Insertion of Phosphoglycerine Kinase (PGK)-Neo 5′ of Jλ1 Dramatically Enhances VJλ1 Rearrangement

By Tianhe Sun and Ursula Storb

From the Department of Molecular Genetics and Cell Biology, University of Chicago, Chicago, Illinois 60637

Abstract
Gene-targeted mice were generated with a loxP-neomycin resistance gene (neo*) cassette inserted upstream of the Jλ1 region and replacement of the glycine 154 codon in the Cλ1 gene with a serine codon. This insertion dramatically increases VA1-Jλ1 recombination. Jλ1 germ-line transcription levels in pre-B cells and thymus cells are also greatly increased, apparently due to the strong housekeeping phosphoglycerine kinase (PGK) promoter driving the neo gene. In contrast, deletion of the neo gene causes a significant decrease in VJλ1 recombination to levels below those in normal mice. This reduction is due to the loxP site left on the chromosome which reduces the Jλ1 germ-line transcription in cis. Thus, the correlation between germline transcription and variable (V), diversity (D), and joining (J) recombination is not just an all or none phenomenon. Rather, the transcription efficiency is directly associated with the recombination efficiency. Furthermore, Jλ1 and VA1 germ-line transcription itself is not sufficient to lead to VJ recombination in T cells or early pre-B cells. The findings may suggest that in vivo: (a) locus and cell type–specific transactivators direct the immunoglobulin or T cell receptor loci, respectively, to a “recombination factory” in the nucleus, and (b) transcription complexes deliver V(D)J recombinase to the recombination signal sequences.

Key words: PGK-neo • immunoglobulin rearrangement • loxP • transcription • B cells

Introduction
The Ig and TCR gene loci are composed of V, D, and J gene segments. During lymphocyte development, these segments somatically recombine to form a functional Ig or TCR gene. This V(D)J recombination is initiated by recombination activating gene (RAG)1/RAG2 proteins and processed by a double strand break repair complex involving DNA-dependent protein kinase catalytic subunit (DNA-PKcs), Ku70/80, x-ray repair complementing defective repair in Chinese hamster cells (XRCC4), and DNA ligase IV (1, 2). The recombination signal sequences (RSSs)1 which mediate this lymphocyte-specific recombination are highly conserved in all the antigen receptor gene segments. Given the conserved RSSs and the same recombination mechanism, the cell type– and stage-specific V(D)J recombination events must be regulated by lineage-specific (B versus T cells) and developmental stage–specific (pre- versus mature cells) accessibility mechanisms (3).

The current model for these lineage- and stage-specific accessibilities holds that the chromatin in different antigen receptor loci is opened in the appropriate cell lineage at the appropriate cell stage so that the V(D)J recombination complexes can be loaded and begin functioning (4, 5).

The transcriptional enhancers in the Igκ, IgH, and TCR genes have been shown to be essential for V(D)J recombination (6–11). Antigen receptor loci generate weak germ-line transcripts at the time when they are recombined (3, 12). For instance, germline transcripts have been observed from Ig heavy chain and κ light chain regions at the same time as their rearrangement (6, 13–16). In a recent paper, the Jκ regions were shown to generate germline transcripts at the small pre-B cell stage when the Igκ genes presumably start to rearrange (17). Even though the germline transcription event is highly correlated with the V(D)J recombination, whether the germline transcription is the cause or a consequence of the chromatin opening is still in debate.

Recent studies have demonstrated that the neomycin resistance gene (neo*), when inserted in the JC intron in the...
Ig heavy chain or κ locus, generally severely inhibits V(D)J recombination (3, 9, 10, 18–20). The inhibition had been explained by postulating that the strong phosphoglycerase kinase (PGK) promoter interacts with Ig enhancer binding transactivating proteins and thus prevents V(D)J recombination either by preventing the transcription of sterile (germ-line) transcripts, preventing chromatin remodeling, preventing the binding of V(D)J recombinase–associated components to the Ig enhancers, or a combination of these (3). The inhibition was clearly due to the PGK-neo gene, as its removal by cre-loxP recombination restored the inhibition. Unfortunately, the effect of PGK-neo on local germ-line DNA transcription was not examined in these studies.

We have created knock-in mice in which a loxP-neo' cassette was inserted upstream of the Jα1 region. The targeted allele can be distinguished from the wild-type allele by a serine codon replacing a glycine codon normally present at position 154 of Ca1. The wild-type Ca1 gene can be cut by KpnI/Asp718 at this position, but the targeted allele cannot. Serine is present in human and chimpanzee λ chains at this position (21). The knock-in mice showed an enormous increase in λ1 B cells which, as we report here, is due to the inserted neo gene driven by the PGK transcriptional promoter.

Materials and Methods

Targeting Vector and Embryonic Stem Cell Screening. The original DNA construct, J3-C1 knock out, was a gift from Dr. D. Huszar, formerly of GenPharm International (San Jose, CA). In this plasmid, the region from the Jα3 exon to part of the Ca1 exon was replaced by a phosphoglycerate kinase promoter–driven puromycin resistance gene (PGK-puro). It was flanked by 4.2 and 4.4 kb of 129S5/SvEvBrd genomic DNA. A PGK promoter–driven thymidine kinase gene (PGK-tk) was located at the 5’ end of the DNA. To make the targeting vector, a 5.3-kb NheI fragment carrying the Jα3 to Ca1 region from a BALB/c genomic clone (22) was first subcloned. A NotI-XhoI fragment from the ploxPneo-1 plasmid (a gift of Dr. A. Nagy, Mount Sinai Hospital, Toronto, Canada [23]) contains a neomycin phosphotransferase (neo) gene, driven by the PGK promoter, terminated by the PGK polyadenylation site, and flanked by two loxP sites. This fragment (loxP-neo) was inserted at the Ndel site, ~100 bp upstream of the R55 of Jα1 in such a way that the neo transcription is toward the J1C1 of the Igα locus. Point mutations were introduced into the Ca1 constant region, changing a glycine codon (GGT) at position 154 of the λ1 chain to a serine codon (AGC). This mutation disrupts a KpnI/Asp718 restriction site. The 7.4-kb NheI fragment containing loxP-neo and the altered Ca1 was then used to replace the NheI fragment containing the PGK-puro in the Jα3-Ca1 knockout plasmid to generate the targeting vector (see Fig. 1 A). The linearized targeting vector was electroporated into AB2.2 embryonic stem (ES) cells (purchased from Lexicon company), and G418- and gancyclovir-resistant clones were selected and further screened by Southern blotting. Two independent clones out of 360 were obtained and injected into C57BL/6 blastocysts to make chimeric mice.

For the screening of positive ES clones, the manual from Lexicon company was followed. The ES cell DNA was digested by HindIII and Asp718 (Boehringer) and the filters were hybridized with a 3’ Ca1 probe (see Fig. 1 A, probe C) first (see Fig. 1 C). They were then stripped and rehybridized with a neo probe (see Fig. 1 A, probe A) to confirm the homologous recombination. The DNAs from positive clones were also digested with Xhol and hybridized with a neo probe for further confirmation of correct targeting.

Mutant Mice. Germline transmitting mice were generated by mating of chimeras with C57BL/6 mice. To generate a κ−/− background, the mice were bred with κ-knockout mice (19, 24). To delete the PGK-neo gene, the κ−/−, λ1-targeted mice were bred with a CRE transgenic mouse line. The Ella-cre transgenic mice (gift from Dr. Westphal, National Institutes of Health, Bethesda, MD [25]) were originally on the FVB background, which has low λ1 expression. They were bred with κ−/−, λ1 wild-type mice first to generate κ−/−, cre transgenic mice with normal λ1 expression. The mice were tested to have normal λ1 expression by serum analysis before breeding with the targeted mice and further verified by FACS® analysis of their spleen cells after the breedings (data not shown). The CRE transgene was screened by PCR using primers: 5’-cre1, 5’-GGTTTCCGCAAGCTT- GAAATG-3’ and 3’-cre1, 5’-CATCTCCACCCGTCAG- TACCTGAG-3’. The κ−/−, cre transgenic, and κ−/−, S-n/S-n mice were bred together to delete the PGK-neo gene. It took two to three breedings to completely eliminate the PGK-neo gene from the germline (25). To analyze the neo-deleted mice, mouse spleens were always biected. Southern blot analysis of DNA from one-half of each spleen was used to check the efficiency of the PGK-neo deletion, while the other half was used to do the FACS® analysis. The results shown in this paper were all done after complete neo deletion. The mouse designations are S-n and S, for neo containing and neo deleted, respectively.

Southern Blot Analysis. DNA isolation from mouse tissues was performed as described (24). The genotype of the κ loci was revealed by hybridizing with an EcoRI-Xhol fragment from pRKU/Xhol (19). To determine the genotypes, rearrangement, and expression levels of the λ1 locus, a 1.0-kb BglII-Smal fragment containing the neo structural gene (neo; Fig. 1 A, probe A) from TN5 neo, a 1.2-kb Xhol-KpnI fragment (JXhol-C1KpnI, probe B) in the Jα1 to Ca1 region and a 0.4-kb EcoRI-HindIII fragment (3’ Ca1, probe C), 4 kb downstream of the Ca1 exon (26), were used. For quantitation of the Southern blotting signals, the filters were scanned on a PhosphorImager and analyzed by ImageQuant v1.2 software (Molecular Dynamics).

Flow Cytometry and Cell Sorting. Bone marrow and spleen tissues were disaggregated using a nylon cell strainer (Becton Dickinson). Red blood cells were then lysed by hypotonic shock. The cell debris was removed by a nylon strainer. Anti-CD16/CD32 monoclonal antibody (clone 2.4G2; BD Pharmingen) was first used to block the Fcy receptors. Cells were then stained (10^6 cells/staining) in the staining buffer (DMEM, free of serum, with 0.1% BSA) with the respective antibodies. Fluorescence analysis was performed on a FACSscan® (Becton Dickinson) and analyzed by CELLQuest™ software. The cells were gated for lymphocytes by forward and side scatter and a total of 10,000 events were collected for each staining. For preparative cell sorting, the stained cells were sorted using a FACStar®Plus™ (Becton Dickinson). Large and small pre-B cells were sorted as B220+IgM−/CD25+ distinguished by cell size (forward scatter; reference 17). Pre-B cells were sorted as B220+ and IgM+ cells.

For the reverse transcription (RT)-PCR of the thymus RNA in Fig. 6, cells from an Igκ-deficient, λ1 G5 S-n mouse thymus were disaggregated as described above. RNA was extracted from these freshly isolated cells. T cell content was verified by staining with

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anti-CD4, CD8 antibodies. Anti-CD19 antibody staining showed that there were <0.5% B cells, if any, in the thymus sample.

The following antibodies were used for staining: Cy-Chrome-conjugated anti-B220 (RA3-6B2), FITC-labeled anti-IgM (Southern Biotechnology Associates, Inc.), PE-labeled anti-CD25 (PC61), PE-labeled anti-CD4, FITC-labeled anti-CD8, and FITC-labeled anti-CD19 (all from BD PharMingen, unless described otherwise). λ1 B cells were revealed by biotinylated L22.18.2 monoclonal antibodies (24, 27) with PE-labeled streptavidin (BD PharMingen) as the secondary reagent. λ2 and λ3 B cells were revealed by FITC-labeled anti-λ2/3 antibody (2B6; references 17 and 28).

RNA Preparation. Freshly harvested cells were homogenized in 1 ml of STAT-60 containing phenol and guanidinium thiocyanate (Tel-Test). The RNA was extracted, precipitated, and suspended in diethylpyrocarbonate-treated water.

PCR, RT-PCR, and Sequence Analysis. The sequences of the primers used for these assays are: 5'-NCRTGCTTCTGAGGTATTATAGTAAACATC-3'; 5'-CACAGCAATGTTCCTTCTGGAGACAGCC-3'; 5'-TCCTCTTTGATTATCAGAACCTGACCG-3'; 5'-GACCTGCAACCAAGCTA-3' (JM000); 5'-AGATGAGTAAGTACCAGGACAGTGAAGGC-3'; 5'-GACCTAGGAACAGTCAGCACGGG-3' (JM01); 5'-GACCTGAAGCTTTCCCAGCCTCTGTGCTGAATGTTC-3'; StA1, 5'-CTTGAGAATAAAATGCATGCAAGG-3'; JM001, 5'-TTTCTTGGAAAGACCTCCTATGAGG-3'; JM005, 5'-TGGAAATTCTGATTAATTCTTCG-3'; JM002, 5'-CCATACCAAGATGGAAATTCTAG-3'; JM006, 5'-TGCTCTCTCTGAGGTCTCCTAGAT-3'; JM015, 5'-GATGTGGATAGATACTGATGACTGTG-3'; JM014, 5'-TTGGGCAATATCTACCTTGTGTAGG-3'; JM012, 5'-GAATGCCCTAGGTATTAATTGAAACATC-3'; JM007, 5'-CCCTATATCATGTTTGAAAGGTCATGCC-3'; JM008, 5'-CACAGCAATGTTCCTTCTGGAGACAGCC-3'; JM009, 5'-CTCTCTCTACTAATCTGAATCCAGTCACGC-3'; JM010, 5'-CAAGGATGACTGTGGTCTGACCTTGCAGG-3'; JM011, 5'-GATGTGGATAGATACTGATGACTGTG-3'; VL1L, 5'-GTTGTTGATTATATTGGCTCTGAT-3'; VL1RS, 5'-GATGGTAGGACCACGAGTGAAGG-3'; SL1, 5'-CTCTCTCTACTAATCTGAATCCAGTCACGC-3'; JM001, 5'-GAATGCCCTAGGTATTAATTGAAACATC-3'; JM007, 5'-CCCTATATCATGTTTGAAAGGTCATGCC-3'; JM008, 5'-CACAGCAATGTTCCTTCTGGAGACAGCC-3'; JM009, 5'-CTCTCTCTACTAATCTGAATCCAGTCACGC-3'; JM010, 5'-CAAGGATGACTGTGGTCTGACCTTGCAGG-3'; JM011, 5'-GATGTGGATAGATACTGATGACTGTG-3'; VL1L, 5'-CTCTCTCTACTAATCTGAATCCAGTCACGC-3'; JM001, 5'-GAATGCCCTAGGTATTAATTGAAACATC-3'; JM007

The positions of PCR and RT-PCR primers in the λ locus are indicated in Figs. 5 A, 6 A, and 7 A. To distinguish the rearrangements in splenic and sorted λ B cells in various mice, PCR with the primers 5'Nreg and 3'J3C1 were used. The cycling conditions for the reactions were optimized to assure linear amplification. PCR conditions were as follows: 94°C for 30 s, 55°C for 30 s, and 72°C for 2 min, with 28–35 cycle times. To clone and sequence the VJ junction in S-n/S-n splenic DNA and sorted G/S samples were always used as the efficient digestion control. Normally, 8 μPCR product were digested with Asp718 at 37°C overnight. To increase the sensitivity of the assay, after the electrophoresis, the samples were transferred onto a nylon filter and hybridized with the specific radiolabeled probes (see Fig. 1 A).

Results

Targeted Insertion of PGK-Neo Upstream of the Jα1 Region. We generated mutant AB2.2 ES cells in which the loxP-flanked PGK-neo cassette was inserted 100 bp upstream of the Jα1 RSS region (Fig. 1, A–C). The PGK-neo was inserted in the same transcriptional orientation as the Jα1 region. Two point mutations were introduced at amino acid position 154 of the CA1 gene, changing a glycine codon into a serine codon. The mutations disrupt a KpnI/Asp718 restriction site so that the targeted allele and the G/G samples were always used as the efficient digestion control. The work described here was done with mice with a κ knockout background unless described otherwise. To eliminate PGK-neo by Cre/loxP recombination, EIIa-cre transgenic mice (25) were bred with the serine-targeted mice to generate PGK-neo deleted mice (Fig. 1 D). Germline transmitting mutant mice were further crossed with Igκ knockout mice (19) to result in a homozygous κ-knockout background. In this way only λ light chains were produced, facilitating their analysis, as normally λ+ B cells represent only 5% of the total B cells. The work described here was done with mice with a κ knockout background unless described otherwise. To eliminate PGK-neo by Cre/loxP recombination, EIIa-cre transgenic mice (25) were bred with the serine-targeted mice to generate PGK-neo deleted mice (Fig. 1 D; S/S). The neo targeted allele is named S-n (serine-neo), and after neo deletion it is named S. The wild-type allele is named G (glycine).

Gene-targeted Mice Have Greatly Increased λ1 B Cell Numbers. Bone marrow and splenic B cells from κ−/− and λ1 wild-type (G/G), heterozygous (G/S-n), or homozygous (S-n/S-n) mice were analyzed by FACS. Surprisingly, compared with wild-type mice, the λ1 B cell numbers in the G/S-n heterozygous mice are greatly increased both in bone marrow and spleen (Fig. 2 A). The homozygous S-n/S-n mice have an even greater enhancement (Fig. 2 A). Although in wild-type mice the ratio of λ1 to A2/3 B cells in the bone marrow and spleen is 1.5 and 1.1, respectively,
the ratio in homozygous knock-in mice, λ1 S-n/S-n, is 14.9 and 11.9. Six independent pairs of mice were analyzed by FACS®; the λ1 B cell percentages in S-n/S-n mice increase dramatically in all cases (Fig. 2 B). On average, the wild-type G/G mice have 54% λ1 B cells, but the mutant S-n/S-n mice have 82%. We also checked the pre-B cell stages and no difference was found between S-n/S-n and G/G (data not shown).

In κ wild-type mice with the S-n/S-n λ1 locus (Table I), the κ B cell number seems to be downregulated at the same time as the λ B cell number increases, altering the κ/λ ratio from 20:1 to 8:1. The total lymphocyte number seems also decreased, which may be caused by the less efficient development of λ B cells than κ B cells due to selection. Indeed, in κ knockout mice with wild-type λ locus, where all B cells are λ producing, the total B cell number decreases to ~5/50% of normal (29).

**High Levels of VJ Recombination in the Targeted Allele.**
We next examined whether rearrangement is increased in the targeted allele. As the serine mutation disrupts a KpnI/Asp718 restriction site, the rearranged targeted allele can be distinguished from the wild-type allele by EcoRI and Asp718 double digestion (Fig. 2 C). In the spleen of the λ1 heterozygous G/S-n mouse, there is a twofold stronger rearranged band from the targeted allele than from the wild-type allele. This finding was made in repeated analyses. This effect is further confirmed by semiquantitative PCR of the splenic DNA from the λ1 wild-type (G/G), heterozygous-targeted (G/S-n), homozygous-targeted (S-n/S-n), heterozygous-targeted with neo deletion (G/S), and homozygous with neo deletion (S/S) mice. After neo deletion the targeted allele (S) gives a 5.9-kb band.

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**Figure 1.** Gene targeting of the Igλ1 locus in ES cells and mice. (A) Strategy for the gene targeting. A general map of the Igα locus is shown at the top. In the targeting vector, a PGK-neo cassette flanked by two loxP sites (shown as black triangles) was inserted ~100 bp upstream of Jα1 RSS region. Two point mutations (*) were introduced into the CA1 exon at amino acid position 154 disrupting a KpnI/Asp718 restriction site. Also indicated with black bars (A, B, or C) are probes used in this study. See Materials and Methods for other details. Cre recombinase was introduced by breeding the targeted mice with the Cre transgenic mice. The restriction sites relevant to the figures in this paper are shown as O (XhoI), X (XbaI), H (HindIII), K (KpnI/Asp718), and E (EcoRI). E2-4, and E3-1 are the λ enhancers. (B) The sequences in the CA1 region where the point mutations were introduced. A glycine codon (GGT) was replaced by a serine codon (AGC) at position 154. (C) Southern blot with the 3′C1 probe (A, probe C) on the HindIII-Asp718-digested ES cell DNA from two targeted cell clones (G/S-n) and wild-type cells (G/G). The targeted allele (S-n) gives a 8-kb band, whereas the wild-type allele (G) gives a 4.7-kb band. (D) Southern blot carried out with the 3′C1 probe on HindIII-Asp718-digested kidney DNA from the λ1 wild-type (G/G), heterozygous-targeted (G/S-n), homozygous-targeted (S-n/S-n), heterozygous-targeted with neo deletion (G/S), and homozygous with neo deletion (S/S) mice. After neo deletion the targeted allele (S) gives a 5.9-kb band.
Figure 2. λ1 B cells in bone marrow and spleen of mice with the Neo insertion. (A) Bone marrow and spleen cells from κ−/− G/G, G/S-n, and S-n/S-n mice were double-stained with FITC-conjugated anti-λ2+λ3 antibody and biotin-coupled anti-λ1 antibody (revealed by PE-coupled streptavidin) and analyzed by flow cytometry. Lymphoid cells were gated by forward and side scatter. The numbers indicate the percentages of the particular cell population among all the gated lymphoid cells. (B) The percentages of λ1 B cells among the total splenic B cells in the κ−/− G/G vs. S-n/S-n mice. The results from six independent FACS® analyses are shown. Three littermates as well as three nonlittermates are included. (C) Southern blot with the J1XbaI-C1KpnI probe (see Fig. 1 A, probe B) on EcoRI-Asp718–digested thymus and spleen DNA from κ−/− G/G, G/S-n, and S-n/S-n mice. The germline bands are indicated as “GL” and the rearranged bands are indicated as “R.” The targeted allele and nontargeted allele are indicated as “S-n” and “G”, respectively. The probe cross-reacts with a germline band from the pseudogene JC14 and therefore the λ4 band was used as the loading control of the experiment.

Table I. λ B Cell Numbers in κ Wild-type Mice

| Genotype | N   | Total nucleated cells | Total lymphocytes | κ B cell numbers | λ B cell numbers | N'   | Percentage of κ B cell | Percentage of λ B cell |
|----------|-----|-----------------------|-------------------|-----------------|-----------------|------|------------------------|------------------------|
| G/G      | 4   | 80 ± 15.5             | 59.20 ± 16.10     | 37.75 ± 13.85   | 1.86 ± 0.43     | 6    | 63.01 ± 8.41           | 3.38 ± 0.59            |
| G/S-n    |     |                       |                   |                 |                 | 3    | 53.70 ± 6.86           | 5.30 ± 0.44            |
| S-n/S-n  | 4   | 70.5 ± 18.41          | 50.48 ± 18.33     | 22.76 ± 4.27    | 3.45 ± 1.38     | 5    | 50.98 ± 13.59          | 6.55 ± 1.24            |

The splenic cell numbers were determined in G/G, G/S-n, and S-n/S-n mice on the κ wild-type background. “N” stands for the total number of mice in which the total spleen cell numbers were counted. “N’” stands for the total number of mice analyzed, including some in which the total nucleated cell numbers were not determined; thus, N’ is greater than N. The unit of the cell number is 10⁶. The average numbers are followed by the SEM. Total lymphocyte numbers were calculated by forward and side scatter gating in FACS® analysis. κ and λ B cells are determined by double-color fluorescence staining using polyclonal antibodies. The percentages of κ and λ B cells are per total lymphocytes.
The PGK-Neo Cassette Is the Cause of the Higher VJ Recombination in the Targeted λ1 Locus. The serine codon mutation, consisting of two single nucleotide changes, is 1.3 kb away from the JA1 RSS making it unlikely that it affects VJ recombination. However, the serine codon could affect λ1 B cell receptor activity and therefore λ1 B cell development. On the other hand, PGK-neo, which is located between the VA1 and JA1 region, could only affect VJ recombination and not λ1 transcription, as PGK-neo will be deleted by VJ recombination. To determine if the PGK-neo leads to the high λ1 expression, the loxP-flanked neo gene was deleted from the germline of targeted mice by breeding with cre transgenic mice (see Materials and Methods). This cre transgene is expressed throughout life in all cells, starting in the early embryo (25). After 2–3 breedings with the cre transgenic mice, the targeted mice had completely lost the PGK-neo cassette from the germline. Southern blotting of the splenic DNA (similar to Fig. 1 D, data not shown) demonstrates that the 8-kb band (S-n, with neo insertion) is replaced by a 5.9-kb band (S, without neo). Flow cytometric analysis of the splenocytes from the cre transgenic mice, the targeted mice had completely lost the PGK-neo cassette from the germline. Southern blotting of the splenic DNA (similar to Fig. 1 D, data not shown) demonstrates that the 8-kb band (S-n, with neo insertion) is replaced by a 5.9-kb band (S, without neo). Flow cytometric analysis of the splenocytes from the cre transgenic mice, the targeted mice had completely lost the PGK-neo cassette from the germline.

The LoxP Site Inhibits VJ Recombination in the Targeted λ1 Locus. The targeted mice with a complete PGK-neo deletion (S/S) have a lower level of λ1 B cells than the wild-type mice: the S/S mice have only 15.8% λ1 B cells (6.07/38.36% of total B) to 38.8% in G/S versus G/S-n mice and from 92.2 to 15.8% in S/S versus S-n/S-n mice (compare Figs. 3 and 2 A). Therefore, the PGK-neo gene in the targeted allele greatly increases VJ recombination at that allele. We also note that the percentage of λ1 B cells is reduced after the PGK-neo deletion (S/S) to even lower levels than in wild-type mice (G/G; Fig. 3; see below).

To check whether the PGK-neo can advance VJ recombination into an earlier B cell stage, we sequenced the Vλ1Jλ1 coding joints from splenic B cells of the S-n/S-n targeted mice (Table II). N region nucleotide addition was found in only 1 of 14 joints, similar to the light chain N addition frequency in λ1 genes of normal mice. The VD and DJ joints of IgH chain genes, which rearrange at the pro-B to early pre-B cell stages, have a high frequency of N additions due to TdT expression (30). Thus, using N additions as a measure of B cell stage, we conclude that PGK-neo cannot drive VJ recombination of λ1 to the early B cell stages at which IgH genes rearrange. As expected from a random sample, only about 1/3 of the rearrangements were in frame (productive; Table II).

Figure 3. λ1 B cells in bone marrow and spleen after neo deletion. The analysis was done as in the legend to Fig. 2 A. A representative analysis from κ−/− G/G, G/S, and S/S mice is shown. Spleens were bisected. One half was used to do a Southern blot (similar to Fig. 1 D; data not shown) confirmed that neo was completely deleted. The other half was used for the FACS® analysis.

Bone Marrow

Spleen

G/G

G/S

S/S

λ1

λ2+λ3
digestion, the wild-type versus targeted rearranged bands from the immature to the mature B cell stage. The ratio of the serine substitution does not inhibit the development (mainly in the bone marrow) with mature B cells (spleen), increases the signaling efficiency of the B cell receptor, B2m, mice, basically on the other hand, the finding that all seven out of seven rearrangements in VJ1 region de-

also, if the wild-type allele, whereas the sequences from wild-type allele), whereas the sequences from genes from each DNA sample (Fig. 4 A). After Asp718 digestion, the wild-type versus targeted rearranged bands can be distinguished (Fig. 4, A and B). The rearranged wild-type \( \lambda 1 \) band is four times stronger than the targeted \( \lambda 1 \) band in \( \lambda 2/\lambda 3 \) B cells, and there is a similar difference in \( \lambda 1 \) B cells (Fig. 4 B). We next sequenced the \( \lambda 1 \) VJ sequences from \( \lambda 2/\lambda 3 \) B cells. All of the sequences (18/18) are nonproductively rearranged (the majority are from the wild-type allele), whereas the sequences from \( \lambda 1 \) B cells are all functional (7/7; Table III). The absence of productive \( \lambda 1 \) rearrangements in \( \lambda 2/3 \) cells has also been found in normal mice (32). It probably indicates that in general, \( \lambda 1 \) is rearranged before \( \lambda 2 \) or \( \lambda 3 \), thus allowing \( \lambda 2/3 \) rearrangement mainly if \( \lambda 1 \) is nonproductively rearranged. On the other hand, the finding that all seven out of seven \( \lambda 1 \) sequences obtained from \( \lambda 1 \) B cells of G/S mice were from the wild-type, G, allele further indicates that the wild-type allele is rearranged much more efficiently than the mutant S allele. Also, if the wild-type \( \lambda 1 \) allele is nonproductively

**Table II. VJ Joints of \( \lambda 1 \) Light Chain Genes**

| V\( \lambda 1 \) region | N region | J\( \lambda 1 \) region | Productivity |
|------------------------|----------|----------------------|-------------|
| AGC AAC CAT TTC        | C TGG GTG TTC |                      | +           |
| 1                      | AGC AAC CAT TT- | - - -G GTG TTC | +           |
| 2                      | AGC AAC CAT TT- | - -GG GTG TTC | -           |
| 3                      | AGC AAC CA-     | - - - -C TGG GTG TTC | +   |
| 4                      | AGC AAC CAT TTC | T                      | -           |
| 5                      | AGC AAC CAT TT- | - - GG GTG TTC | +           |
| 6                      | AGC AAC CAT T-- | - - GG GTG TTC | +           |
| 7                      | AGC AAC CAT T-- | - - GG GTG TTC | +           |
| 8                      | AGC AAC CAT TT- | - - GG GTG TTC | -           |
| 9                      | AGC AAC CAT TT- | - - GG GTG TTC | -           |
| 10                     | AGC AAC CAT TTC | - - GG GTG TTC | -           |
| 11                     | AGC AAC CAT TTC | - - GG GTG TTC | -           |
| 12                     | AGC AAC CAT TT- | - - GG GTG TTC | -           |
| 13                     | AGC AAC CAT TT- | - - GG GTG TTC | -           |
| 14                     | AGC AAC CAT TT- | - - GG GTG TTC | -           |
Table III. Sequence Analysis of the Rearranged Λ1 Gene

| Germine sequences | Productivity |
|-------------------|--------------|
| VA1-AAC CAT TTC   |              |

From λ2/3 B cells

1  AAC CAT -GG GGT TTC GGT -
2  AAC CAT -GG GGT TTC GGT -
3  AAC CAT -GG GGT TTC GGT -
4  AAC CAT TT- -GG GGT TTC GGT -
5  AAC CAT TT- -GG GGT TTC GGT -
6  AAC CAT TT- -GG GGT TTC GGT -
7  AAC CAT TT- -GG GGT TTC GGT -
8  AAC CAT TT- -GG GGT TTC GGT -
9  AAC CAT TT- -GG GGT TTC GGT -
10 AAC CAT TT- -GG GGT TTC GGT -
11 AAC CAT TT- -GG GGT TTC GGT -
12 AAC CAT TTC GGA -GG GGT TTC GGT -
13 AAC CAT TTC -GG GGT TTC GGT -
14 AAC CAT TT- -GG GGT TTC GGT -
15 AAC CAT TT- -GG GGT TTC GGT -
16 AAC CAT TT- -GG GGT TTC GGT -
17 AAC CAT TT- -GG GGT TTC GGT -
18 AAC CAT TT- -GG GGT TTC GGT -

From λ1 B cells

1  AAC CAT TT- -GG GGT TTC GGT +
2  AAC CAT TT- -GG GGT TTC GGT +
3  AAC CAT TT- -GG GGT TTC GGT +
4  AAC CAT TT- -GG GGT TTC GGT +
5  AAC CAT TT- -GG GGT TTC GGT +
6  AAC CAT TT- -GG GGT TTC GGT +
7  AAC CAT TT- -GG GGT TTC GGT +

Sequence analysis of the rearranged Λ1 gene in either sorted Λ1 or sorted λ2+Λ3 B cells from a κ<sup>-/-</sup> G/S mouse. The germine sequences of VA1 and Λ1 coding ends are shown at the top. Independent clones from the sorted B cells are shown below. Nucleotides deleted at the coding ends are indicated by dashes. P (palindromic) elements are shown in italics. N nucleotides are shown as underlined italics. Productivity: + (−), the gene is productively (nonproductively) rearranged.

The levels of Λ1 germline transcripts are correlated with the efficiency of VJ recombination in S-n, S, and G alleles. To determine the relationship between VJ recombination efficiencies and germline transcription activity, we measured the levels of Λ1 germline transcripts in heterozygous mice before and after deletion of PGK-neo and in the wild-type Λ1 locus. The Igα light chain genes normally start to rearrange at the small pre-B cell stage, at which time Λ1 germline transcription is found (17). We sorted small pre-B (IgM<sup>−</sup>B220<sup>+</sup>CD25<sup>+</sup>, small) as well as large (less mature) pre-B cells (IgM<sup>−</sup>B220<sup>+</sup>CD25<sup>+</sup>, large) from the bone marrow of two G/S-n heterozygous mice. RT-PCR and restriction digestion were performed to distinguish the germline transcript levels from the two different alleles (Fig. 5). At the small pre-B cell stage, the level of germline transcripts is 1.6 to 5.9 times higher from the targeted allele (S-n) than from the wild-type allele (G). A much greater difference is observed in less mature large pre-B cells. Transcripts from the targeted allele are present at similar levels as in small pre-B cells by semiquantitative estimation relative to β-actin, but no transcripts from the

Figure 5. Germline transcription levels of Λ1 in κ<sup>-/-</sup> G/S-n and G/S mice. (A) Schematic structure of the Λ1 germline transcripts. The germine transcripts are initiated 5’ of the J region and spliced to the C region. Location of primers used for RT-PCR are shown by small arrows. The gray box indicates the probe (see Fig. 1 A, probe B) used in the Southern blot of the RT-PCR products. (B) Southern blot of the RT-PCR products of the germine transcripts from the indicated cell stage/lineage. The RT-PCR products were digested by Asp718 (indicated as “Cut”) to distinguish products from the wild-type and targeted alleles. The blots were hybridized with the J1Xhol-C1Kpol probe. Germine transcripts corresponding to the targeted or the wild-type alleles are indicated. The experiments from two dilutions of thymus sample are shown. No amplification products were obtained when reverse transcriptase was omitted from the reaction of all the samples (two are shown as “RNA,” others not shown).
alleles after neo deletion we sorted pre-B cells (IgM^+2, B220^+) and T cells (Fig. 2 C; see Discussion).

To compare the expression of the wild-type and targeted alleles after neo deletion we sorted pre-B cells (IgM^+2, B220^+) from the heterozygous G/S mouse bone marrow. The germline transcripts from these cells are clearly biased to the wild-type allele (Fig. 5 B). Coincidentally, in these mice, VJ recombination also preferably occurs at the wild-type allele (Fig. 4). Thus, the germline transcript levels exactly parallel the recombination efficiencies from the highest in PGK-neo containing (S-n), through the intermediate in wild-type (G), to the lowest in PGK-neo deleted (S) α1 alleles.

Mapping of the Germline Jα1 Promoter. To understand the relationship between the inserted PGK-neo gene and the germline Jα1 promoter we determined the position of this promoter. Jα1 germline transcription was found previously in wild-type mice, but the transcript initiation site was not defined in mice (17). To map the promoter, a series of 13 5' primers were used that are located 5' of Jα1, together with a 3' primer in CA1 to amplify oligo-dT-primed cDNAs from pre-B cells of λ wild-type mice. As shown in Fig. 6 A, primers located up to 269 nucleotides upstream of the NdeI site gave a PCR product, whereas primers located further upstream did not. The most 3' two of the latter primers were found to amplify a correct sized and sequenced product from genomic DNA; therefore, the lack of cDNAs extending into this region is not due to inefficient primers. Thus, this analysis places the initiation of the germline Jα1 transcript between primers JM014 and JM015. There is a good candidate for a TATA box just upstream of primer JM014 which would place the transcription start site at about the G located 29 nucleotides 3' of the putative TATA box (Fig. 6 B). This start site is six nucleotides 5' of the 3' end of primer JM014, making the primer overlap too short for stable annealing. Thus, the start of the Jα1 germline transcript is located 420 nucleotides 5' of Jα1 in the untargeted λ locus (Fig. 6 B). In the targeted locus it resides 336 nucleotides 5' of the inserted PGK-neo gene and 1 kb 5' of the start of the neo gene transcripts.

Do All Jα1 Germline Transcripts from the Targeted Allele Initiate from the PGK Promoter? Jα1-containing transcripts in thymus and large pre-B cells of G/S-n mice are all from the targeted allele (Fig. 5 B). These transcripts must initiate from the PGK promoter, as none are found that initiate upstream of this promoter in the targeted allele of G/S-n mice (Fig. 6 C). On the other hand, in small pre-B cells, Jα1-containing transcripts arise from both the targeted and the wild-type allele in G/S-n mice (Fig. 5 B). Interestingly, in these cells transcripts are found that start upstream of the PGK promoter and most likely initiate at the germline Jκ1 promoter. Thus, this promoter is only active in small pre-B cells when the λ enhancers are activated and VJ recombination takes place, even in the presence of PGK-neo.

Discussion

Jα1 Germline Transcription and VJ Recombination Is Promoted by the PGK-Neo Insertion. When PGK-neo is inserted in the κ or heavy chain intron, in most cases it causes strong inhibition of V(D)J recombination (3, 9, 10, 18–20). A similar phenotype has also been observed in the TCR loci (33). This was postulated to be due to the higher density of CpGs in the coding sequence of the neo gene (18) or to the neo coding region itself (34). Alternatively, it may be due to a squelching effect, in which the PGK promoter interacts with the Ig enhancer at the expense of an Ig promoter/enhancer interaction (9, 10). In our experiment, the location of the PGK promoter relative to the germline Jα1 promoter and the EA3-1 enhancer is such that the PGK promoter initiates transcripts that pass through the RSS. Therefore, instead of possibly interfering with the germline transcription of the JC κ1 region, a strong PGK promoter in this case promotes transcription through Jα1. However, similar to findings by others (35) transcription alone is not sufficient for rearrangement (see below).

The Jα1 Germline Transcription Is Inhibited by the LoxP Site. A single remnant loxP site with its short flanking region dramatically inhibits VJ recombination. This is probably a consequence of the severe inhibition of Jα1 germline transcription by this loxP remnant (Fig. 5). We consider two possible reasons. The remnant loxP site is very rich in CpG dinucleotides. As CpGs are substrates for DNA methylation, methylated CpGs may be involved in inactivating the region, leading to the repression of Jα1 germline transcription. It has been shown that hypomethylation is correlated with V(D)J recombination activity (36–38).

Alternatively, the additional 150 nucleotides of the remnant loxP site might directly interfere with transcription initiation from the Jα1 promoter. In another study, a loxP remnant was reported to have no effect on VJ recombination in the Igκ locus, nor to affect Jκ germline transcription (39). In that study, the loxP site was located farther away from the Jκ RSS (2 kb). Also, the flanking region of the loxP site was different from ours.

Finally, the loxP site remaining in the “S” mutation may disrupt a Jκ proximal regulatory element similar to the κI/κII sequences which are important for Jκ gene rearrangement (39).

As the cre-loxP system is extensively used to eliminate potential effects of the selectable marker in targeted loci, it is important to consider possible consequences of retaining a loxP site. The loxP flanked marker gene should be placed at a safe distance from any essential control element, and the constitution of the polylinker region should be considered.
The Levels of Jα1 Germline Transcription Are Highly Correlated with VJ Recombination. What is the molecular mechanism for the PGK-neo effect? As the PGK promoter is a strong housekeeping promoter, it is presumably active in all cells. Clearly, transcripts from the S-n allele initiating upstream of Jα1 can be detected in both large pro-, pre-B cells (stages A–C of the Hardy nomenclature [40]) and in T cells, where germline transcripts from the wild-type allele are not found (Figs. 5 and 6). Nevertheless, Jα1 germline transcription is not sufficient to promote α1 rearrangement in either T cells or in the early B cell stage (Fig. 2 C).

We find no rearranged Jα1 band in the thymus from an S-n/S-n mouse (Fig. 2 C). The Vα11Jα1 joints found in the S-n/S-n homozygous targeted mice do not show an increase in N additions (Table II), as would be expected if the rearrangement took place as early as Ig heavy chain gene rearrangement in B cell development when TdT is expressed (41).

Figure 6. The germline Jα1 promoter. (A) Map and RT-PCR primers used to identify the endogenous Jα1 germline transcription initiation site. The endogenous Jα1 locus is shown in the top three lanes, with the position of the NdeI site, where PGK-neo is inserted in the targeted S-n allele. The position of each upstream primer used is shown, with the number indicating the distance (in basepairs) between the 5' end of the primer and the NdeI site. The downstream primer used for RT-PCR was 3'J3C1, located in the Cα1 exon. The names of the primers are in brackets. The position of the potential initiation site is marked with a black arrow. DNA PCR: four primers tested and found positive in PCR amplification of genomic DNA with the 3'J3C1 downstream primer. The position of the primer used in C, 3'PNR-2, is shown. (B) Sequences 461 to 1,952 between Cα3 and Jα1 were not previously sequenced. Exons and nonamer/heptamer are marked by solid and dotted underlines, respectively. The primers JM014 and JM015 are indicated by dotted lines with arrows. The potential TATA box and the transcriptional start site are indicated. (C) RT-PCR to detect germline transcripts from the endogenous promoter in the S-n targeted allele. The primers, JM001 and 3'PNR-2 (see A), were used for PCR reactions with cDNA or RNA of the indicated cell types from a G/S-n mouse. The germline transcription products (248 bp) are indicated as “PGK junction.” G/S-n thymus DNA sample and positive ES clone DNA were included as positive PCR controls. β-actin was included as the cDNA quantity control.
The lack of VJA1 rearrangement in thymus and large pre-B cells of S-n/S-n mice is not due to lack of VA1 germ-line transcription. Surprisingly, VA1 germ-line transcripts are apparently present in both large pre-B cells and thymus RNA (not shown). We also found VA1 germ-line transcripts in thymus RNA from wild-type mice (data not shown). We consider two possibilities: The germ-line transcription of VA1 might not reflect the same chromatin structure alteration required for V(D)J recombination in the J region. Indeed, a recent study showed an unusual profile of germ-line transcription, nucleosome sensitivity, and methylation of the V regions in the TCR-β locus, suggesting that the V region might use a unique structural conformation to be accessed by the V(D)J recombinase complex (42). Second, the germ-line transcription of VA1 might suggest constitutively open chromatin at that region in B and T cells. Therefore, the V(D)J recombination complex may be able to access VA1 region all the time. Thus, the lack of rearrangement of the S-n allele in large pre-B and T cells most likely indicates that rearrangement also requires activation of the λ enhancers by transactivating factors and that this does not occur until the small pre-B stage, when λ1 rearrangement normally takes place. At this stage, the germ-line J1 promoter is active (Fig. 6) and all other required elements for λ1 rearrangement are functional.

In small pre-B cells, the level of the germ-line transcripts from the S-n allele is clearly higher than that from the wild-type allele (Fig. 5 B). It is possible that there is a difference in the stability between the endogenous and the PGK-neo inserted transcripts. However, as Ig transcripts, generally, are very stable, there is no reason to expect that the wild-type transcripts would be less stable than the transcripts from the targeted locus. Therefore, the presence of PGK-neo upstream of the J1 RSS most likely causes up-regulation of germ-line transcription. There are at least two possible models how the levels of germ-line transcription may affect λ1 rearrangement efficiency. In G/S-n mice, the increased germ-line transcription of the targeted allele compared with the wild-type allele could represent a more completely opened chromatin around the J1 RSS via ATP-dependent remodeling and acetylation of histones (43, 44). The more open RSS configuration would be more accessible to limiting quantities of V(D)J recombination components. Conversely, in the neo-deleted mice, in the presence of the CpG-rich loxP remnant, chromatin condensation could persist via the binding of methylated DNA binding proteins and possible recruitment of histone deacetylases (43, 45). The chromatin condensation could make the RSS directly inaccessible or prevent transcription from the germ-line J1 promoter.

Alternatively, the germ-line transcription complexes themselves could be involved in the up-regulation of VJ recombination. In light of the previous findings of inhibition of rearrangement by PGK-neo when inserted into the JC intron and transcribing away from the RSS, our findings would be well explained by an enhancement of V(D)J recombination by transcription through the RSS. Perhaps components of the V(D)J recombinase could be associated with transcription complexes during transcript elongation and be more efficiently deposited at the RSS. It is likely that open chromatin exists in small pre-B cells at both the wild-type and S-n J1 RSSs, as transcripts are seen. However, in an S-n allele with the PGK promoter, transcript initiation occurs at a higher rate allowing more transcription complexes to clear the promoter and possibly to carry V(D)J recombination factors to the RSS during transcript elongation. It would be interesting to turn around the PGK promoter both in the λ and κ loci. Perhaps, no more enhancement of J1 rearrangement would be found, because the PGK initiated transcripts do not pass through this RSS anymore. Further, if directionality is not an issue, κ gene rearrangement may actually be enhanced by transcripts passing through the κ RSS in the opposite orientation.

However, to boost rearrangement, the transcripts may have to initiate at an Ig germ-line promoter. The presence of enhancer elements in PGK-neo may nonspecifically up-regulate transcription from the germ-line J1 promoter when the λ enhancers are active. It appears that there are more transcripts initiating at the PGK promoter than at the germ-line J1 promoter in small pre-B cells (not shown). However, it is possible that the majority of the transcripts are from small pre-B cells that are unable to rearrange VJA1.

It is clear that the accessibility to V(D)J recombination is controlled at several levels besides the requirement for the recombinase complex. Every Ig or TCR gene segment has not only its own RSS but also its own germ-line promoter whose activity is required for V(D)J recombination. RSS strength has been shown to affect the recombination frequencies (46–48). Our findings suggest that there is a potential secondary level of control in which the absolute activity of a germ-line promoter determines the absolute efficiency of recombination. The gene segment carrying a strong promoter will have a greater chance to be rearranged than the one with a weak promoter in the same locus. This, besides RSS quality, may explain the different intrinsic rearrangement efficiencies of different V, (D), and J regions in the Ig and TCR loci.

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