Altered kinetics of nonhomologous end joining and class switch recombination in ligase IV–deficient B cells

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Immunoglobulin heavy chain class switch recombination (CSR) is believed to occur through the generation and repair of DNA double-strand breaks (DSBs) in the long and repetitive switch regions. Although implied, the role of the major vertebrate DSB repair pathway, nonhomologous end joining (NHEJ), in CSR has been controversial. By somatic gene targeting of DNA ligase IV (Lig4; a key component of NHEJ) in a B cell line (CH12F3) capable of highly efficient CSR in vitro, we found that NHEJ is required for efficient CSR. Disruption of the Lig4 gene in CH12F3 cells severely inhibits the initial rate of CSR and causes a late cell proliferation defect under cytokine stimulation. However, unlike V(DJ) recombination, which absolutely requires NHEJ, CSR accumulates to a substantial level in Lig4-null cells. The data revealed a fast-acting NHEJ and a slow-acting alternative end joining of switch region breaks during CSR.
or no homology, consistent with direct joining of switch region breaks through NHEJ. Although the exact mechanism for switch region breaks remains unclear, the current view is that AID catalyzes cytidine deamination (converts cytidines to uracils) in the switch regions, followed by the repair of uracils that ultimately results in DSBs (3).

In contrast to V(D)J recombination, which absolutely requires NHEJ, direct testing of the role of NHEJ in CSR has met with considerable challenge. One obvious obstacle is that NHEJ-deficient animals cannot generate mature lymphocytes necessary for a CSR assay because the V(D)J recombination defect blocks T and B cell development. To circumvent the developmental block on B cells, Ku- or DNA-PKcs-deficient mice have been crossed with mice harboring preassembled Ig heavy and light chain genes to generate monoclonal NHEJ-deficient B cells (4–7). By this method, CSR was found completely abolished in the absence of Ku70 or Ku80 (4, 5). However, this conclusion must be interpreted with caution, because Ku may have non-NHEJ-related functions (e.g., telomere maintenance) and Ku deficiency causes defective cell proliferation (4, 5). As for DNA-PKcs, controversial results were obtained between a point mutation that inactivates DNA-PKcs and a complete deletion of the gene (6, 7). Although catalytically inactive DNA-PKcs allows normal CSR (7), deletion of the DNA-PKcs gene inhibited CSR to all isotypes except IgG1 (6). A caveat of this approach is that there are no T cells in the B cell–reconstituted mice. As a result, these mice have greatly reduced B cell numbers (4–6) and the B cells are inactive (8). The importance of T cell regulation was demonstrated by a later study showing restored CSR in DNA-PKcs–null mice upon T cell transplantation from a B cell–deficient host (8).

In contrast to Ku and DNA-PKcs, XRCC4 and Lig4 have no identified function outside NHEJ and their inactivation results in the most severe form of NHEJ deficiency. Therefore, XRCC4 and Lig4 are the most suitable and specific targets for abolishing NHEJ. However, the B cell reconstitution strategy is not feasible for XRCC4 or Lig4 because deletion of either gene in mice results in embryonic lethality (9, 10). Although p53 deficiency can extend the life of XRCC4 and Lig4 knockout mice, they all succumb to pro-B cell lymphomas shortly after birth (11, 12). Recently, two independent studies used conditional knockout methods to delete XRCC4 in mature B cells (13, 14). Both studies found that CSR was reduced but not abolished in the absence of XRCC4. It is known that XRCC4-deficient cells have a very low level of Ligg4. However, it was also reported that even a very low level of Ligg4 is still sufficient for NHEJ (15).

To unequivocally determine the role of NHEJ in CSR while avoiding the complications associated with some of the animal models, we disrupted the Lig4 gene in a B cell line (CH12F3) capable of highly efficient cytokine-induced CSR in vitro (16). Like its parental line CH12.LX (17, 18), CH12F3 has a mostly stable diploid genome and is thus suitable for gene targeting. We chose to disrupt the Lig4 gene not only because of its central role in NHEJ but also because it is dispensable for somatic cell growth (19). We found that deletion of Lig4 in CH12F3 cells mildly reduced CSR after 3 d of cytokine stimulation, which is qualitatively and quantitatively similar to what was observed with XRCC4-deficient B cells, as well as the Lig4<sup>−/−</sup> p53<sup>−/−</sup> B cells (13, 14). However, we also found a proliferation defect of Lig4-null CH12F3 cells in the presence of cytokine stimulation. This led us to another important finding that the initial rate of CSR is severely inhibited in Lig4-null cells. Thus, the initially observed mild reduction appears to be an underestimation of a marked CSR defect, which is masked by an enrichment of switched cells. These observations reveal the kinetics of a fast-acting NHEJ and a slow-acting alternative end joining during CSR.

RESULTS AND DISCUSSION

Gene targeting of the Lig4 gene in CH12F3 cells

Lig4 is essential for mouse embryo development but dispensable for somatic cell growth. To determine the role of Lig4 in CSR, we disrupted the Lig4 gene on both alleles in CH12F3 cells. Mouse Lig4 has a total of 911 amino acids. The entire coding sequence is located within the second exon. We replaced 78% of the Lig4 coding region with a floxed puromycin marker (Fig. 1 A). The deleted region encodes amino acids 220–911, which contain the catalytic domain and the two BRCT motifs. This created a larger deletion than a previously defined null allele (10).

Six independent clones harboring a correctly targeted allele (designated L4<sup>+/Δ</sup>) were obtained from a total of 192 puromycin-resistant clones. One of these was randomly selected for gene targeting of the second allele. First, the puromycin marker was excised by the Cre-LoxP reaction (designated L4<sup>+/Δ</sup>). Then, the same targeting vector was used again to disrupt the second allele. One correctly targeted clone (designated L4<sup>P/Δ</sup>) was obtained from a total of 34 puromycin-resistant clones (Fig. 1).

Because NHEJ deficiency is known to result in cellular hypersensitivity to DSB-inducing agents such as ionizing radiation and bleomycin, we tested the Lig4-null CH12F3 cells for this property. As expected, Lig4-null cells exhibited a marked sensitivity to zeocin (a bleomycin analogue; Fig. 1 C), based on a colorimetric assay that measures the metabolic conversion of thiazolyl blue tetrazolium bromide (MTT) in the mitochondria of living cells (20).

Reduced CSR in Lig4-null cells

To determine the CSR capacity of Lig4-null CH12F3 cells, cells were seeded at a low density (5 × 10<sup>4</sup> cells/ml) and grown in the presence of cytokines for 72 h, as previously described (see Materials and methods) (16). CSR in Lig4-null cells is reduced to ~40–50% of that of the Lig4-heterozygous cells (Fig. 2 A and B), which are indistinguishable from wild-type cells (not depicted). This is consistent with the two recent studies (13, 14) showing a similar reduction of CSR in XRCC4-deficient primary B cells and in Lig4<sup>−/−</sup> p53<sup>−/−</sup> B cells, indicating the existence of alternative end joining that can resolve switch region breaks.
To confirm that the reduction was indeed caused by a Lig4 deficiency rather than a clonal effect, a genetic complementation assay was performed. First, the floxed puromycin marker in L4 P/Δ cells was excised by the Cre-LoxP reaction to generate puromycin-sensitive Lig4-null cells (designated L4 Δ/Δ; Fig. 2 C). Then, L4 Δ/Δ cells were infected with recombinant retroviruses expressing the mouse Lig4 or control virus made from the empty vector. Infected cells were selected by puromycin and subjected to the CSR analysis. As shown in Fig. 2 D, Lig4 virus–transduced cells had restored normal levels of CSR, but the cells transduced with the control virus did not.

**Altered kinetics of CSR in Lig4-null cells**

Because CSR is regulated by cell division, we analyzed the effect of Lig4 deletion on cell proliferation. Lig4-null CH12F3 cells propagated normally without cytokine stimulation (Fig. 3 A, left). However, slower propagation of Lig4-null cells was detected at 48 and 72 h in the presence of cytokines (Fig. 3 A, right). To distinguish whether this was caused by a reduced proliferation rate or increased cell death, we stained the cells with CFSE and monitored cell divisions through the passive dilution of CFSE. Although the CFSE fluorescence profiles of Lig4-heterozygous and Lig4-null cells completely overlap in the absence of cytokines (Fig. 3 B, left column), a shift toward brighter CFSE was observed for Lig4-null cells in the presence of cytokines at 48 and 72 h (Fig. 3 B, right column), indicating reduced proliferation. We next analyzed cell death using FITC-conjugated annexin V and propidium iodide (PI), which stain early apoptotic and dead cells, respectively. No significant difference in apoptosis or cell death was observed for the two genotypes in the absence of cytokines (Fig. 3 C). However, an increase of apoptosis or cell death

![Figure 1. Gene targeting of Lig4 in CH12F3 cells. (A) Genomic organization of wild-type and targeted mouse Lig4 locus. Closed boxes indicate exons. Small triangles indicate loxP sites. Restriction enzyme sites: B, Bam HI; Bg, Bgl II; C, Cla I; E, EcoR V. DTA, diphtheria toxin A chain; PGK, phosphoglycerate kinase promoter; Puro, puromycin resistance gene; SV40, SV40 early promoter. (B) Southern blot analysis of Bam HI– and EcoR V–digested genomic DNA. Genotype symbols: +, wild-type allele; P and Δ, targeted allele with or without the puro-selectable marker, respectively. (C) Cellular sensitivity to zeocin as determined by the MTT assay. Higher absorbance indicates more live cells in the culture.](image-url)
was observed in *Lig4*-null culture in the presence of cytokines at 48 and 72 h (Fig. 3 C). This suggests that diminished repair capacity in *Lig4*-null cells leads to elevated apoptosis triggered by CSR-associated DSB.

The growth difference of *Lig4*-heterozygous and *Lig4*-null cells in the presence of cytokines prompted us to reevaluate the mild effect of *Lig4* deletion initially observed on CSR. Two possible mechanisms could account for the reduction of CSR in the absence of Lig4. One possibility is that Lig4 affects cell proliferation in a way unrelated to NHEJ, which in turn affects CSR. The other explanation is that Lig4 is essential for efficient repair of switch region breaks. Without Lig4, accumulation of switch region breaks cause cell-cycle arrest, which is manifested as a cell proliferation defect. If the former were true, CSR level between the two genotypes (L4+/Δ vs. P/Δ) should be similar at 24 h, as cell proliferation is apparently the same (Fig. 3 A, right). However, if the latter were true, CSR difference between the two genotypes (L4+/Δ vs. P/Δ) should be more significant at the beginning before the elimination of CSR-failed cells (and the enrichment of switched cells).

To distinguish these two possibilities, a time course of CSR was performed. A representative experiment is shown in Fig. 4 A (left). Although the numerical value of CSR varies slightly in different experiments, the relative ratio of CSR between the two genotypes (L4+/Δ vs. P/Δ) is consistent. At 24, 48, and 72 h, CSR of the *Lig4*-null cell was at 13, 29, and 48%, respectively, of that of the *Lig4*-heterozygous cell (Fig. 4 A, right). Continued subculturing (>7 d) of *Lig4*-null cells in the presence of cytokines eventually led to ~80–90% of the wild-type CSR level (unpublished data). CH12F3 cells grow rapidly in culture with an estimated doubling time of ~8–10 h. Seeding at a very low density is necessary to achieve maximum levels of CSR without subculturing. However, this condition is less optimal for accurately measuring the initial rate of CSR. Therefore, we also seeded the cells at 3 × 10⁵ cells/ml (early log phase) and measured CSR after 24 h. Again, a marked reduction of CSR in *Lig4*-null cells was observed (15% of wild-type level; Fig. 4 B, right) despite a comparable rate of cell proliferation (Fig. 4 B, left). Thus, the optimal rate of CSR requires Lig4 regardless of stimulation conditions.

These data support the hypothesis that NHEJ is the primary repair mechanism for switch region breaks. The small endpoint reduction observed initially is likely an underestimation of a marked CSR defect, which is probably masked by an enrichment of IgA cells that have managed to repair their switch region breaks through alternative end joining.

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**Figure 2. Reduced CSR in *Lig4*-null cells.** (A) FACS analysis of CSR by surface staining of IgA after 72 h of cell growth with or without cytokines. Numbers in boxed areas indicate percentages. CIT, anti-CD40 antibody (IL-4 and TGF-β1). (B) Reduced CSR in *Lig4*-null B cells after 72 h of cytokine stimulation. Error bars represent standard deviations from three independent experiments. (C) Southern blot analysis of Cre-mediated excision of puro resistance marker in L4 P/Δ cells. (D) Genetic complementation of CSR defect in *Lig4*-null cells by retroviral transduction of *Lig4*. Error bars represent standard deviations from three independent experiments.
Nevertheless, in the absence of NHEJ, alternative end joining can effectively repair switch region breaks, albeit at a substantially slower rate.

**Switch junction sequences**

Numerous studies have shown increased use of microhomology at the junctions in NHEJ-deficient cells (14, 21, 22). To determine whether switch junctions in Lig4-null cells are microhomology mediated, we cloned and sequenced Sμ–Sα junctions from stimulated Lig4-heterozygous and Lig4-null cells. Because Sα is highly homologous to Sμ, we chose to PCR amplify switch junctions from individual cell clones. Cloning individual switch junctions instead of amplifying from a pool of cells helps to eliminate false junctions produced by

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**Figure 3.** Proliferation of Lig4-null cells. (A) Normal and reduced propagation of Lig4-null cells in the absence (top) and presence (bottom) of cytokine stimulation, respectively. Error bars represent standard deviations from three independent experiments. (B) CFSE profiles of Lig4-heterozygous (red) and Lig4-null (blue) cells at every 24 h in the absence (left) and presence (right) of cytokines (CIT). (C) Early apoptosis (annexin V-FITC) and cell death (PI) at every 24 h in the absence and presence of cytokines (CIT). Numbers in boxed areas indicate percentages.
template switching of stalled PCR intermediates, as well as the bias against large amplicons. In addition, the distinctive size of each PCR product was used to reference the authenticity of the sequenced junction. We found that 27% of the junctions from Lig4-heterozygous cells were direct joins, but there were none for the Lig4-null cells (Fig. 5). The difference is statistically significant (7 out of 26 vs. 0 out of 23; two-tail p-value of 0.01 by Fisher’s exact test) and consistent with what was observed for the Sμ–Sy junctions in XRCC4-deficient B cells (14). The distribution of recombination break points is indistinguishable between the two genotypes (unpublished data).

DNA end joining in CSR

This study highlights the importance of NHEJ as the primary mechanism of DSBRD repair in CSR but also reveals the flexibility of using alternative end joining, which is in sharp contrast to NHEJ-dependent V(D)J recombination. The mild endpoint reduction of CSR in Lig4-null CH12F3 cells is in agreement with two recent studies using a conditional knockout of XRCC4 in mature B cells (13, 14). However, our study differs from those two regarding cell proliferation under cytokine stimulation. We found a reduced proliferation of Lig4–null cells at the late stage of stimulation, which was not obvious in the two XRCC4 studies. Our observation would argue for a dominant role of NHEJ in CSR, whereas others would argue for the robustness of the alternative end joining. It should be noted that in one of the studies, deletion of XRCC4 is incomplete (13). In the other study, cell proliferation was measured on a p53-deficient background, which lacks the proper cell-cycle checkpoint control (14).

Although alternative NHEJ can efficiently join linearized plasmids, it clearly cannot replace NHEJ for V(D)J recombination or cellular resistance to ionizing radiation. One speculation has been that alternative end joining is not optimized for repair chromosomal breaks, which is not applicable in this case. Another view is that DSBs are repaired in a manner reflecting how they are generated. It is widely believed that the strict dependence of V(D)J recombination on NHEJ stems from the active guidance of the RAG proteins in the postcleavage complex (14, 22). Indeed, fusion of the RAG proteins to the I-SceI endonuclease can direct I-SceI breaks solely to NHEJ (23). On the other hand, certain RAG mutations can significantly alleviate the dependence of V(D)J recombination on NHEJ (22). Therefore, the flexibility of using alternative end joining in CSR might originate from the complete accessibility of AID-initiated breaks. Interestingly, a single I-SceI I break engineered at an acceptor switch region can efficiently support CSR (24), strongly suggesting that CSR does not require the type of synaptic complex presumed to exist in V(D)J recombination.

Alternative end joining was found to excessively use microhomology. Because all switch regions are abundant in conserved pentamer sequences (e.g., GAGCT and GGG G/C T; underlining indicates one nucleotide position), these short sequences could potentially serve as homology blocks and promote the utilization of alternative end joining. Indeed, there was a marked shift toward microhomology-mediated joining in Lig4–null cells. The most remarkable observation was that there was no direct join in the absence of Lig4. This is in complete agreement with what has been observed for Sμ–Sy junctions in XRCC4-deficient mouse B cells (13, 14) and in human patients harboring hypomorphic Lig4 mutations (21).

Mammalian cells have three ATP-dependent DNA ligases (Lig1, 3, and 4). As a dedicated DSB repair enzyme, The Lig4 complex (XLF–XRCC4–Lig4) has a remarkable flexibility of joining blunt, compatible, and incompatible ends, and even ligating across a gap (25). In contrast, Lig1 and Lig3 are considered as nick ligases. However, recent studies have shown that Lig3 is also capable of intermolecular ligation of blunt and short compatible ends (25, 26). Although the exact architecture of alternative end joining remains unclear, biochemical evidence has suggested the XRCC1–Lig3 complex as a primary candidate for the ligation step (27, 28). Disruption of Lig3 or XRCC1 in mice causes embryonic lethality, thereby limiting their genetic characterizations in alternative end joining. However, Lig3 deficiency might not be cellulary lethal given that XRCC1-deficient cells are viable. In this regard, somatic gene targeting of Lig3 might be a suitable way of dissecting the role of alternative end joining in CSR, as well as DSBRD repair in general in mammalian cells.

MATERIALS AND METHODS

Cell culture and CSR assay. CH12F3 cells were obtained from T. Honjo (Kyoto University, Kyoto, Japan). Cells were cultured as previously described (16). Cell propagation was analyzed by counting live cells every 24 h.
Figure 5. Sμ–Sα junction sequences. (A) Percentage of switch junctions with the indicated length of microhomology (excluding nucleotide additions). (B and C) Alignment of switch junctions with germline sequences. Germline Sμ (black) and Sα (gray) sequences are listed on the top and bottom, respectively, of each junction sequence (blue). Microhomologies (boxes) are identified as the largest perfect matches to the germline sequences. Nucleotide additions are underlined. Long vertical lines indicate direct joins. Small vertical lines indicate identity between the junction and germline sequences. Several junctions containing inverted Sμ are indicated by arrows.
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Correction

The authors regret that their paper incorrectly presented data from another study.

The authors state:

“In the Discussion of our recently published paper, we made an error regarding the study by Yan et al. (Yan, C.T., C. Boboila, E.K. Souza, S. Franco, T.R. Hickernell, M. Murphy, S. Gumaste, M. Geyer, A.A. Zarrin, J.P. Manis, K. Rajewsky, and F.W. Alt. Nature. 2007. 449: 478–482; reference 14). We stated that the XP-T/HL cell proliferation curve shown in Fig. 1 B of that paper was done using cells from p53-deficient mice. However, that experiment was performed with cells from mice on a p53+/− background. The authors of that paper also cited unpublished data showing that cells from CD21-cre-Xrcc(c/-) mice (i.e., p53+/−) proliferate normally. Thus, our interpretation that the normal proliferation and class switch kinetics observed with XRCC4-deficient primary B cells was caused by a lack of cell cycle check point control was incorrect.”