts [7AAD<sup>low</sup>, FSC<sup>high</sup>], bottom). Results are representative of those obtained from >40 CD22<sup>−/−</sup>[N1] littermates assessed in 10 independent experiments. (C) Survival analysis of BCR-stimulated B cells from CD22<sup>−/−</sup>[N1] littermates used for genomic SNP genotyping. Results are pooled from 10 independent CD22<sup>−/−</sup>[N1] mouse B cell blast assays (open squares), as described in B. Each value represents a single mouse, with dashed boxes distinguishing between CD22<sup>−/−</sup>[N1] littermates having B cells sufficient in blast development (top box), or insufficient in blast development (bottom box). Shown for comparison are the parental mouse B cells assessed in each of the assays as controls: CD22<sup>−/−</sup>[B6] B cells (apo) and CD22<sup>−/−</sup>[N1] B cells (viable, gray circles). (D) QTL mapping results from the 44 CD22<sup>−/−</sup>[N1] experimental mice shown in C, revealing a Chr 15 locus with a LOD score peaking at >12 (P < 0.000001, χ<sup>2</sup> statistic, MapManager QTX permutation test). The Cd22 gene location (Chr 7) is indicated. (E) Detailed heat map display of the Chr 15 mapping data shown in D. The locus of significance was defined by four consecutive SNPs positioned at 68, 71, 78, and 91 Mbp with LOD scores of 8.3–12.6. Columns represent individual CD22<sup>−/−</sup>[N1] mice, with blue squares indicating homozygous B6:B6 SNP loci and red squares indicating heterozygous B6:129 SNP loci. Representative genotypes of parental CD22<sup>−/−</sup>[B6] and CD22<sup>−/−</sup>[N1] mice are shown at the right, with green squares indicating regions of 129 homozygosity. (F) Refined PCR-based SNP mapping of the Chr 15 locus of CD22<sup>−/−</sup>[N1] mice at 1 Mbp intervals from 78 to 102 Mbp. Open squares represent SNPs that were not assessed. (G) B cells from an additional 59 CD22<sup>−/−</sup>[N1] mice were identified as viable or apoptotic in addition to the 44 CD22<sup>−/−</sup>[N1] mice shown in C–F. All 103 mice (55 with viable B cells and 48 with apoptotic B cells) were then assessed at 0.25–1 Mbp resolution by PCR-based SNP genotyping within the proximal (93.3 Mbp) to distal (98.3 Mbp) region of Chr 15.

To narrow the 78 to 102 Mbp Chr 15 locus, 23 additional SNPs within this region at 1 Mbp intervals were genotyped in the original 44 N1 mice. Using this method, the AICD phenotype mapped to the 94–98 Mbp region for all 44 CD22<sup>−/−</sup>[N1] mice (Fig. 2 F), including refined border regions for mice #14 and #28. Mice having apparently discordant
EndoU, a gene within the Chr 15 locus that encodes a novel ssRNA binding protein, is overexpressed in anergic B cells. (A) Relative transcript expression of the 28 genes within the Chr 15 locus in BCR-stimulated spleen B cells from CD22^{−/−} [B6] (closed bars) or CD22^{−/−} [inbr] parental mice. A single gene, EndoU (arrows), was differentially expressed by B cells from CD22^{−/−} and CD22^{−/−}[inbr] parental mice. (B) RNA was isolated from purified spleen B cells that were untreated (open bars) or were stimulated with F(ab')_{2} anti-IgM Abs for 18 h (closed bars), followed by real-time RT-PCR analysis of EndoU gene expression. Values represent mean (±SEM) fold changes in EndoU expression with Cd20 expression as the internal control gene in B cells from three mice of each genotype. All values were normalized relative to WT [129] B cells after BCR stimulation (dashed line). The diploid genotypes of each mouse line at the Chr 15 SNP-defined locus are indicated. (C) Purified spleen B cells or CD4^{+} T cells were cultured alone or with F(ab')_{2} anti-IgM Abs or mitogenic CD3 mAb, respectively, for 18 h. Total cellular proteins were separated by SDS-PAGE, followed by transfer to nitrocellulose membranes and Western blot analysis using polyclonal anti-EndoU Ab. The arrow indicates the position of the major EndoU band, which was dominant in CD22^{−/−} [B6] B cells. A nonspecific band (n.s.) is present in all lanes that is also observed when the secondary detection Ab is used alone as a control to probe CD22^{−/−} [B6] B cell lysates (far right lane). Membranes were reprobed with a polyclonal ERK2 Ab as a control for total protein loading. Bar graphs indicate relative B cell EndoU band densities (±SEM) from three each CD22^{−/−} [B6] and CD22^{−/−} [inbr] mice assessed in independent experiments. **, P < 0.01, Student's t test. (D) Purified spleen B cells from the indicated mice were cultured alone or with F(ab')_{2} anti-IgM Abs for 18 h. Total cellular proteins were separated by SDS-PAGE, followed by transfer to nitrocellulose membranes and Western blot analysis using polyclonal anti-EndoU Ab as described in C. Bar graphs indicate mean EndoU band densities from three each Ig^{Tg} and Ig^{Tg}sHEL mice as assessed in two independent experiments producing similar results. **, P < 0.01, Student's t test. (E) \(^{[32}P\)-labeled ssRNA substrates described for XendoU (Laneve et al., 2008) were incubated with recombinant EndoU protein in the presence or absence of Mn\(^{2+}\) or EDTA as indicated. Reactions were then separated by 15% PAGE and visualized by phosphorimager. The migration positions of EndoU-RNA complexes and of unbound RNA are indicated. Results are representative of those obtained in three experiments producing similar results. (F) c-Myc mRNA (sequence at left) shows an enrichment for poly(U) sequences. Regions of three or more consecutive uridines are highlighted. Bar graph at right shows the binding of in vitro transcribed, \(^{[32}P\)-labeled c-Myc mRNA by recombinant EndoU protein bound to IgG-coated magnetic beads as quantified by scintillation counting. A >100-fold molar excess of nonspecific protein (BSA) and RNA (tRNAs) were used as carriers in these assays. Background c-Myc RNA binding to magnetic beads without EndoU (Carrier) is shown. Bar graphs represent means (±SEM) from three binding assays per group. Results represent one of two experiments producing similar results. (G) NIH-3T3 cells were stably transfected with a plasmid encoding EndoU fused with CFP reporter, or with plasmid expressing a reporter protein alone as a control. Shown are representative histogram overlays of immunofluorescence staining of intracellular c-Myc protein in reporter-positive cells as determined by flow cytometry. The percent decrease in and p-value for the reduction of mean c-Myc MFI values (minus isotype control [Iso ctl] Ab staining) in EndoU-transfected cells (EndoU) relative to control-transfected cells (Vector) from four independent experiments are indicated. The significance level was determined using the Student's t test.

Abs for 18 h. Total cellular proteins were separated by SDS-PAGE, followed by transfer to nitrocellulose membranes and Western blot analysis using polyclonal anti-EndoU Ab as described in C. Bar graphs indicate mean EndoU band densities from three each Ig^{9} and Ig^{sHEL} mice as assessed in two independent experiments producing similar results. **, P < 0.01, Student's t test. (E) \(^{[32}P\)-labeled ssRNA substrates described for XendoU (Laneve et al., 2008) were incubated with recombinant EndoU protein in the presence or absence of Mn\(^{2+}\) or EDTA as indicated. Reactions were then separated by 15% PAGE and visualized by phosphorimager. The migration positions of EndoU-RNA complexes and of unbound RNA are indicated. Results are representative of those obtained in three experiments producing similar results. (F) c-Myc mRNA (sequence at left) shows an enrichment for poly(U) sequences. Regions of three or more consecutive uridines are highlighted. Bar graph at right shows the binding of in vitro transcribed, \(^{[32}P\)-labeled c-Myc mRNA by recombinant EndoU protein bound to IgG-coated magnetic beads as quantified by scintillation counting. A >100-fold molar excess of nonspecific protein (BSA) and RNA (tRNAs) were used as carriers in these assays. Background c-Myc RNA binding to magnetic beads without EndoU (Carrier) is shown. Bar graphs represent means (±SEM) from three binding assays per group. Results represent one of two experiments producing similar results. (G) NIH-3T3 cells were stably transfected with a plasmid encoding EndoU fused with CFP reporter, or with plasmid expressing a reporter protein alone as a control. Shown are representative histogram overlays of immunofluorescence staining of intracellular c-Myc protein in reporter-positive cells as determined by flow cytometry. The percent decrease in and p-value for the reduction of mean c-Myc MFI values (minus isotype control [Iso ctl] Ab staining) in EndoU-transfected cells (EndoU) relative to control-transfected cells (Vector) from four independent experiments are indicated. The significance level was determined using the Student's t test.

Figure 3. EndoU, a gene within the Chr 15 locus that encodes a novel ssRNA binding protein, is overexpressed in anergic B cells. (A) Relative transcript expression of the 28 genes within the Chr 15 locus in BCR-stimulated spleen B cells from CD22^{−/−} [B6] (closed bars) or CD22^{−/−}[inbr] (open bars) mice. A single gene, EndoU (arrows), was differentially expressed by B cells from CD22^{−/−} and CD22^{−/−}[inbr] parental mice. (B) RNA was isolated from purified spleen B cells that were untreated (open bars) or were stimulated with F(ab')_{2}, anti-IgM Abs for 18 h (closed bars), followed by real-time RT-PCR analysis of EndoU gene expression. Values represent mean (±SEM) fold changes in EndoU expression with Cd20 expression as the internal control gene in B cells from three mice of each genotype. All values were normalized relative to WT [129] B cells after BCR stimulation (dashed line). The diploid genotypes of each mouse line at the Chr 15 SNP-defined locus are indicated. (C) Purified spleen B cells or CD4^{+} T cells were cultured alone or with F(ab')_{2}, anti-IgM Abs or mitogenic CD3 mAb, respectively, for 18 h. Total cellular proteins were separated by SDS-PAGE, followed by transfer to nitrocellulose membranes and Western blot analysis using polyclonal anti-EndoU Ab. The arrow indicates the position of the major EndoU band, which was dominant in CD22^{−/−} [B6] B cells. A nonspecific band (n.s.) is present in all lanes that is also observed when the secondary detection Ab is used alone as a control to probe CD22^{−/−} [B6] B cell lysates (far right lane). Membranes were reprobed with a polyclonal ERK2 Ab as a control for total protein loading. Bar graphs indicate mean EndoU band densities (±SEM) from three each CD22^{−/−} [B6] and CD22^{−/−}[inbr] mice assessed in independent experiments. **, P < 0.01, Student's t test. (D) Purified spleen B cells from the indicated mice were cultured alone or with F(ab')_{2}, anti-IgM Abs for 18 h. Total cellular proteins were separated by SDS-PAGE, followed by transfer to nitrocellulose membranes and Western blot analysis using polyclonal anti-EndoU Ab as described in C. Bar graphs indicate mean EndoU band densities from three each Ig^{Tg} and Ig^{Tg}sHEL mice as assessed in two independent experiments producing similar results. **, P < 0.01, Student's t test. (E) \(^{[32}P\)-labeled ssRNA substrates described for XendoU (Laneve et al., 2008) were incubated with recombinant EndoU protein in the presence or absence of Mn\(^{2+}\) or EDTA as indicated. Reactions were then separated by 15% PAGE and visualized by phosphorimager. The migration positions of EndoU-RNA complexes and of unbound RNA are indicated. Results are representative of those obtained in three experiments producing similar results. (F) c-Myc mRNA (sequence at left) shows an enrichment for poly(U) sequences. Regions of three or more consecutive uridines are highlighted. Bar graph at right shows the binding of in vitro transcribed, \(^{[32}P\)-labeled c-Myc mRNA by recombinant EndoU protein bound to IgG-coated magnetic beads as quantified by scintillation counting. A >100-fold molar excess of nonspecific protein (BSA) and RNA (tRNAs) were used as carriers in these assays. Background c-Myc RNA binding to magnetic beads without EndoU (Carrier) is shown. Bar graphs represent means (±SEM) from three binding assays per group. Results represent one of two experiments producing similar results. (G) NIH-3T3 cells were stably transfected with a plasmid encoding EndoU fused with CFP reporter, or with plasmid expressing a reporter protein alone as a control. Shown are representative histogram overlays of immunofluorescence staining of intracellular c-Myc protein in reporter-positive cells as determined by flow cytometry. The percent decrease in and p-value for the reduction of mean c-Myc MFI values (minus isotype control [Iso ctl] Ab staining) in EndoU-transfected cells (EndoU) relative to control-transfected cells (Vector) from four independent experiments are indicated. The significance level was determined using the Student's t test.

This locus as harboring the genetic element controlling AICD, additional SNPs and a total of 103 CD22^{−/−}[N1] mice were analyzed (Fig. 2 G). Genotype and observed phenotypes were...
98% (101/103) concordant in the mice, with additional crossovers observed in CD22^{+/−} [N1] mice #21 and 64. Thus, the CD22^{+/−} [N1] survival locus mapped to a refined 5 Mbp region spanning from 93.3 to 98.3 Mbp on Chr 15.

**EndoU is overexpressed in B cells that undergo AICD**

The 93.3–98.3 Mbp region encodes 17 pseudogenes and 38 known genes, including 10 olfactory receptor genes not expressed in hematopoietic cells. Therefore, the differential expression of transcripts from these genes was compared using BCR-stimulated CD22^{−/−} [B6] and CD22^{+/−} [B6] B cells that remained similarly viable for 18 h. A single gene was differentially expressed between mouse lines, with 4.4-fold higher EndoU (formerly Ppt1r/Tol-30) levels in CD22^{−/−} [B6] B cells compared with CD22^{+/−} [B6] B cells (Fig. 3 A). B cells from WT [B6] mice expressed EndoU at lower levels than B cells from WT 129 (WT[129]) mice before and after BCR ligation (Fig. 3 B). EndoU transcript levels in CD22^{−/−} [B6] B cells remained uniquely higher relative to CD22^{+/−} [B6] mice and all other genotypes after BCR ligation, whereas CD22^{+/−} [B6] B cells expressed intermediate transcript levels. Thus, EndoU overexpression required CD22 deficiency in the context of a B6 genetic background.

To ensure that EndoU protein levels correlated with gene overexpression in CD22^{−/−} [B6] B cells, an affinity-purified rabbit polyclonal Ab specific for recombinant EndoU was generated. Western blot analysis of whole cell lysates confirmed increased EndoU expression in both resting and BCR-stimulated CD22^{−/−} [B6] B cells relative to CD22^{+/−} [B6] B cells (Fig. 3 C). EndoU overexpression in CD22^{−/−} [B6] mice was B cell specific, as EndoU levels were modest in activated spleen T cells from CD22^{+/−} [B6] and CD22^{−/−} [B6] mice. Remarkably, EndoU was also overexpressed in B cells from Ig<sup>κ</sup>κsHEL mice both before and after BCR ligation but was nearly undetectable in B cells from Ig<sup>κ</sup>κ luternate mice (Fig. 3 D). Thus, B cell EndoU overexpression required both a homozygous B6:B6 EndoU locus in CD22^{−/−} [B6] and Ig<sup>κ</sup>κsHEL mice that correlated with their characteristic AICD phenotypes.

**Mammalian EndoU is a uridine-directed ssRNA binding protein**

EndoU function in mammals is unknown, although EndoU transcripts are abundant in glucocorticoid-sensitive thymocytes undergoing apoptosis (Baughman et al., 1992). Highly conserved EndoU orthologs are found in many advanced and primitive species (see Fig. 8 A; Renzi et al., 2006), but EndoU paralogs derived from ancestral gene duplications do not exist in mice. The *Xenopus* ortholog of EndoU, XendoU, has been defined as an endonuclease targeting ssRNA molecules harboring polyuridine (poly(U)) regions (Renzi et al., 2006). Because EndoU substrates have not been defined in any species and EndoU-related molecules that could share redundant activities are not found in mice, the ability of EndoU to bind and/or process XendoU ssRNA substrates was examined. Recombinant EndoU protein specifically bound poly(U) ssRNAs, but not ssRNA species lacking these sites (Fig. 5 E). However, EndoU did not cleave these poly(U) ssRNAs, even

![Figure 4. Generation of EndoU-deficient (EndoU<sup>−/−</sup>) mice.](image-reference)

(A) EndoU gene targeting strategy. The EndoU gene contains nine exons. Positions encoding a predicted transmembrane (TM) domain, the somatomedin B-like (SMB) domain, and the XendoU superfamily domain are indicated. A targeting vector was generated that contained a neomycin-resistance gene (<sup>neo</sup>) introduced into two unique *BsrBI* restriction sites, which introduced an in-frame termination codon. A thymidine kinase gene (<sup>tk</sup>) was inserted at the 5′ end of the EndoU gene sequence. (B) Homologous recombination of the targeting vector into the EndoU gene of heterozygous and homozygous offspring expanded the endogenous 7.5 kb *HindIII* digested genomic DNA fragment into an 8.5 kb fragment as assessed by Southern blot. (C) Appropriate targeting of the EndoU gene was further confirmed by PCR analysis (primer locations shown in A), whereby a common forward primer (PCR1) was used in combination with reverse primers either adjacent to (PCR2) or within (PCR3) the <sup>neo</sup> gene. Results show amplification of genomic DNA from WT, heterozygous, and homozygous offspring. (D) Western blot analysis of thymocyte lysates confirmed the absence of EndoU protein in EndoU<sup>−/−</sup> mice. The presence of a nonspecific band (n.s.) as also described above is indicated. (E) The major EndoU protein band in CD22^{−/−} [B6] thymocytes (lane 2) migrates slower than the EndoU protein from CD22^{+/−} [B6] thymocytes (lane 3). A 1:1 mixture of lysates from CD22^{+/−} [B6] and CD22^{−/−} [B6] thymocytes generates a double band (lane 4). The presence of a nonspecific band (n.s.) also described above is indicated. (F) The difference in EndoU protein migration results from a nonsynonymous SNP in the second exon of the *EndoU* gene. EndoU protein of CD22^{−/−} [B6] and AU mice are identical in sequence, but different from that of CD22^{−/−} [B6] mice as indicated by nucleotide sequence shown and protein migration by SDS-PAGE. The presence of a nonspecific band (n.s.) as also described above is indicated.
in the presence of Mn\(^{2+}\), a condition favorable for XendoU endonuclease activity (Laneve et al., 2003). Mn\(^{2+}\) chelation did not interfere with EndoU binding of ssRNA molecules, as described for XendoU (Gioia et al., 2005). Thus, EndoU binds ssRNAs, likely through its predicted XendoU Superfamily domain (see Fig. 8 A).

Because c-Myc expression is reduced in CD22\(^{-/-}\)[B6] and Ig\(^{16}\)sHEL B cells after BCR ligation and c-Myc transcripts contain numerous stretches of poly(U) residues within the 3’ untranslated region (3’-UTR, Fig. 3 F), c-Myc mRNA interactions with EndoU were examined. Like the XendoU ssRNA substrate, c-Myc mRNA was also bound by recombinant EndoU (Fig. 3 F), but mRNA cleavage was not detectable (not depicted). Thus, c-Myc mRNA is a potential EndoU substrate, which may explain reduced c-Myc expression when B cells overexpress EndoU. Further supporting this notion, ectopic EndoU expression in mouse NIH-3T3 fibroblasts significantly reduced c-Myc protein levels (Fig. 3 G; 32% decrease, P < 0.05).

**EndoU deficiency reverses CD22\(^{-/-}\)[B6] B cell AICD**

Because EndoU expression is low in WT and CD22\(^{-/-}\)[B6] B cells after BCR ligation (Fig. 3, B and C), the effect of reduced EndoU expression on CD22\(^{-/-}\)[B6] B cell AICD was assessed. EndoU-deficient mice were generated by appropriately targeting the EndoU gene on B6 Chr 15 (Fig. 4, A–C). PCR-based SNP genotyping of genomic DNA from targeted mice confirmed that the targeted EndoU locus was B6 in origin. The chimeric mice were then bred onto a B6 genetic background to generate EndoU\(^{-/-}\)[B6] mice that were homozygous B6:B6 throughout the 93.3–98.3 Mbp Chr 15 region (unpublished data). The absence of EndoU protein in homozygous gene-targeted mice was confirmed by Western blot analysis, which identified an appropriately sized protein in WT mice (Fig. 4 D). EndoU from CD22\(^{-/-}\)[B6] B cells and thymocytes migrated more slowly on SDS-PAGE gels (~50 kD band) than EndoU from CD22\(^{-/-}\)[B6] thymocytes (~48 kD band, Figs. 3 C and 4 E), due to a nonsynonymous SNP present within EndoU exon 2 which switches a single Glu (B6 mice) to Lys (129 mice). This was verified using A/J mice that share the EndoU SNP sequence with 129 mice, with EndoU from both CD22\(^{-/-}\)[B6] and A/J mice migrating identically (Fig. 4 F). Splenocytes from CD22\(^{-/-}\)[B6] and CD22\(^{-/-}\)[A/J] mice expressed a single EndoU cDNA species as confirmed using 5’ RACE procedures (unpublished data). Thus, EndoU\(^{-/-}\)[B6] mice do not express EndoU protein.

EndoU\(^{-/-}\)[B6] and CD22\(^{-/-}\)[B6] mice were crossed to generate double-deficient (EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6]) progeny. A modest but significant reduction of total spleen B cell numbers is observed in CD22\(^{-/-}\)[B6] mice, which normalized in EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] mice (Fig. 5 A). AICD was also reversed in B cells from EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] mice with normal BCR-induced blast development (Fig. 5 B) and proliferation (Fig. 5 C). c-Myc expression was also normalized in BCR-stimulated EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] B cells relative to CD22\(^{-/-}\)[B6] B cells (Fig. 5, D and E). EndoU may preferentially regulate c-Myc expression in CD22\(^{-/-}\)[B6] B cells after BCR ligation because proliferating cell nuclear Ag (PCNA) was expressed at normal levels (Fig. 5 E), as reported for other molecules critical for cell cycle progression (Poe et al., 2004). The rescue of EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] B cells from AICD appeared to be B cell intrinsic because lymphoid tissue and thymic development and cellularity were normal in EndoU\(^{-/-}\) mice (unpublished data).

B cells from both CD22\(^{-/-}\)[B6] and EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] mice shared a surface IgM\(^{low}\) phenotype (Fig. 5 F) that results from enhanced BCR tonic signals in the absence of CD22 negative regulation (Doody et al., 1995; O’Keefe et al., 1996; Otipoby et al., 1996; Sato et al., 1996; Nitschke et al., 1997; Tedder et al., 1997). Otherwise, the modest reductions in surface IgD and CD21 expression that characterize CD22\(^{-/-}\)[B6] B cells (Poe et al., 2004) were normalized in EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] B cells (Fig. 5 F). Enhanced heat stable Ag (HSA/CD24) expression is commonly used as a marker for immature/transitional B cells, but as with CD5, enhanced HSA expression is also characteristic of chronically stimulated or anergic B cells (Noorchashm et al., 1999). Regardless, the characteristic CD5\(^{high}\)/HSA\(^{high}\) phenotype of CD22\(^{-/-}\)[B6] B cells (Poe et al., 2004) was largely normalized in EndoU\(^{-/-}\)CD22\(^{-/-}\)[B6] B cells (Fig. 5 F), and CD22\(^{-/-}\)[B6] mice have a normal ratio of mature to immature/transitional peripheral B cells (Poe et al., 2004). Thereby, elevated cell surface CD5 and HSA densities were phenotypic indicators for B cell AICD after BCR ligation.

**EndoU-deficient Ig\(^{16}\)sHEL mice produce auto-Abs**

A defining feature of Ig\(^{16}\)sHEL mice is the paucity of anti-HEL auto-Ab production. To examine EndoU function in this process, EndoU\(^{-/-}\)Ig\(^{16}\)sHEL mice were generated. Serum anti-HEL auto-Ab levels were strikingly high in some EndoU\(^{-/-}\)Ig\(^{16}\)sHEL mice, which were labeled “high responders” (Fig. 6 A). Anti-HEL auto-Ab levels in high-responder EndoU\(^{-/-}\)Ig\(^{16}\)sHEL mice were even greater than the levels of spontaneous anti-HEL Ab produced in Ig\(^{16}\) mice (1.9-fold increase, P = 0.02) and EndoU\(^{-/-}\)Ig\(^{16}\) mice (1.8-fold increase, P < 0.01) lacking auto-Ag. High-responder EndoU\(^{-/-}\)Ig\(^{16}\)sHEL mice were present in both sexes, and carried both the Ig\(^{16}\) and sHEL transgenes (Fig. 6 B). High serum anti-HEL auto-Ab levels developed in most high-responder mice after 5 wk of age and were maximal by 9–12 wk of age (Fig. 6 A, B and C). In addition to high-responder EndoU\(^{-/-}\)Ig\(^{16}\)sHEL mice, their age-matched littermates had augmented anti-HEL auto-Ab levels that were 2.4-fold higher than Ig\(^{16}\)sHEL mice (P = 0.01, Fig. 6 A). Anti-HEL auto-Ab levels in augmented-responder mice continued to increase with age (Fig. 6 D), whereas anti-HEL auto-Abs were not enhanced in old Ig\(^{16}\)sHEL mice (not depicted). Thus, EndoU overexpression regulates B cell auto-Ag responses in Ig\(^{16}\)sHEL mice.

**EndoU deficiency normalizes the phenotype of Ig\(^{16}\)sHEL mice**

Compared with Ig\(^{16}\)sHEL mice, high-responder EndoU\(^{-/-}\)Ig\(^{16}\)sHEL mice had a marked increase in B cell blast...
development and c-Myc expression after BCR ligation, increased spleen B cell numbers and surface IgM expression, and reduced CD5 and HSA expression (Fig. 6, E and F). In fact, high-responder mice were comparable to IgTg and EndoU−/− IgTg littermates lacking sHEL auto-Ag, except that surface IgM levels were modestly decreased. In contrast, augmented-responder EndoU−/− IgTg sHEL mice remained phenotypically similar to IgTg sHEL mice. Some B cells in augmented-responder mice exhibited surface IgM levels comparable to high-responder and IgTg B cells (Fig. 6 F, small peak), but otherwise their B cells had reduced IgM expression, modest blast development and c-Myc expression after BCR ligation, and high CD5 and HSA expression, with low spleen B cell numbers (Fig. 6, E and F). The frequency of IgMhigh B cells in augmented-responder mice (11% ± 2.1%) was significantly greater than the frequency of IgMhigh B cells present in IgTg sHEL mice (2.2 ± 0.5%, P < 0.01). Thus, the normalized phenotype of high-responder EndoU−/− IgTg sHEL mice and the IgMlow CD5high HSAhigh phenotype of augmented-responder EndoU−/− IgTg sHEL B cells suggests differential sensitivities to auto-Ag exposure.

Whether reduced B cell AICD led to the depletion of sHEL auto-Ag in EndoU−/− IgTg sHEL mice with age was examined. Blood B220+ cells isolated from IgTg and high-responder
EndoU−/−IgTβsHEL mice expressed relatively high surface IgMα levels and bound fluorescently labeled HEL protein at similar high levels (Fig. 7 A), suggesting that their BCRs were not occupied by sHEL. In contrast, B cells from both IgTβsHEL and augmented-responder EndoU−/−IgTβsHEL mice maintained an IgMα-low phenotype and did not bind labeled HEL protein, indicating BCR occupancy by sHEL. Dye-labeled spleen B cells from high-responder EndoU−/−IgTβsHEL mice were also adoptively transferred into WT or sHEL transgenic mice (Fig. 7 B). There was a dramatic reduction in IgM levels and HEL-binding capacity for high-responder B cells recovered from sHEL recipients in comparison with WT recipients (Fig. 7, B and C). Thus, the absence of AICD in high-responder EndoU−/−IgTβsHEL mice promoted auto-Ab production and endogenous sHEL clearance in vivo.

**DISCUSSION**

EndoU was identified in a gene modifier screen as a unique and potent regulator of B cell AICD. In WT [B6] mice, EndoU transcription and protein expression were suppressed by the balance between tonic BCR signals and their modulation by CD22, permitting normal BCR-mediated responses without blatant AICD after BCR ligation. In contrast, heightened BCR signals in peripheral B cells of CD22−/−[B6] mice drove EndoU overexpression and the CD5hiHEL− phenotype with predominant AICD after BCR ligation. Genetic deletion of EndoU expression reversed the CD5hiHEL− phenotype of B cells from CD22−/−[B6] mice and normalized AICD, confirming its important regulatory function in these processes. Bona fide EndoU substrates are likely to include c-Myc and other regulatory transcripts in vivo because c-Myc...
up-regulation is uniquely impaired in CD22−/−[B6] B cells (Poe et al., 2004). c-Myc expression was normalized in CD22−/−[B6] B cells by EndoU deficiency, EndoU bound c-Myc mRNA in vitro, and ectopic EndoU expression in a fibroblast cell line reduced c-Myc levels. A regulatory role for c-Myc in the mouse WEHI-231 B cell model of AICD has been previously described (Sonenshein, 1997; Donjerković and Scott, 2000). In CD22−/− mice, EndoU overexpression and AICD required B6 homozygosity at the EndoU locus. Therefore, genetic alterations must exist within the EndoU locus between the B6 and 129 genetic backgrounds that positively influences EndoU[B6] allele transcription in B cells, but not T cells. Although the EndoU locus has yet to be fine-mapped or sequenced in 129 mice, these important regulatory elements are likely to be located within the numerous stretches of conserved noncoding DNA present within the EndoU 5′ promoter region and 3′-UTR between B6 mice and humans (Fig. 8 C).

These collective observations support a model in which a CD22, EndoU, and c-Myc expression axis controls B cell fate after BCR ligation in B6 mice. EndoU levels remain low in WT[B6] B cells, allowing robust c-Myc expression and cell cycle progression after BCR engagement. In contrast, chronically high EndoU expression by CD22−/−[B6] B cells prevents c-Myc up-regulation after BCR ligation, resulting in AICD. EndoU was a major regulator of AICD in CD22−/−[B6] and Ig5+ψHEL mice, although additional unknown mechanisms undoubtedly also contribute to this complex regulatory process.

EndoU overexpression in vivo also occurred in response to BCR signals generated in Ig5+ψHEL mice, where B cells displayed a CD5highHSAhighIgMlow phenotype and failed to up-regulate c-Myc expression as was observed in CD22−/−[B6] mice. Genetic deletion of EndoU normalized this B cell phenotype in a substantial number of Ig5+ψHEL mice, which correlated with robust anti-HEL auto-Ab responses at an early age in mice that were defined as high responders. In augmented-responder EndoU−/−Ig5+ψHEL mice, some B cells expressed normalized phenotypes, suggesting that some B cells escape from an AICD phenotype with age to generate higher anti-HEL auto-Ab levels than Ig5+ψHEL mice. Given the link-age between EndoU, c-Myc up-regulation and AICD, it is likely that decreased B cell AICD in EndoU−/−Ig5+ψHEL mice leads to increased numbers of B cells that eventually consume endogenous HEL auto-Ag, permitting clonal expansion with a resultant increase in anti-HEL auto-Ab production.

The divergence of phenotypes between high-responder and augmented-responder mice has preceded based on previous studies. For example, Ig5+ψHEL mice deficient in the inositol phosphatase PTEN have reduced BCR occupancy by HEL and produce anti-HEL auto-Ab levels at a higher level than Ig5+ψHEL mice. This results in a subsequent break in tolerance in some mice caused by depletion of available HEL-auto-Ag (Browne et al., 2009). Likewise, Ig5+ψHEL mice transgenic for CD19 overexpression progressively break tolerance as they age and eventually produce high anti-HEL auto-Ab levels (Inaoki et al., 1997). In fact, remarkably similar to EndoU−/−Ig5+ψHEL mice, CD19+ψIg5+ψHEL mice develop as both high- and augmented-responder mice by 2 mo of age. Thus, modest increases in B cell numbers or enhancement in auto-Ab production resulting from altered B cell signaling can lead to auto-Ag depletion. Once this initial reduction in HEL levels occurs, the resulting lack of B cell anergy enables peripheral B cell numbers to be maintained at a level that exceeds HEL production, resulting in sustained and high-level anti-HEL auto-Ab production. Thereby, the EndoU regulatory pathway is likely to be an important component of B cell tolerance regulation, which is consistent with the known polygenic origins of autoimmunity.

The EndoU gene has an ancient origin, with conserved orthologs found in lower vertebrates (Suzuki et al., 2002; Gioia et al., 2005) and in invertebrates (Fig. 8, A and B).
Figure 8. **EndoU orthologs are conserved among diverse species.** (A) Protein sequences for EndoU and representative vertebrate orthologs were extracted from Ensembl or NCBI databases (human #ENST00000229003; mouse #ENSMUST00000023105; chicken #ENSGALT000000012388; opossum #ENSMO0100000012388; anole lizard #ENSAACAT00000014936; Xenopus tropicalis #XM_002935652.1; Fugu rubripes #ENSTRUT00000035070). Yellow highlighted residues with red font are conserved among all species. The consensus sequence at the bottom shows residues conserved in six out of seven species.

(B) Phylogenetic tree showing the evolutionary relationships among the orthologs of EndoU from various species.

(C) Graphical representation of amino acid substitutions and % homology for the 5' and 3' ends of the EndoU orthologs.
EndoU transcripts expressed in mouse B cells encoded a protein of 454 amino acids, which includes an N-terminal somatomedin B-like domain believed to be involved in protein–protein interactions (Gijssbers et al., 2003), a XendoU Superfamily ssRNA-binding domain, and a predicted transmembrane domain. EndoU specifically interacted with an ssRNA substrate harboring U-rich sequences that is also bound by XendoU, which is implicated in small nucleolar RNA processing in Xenopus (Gioia et al., 2005; Renzi et al., 2006). XendoU apparently generates snoRNAs through the cleavage of premiRNAs encoded within introns (Laneve et al., 2003). Human EndoU may exhibit low-level U-directed ssRNA endoribonuclease activity against the same in vitro ssRNA substrates recognized by XendoU (Laneve et al., 2008). However, mouse EndoU did not demonstrate measurable endonuclease activity against XendoU substrates. Mammalian EndoU may thus be functionally distinct from its Xenopus ortholog because human EndoU is excluded from the nucleolus of multiple cell lines (http://www.proteinatlas.org/ENSG00000111405/subcellular).

Viral orthologs of EndoU also exist (Fig. 8 B). Nsp15 (NendoU) is unique to the nidovirus family of ssRNA viruses, which includes the SARS coronavirus (Laneve et al., 2003; Bhardwaj et al., 2004; Ivanov et al., 2004; Gioia et al., 2005). Although Nsp15 is required for viral replication, specific Nsp15 substrates have not been identified and, surprisingly, appear to exclude the viral genome itself (Ivanov et al., 2004). Nsp15 has been postulated to suppress host immune responses (Ricagno et al., 2006) and to facilitate apoptosis in cells expressing the MAVS (mitochondrial anti-viral signaling) adapter protein that induces host anti-viral responses (Lei et al., 2009). Mouse hepatitis virus is also a member of the nidovirus family, is lymphotropic, and induces immunosuppression at least in part through the elimination of developing B cells (Lamontagne et al., 1989; Jolicoeur and Lamontagne, 1994). It therefore remains possible that mouse EndoU and hepatitis virus Nsp15 may target similar ssRNA substrates in B cells despite their evolutionary divergence. If true, each would have the capacity to suppress immune responses by inducing B cell AICD.

The current studies demonstrate for the first time that an RNA-binding protein plays a major role in regulating AICD, c-Myc expression, and the phenotype of peripheral B cells. Furthermore, these studies suggest an EndoU, CD22, c-Myc, and CD19 regulatory loop that directs normal B cell survival and malignant transformation (Fujimoto et al., 1999; Poe et al., 2012). EndoU likely regulates c-Myc and potentially other crucial ssRNA (miRNA or pre-miRNA) transcripts and nonclassical ssRNAs that directly regulate or fine-tune specific subsets of genes required for optimal B cell proliferation and survival. For autoreactive B cells that survive central selection and inhabit peripheral tissues, EndoU overexpression has nonetheless evolved as a checkpoint protein within the complex pathway that keeps potentially pathogenic B cells at bay through deletion via AICD.

**MATERIALS AND METHODS**

**Mice.** All procedures were approved by the Duke University Institutional Animal Care and Use Committee. CD22<sup>−/−</sup>[B6] and CD22<sup>−/−</sup>[inbr] mice were as previously described (Poe et al., 2004). EndoU<sup>−/−</sup>CD22<sup>−/−</sup>[B6] mice were produced by breeding EndoU<sup>−/−</sup> mice with CD22<sup>−/−</sup>[B6] mice, and then crossing heterozygous F1 animals to generate homozygous mice. WT littermates generated from the same F1 intercross served as controls. Ig<sup>κ</sup> and Ig<sup>κ</sup>/HEL mice (B6 background) were as previously described (Goodnow et al., 1989), with the Ig<sup>κ</sup>/HEL transgene confirmed using JAX Mouse Genotyping Protocols (The Jackson Laboratory).

To disrupt the EndoU gene by homologous recombination, an 8 kb DNA fragment containing exons 1–7 was isolated from a B6 genomic BAC clone (RP24-160P4). Flanking BstBI restriction sites were used to remove the distal end of exon 4, a portion of intron 4 and the splice donor site, which was replaced with a neomycin-resistance gene (neo) that introduced an intron in-frame termination codon. A thymidine kinase (tk) gene was inserted at the 5′ end of the EndoU targeting sequence. Appropriate homologous recombination in EFG1 (B6:129 hybrid) embryonic stem cells (Duke Transgenic Mouse Facility) was confirmed by Southern blot analysis of HindIII digested genomic DNA using a probe outside of the targeting sequence. The targeted allele generated an expected 8.5 kb band, compared with the WT 7.5 kb fragment. Correct EndoU gene targeting in subsequent offspring was verified by PCR analysis using a common forward primer (PCR1) in combination with reverse primers either 3′ of (PCR2) or within (PCR3) the inserted neo gene.

**B cell blast development and proliferation.** Purified spleen B cells (B cell isolation kit; Miltenyi Biotec) were analyzed by flow cytometry for blast development after 48 h of culture with F(ab′)<sub>2</sub> goat anti–mouse IgM Abs as previously described (Poe et al., 2004). Alternatively, B cell proliferation during 72-h cultures was assessed by λ-thymidine incorporation or CFSE labeling and dilution as previously described (Poe et al., 2004).

**Flow cytometry analysis.** Assessment of cell surface Ags was performed by flow cytometry using fluorescein-conjugated Abs as previously described (Poe et al., 2004). For intracellular c-Myc and PCNA expression, purified spleen B cells were cultured for 18 h in the presence or absence of F(ab′)<sub>2</sub> anti–mouse IgM Ab and then fixed and permeabilized using a Cytofix/Cytoperm Plus kit (BD). Intracellular protein staining used either c-Myc Ab (IgG1 clone 9E10; BioLegend), followed by R-PE-conjugated goat anti–mouse IgG1 secondary Abs (SouthernBiotech), or PCNA Ab (FITC-conjugated mouse IgG2a clone PC10; eBioscience). Immunofluorescence staining was assessed by flow cytometry with background staining levels determined using nonreactive isotype-matched, fluorescein-conjugated control Abs.

**SNP genotyping and QTL analysis.** CD22<sup>−/−</sup>[B6] mice were crossed with CD22<sup>−/−</sup>[inbr] mice to generate syngeneic F1 progeny (CD22<sup>−/−</sup>[F1]), CD22<sup>−/−</sup>[F1] mice were backcrossed to CD22<sup>−/−</sup>[B6] mice to generate CD22<sup>−/−</sup>[NI].
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littermates. Blast development after BCR ligation was assessed for purified spleen B cells from CD22-/-[N1] littermates, along with B cells from single parental CD22-/-[B6] and CD22-/-[inbr] mice as controls. The results of 10 independent experiments were pooled. To reduce variability between mice and experiments, CD22-/-[N1] mice B cell blast frequencies >1 SD above the mean percentage blasts were considered viable, whereas B cell blast frequencies <1 SD below the mean were considered nonviable. Genomic DNA from 22 viable and 22 nonviable CD22-/-[N1] mice was used for genome-wide genotyping of 250 relevant SNPs between B6 and 129 mice (Illumina BeadArray; Duke University Genotyping Facility). QTL mapping used Map Manager QTX regression analysis software (version QTXb20; Manly et al., 2001). Informative SNPs between B6 and 129 mice were identified using the Mouse Genome Informatics and dbSNP databases. DNA primer pairs were used to PCR amplify these SNPs and their surrounding sequences, which were sequenced (Duke Cancer Center DNA Analysis Facility) to identify single nucleotide (homozygous B6:B6) or double nucleotide (heterozygous B6:129) peaks at SNP sites.

Gene expression analysis. Spleen B cells from mice of the indicated genotypes were cultured in triplicate for 18 h in medium alone or with F(ab')2 anti-mouse IgM Ab, and then total RNA was isolated from pooled cells (R.Neasy Plus; Qiagen). For analysis of c-Myc and other signaling molecules, [32P]–radiolabeled cdNA was generated and hybridized to GEArray NF-kB signaling pathway gene arrays (SABiosciences) according to the manufacturer's instructions, with hybridization assessed using x-ray film. Gene expression within the Chr 15 locus was analyzed using Mouse 430.2 RNA arrays (Affymetrix; Duke Cancer Inst. DNA Analysis Facility) with GeneSpring GX software (Agilent Technologies) analysis. For real-time PCR analysis, cdNA was synthesized using random primers and analyzed using a LightCycler Instrument (software version 3) and FastStart DNA MasterPLUS SYBR Green 1 kit (Roche Diagnostics Corp), with EndoU expression quantified using EndoU exon 5 (forward) 5'-CGTCAAGAGAAGCTGTTCTCCAAG-3' and EndoU exon 6 (reverse) 5'-CCACATGTTCTTCAAATCGTCCAC-3' primers. Cd20 expression was the internal control. Relative EndoU expression was quantified by using the REST program (version 2).

Western blot analysis. Polyclonal antiserum reactive with EndoU was transfected (FuGENE HD; Promega) with a plasmid containing full length EndoU fused at the N terminal with CFP using the pEFP-C1 expression vector backbone (Takara Bio Inc.). Control-transfected reporter-positive NIH-3T3 cells were generated independently using the CFP (pEFP-C1) vector alone, or by co-transfection using the vector alone expressing GFP (pEGFP-C1), with similar results. Cells with stable plasmid incorporation were selected for appropriate reporter expression in the presence of 1 µg/ml Genetecin (Life Technologies).

ELISAs. Serum anti-HEL Ab levels in mice expressing the IgG3 transgene were measured using serum diluted 1:100 in Tris-buffered saline containing 1% BSA as previously described (Iwasaki et al., 1997).

Fluorescent labeling of HEL protein for flow cytometry analysis. Fluorescently labeled HEL protein (Sigma-Aldrich) was used to detect c-Myc surface expression levels and HEL–binding capacity.

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For XendoU substrates, the reactions were performed for 45 min in the presence or absence of 5 mM Mn2+ or 20 mM EDTA before analysis by 15% PAGE, with phosphorimager visualization. For c-Myc RNA, binding was assessed by scintillation counting.

To induce EndoU expression in mouse NIH-3T3 fibroblasts, cells were transfected (FuGENE HD; Promega) with a plasmid containing full length EndoU fused at the N terminus with CFP using the pEFP-N1 expression vector backbone (Takara Bio Inc.). Control-transfected reporter-positive NIH-3T3 cells were generated independently using the CFP (pEFP-N1) vector alone, or by co-transfection using the vector alone expressing GFP (pEGFP-C1), with similar results. Cells with stable plasmid incorporation were selected for appropriate reporter expression in the presence of 1 µg/ml Genetecin (Life Technologies).

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