Polζ Ablation in B Cells Impairs the Germinal Center Reaction, Class Switch Recombination, DNA Break Repair, and Genome Stability

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters.

Citation

Published Version
doi:10.1084/jem.20080669

Citable link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:4777439

Terms of Use
This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA
Polζ ablation in B cells impairs the germinal center reaction, class switch recombination, DNA break repair, and genome stability

Dominik Schenten,1,2 Sven Kracker,1,2 Gloria Esposito,3 Sonia Franco,1,2,4 Ulf Klein,3 Michael Murphy,1 Frederick W. Alt,1,2,4,5 and Klaus Rajewsky1,2,3

Polζ is an error-prone DNA polymerase that is critical for embryonic development and maintenance of genome stability. To analyze its suggested role in somatic hypermutation (SHM) and possible contribution to DNA double-strand break (DSB) repair in class switch recombination (CSR), we ablated Rev3, the catalytic subunit of Polζ, selectively in mature B cells in vivo. The frequency of somatic mutation was reduced in the mutant cells but the pattern of SHM was unaffected. Rev3-deficient B cells also exhibited pronounced chromosomal instability and impaired proliferation capacity. Although the data thus argue against a direct role of Polζ in SHM, Polζ deficiency directly interfered with CSR in that activated Rev3-deficient B cells exhibited a reduced efficiency of CSR and an increased frequency of DNA breaks in the immunoglobulin H locus. Based on our results, we suggest a nonredundant role of Polζ in DNA DSB repair through nonhomologous end joining.

© 2009 Schenten et al. This article is distributed under the terms of an Attribution–Noncommercial–Share Alike–No Mirror Sites license for the first six months after the publication date (see http://www.jem.org/misc/terms.shtml). After six months it is available under a Creative Commons License (Attribution–Noncommercial–Share Alike 3.0 Unported license, as described at http://creativecommons.org/licenses/by-nc-sa/3.0/).

CORRESPONDENCE
Klaus Rajewsky; rajewsky@idi.med.harvard.edu

Abbreviations used: AID, activation-induced deaminase; CSR, class switch recombination; DSB, double-strand break; ES, embryonic stem; FISH, fluorescence in situ hybridization; GC, germinal center; NHEJ, nonhomologous end joining; SHM, somatic hypermutation; SKY, spectral karyotyping; V, variable.

In T cell–dependent antibody responses, B cells are triggered to undergo a second round of antibody diversification in germinal centers (GCs) (1). Somatic hypermutation (SHM) introduces mutations into rearranged variable (V) regions of Ig genes allowing antibody affinity maturation (2, 3), whereas class switch recombination (CSR) exchanges the Ig constant (C) region to modify the effector function of the antibody (4). Both SHM and CSR rely on activation-induced deaminase (AID), an enzyme which deaminates cytidine residues in single-stranded DNA (5). The DNA deamination model of SHM suggests the conversion of G-C basepairs into G-U mismatches within the V region by AID (6, 7), which are subsequently processed in one of three ways: direct replication across the G-U mismatches results in G-C to A-T mutations; the removal of the uracil residues by the uracil degycosylaseUNG creates abasic sites, and DNA synthesis by error-prone DNA polymerases generates additional mutations; or the recognition of the G-U mismatches by the mismatch repair enzymes MSH2 and MSH6 leads to subsequent error-prone short-patch DNA synthesis, which introduces mutations outside the initial site of the lesion.

In the case of CSR, it is widely believed that upon cytidine deamination by AID, staggered DNA double-strand breaks (DSBs) are generated by the removal of the uracil residues by UNG, followed by the cleavage of the abasic sites by APE1/2 during the G1 phase of the cell cycle (5, 8–13). Alternatively, the mismatch-repair pathway can lead to the generation of staggered DSBs via the recognition of uracil by
MSH2 and MSH6 (8, 14, 15). The DSBs are then resolved by a process that includes DNA damage response proteins such as H2AX, MDC1, ATM, 53BP1, and the Nibrin–Mre11–Rad50 complex, the mismatch repair enzymes Pms2 and Mlh1, the exonuclease Exo1, and the classical and alternative nonhomologous end-joining (NHEJ) machinery (16–28).

Although the DNA polymerases required for filling in the staggered DNA breaks generated in CSR have not been identified, a possible involvement of several error-prone DNA polymerases in SHM has been tested using both hypermutating cell lines and KO mice deficient of these enzymes. From this work, Poλη, Poλθ, and perhaps Poλκ have emerged as important components of the SHM mechanism, whereas Poλβ, Poλκ, Poλλ, and Poλμ do not appear to play a significant role (29–35). Poλκ is an error-prone DNA polymerase that is characterized by its ability to extend mismatched primer-template termini (36). Poλκ, together with Poλλ, has been suggested as the prime example of the two-step inserter-extender model of translesion synthesis, in which a first DNA polymerase (Poλk) synthesizes across the DNA lesion and a second polymerase (Poλκ) extends the resulting mismatch (36). These features highlighted Poλκ as an additional candidate enzyme of the SHM machinery. Indeed, studies in a hypermutating cell line and a transgenic mouse strain that express antisense RNA against Rev3, the catalytic subunit of Poλκ, demonstrated a reduction of the frequency of somatic mutations in rearranged Ig V region genes, suggesting an involvement of Poλκ in SHM (37, 38).

Attempts to address this issue in vivo by genetically ablating Poλκ in mice have been hampered by the embryonic lethality observed upon deletion of the Poλκ gene in the mouse germ line (39–41). This embryonic lethality could be a result of the pronounced genomic instability observed upon Poλκ ablation in a wide variety of cellular systems (42), in turn suggesting a role of this enzyme in DNA repair and thus, potentially, CSR (16). To assess a possible contribution of Poλκ to SHM and CSR in vivo, we generated mice that carry a deletion of Rev3 selectively in mature B cells (subsequently called Poλκ+/CD21-cre mice). In this paper, we show that Poλκ-deficient B cells are impaired in their ability to proliferate and to maintain a stable genome. The mutant cells fail to undergo an efficient GC reaction and exhibit a reduced frequency of SHM and impaired CSR. The CSR defect is associated with an increased frequency of aberrantly or unrepaird DNA breaks within the IgH locus, suggesting that Poλκ plays a non-redundant role in DNA DSB repair through NHEJ.

RESULTS

Generation of a conditional Poλκ allele

This was achieved by flanking exon 2 of the Rev3 gene by loxP sites, to allow its deletion by Cre-mediated recombination (Fig. 1, A and B). Exon 2 is the second coding exon of Rev3, and messenger RNA splicing from exon 1 to exon 3 results in a frameshift mutation. Some Rev3 transcripts incorporate an additional exon downstream of exon 1. RNA splicing from this exon to exon 3 also results in a frameshift mutation (43). Two targeted embryonic stem (ES) cell clones were injected into blastocysts. Mice carrying the floxed Rev3 allele (subsequently called Poλκf/f) were viable and born at Mendelian ratios. The correct targeting of the Rev3 locus was confirmed by a nested PCR assay on tail DNA from Poλκf/f mice and wild-type controls. The PCR relied on a forward primer located outside the targeting vector to ensure the integration of the latter into the Rev3 locus, and a reverse primer located 3′ of the downstream loxP site. The amplification of both the wild-type and floxed alleles migrated at the expected size of ~5.2 kb (unpublished data). The PCR products were then gel purified and used as templates for two separate PCRs that amplified across the first and second loxP site, respectively. Each PCR amplified only one band for wild-type mice. In Poλκf/f mice, however, an additional band appeared in both cases whose larger size was consistent with the presence of an inserted loxP site (Fig. 1 C). Mice carrying the Poλκ allele were intercrossed with CD21-cre or deleter mice to generate Poλκ+/CD21-cre compound mutants, in which Poλκ function was specifically inactivated in mature B cells (44, 45). The deletion of exon 2 in the Poλκf allele was confirmed by gene amplification and subsequent sequencing of the PCR product (unpublished data).

Immune response and GC formation in Poλκ+/CD21-cre mice

No defects in B cell development were observed in Poλκ+/CD21-cre mice (unpublished data). Likewise, the number and proportions of resting B cells in the periphery was indistinguishable from those in control mice (unpublished data). We then characterized the antibody response of Poλκ+/CD21-cre mice against the T cell–dependent antigen NP-CG (3-hydroxy-nitrophenyl acetyl coupled to chicken γ-globulin). We injected Poλκ+/CD21-cre mice and control animals with 100 μg NP-CG in alum intraperitoneally and measured the serum titers of the NP-specific IgG1 and IgA antibodies at days 7 and 14 after immunization. At both time points, the response of Poλκ+/CD21-cre mice was reduced by 50% compared with that of control mice (unpublished data). Poλκ deficiency may affect cell viability of highly proliferating cells such as GC B cells. We therefore compared the proportions of CD19+ Fas+ PNA high GC B cells and CD19+ Fas+ PNA high GC B cells and CD19+ Fas+ PNA low naive B cells from control mice 14 d after immunization by flow cytometry and determined the deletion efficiency of the Poλκ allele in these cells by PCR. Competitive amplification of the Poλκ and Poλκ alleles and comparison to a standard curve with known ratios of the two alleles showed that ~90% of the PCR products in the CD19+ Fas+ PNAlow B cell
fraction and 80% in the CD19<sup>+</sup>Fas<sup>+</sup>PNA<sup>high</sup> B cell fraction were derived from the Pol<sup>−/−</sup> allele in Pol<sup>−/−</sup>/CD21-cre mice (Fig. 2B). This corresponds to a deletion efficiency of 80% in naive and 60% in GC B cells, as the mice were heterozygous for the Pol<sup>−/−</sup> allele. Thus, there was a moderate enrichment of cells that failed to inactivate Pol<sup>−</sup> in the GCs.

Reduced mutation frequency in Pol<sup>−/−</sup>/CD21-cre mice

To examine the impact of Pol<sup>−</sup> deficiency on SHM, we isolated naive and GC B cells from Pol<sup>−/−</sup>/CD21-cre and control mice that had been immunized with NP-CG 14 d earlier. As in the mutants, a substantial fraction of these cells had escaped Cre-mediated recombination. An analysis of VDJ joints amplified from populations of such cells is a priori compromised. Indeed, when we analyzed the accumulation of somatic mutations by bulk PCR using a primer pair that anneals in the framework region 3 of most V<sub>H</sub>J<sub>558</sub> gene segments and in the intron downstream of the J<sub>H</sub>4 gene segment (46), we found only a moderate reduction of the frequency of SHM in the mutants compared with the controls (Fig. S1A, available at http://www.jem.org/cgi/content/full/jem.20080669/DC1). 76 and 79% of the mutations in Pol<sup>−/−</sup>/CD21-cre mice occurred at...
A-T basepairs, compared with 56 and 65% in the controls. There was no difference in the ratio of transitions to transversions between mutants and controls (Fig. S1 B).

To approach the impact of Polζ deficiency on SHM at a higher level of resolution, we analyzed the accumulation of somatic mutations in Polζ+/CD21-cre mice at the level of single GC B cells, which we simultaneously genotyped for the inactivation of the Polζ gene. This approach offered an additional advantage in that it is not sensitive to a different rate of PCR errors across the individual samples. We isolated single GC B cells by flow cytometry and performed a PCR using a primer pair that anneals in the framework region 3 of most Vβ558 gene segments and in the intron downstream of the JH4 gene segment together with primer pairs that allow the genotyping of the Polζ and Polζ+ alleles. The amplification efficiency of both the Polζ and the Polζ+ allele in sorted GC Polζ+/CD21-cre B cells was 80%. We successfully amplified the rearranged JH4 gene segments in 66% of Polζ+/CD21-cre GC B cells from mice without the CD21-cre allele, in 42% of Polζ+/CD21-cre GC B cells from Polζ+/CD21-cre mice, and in 66% of Polζ+/CD21-cre GC B cells from Polζ+/CD21-cre mice. These differences in amplification efficiencies likely resulted from different extents of contamination of the sorted cells by non-B cells. The PCR products corresponding to the four introns downstream of the rearranged JH gene segments were sequenced and analyzed for the presence of somatic mutations. GC B cells that had ablated Polζ mutated

Figure 2. GC formation in Polζ+/CD21-cre mice. (A) Generation of GC B cells. Mice were immunized with 100 μg NP-CG in alum and analyzed for the presence of CD19+PNA+Fas+ GC B cells 14 d after immunization. Only CD19+ cells are shown. SPL, spleen; PP, Peyer’s patches; MLN, mesenteric lymph nodes. Data represent one of two independent experiments. (B) Deletion efficiency of the Polζ allele. The Polζ+ allele in B cells of Polζ+/CD21-cre and control mice were amplified in a competitive PCR using primers that anneal 5’, within, and 3’ of the floxed region and compared with a standard with known ratios of the two alleles. The fragments corresponding to the floxed and deleted allele migrate at around 300 and 450 bp, respectively.
their Ig genes with a frequency of 0.19% (Fig. 3 A). In contrast, the mutation frequency in GC B cells that had retained the Pol\(\gamma\) allele was 0.68%. Moreover, almost 50% of all sequences derived from Pol\(\gamma^{\Delta}\) cells contained only one mutation, whereas close to 80% of the sequences derived from Pol\(\gamma^{\Delta}\) cells had multiple mutations. Likewise, 38% of the sequences derived from Pol\(\gamma^{\Delta}\) cells remained unmutated.

To compare the pattern of somatic mutations in mutants and controls, we first established the mutation patterns in the individual introns downstream of each rearranged J\(H\) gene segment. Subsequently, we normalized the mutation patterns to the base composition of the respective intron and then combined the mutation patterns of all four introns. There was no significant difference in the mutation pattern between

**Figure 3. Single cell analysis of S\(\text{H}M\) in Pol\(\gamma^{\Delta}/\text{CD}21\)-cre mice.** (A) Distribution of mutations per B cell and mutation frequency in Pol\(\gamma^{\Delta}\)-deficient and Pol\(\gamma^{\Delta}\)-proficient B cells. Single GC B cells were sorted by FACS 14 d after immunization with NP-CG. PCR fragments containing the introns downstream of the rearranged VDJ elements were amplified in a seminested PCR and subsequently sequenced. Simultaneously, the single cells were genotyped for the presence of the Pol\(\gamma^{\Delta}\) allele. The mutation frequency was determined by the number of mutations present in the sequence of 500 bp, downstream of the individual VDJ rearrangement. (B) To compare the patterns of mutations of Pol\(\gamma^{\Delta}\)-deficient and -proficient cells, sequences derived from B cells using the same rearranged J\(\gamma\) element were grouped together and the mutations in each group were counted. The emerging mutation pattern in each group was corrected for the base composition of that particular intron before the patterns of the four J\(\gamma\) introns were compiled together. All values are shown in percentages and were rounded to the nearest whole number. n = the number of mutations; Tr./Tv., the transitions (Tr.) over transversions (Tv.) ratio. Shown are the combined mutation patterns corrected for the base composition of the sequences. (C) Percentage of mutations at A-T versus C-G basepairs in the same analysis.
Polζ and Polδ-α GC B cells. In Polζ-α cells, 62% of all mutations accumulated at A-T basepairs, which compared with 61% for Polδ-α cells (Fig. 3, B and C). Likewise, the ratio of transitions to transversions was not altered. We conclude from these data that Polζ deficiency has no impact on the pattern of SHM and that the differences in the A-T mutations observed in the bulk analysis must have been a result of differences in the fidelity of gene amplification between the experimental groups, a possibility which is excluded in the single cell analysis. With respect to the frequency of SHM, two interpretations offer themselves to explain the results: Polζ deficiency either directly impacts the SHM mechanism or it results in a rapid arrest of cellular proliferation preventing further SHM. The absence of a change of mutation pattern favors the latter interpretation.

Impaired CSR in Polζ-α/CD21-cre mice
To study a possible effect of Polζ deficiency on CSR, splenic B cells from Polζ-α/CD21-cre and control mice were stimulated with LPS in the presence or absence of IL-4, and the percentages of class-switched cells were measured by flow cytometry 3 or 4 d later. Compared with wild-type B cells, B cells from the mutant mice exhibited 2–3-fold reduced frequencies of cells switched to the expression of IgG1 or IgG3, respectively (Fig. 4 A and Table I). As CSR and cell proliferation are linked (47), we labeled Polζ-α/CD21-cre B cells and control cells with CFSE and measured their ability to divide. Proliferation of Polζ-α/CD21-cre B cells was reduced compared with controls (Fig. 4 A). Moreover, cell viability of Polζ-α/CD21-cre B cells dropped significantly 3 d after activation (unpublished data). Defective proliferation and decreased viability could explain the reduced frequency of Polζ-deficient cells switched to the expression of IgG1 and IgG3. However, the defect in CSR did not appear to be linked to the reduced ability of Polζ-α/CD21-cre B cells to class-switched cells was determined 4 d later by flow cytometry. Numbers in the graphs represent the percentage of switched cells. Numbers next to the red brackets below the graphs represent the percentage of class-switched cells among cells that have proliferated similarly, as indicated by the red brackets in the graphs. The histograms depict the proliferation of all cells in the culture. Black lines, Polζ-α+/ CD21-cre cells; red lines, Polζ-α−/ CD21-cre cells. Data represent one of three independent experiments.

(B) Germline transcription of the S regions in Polζ-α/CD21-cre mice and control mice. B cells were stimulated with LPS or LPS + IL-4 to undergo CSR in vitro for 2 d, after which germline transcripts of the Sμ, Sγ1, and Sγ3 regions and messenger RNA transcripts of the β-actin locus were amplified by RT-PCR. (C) Microhomologies at the junction between the Sμ and Sγ3 regions in IgG3+ Polζ-α−/CD21-cre B cells. B cells were stimulated with LPS to undergo CSR to IgG3+ for 3 d. After isolation of the DNA, the switch junctions were amplified by a nested PCR using primers that anneal in Sμ and Sγ3, subcloned, and sequenced. Black bars, Polζ-α+/CD21-cre cells (64 sequences); gray bars, Polζ-α−/CD21-cre cells (41 sequences). Statistical significance for changes in the frequency of sequences with insertions, no or short microhomologies (0–2 nt), and longer microhomologies (3–7 nt) in Polζ-α/CD21-cre cells, P = 0.04.
We then analyzed the sequences by BLAST alignment to the subcloned, and sequenced, excluding clonally related cells. Other S regions in class-switched cells (15, 19, 48, 49). PCR amplification and sequenced S genes involved in DNA repair, such as MSH2, PMS2, or Polβ, to test whether Polβ amplitudes and sequenced S genes. The percentages of class-switched cells were also 2–3-fold reduced when cells having undergone a similar number of divisions were analyzed for CSR (Fig. 4 A, gate indicated by red bracket). The latter result was independent of the location or width of the gate.

Germline transcription of the S regions is required for the initiation of CSR. Hence, we asked whether differences of germline transcription could account for the impairment of CSR in Polβ−/−/CD21-cre B cells. We stimulated splenic B cells with either LPS + IL-4 or LPS alone for 2 d and then amplified germline transcripts of Sμ, Sγ1, and Sγ3 by RT-PCR. Polβ−/−/CD21-cre and control B cells produced similar levels of Sμ, Sγ1, and Sγ3 transcripts (Fig. 4 B). To test whether Polβ is required for the repair phase of CSR, we amplified and sequenced Sμ–Sγ3 junctions from mutant and control cells, as it is known that the inactivation of some genes involved in DNA repair, such as MSH2, PMS2, or Polη, can lead to changes in the nature of the microhomologies or mutations at the junctions between the Sμ and Sγ3 or other S regions in class-switched cells (15, 19, 48, 49). PCR products between 500 and 800 nt were selected for analysis, subcloned, and sequenced, excluding clonally related cells. We then analyzed the sequences by BLAST alignment to the germline Sμ and Sγ3 sequences and counted the number of nucleotides at the S junctions that were identical to both the Sμ and Sγ3 region. The majority (53%) of the sequences derived from Polβ−/−/CD21-cre B cells displayed little (1–2 nt) or no microhomology at the S junctions, whereas a smaller fraction (21%) of sequences exhibited a longer stretch (3–7 nt) of nucleotide overlap (Fig. 4 C). The remaining fraction (25%) had insertions of 1 nt or more at the S junctions. Similar to the control cells, the majority of sequences (56%) derived from Polβ−/−/CD21-cre B cells contained no or only small microhomologies. In contrast, the frequency of sequences containing microhomologies of 3–7 nt was increased (34%), whereas the frequency of sequences containing insertions was reduced (10%). Insertions of 2 nt or longer were absent in sequences from the mutant B cells in contrast with control sequences. The changes in the nature of the Sμ–Sγ3 junctions, albeit modest, are consistent with a possible requirement for Polβ in the efficient processing of free DNA ends during CSR.

We also analyzed the switch junctions for somatic point mutations, which are known to occur in these regions through the recruitment of AID in the initiation of CSR (13, 50). In cells initiating but not completing CSR in the course of proliferation.

### Table I. General and IgH locus-specific genomic instability in stimulated Polβ−/−/CD21-cre B cells measured by FISH

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Cytokine stimulation</th>
<th>Percentage of IgG1+ cells on day 4</th>
<th>General genome instability</th>
<th>IgH locus-specific instability</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Number of metaphases analyzed</td>
<td>Number of chromosomal aberrations</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>30.3</td>
<td>30</td>
<td>1</td>
</tr>
<tr>
<td>CD21-cre</td>
<td>α-CD40 + IL-4</td>
<td>36.1</td>
<td>30</td>
<td>2</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>28.4</td>
<td>30</td>
<td>8</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>26.1</td>
<td>30</td>
<td>9</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>32.6</td>
<td>30</td>
<td>5</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>34.7</td>
<td>30</td>
<td>2</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>23.7</td>
<td>30</td>
<td>3</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>1.0</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40</td>
<td>0.1</td>
<td>28</td>
<td>1</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-RP105</td>
<td>0.2</td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>16.7</td>
<td>30</td>
<td>47</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>13.0</td>
<td>30</td>
<td>41</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>19.5</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>16.6</td>
<td>30</td>
<td>35</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>16.8</td>
<td>30</td>
<td>16</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40 + IL-4</td>
<td>18.0</td>
<td>30</td>
<td>19</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40</td>
<td>0.5</td>
<td>30</td>
<td>18</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40</td>
<td>0.1</td>
<td>20</td>
<td>10</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-CD40</td>
<td>0.1</td>
<td>23</td>
<td>1</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-RP105</td>
<td>0.1</td>
<td>30</td>
<td>29</td>
</tr>
<tr>
<td>Polβ+/+</td>
<td>α-RP105</td>
<td>0.1</td>
<td>30</td>
<td>31</td>
</tr>
</tbody>
</table>

Each row represents the results from one individual mouse.
proliferation, such mutations are introduced into Sμ but are rarely found in Sγ1 or Sγ3. It is believed that this explains the higher load of mutations in the Sμ parts of Sμ–Sγ1 or Sμ–Sγ3 switch junctions (24, 49, 50). Comparing the sequences of Sμ–Sγ1 junctions within a 200-bp window, we found that in wild-type cells the mutation frequencies in the Sμ portions of the junctions were indeed higher than in Sγ3 portions (0.53 and 0.27%, respectively), whereas in the mutants they were similarly low (0.21 and 0.22%, respectively; Table S1, available at http://www.jem.org/cgi/content/full/jem.20080669/DC1). The limited number of mutations did not allow a meaningful comparison of the mutation pattern. These results indicate that somatic mutations were equally introduced in mutant and control cells into switch regions undergoing switch recombination but that because of impaired proliferation and survival, the mutant cells had less opportunity to accumulate mutations in Sμ before switching.

Chromosomal aberrations in Polζ/CD21-cre B cells
Polζ had been previously shown to be required for genome stability in the DT40 chicken B cell line and mouse embryonic fibroblasts (51, 52). As genomic instability might explain the defective proliferation and CSR in LPS-activated Polζ-deficient B cells, as well as the counterselection of Polζ-deficient GC B cells, we stimulated B cells from two Polζf/CD21-cre and two control (Polζf/+ /CD21-cre) mice with LPS + IL-4 or LPS alone in vitro, prepared metaphase spreads 3 d later, stained the chromosomes with DAPI, and analyzed the metaphases by microscopy (Fig. 5 A). Chromosomal abnormalities were observed in 3% of the metaphases.

Figure 5. Chromosome stability in Polζ-deficient B cells. (A) Chromosome aberrations in B cells of Polζf/CD21-cre mice. B cells of Polζf/CD21-cre mice and control mice were activated to proliferate by the stimulation with LPS + IL-4 or LPS alone for 3 d. After induction of chromosome condensation and fixation of the cells, metaphase spreads were prepared and stained with DAPI for the visualization by fluorescence microscopy. Shown are the combined data of two mice per genotype. Bar, ~1 μm. (B) SKY of B cells of Polζf/CD21-cre mice. After stimulation of B cells with LPS + IL-4, metaphases were prepared, hybridized with suitable probes, and examined by fluorescence microscopy. Shown are the combined data of two mice per genotype. Red arrows indicate chromosome abnormalities. Bar, ~5 μm.
from B cells of the control mice in contrast with 39% of the metaphases from B cells of Polζ−/−/CD21-cre mice. We saw approximately equal frequencies of broken chromosomes and chromatids versus fused chromosomes in the affected metaphases of Polζ-deficient B cells. The occurrence of broken chromosomes and broken chromatids (which are present in mitotic cells as a result of unrepaired DSBs introduced before or after replication, respectively) suggested a DNA repair defect throughout the cell cycle (Fig. 5 A, Table S2 [available at http://www.jem.org/cgi/content/full/jem.20080669/DC1], and see Fig. 7). Spectral karyotyping (SKY) confirmed the general chromosomal instability detected in Polζ-deficient B cells (Fig. 5 B). Consistent with the previous analysis, only a modest number (5%) of metaphases of B cells from control mice contained irregular chromosomes, whereas metaphases containing broken chromosomes, chromosome fusions, and translocations were much more frequent (23%) in B cells of Polζ−/−/CD21-cre mice stimulated in vitro with LPS + IL-4 for 3 d. Even though Polζ-deficient metaphases were present with a broken chromosome 12, which carries the IgH locus in the mouse and where CSR occurs, other chromosomes (namely chromosomes 1, 3, and 11) were affected at a similar frequency. However, the sensitivity of either DAPI staining or SKY in the detection of CSR-specific breaks is limited, as the IgH locus is located in the subtelomeric region of chromosome 12, and such techniques may fail to distinguish an intact chromosome 12 from one lacking a small telomeric fragment as a result of a break within the IgH locus. We therefore proceeded to analyze CSR-specific chromosome breaks by fluorescence in situ hybridization (FISH) using IgH locus-specific probes.

**Increase of IgH locus-specific breaks in B cells of Polζ−/−/CD21-cre mice undergoing CSR**

In these experiments, we induced CSR in B cells of Polζ−/−/CD21-cre mice and of control mice by stimulation with an α-CD40 antibody in the presence of LPS + IL-4. This treatment efficiently induces CSR to IgG1 and allows comparison to stimulation with α-CD40 mAb alone, which activates the cells by itself and, in contrast with LPS, does not induce CSR. On day 4 after stimulation, metaphases were analyzed by FISH. Using a telomere-specific probe, we confirmed the increased general genome instability in Polζ−/−/CD21-cre mice in that we observed chromosomal breaks in B cells of the control mice in contrast with 39% of the metaphases from B cells of Polζ−/−/CD21-cre mice. We saw approximately equal frequencies of broken chromosomes and chromatids versus fused chromosomes in the affected metaphases of Polζ−/−/CD21-cre mice stimulated in vitro with LPS + IL-4 for 3 d. Even though Polζ-deficient metaphases were present with a broken chromosome 12, which carries the IgH locus in the mouse and where CSR occurs, other chromosomes (namely chromosomes 1, 3, and 11) were affected at a similar frequency. However, the sensitivity of either DAPI staining or SKY in the detection of CSR-specific breaks is limited, as the IgH locus is located in the subtelomeric region of chromosome 12, and such techniques may fail to distinguish an intact chromosome 12 from one lacking a small telomeric fragment as a result of a break within the IgH locus. We therefore proceeded to analyze CSR-specific chromosome breaks by fluorescence in situ hybridization (FISH) using IgH locus-specific probes.
inhibitors Xrcc4 and ligase 4 harbor predominantly chromosome lesions and may facilitate postreplication DNA repair via homologous recombination or translesion synthesis and mismatch extension (42, 51). However, the role of Polζ in the maintenance of genome stability has not been determined with precision. Many components of the NHEJ or DNA damage response machinery do not affect proliferation and SHM (24, 28, 54). B cells deficient of the classical NHEJ factors Xrcc4 and ligase 4 harbor predominantly chromosome breaks, which are indicative of failed DSB repair in the G1 phase of the cell cycle (23). In contrast, we found high frequencies of both chromosome and chromatid breaks in the Polζ-deficient B cells, indicating pre- and postreplication repair defects, respectively. It is thus also conceivable that the impaired proliferation and reduced cell viability of Polζ-deficient B cells are a result of a function of Polζ in NHEJ, translesion synthesis, and homologous recombination.

Genomic instability and impaired proliferative capacity of Polζ-deficient B cells are likely the main reasons for the impaired GC and IgG1 antibody response in our mutant animals, in which Polζ was selectively ablated in mature B cells. GCs were reduced in size, the frequency of somatic mutations introduced into V region genes was reduced, and CSR was impaired. We also observed counterselection of cells that had undergone deletion of the Rev3 gene in GCs. Analyzing the extent and pattern of SHM in Polζ-proficient and -deficient GC B cells by single cell PCR, we found a >3.5-fold reduction of the frequency of somatic mutations in the mutant cells but no change in the mutational pattern. Given that Polη has been identified as the sole polymerase responsible for mutations at A-T basepairs (55), a contribution of Polζ to SHM should have resulted in a change of the mutational pattern in Polζ-deficient B cells. In the absence of such a change, our results are consistent with the interpretation that the

under these conditions (Fig. 6, B and C; and Table I). We conclude that the increased frequency of chromosomal breaks in the IgH loci in B cells from Polζ/Δ/CD21-cre mice after α-CD40 mAb + IL-4 stimulation is a result of defective repair of CSR-induced DSBs.

DISCUSSION

Ubiquitous Polζ deficiency in the mouse is embryonic lethal (39–41). We have overcome this lethality by generating a conditional allele of Rev3 and used this allele to render mature resting CD21-expressing B cells deficient of Polζ. Using this system, we show that Polζ is not essential for B cell maintenance but is required for proper B cell proliferation, presumably because of its role in the maintenance of genome stability. The genomic instability and proliferation defect of Polζ-deficient B cells observed in the present study is reminiscent of the phenotype of Polζ-deficient chicken DT40 B cells, which also accumulate genomic aberrations and are impaired in their ability to proliferate (52). Genomic instability was also observed in Polζ-deficient mouse embryonic fibroblasts, which barely divide and require inactivation of p53 to proliferate (51). It has been speculated that Polζ may prevent the collapse of replication forks that have stalled at DNA lesions and may facilitate postreplication DNA repair via homologous recombination or translesion synthesis and mismatch extension (42, 51). However, the role of Polζ in the maintenance of genome stability has not been determined with precision. Many components of the NHEJ or DNA damage response machinery do not affect proliferation and SHM (24, 28, 54). B cells deficient of the classical NHEJ factors Xrcc4 and ligase 4 harbor predominantly chromosome breaks, which are indicative of failed DSB repair in the G1 phase of the cell cycle (23). In contrast, we found high frequencies of both chromosome and chromatid breaks in the Polζ-deficient B cells, indicating pre- and postreplication repair defects, respectively. It is thus also conceivable that the impaired proliferation and reduced cell viability of Polζ-deficient B cells are a result of a function of Polζ in NHEJ, translesion synthesis, and homologous recombination.

Genomic instability and impaired proliferative capacity of Polζ-deficient B cells are likely the main reasons for the impaired GC and IgG1 antibody response in our mutant animals, in which Polζ was selectively ablated in mature B cells. GCs were reduced in size, the frequency of somatic mutations introduced into V region genes was reduced, and CSR was impaired. We also observed counterselection of cells that had undergone deletion of the Rev3 gene in GCs. Analyzing the extent and pattern of SHM in Polζ-proficient and -deficient GC B cells by single cell PCR, we found a >3.5-fold reduction of the frequency of somatic mutations in the mutant cells but no change in the mutational pattern. Given that Polη has been identified as the sole polymerase responsible for mutations at A-T basepairs (55), a contribution of Polζ to SHM should have resulted in a change of the mutational pattern in Polζ-deficient B cells. In the absence of such a change, our results are consistent with the interpretation that the

53 versus 13% of all metaphases of B cells from Polζ/Δ/CD21-cre and control (Polζ+/+ /CD21-cre) mice, respectively (Fig. 6 and Table I). The chromosomal breaks in B cells from both Polζ/Δ/CD21-cre and control (Polζ+/+ /CD21-cre) mice consisted of approximately equal proportions of chromatid and chromosome breaks (Fig. 7 and Table S2). The results in Table S2 also raise the possibility that Polζ haploinsufficiency affects genome stability in cells activated to undergo CSR, but more data would be required to firmly establish this point.

To assay for breaks or translocations within the IgH locus, we used a FISH assay in which the first probe recognizes sequences just upstream of the IgH VH domain on the telomeric region of chromosome 12 (5′ IgH probe) and the second probe sequences immediately downstream of the C H locus (3′ IgH probe) (26). This assay revealed chromosomal breaks within the IgH locus, visualized as separated chromatid and Robertsonian chromosomes. See also Table S2 (available at http://www.jem.org/cgi/content/full/jem.20080669/DC1).

Figure 7. Types of genomic instabilities in stimulated Polζ/Δ/CD21-cre B cells measured by telomere-specific FISH. Purified B cells were stimulated with α-CD40 + IL-4 for 4 d. Subsequently, metaphase spreads were hybridized with a telomere probe to measure chromosomal aberrations. Shown is the mean number of chromosomal aberrations per 30 metaphases from Polζ/Δ/CD21-cre and control B cells plus the SD. cb, chromatid break; CB, chromosome break; other includes chromosomal rearrangements such as dicentric and Robertsonian chromosomes. See also Table S2 (available at http://www.jem.org/cgi/content/full/jem.20080669/DC1).

To ensure that the enhancement of IgH breaks in activated B cells from Polζ/Δ/CD21-cre mice after α-CD40 mAb + IL-4 stimulation is a result of defective repair of CSR-induced DSBs.

polζ /Δ/CD21-cre mice and control (Polζ+/+/ CD21-cre) mice, respectively (Fig. 6 and Table I). The chromosomal breaks in B cells from both Polζ/Δ/CD21-cre and control (Polζ+/+ /CD21-cre) mice consisted of approximately equal proportions of chromatid and chromosome breaks (Fig. 7 and Table S2). The results in Table S2 also raise the possibility that Polζ haploinsufficiency affects genome stability in cells activated to undergo CSR, but more data would be required to firmly establish this point.

To assay for breaks or translocations within the IgH locus, we used a FISH assay in which the first probe recognizes sequences just upstream of the IgH VH domain on the telomeric region of chromosome 12 (5′ IgH probe) and the second probe sequences immediately downstream of the C H locus (3′ IgH probe) (26). This assay revealed chromosomal breaks within the IgH locus, visualized as separated chromatid and Robertsonian chromosomes. See also Table S2 (available at http://www.jem.org/cgi/content/full/jem.20080669/DC1).

To ensure that the enhancement of IgH breaks in activated Polζ/Δ/CD21-cre B cells is indeed caused by CSR, we also stimulated B cells from Polζ/Δ/CD21-cre and control mice with an α-CD40 or α-RP105 mAb alone because such treatments induce proliferation but not CSR (53). Although B cells of Polζ/Δ/CD21-cre mice again accumulated chromosomal aberrations at a higher level than the control cells, IgH locus-specific breaks were not detected in either cell type
impairment of SHM by Polζ deficiency is caused by the im-
paired cell viability and consequent counterselection of the
mutant cells in the GC reaction.

In two earlier papers a direct involvement of Polζ in the
mechanism of SHM had been suggested (37, 38). In one ap-
proach, Diaz et al. (37) constitutively expressed antisense
RNA against Polζ in transgenic mice and observed compro-
mised SHM in the absence of a change of mutational pattern,
which is similar to the present data. Although these results
were difficult to interpret because of the non–B cell–autono-
mous expression of the transgenic antisense RNA and posi-
ble genomic instability also in that system, a study of SHM in
a Burkitt lymphoma cell line again demonstrated a reduced
frequency of somatic mutation upon transfection of the cells
with Rev3-specific antisense oligonucleotides, without an
effect on the mutation pattern (37, 38). Rev3 expression was
only partially inhibited in these experiments, and cell prolif-
eration was not detectably affected. It remains to be seen to
which extent these latter experiments reflect the require-
ments for SHM in GC B cells in vivo, where a direct partici-
pation of Polζ in its control clearly remains tentative.

In contrast with SHM, where our results suggest indirect
effects of Polζ deficiency, our data indicate a direct involve-
ment of Polζ in CSR. We observed a reduction in CSR effi-
ciency of Polζ-deficient B cells compared with control B
cells even when we compared cells that had gone through the
same number of cell divisions. Together with the observation
that germline transcription of the S regions remains unaf-
fected in Polζ/+/CD21-cre mice, this suggested an involve-
ment of Polζ in the repair phase of CSR, a notion which was
directly supported by the increased frequency of IgH locus-
specific DNA breaks in Polζ-deficient B cells undergoing
CSR in response to anti-CD40 stimulation in the presence of
IL-4. Our evidence clearly indicates that these breaks were
cause by a defective CSR process because they were absent in
Polζ-deficient B cells activated with α-CD40 or α-RP105
mAbs alone, an activation protocol which induces prolifer-
ation but not CSR. Among 1,200 chromosomes analyzed by
telomere FISH from individual α-CD40 + IL-4 cultures, 
3% of the chromosomes from Polζ-deficient B cells showed
abnormalities including chromatid breaks, chromosome breaks,
and chromosome fusions (Fig. 7, Table 1, and Table S2).
However, telomere FISH does not distinguish between
CSR-specific and –unspecific chromosomal aberrations. To
approach this problem, we determined the frequency of such
CSR-specific breaks in the activated Polζ-deficient cells
within the IgH locus on chromosome 12, where CSR oc-
curs. Based on the genome instability detected by telomere
FISH, we estimated that if chromosome 12 was targeted ran-
domly, three breaks in this chromosome should be found
among 50 metaphases and these breaks should be randomly
distributed along the chromosome. In contrast with the latter
prediction, we detected three to five IgH locus–specific chro-
mosome breaks in 50 metaphases (Table 1). Considering that
the IgH locus covers only ~3% of chromosome 12, this im-
plies that IgH locus-specific breaks in Polζ-deficient B cells
stimulated to undergo CSR are the result of a targeted rather
than a random process. In further support of this notion, we
failed to observe chromatid breaks at the IgH locus in meta-
phases of Polζ-deficient cells (in contrast with their frequent
occurrence at other loci), suggesting that, specifically within
the IgH locus, Polζ functions to repair a subset of DSBs re-
stricted to prereplicative stages of the cell cycle (G1) precisely
when AID is known to target S regions (13, 56). In principle,
an increased frequency of IgH locus–specific DNA breaks
could also result from defective V(D)J recombination. This
has been recently observed in ATM-deficient B cells, which
accumulate such breaks in an AID-independent and RAG-
dependent manner (53). Because the CD21-cre allele causes
the deletion of the floxed Polζ allele in mature but not devel-
oping B cells (44) and no IgH locus–specific DNA breaks
were detected in Polζ-deficient B cells stimulated with α-
CD40 or α-RP105 mAbs, such a possibility seems to be ex-
cluded in the present context.

CSR involves the generation of staggered DSBs during
the G1 phase of the cell cycle, which need to be processed
to create blunt DNA ends that serve as substrates for the alter-
native and classical NHEJ machinery (13, 23, 56–58). This
processing requires the activity of exonucleases and DNA
polymerases, with the former being provided by exonuclease 1
among perhaps other enzymes (20). The present results
suggest that Polζ catalyzes fill-in reactions in the course of
CSR–associated DSB repair by NHEJ. This is in line with
observations in Saccharomyces cerevisiae that ATR homologue
Mec1-dependent phosphorylation, which plays a major role
in DNA damage checkpoint responses, promotes the associa-
tion of Polζ/Rev1 with DSBs (59, 60). Our observation that
CSR is impaired in the absence of Polζ, yet not completely
abolished, points to the participation of additional polymer-
ases in the repair phase of CSR. These could include Polμ
and Polκ, both of which have been suggested to operate in
classical NHEJ (61, 62).

In normal B cells undergoing CSR, strict dosing of AID
and efficient repair by NHEJ prevents aberrant rejoining of
IgH locus–specific DSBs to DSBs within other loci. Dysregu-
lation of AID activity or deficiency of specific repair factors has
been shown to increase the frequency of oncogenic IgH–myc
translocations in in vitro cultured primary B cells undergoing
CSR (27). In this context, we report frequent chromosomal
translocations involving the IgH locus (and multiple uncharac-
terized partners) in primary Polζ-deficient B cells undergoing
CSR in vitro. We also observed a slight trend toward increased
genomic instability in Polζ/+/CD21-cre heterozygous B cells
compared with wild-type B cells (Table S2). Thus, genetic or
epigenic alterations that result in Polζ (haplo) insufficiency
might act in vivo to increase the frequency of oncogenic IgH
locus translocations.

MATERIALS AND METHODS
Targeting vector and generation of gene-targeted mice. A 25-kb
clon comprising 4 exons of rev3 was obtained by screening a P1 library
generated from C57BL/6 genomic DNA. The clone was used to construct
a targeting vector containing a loxP-flanked exon 2. The culture and transfection of C57BL/6-derived Bruce4 ES cells has been described previously. Homologous recombination was determined by Southern hybridization of probe A with EcoRI-digested genomic DNA or of probe B with BamHI-digested genomic DNA. The latter assay was also used to ensure the integration of the third loxP site. After homologous recombination, the neomycin resistance gene was removed in vitro by transfection of ES cells with the Cre recombinase–expressing plasmid pIC-Cre. Two independently targeted ES cell clones were injected into CB.20-derived blastocysts, which were implanted into pseudo-pregnant CB.20 foster mice to generate chimeric mice. Both clones transmitted the targeted allelle into the germline. The correct targeting was confirmed by a nested PCR. The first PCR round used the primers polz-1 and polz-2 [5'-GGAGCCCTTGAATTCTCATGCAGCTTTGACAGTCTTGCTCATGAGGCAG-3'], which anneal 5' of the region of homology and 3' of the second loxP site, respectively (Fig. 1B). Both the Polδ- and Polγ- allele yielded a PCR product of ~5.2 kb. This was followed by an amplification step of the gel-purified PCR products using primers that flank each loxP site (first loxP site: primers polz-3 [5'-GCACGTGTCAAGGTGAGGGATGATCAC-3'] and polz-4 [5'-TACCACTGATCCACTATTGGACCTGTTA-3'] and second loxP site: primers polz-5 [5'-CTCTGCACTATGTTAGGACACCTTTTGGAG-3'] and polz-6 [5'-CTATGTTACTCACCCTATTGCATCTCTGC-3']). The primers polz-1 and polz-2 were also used for the amplification of the Polδ- allele from tail DNA of Polδ-1 mice. Parts of the PCR product (3.2 kb) were sequenced using the primers polz-1, polz-3, and polz-6 to confirm the integration of the targeting vector into the Rev3 locus and the excision of exon 2.

Mice. Polδ-/- mice were intercrossed with deleter mice to obtain mice carrying a nonfunctional Polδ allele (Polδ-/- mice) (45). Subsequent intercrosses between Polδ-/- mice, Polδ-/- mice, and CD21-/- mice generated Polδ-/-/CD21-/- mice (44). The mice were kept on a C57BL/6 background. All mouse experiments were approved by the Institutional Animal Care and Use Committee of Harvard University and by the Immune Disease Institute.

Flow cytometry. Single-cell suspensions prepared from lymphoid organs were stained with the following monoclonal antibodies conjugated to FITC, PE, CyC, APC, or biotin: anti-CD3 (145-2C11), anti-CD4 (L3T4), anti-CD8 (53-67), anti-CD19 (ID3), anti-CD21, anti-CD23 (B3B4), anti-CD43 (57), and anti-Fas (Jo2; BD). Monoclonal antibodies against IgM (R3.3-2412) and B220 (RA3-6B2) were grouped and compared with cells from the same mouse that retained a floxed Polδ allele (Polδ-/-). To compare the pattern of somatic mutations, the mutations were analyzed in each JH intron and subsequently normalized according to the base composition of the individual JH introns. The isolation of the intron upstream of JH4, which is in the 5'-flanking region of the gene, was achieved by transfection of ES cells with the Cre allele. The deletion of the Polδ allele was determined with the primers zeta4 (5'-AAGAATACAGTTGAAAGGACGGCATCGG-3') and zeta5 (5'-CCATGAGGAGGCTTGATCTCAGGAGGC-3') detecting the Polδ allele. The IgH immunoglobulins were amplified with the primers J588Fr and JH4Hint-1 (5'-GGCTTCTCCTGACTATCAC-3'). The annealing temperature was 55°C followed by 5 cycles at 60°C and 3 cycles at 55°C. The second PCR reactions for the amplification of the Polδ alleles and JH introns were performed using anti-CD40 mAb (clone 11C10) as the initial PCR reaction mix. The deletion of the Polδ allele was determined with the primers zeta4 (5'-AAGAATACAGTTGAAAGGACGGCATCGG-3') and zeta5 (5'-CCATGAGGAGGCTTGATCTCAGGAGGC-3') detecting the Polδ allele. The IgH introns were directly sequenced with the primers J588Fr and JH4Hint-1 (5'.1.1, followed by C.1.147, G.1.104; T.0.87, intronic region JH1 [A.0.97; C.1.147, G.1.04; T.0.87], intronic region JH2 [A.1.15; C.0.94; G.1.06; T.0.88; intronic region JH3 [A.1.08; C.1.28; G.0.83; T.0.91], and intronic region JH4 [A.0.94; C.1.156; G.0.92; T.0.82]).

The analysis of SHM in GC B cells by bulk PCR has been published previously (32, 33). Splenic GC B cells were sorted into a naive fraction (CD19+PNA-CD4+CD8-), a GC fraction (CD19+PNA+CD4+CD8-), and a GC fraction (CD19+PNA+CD4+CD8-). PCR fragments corresponding to the intron downstream of JH4 were obtained from DNA purified cell populations using the primers J588Fr and JH4Hint-1 (46). PCR fragments containing VDJ rearrangements involving a JH4 element were selected for further analysis, subcloned, and sequenced. A stretch of 500 bp of intron sequence immediately downstream of the JH4 element was analyzed for somatic mutations.

Class switch analysis. B cells were purified from spleen single cell suspensions by MACS depletion using anti-CD43 microbeads (Miltenyi Biotec). The cells were stimulated with either 20 μg/ml LPS alone, 25 ng/ml IL-4 (R&D Systems), or 1 μg/ml anti-CD40 mAb (clone 11C10) for 5 days and then analyzed by flow cytometry. The preparation of metaphases has been described elsewhere. In brief, the cells were

Analysis of SHM. Single GC B cells (CD19+PNA+CD4+CD8-) were sorted by flow cytometry into 96-well plates and analyzed individually using a nested PCR for the genotyping of the cells and a semitone PCR for the amplification of DNA fragments corresponding to the introns downstream of the rearranged VDJH elements. After cell lysis with proteinase K in PCR buffer, the PCR was performed amplifying DNA fragments corresponding to the Polδ and Polγ alleles using the primers zeta1 (5'-ACCAAAAGAATCAGTTGAAAGGAGGGG-3'), zeta2 (5'-TTTAGATGTTGACAGTTGAAAGGAGGGG-3'), zeta3 (5'-TTTAGATGTTGACAGTTGAAAGGAGGGG-3'), and zeta6 (5'-TTGCTTCTTACTACAGAGTTGACAGTTGAAAGGAGGGG-3') and DNA fragments containing the introns downstream of the rearranged VDJH elements using the primers J588Fr (5'-CAGCCCTGACATCGGAGTCTCGC-3') and JH4-int2 (5'-GTCTGACCATGACAGTTGAAAGGAGGGG-3'), which anneal in the framework 3 region of most VDJH elements and the intron downstream of JH4, respectively (46). The annealing temperature was 55°C followed by 5 cycles at 60°C and 3 cycles at 55°C.

The second PCR reactions for the amplification of the Polδ alleles and JH introns were performed using anti-CD40 mAb (clone 11C10) as the initial PCR reaction mix. The deletion of the Polδ allele was determined with the primers zeta4 (5'-AAGAATACAGTTGAAAGGACGGCATCGG-3') and zeta5 (5'-CCATGAGGAGGCTTGATCTCAGGAGGC-3') detecting the Polδ allele. The IgH introns were directly sequenced with the primers J588Fr and JH4Hint-1 (5'-GGCTTCTCCTGACTATCAC-3'). The annealing temperature of the secondary PCR reactions was 58°C. DNA products corresponding to the JH introns were directly sequenced with the primers J588Fr and JH4Hint-1 (5'.1.1, followed by C.1.147, G.1.04; T.0.87, intronic region JH1 [A.0.97; C.1.147, G.1.04; T.0.87], intronic region JH2 [A.1.15; C.0.94; G.1.06; T.0.88; intronic region JH3 [A.1.08; C.1.28; G.0.83; T.0.91], and intronic region JH4 [A.0.94; C.1.156; G.0.92; T.0.82]).

The analysis of SHM in GC B cells by bulk PCR has been published previously (32, 33). Splenic GC B cells were sorted into a naive fraction (CD19+PNA-CD4+CD8-) and a GC fraction (CD19+PNA+CD4+CD8-). PCR fragments corresponding to the intron downstream of JH4 were obtained from DNA purified cell populations using the primers J588Fr and JH4Hint-1 (46). PCR fragments containing VDJ rearrangements involving a JH4 element were selected for further analysis, subcloned, and sequenced. A stretch of 500 bp of intron sequence immediately downstream of the JH4 element was analyzed for somatic mutations.

Class switch analysis. B cells were purified from spleen single cell suspensions by MACS depletion using anti-CD43 microbeads (Miltenyi Biotec). The cells were stimulated with either 20 μg/ml LPS alone, 25 ng/ml IL-4 (R&D Systems), or 1 μg/ml anti-CD40 mAb (clone 11C10) for 5 days and then analyzed by flow cytometry. The preparation of metaphases has been described elsewhere. In brief, the cells were

Cyto genetic analysis. MACS-purified B cells were stimulated with 20 μg/ml LPS and 25 ng/ml IL-4 and cultured for 3 d, at which time 50 ng/ml cocolcemid (KaryoMAX Solution; Invitrogen) was added to the medium for 2 h. The preparation of metaphases has been described elsewhere. In brief, the cells were
were swelled in 0.4% potassium chloride solution for 10 min at 37°C, incubated four times in 3:1 methanol/acidic acid solution, dropped onto microscope slides, and subsequently passed through hot steam and air dried. The metaphases were either covered with DAPI-containing mounting medium (Vectorshield; Vector Laboratories) or further processed for SKY using the Sky-Paint DNA kit (Applied Spectral Imaging) according to the manufacturer’s instructions. The analyses were performed on a microscope (Eclipse; Nikon) using a black and white 12-bit charge-coupled device camera (Applied Spectral Imaging) and a 63× objective lens. The cytogenetic analysis by FISH has been published elsewhere (26). In brief, the cells were stimulated with 1 μg/ml of an α-CD40 mAb (clone HM1-3; BD), 2.5 μg/ml of an α-CD180 mAb (anti-CD180; clone RP/14; eBioscience), or an α-CD40 mAb and 25 ng/ml IL-4 for 4 d. Subsequently, metaphases were prepared as described and hybridized with the fluorescence-labeled BAC207 and BAC199 for the detection of the 5′ and 3′ ends of the IgH locus, respectively. The telomeres were stained with an Cy3-conjugated (TTagGg), PNA probe (Applied Biosystems).

Online supplemental material. Fig. S1 contains the analysis of SHM in GC B cells of Polκ−/−CD21−/− or control mice by bulk PCR. Table S1 shows the frequency of mutations in Sp and Sy3 regions near Spα-Sy3 junctions in Polκ-deficient and control B cells, and Table S2 describes the types of genomic abnormalities observed in stimulated Polκ-deficient and control B cells. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20080669/DC1.

We thank Angela Egert, Anthony Monti, Margaret Curnutte, and Victoria Dreier for expert mouse work. We also appreciate the helpful advice and comments of Sergei Neuberger, M.S., R.S. Harris, J. Di Noia, and S.K. Petersen-Mahrt. We thank Angela Egert, Anthony Monti, Margaret Curnutte, and Victoria Dreier for expert mouse work. We also appreciate the helpful advice and comments of Sergei Neuberger, M.S., R.S. Harris, J. Di Noia, and S.K. Petersen-Mahrt.

Submitted: 31 March 2008
Accepted: 21 January 2009

REFERENCES

et al. 2006. Role of genomic instability and p53 in AID-induced c-myc- 
Carpenter. 2004. 53BP1 links DNA damage-response pathways to im-
5:481–487.
Reyraud. 2005. Contribution of DNA polymerase η to immunoglobu-
30. Bertoci, B., A. De Smet, E. Flatter, A. Dahan, J.C. Bories, C. Landreau, 
u and lambda are dispensable for Ig gene hypermutation. J. Immunol. 
168:3702–3706.
31. Faili, A., S. Aoufouchi, E. Flatter, Q. Gueranger, C.A. Reynaud, and 
DNA polymerase theta contributes to the generation of C/G mutations 
during somatic hypermutation of Ig genes. Proc. Natl. Acad. Sci. USA. 
102:13986–13991.
33. Esposto, G., G. Tezido, U.A. Betz, H. Gu, W. Muller, U. Klein, and 
K. Rajewsky. 2000. Mice reconstituted with DNA polymerase beta-
deficient fetal liver cells are able to mount a T cell-dependent immune 
response and mutate their Ig genes normally. Proc. Natl. Acad. Sci. USA. 
97:1166–1171.
34. Johnson, R.E., M.T. Washington, L. Haracska, S. Prakash, and L. 
Carpenter. 2004. 53BP1 links DNA damage-response pathways to im-
5:481–487.
Carpenter. 2004. 53BP1 links DNA damage-response pathways to im-
5:481–487.
Carpenter. 2004. 53BP1 links DNA damage-response pathways to im-
5:481–487.
The translesion DNA polymerase ζ plays a dominant role in immunoglobu-
in gene somatic hypermu-
DNA polymerase η-mediated repair of DNA substrates mimicking non-
homologous end joining and inter-
2004. Absence of DNA polymerase γ reveals targeting of C mutations on 
the nontranscribed strand in immunoglobulin switch regions. J. Exp. 
Med. 199:917–924.
Reyraud, and J.C. Weill. 2004. DNA polymerase η is involved in hy-
permutation occurring during immunoglobulin class switch recombin-
41. Schrader, C.E., S.P. Bradley, J. Vardo, S.N. Mochegevo, E. Flaman, 
and J. Stavnezer. 2003. Mutations occur in the Ig S mu region but 
are rare in Igμ regions prior to class switch recombination. EMBO J. 
Loss of DNA polymerase θ causes chromosomal instability in mam-
43. Sonoda, E., T. Okada, G.Y. Zhao, S. Tatemichi, K. Araki, M. Yama-
2003. Multiple roles of Rev3, the catalytic subunit of polzeta in main-
2004. Absence of DNA polymerase γ reveals targeting of C mutations on 
the nontranscribed strand in immunoglobulin switch regions. J. Exp. 
Med. 199:917–924.
43. Sonoda, E., T. Okada, G.Y. Zhao, S. Tatemichi, K. Araki, M. Yama-
2003. Multiple roles of Rev3, the catalytic subunit of polzeta in main-
2004. Absence of DNA polymerase γ reveals targeting of C mutations on 
the nontranscribed strand in immunoglobulin switch regions. J. Exp. 
Med. 199:917–924.
43. Sonoda, E., T. Okada, G.Y. Zhao, S. Tatemichi, K. Araki, M. Yama-
2003. Multiple roles of Rev3, the catalytic subunit of polzeta in main-
2004. Absence of DNA polymerase γ reveals targeting of C mutations on 
the nontranscribed strand in immunoglobulin switch regions. J. Exp. 
Med. 199:917–924.
43. Sonoda, E., T. Okada, G.Y. Zhao, S. Tatemichi, K. Araki, M. Yama-
2003. Multiple roles of Rev3, the catalytic subunit of polzeta in main-
2004. Absence of DNA polymerase γ reveals targeting of C mutations on 
the nontranscribed strand in immunoglobulin switch regions. J. Exp. 
Med. 199:917–924.
43. Sonoda, E., T. Okada, G.Y. Zhao, S. Tatemichi, K. Araki, M. Yama-
2003. Multiple roles of Rev3, the catalytic subunit of polzeta in main-
2004. Absence of DNA polymerase γ reveals targeting of C mutations on 
the nontranscribed strand in immunoglobulin switch regions. J. Exp. 
Med. 199:917–924.