Biallelic, ubiquitous transcription from the distal germline $\lg \kappa$ locus promoter during B cell development

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Allelic exclusion of Ig gene expression is necessary to limit the number of functional receptors to one per B cell. The mechanism underlying allelic exclusion is unknown. Because germline transcription of Ig and TCR loci is tightly correlated with rearrangement, we created two novel knock-in mice that report transcriptional activity of the J $_{\rm K}$ germline promoters in the $Ig_{\rm K}$ locus. Analysis of these mice revealed that germline transcription is biallelic and occurs in all pre-B cells. Moreover, we found that the two germline promoters in this region are not equivalent but that the distal promoter accounts for the vast majority of observed germline transcript in pre-B cells while the activity of the proximal promoter increases later in development. Allelic exclusion of the $Ig_{\rm K}$ locus thus occurs at the level of rearrangement, but not germline transcription.

allelic exclusion | knock-in mouse

The V(D)J recombinase and various DNA repair proteins catalyze the assembly of Ig heavy- and light-chain variable region exons from dispersed gene-segments during B cell development (1). The presence of two alleles for each gene and two light chain loci ($Ig\kappa$ and $Ig\lambda$), suggests that, in theory, each B cell could produce eight different antibody specificities. However, as postulated by Burnet 50 years ago (2), the vast majority of B cells are monospecific. The underlying basis for the "one cell, one receptor" phenomenon is the allelic and isotypic exclusion of Ig gene expression such that the vast majority of mature B cells possess only one functional heavy and one functional light chain gene rearrangement. Despite intensive effort, the mechanism underlying allelic exclusion is unknown.

Two general models have been proposed to account for allelic exclusion (3). The first is an instructive model in which an event early in development differentially marks the two alleles such that one allele is the preferred substrate for recombinase activity. Once established, this differential mark propagates in a clonal manner. The second model proposes a stochastic mechanism where each allele is equally likely to recombine, but the probability of recombination is low so that the percentage of cells with two functional rearrangements is vanishingly small. In support of the first model, it was observed that the two mouse $Ig\kappa$ alleles replicate asynchronously in S-phase beginning at a very early stage of development (4). However, it has not been proven conclusively that the early replicating allele is the preferential target of recombination, nor what other marks might be clonally propagated to dictate allele-specific recombination. In favor of the second model, we previously reported that a GFP cDNA knocked-in to the mouse $Ig\kappa$ locus is expressed in only 1-5% of pre-B cells (5). As germline transcription is tightly correlated with recombination, we interpreted this to indicate that only a small fraction of unrearranged Igk alleles are transcriptionally active and therefore suitable targets for the recombinase at any given time. Neither model fully accounts for observed recombination frequencies or patterns, in particular

receptor editing of the $Ig\kappa$ locus (6). If the preferred allele is dictated, it is unclear how the second allele becomes a substrate for recombination if the first fails to encode a suitable protein. Similarly, if recombination is rare and stochastic, it is unlikely that a cell would ever recombine both alleles; yet a significant fraction of splenic B cells contain two rearranged $Ig\kappa$ alleles (6).

All rearranging loci undergo sterile or germline transcription. Sterile transcripts originate from promoters upstream of the recombining segments in a developmental and tissue-specific manner that correlates tightly with recombination of the associated loci (7, 8). A recent report demonstrated that germline transcription through a cluster of $J\alpha$ segments is necessary for their recombination (9). Insertion of a transcription termination sequence into the J α cluster prevented local V α -to-J α rearrangement. These workers concluded that RNA polymerase II transit across the recombining segments recruits chromatin modifications that make the locus accessible to the recombinase. The J κ cluster of gene segments contains two germline transcript promoters, located approximately 100 bp and 3.5 kb upstream of the J κ 1 gene segment (10–12). Deletion of both promoters abolishes κ locus recombination in AMuLV transformed pro-B cell lines, although the identical experiment has not been done in mice nor has either promoter been deleted individually (13, 14). Although germline transcription appears necessary for recombination, it is unclear if it is sufficient. If germline transcription is sufficient to target the recombinase, monoallelic expression of germline transcripts may underlie antigen receptor allelic exclusion. However, a previous report demonstrated by single cell RT-PCR and RNA FISH that germline transcription of the J κ cluster is biallelic (15). This is in contrast to our report of variegated, predominantly monoallelic expression of this locus (5). Because these results are at odds with one another, it remains unclear how the J κ germline transcripts are expressed and how this might regulate locus recombination.

To further probe the mechanism of allelic exclusion and its relation to germline transcription, we created two new knock-in mouse strains that report transcriptional activity of the germline $Ig\kappa$ locus. We found, contrary to expectation, that germline transcription of the $J\kappa$ cluster is biallelic and occurs in all pre-B cells. In reconciling this data with previous literature, we discovered that the two $J\kappa$ germline promoters in the $Ig\kappa$ locus are

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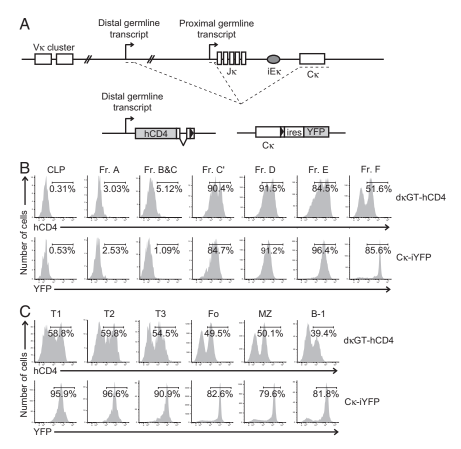


Fig. 1. Marker protein expression in d_KGT -hCD4 and C_K -iYFP knock-in mice. (A) Schematic of the d_KGT -hCD4 and C_K -iYFP knock-in mutations at the lg_K locus. Both the proximal and distal germline transcript promoters are shown as arrows. Previously defined splicing patterns of both transcripts are depicted as dashed lines. The hCD4 cDNA is followed by an SV40 intron and polyA sequence. Black triangles represent the positions of loxP sites remaining in the locus after Cre recombinase-mediated deletion of a neomycin resistance gene. (B) Flow cytometric analysis of hCD4 or YFP marker expression in developing bone marrow B cells from heterozygous d_KGT -hCD4 or C_K -iYFP knock-in mice. Harvested bone marrow was labeled with antibodies to delineate B cell developmental subsets and mark hCD4-expressing cells (in d_KGT -hCD4 mice only). CLP through fraction F are shown. The percentage of marker-positive cells at each developmental stage is shown. Control C57/BL6 mice had no detectable expression of either marker (data not shown). Data are representative of at least five independent experiments analyzing two to four mice per experiment. (C) Flow cytometric analysis of hCD4 or YFP marker expression in splenic B cells from heterozygous d_KGT -hCD4 or C_K -iYFP knock-in mice. Harvested splenocytes were labeled with antibodies to delineate B cell developmental subsets and mark hCD4-expressing cells (in d_KGT -hCD4 mice only). Transitional 1 (T1), transitional 2 (T2), transitional 3 (T3), follicular (Fo), marginal zone (MZ), and B-1 B cell subsets are shown along with the percentage of marker positive cells at each developmental stage. Data are representative of at least two independent experiments analyzing two to four mice per experiment.

not equivalent and that the bulk of germline transcription in this locus derives from the distal promoter in pre-B cells.

Results

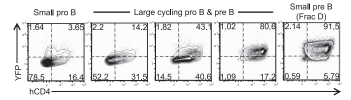
New Igk Knock-In Mice Reveal Germline Transcription in All Developing Pre-B Cells. Our previously studied Igk reporter mouse strain contained a GFP cDNA, heterologous intron and polyadenylation (polyA) sequence inserted into the first Jκ gene segment (referred to as κ^0 GFP) (5). Although this strain reported proximal promoter activity, the polyA sequence decreased recombination to gene segments 3' of the knock-in (data not shown) and resulted in skewed allele usage in mature B cells in heterozygous mice. To confirm our previous results in a more rigorous manner, we sought to create additional reporter mice which would not affect recombination of the locus. Because the $J\kappa$ cluster contains two germline promoters that were thought to be equivalent, we reasoned that placement of a reporter cassette and polyA sequence downstream of the distal promoter was less likely to affect recombination as the proximal promoter was left intact (the polyA sequence is necessary for proper transcript processing and expression of the reporter cassette). We created a mouse in which a human CD4 (hCD4) cDNA, intron and polyA sequence is inserted downstream of the distal J κ germline promoter, approximately 3.5 kb upstream of the J κ 1 gene segment (referred to subsequently as d κ GT-hCD4, Fig. 1A and supporting information (SI) Fig. S1a). The hCD4 was mutated to prevent intracellular signaling and interaction with class II MHC (16). Because the d κ GT-hCD4 insertion could potentially disrupt $Ig\kappa$ control elements, we created a second mouse with an internal ribosome entry site (IRES) and yellow fluorescent protein (YFP) cDNA inserted after the C κ stop codon and before the endogenous polyA sequence (referred to as C κ -iYFP, Fig. 1A and Fig. S1b). The C κ -iYFP insertion has the additional advantage of serving as a reporter for both germline and rearranged transcripts since both include the C κ exon. Both strains of mice were born in equal proportion to wild-type litter mates and revealed no gross physiological abnormalities (data not shown).

We predicted the pattern of marker protein expression from the new knock-in strains would be similar to the κ^0 GFP mouse. However, flow cytometric analysis of heterozygous d κ GT-hCD4 and C κ -iYFP mice revealed a very different pattern of marker expression during B cell development (Fig. 1 B and C). Compared to wild-type mice, heterozygous animals from both knock-in strains displayed very little marker expression in the early stages of B cell development (common lymphoid progen-

itor (CLP) to fraction C). This was expected since the Igk locus does not begin to rearrange until the pre-B cell stage (fraction D) of development. The small percentage of marker-positive cells in fractions B and C derive mainly from cells transitioning from C to C' (data not shown). At the large cycling and small resting pre-B cell stages (fractions C' and D) almost all B cells became hCD4 or YFP positive. This is in contrast to the κ^0 GFP mice, in which <1% of small resting pre-B cells from heterozygotes are GFP positive (data not shown). In the dkGT-hCD4 strain, the marker is in the $V\kappa$ -J κ genomic interval and thus is deleted from the locus upon V_{κ} -to- J_{κ} rearrangement. As such, we found hCD4 marker expression began to decline in immature B cells (fraction E) and was reduced to approximately 50% in transitional and splenic B cells (Fig. 1C). In contrast, the YFP marker is downstream of Cκ, remains in the locus after rearrangement and continues to be transcribed from the incoming $V\kappa$ promoter. Because $V\kappa$ promoters are stronger than the germline J κ promoters, we found the mean fluorescence intensity of YFP to steadily increase as cells progressed from pre-B to splenic B cells. We did detect a fraction of YFP negative cells in IgD+ bone marrow and splenic B cells. These correspond to alleles that have been deleted by recombination to the recombining sequence (RS) element (data not shown).

The κ^0 GFP knock-in disrupted normal recombination of the mutant allele and led to a 4:1 skew in $Ig\kappa$ usage in favor of the wild-type allele in IgM+ B cells (5). To determine if either dκGT-hCD4 or Cκ-iYFP insertions cause similar recombination defects, we mated both strains to h $C\kappa$ knock-in mice which carry a human $Ig\kappa$ constant region exon in place of the mouse exon. Analysis of immature B and splenic B cells in hCκ/dκGT-hCD4 heterozygous mice revealed a 2:1 skew in favor of expression of the human C_{κ} allele (Fig. S2) (17). This agreed with the higher than expected percentage of hCD4 positive splenic B cells (50%) in heterozygous dkGT-hCD4 mice. If the hCD4 knock-in were truly innocuous, approximately 30% of these cells would retain the hCD4 allele in the germline configuration and hence be hCD4 surface positive. In contrast, immature B cells from $hC\kappa/C\kappa$ -iYFP heterozygous mice had a 1:1 ratio of mouse to human $C\kappa$, similar to what is observed in h $C\kappa$ /C57BL6 heterozygous mice. Homozygous dκGT-hCD4 and Cκ-iYFP mice had normal numbers of B cells at all developmental stages compared to 129/SvJ wild-type littermates (data not shown). Thus, while the hCD4 insertion does cause a slight recombination defect, it does not grossly disrupt B cell development. As predicted, the $C\kappa$ -iYFP insertion appeared completely innocuous.

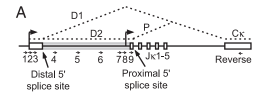
Germline Transcription of the Jk Cluster Is Bialleic. Essentially all small resting pre-B cells appeared positive for either the hCD4 or YFP markers in heterozygotes from the respective knock-in strains. This result suggested that germline transcription may be biallelic. To confirm this interpretation, we mated dκGT-hCD4 and Cκ-iYFP mice and analyzed the resulting compound heterozygotes for expression of both markers during B cell development (Fig. 2). We found that expression of both markers increased as cells progressed from pro-B to the pre-B stage, with almost all small resting pre-B cells being positive for both markers. The hCD4 marker appeared earlier during the pro-topre-B cell transition than YFP. This might be due to the fact that the hCD4 marker is a few hundred base pairs from the distal promoter and thus is sensitive to any transcription initiating from this region while the YFP lies at the end of the transcript and requires that RNA polymerase II transcribes the entire 8 kb region; any stalled or aborted transcripts would not give rise to YFP protein. Alternatively, the sensitivity of the assay for hCD4 may exceed that for YFP. Regardless, all pre-B cells were double positive, confirming that on a single cell level, both $Ig\kappa$ alleles are transcriptionally active during recombination of this locus. This is in agreement with a previous report which demonstrated

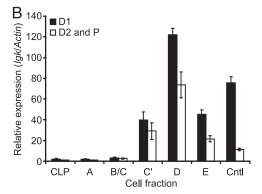


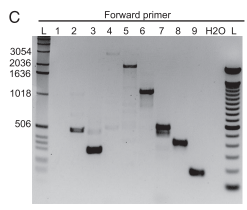
Germline transcription at the J_K cluster is biallelic. Flow cytometric analysis of marker expression in developing B cells from dκGT-hCD4/Cκ-iYFP compound heterozygous mice. Harvested bone marrow was labeled with antibodies to delineate developing B cell subsets and mark hCD4 expression. The progression of cells from the pro-B to the small resting pre-B cell stages is shown. Small pro-B cells were defined as forward scatter low CD43⁺B220⁺ while small pre-B cells were forward scatter low CD43⁻B220⁺. The intermediate stages represent forward scatter high cells (cycling) with intermediate levels of CD43 progressing from high to low. Data are representative of two independent experiments analyzing three mice each.

biallelic Jκ transcription by single-cell RT-PCR analysis and FISH (15).

The Two Jk Germline Promoters Are Not Equivalent. To confirm the validity of the new reporter mice, we fractionated bone marrow B cells from wild-type mice into successive developmental stages by flow cytometry and measured abundance of both the proximal and distal germline transcripts by real-time quantitative RT-PCR (Fig. 3 A and B). In agreement with the reporter mice, the RT-PCR assay demonstrated that J κ germline transcription begins in fraction C' and continues through the rest of B cell development, reaching a peak in small resting pre-B cells. This is a very different pattern of expression than we observed using our previous κ^0 GFP knock-in. Although only a small percentage of pre-B cells are GFP positive in κ^0 GFP mice (5), by conventional RT-PCR the proximal $J\kappa$ promoter appeared to be extremely active. It was possible that the κ^0 GFP mutation is not a faithful reporter of proximal promoter activity, perhaps due to silencing of the targeted allele. However, it is unlikely that a reporter would be systematically silenced early in development only to loose that silencing later (100% of cells that have a germline κ^0 GFP allele express GFP in peripheral B cells). In other reported cases of transgene silencing, the silencing is stable throughout development (18). Another possibility was that the κ^0 GFP insertion was accurately reporting proximal promoter activity, but that the RT-PCR analysis used to assess the proximal promoter-derived transcript was flawed. We hypothesized that the transcript originating from the distal J κ germline promoter may have an alternatively spliced variant that utilizes a splice donor site identical to the proximal transcript splice donor site (Fig. 3A, transcript D2). Such a transcript would wholly encompass the proximal J κ germline transcript in extent and make it impossible to distinguish the proximal transcript from this distal-promoter-derived transcript by conventional RT-PCR (Fig. 3A). To test if such a transcript exists, we performed RT-PCR on wild-type pre-B cell cDNA using a common anchor primer in the $C\kappa$ exon paired with various primers arrayed across the region between the distal germline transcript promoter and the J κ cluster. As shown in Fig. 3C, we observed bands of the predicted sizes for a transcript that initiates from the distal germline promoter, but that utilizes a 5' splice site downstream of $J\kappa 1$ (lanes 4–6). These were not primary transcripts as the intron between $J\kappa 1$ and $C\kappa$ was correctly spliced from all observed products. Primers placed upstream of the previously reported distal transcript splice donor site also gave bands of the predicted size for transcript D1 (lanes 2 and 3). A primer placed upstream of the distal germline promoter gave no product, indicating the absence of transcription originating further upstream (lane 1). We confirmed this







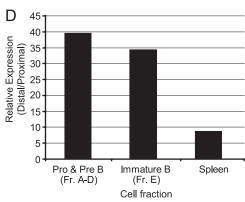


Fig. 3. The distal germline J_K promoter is far more active than the proximal germline Jk promoter in developing pre-B cells. (A) Schematic of the Jk cluster. The transcription start sites for both the distal and proximal promoters are shown as arrows above the diagram along with the deduced splicing pattern of the germline transcripts in dashed lines. The unfilled box at the left represents the first exon of one of the distal-promoter transcripts (D1), and the gray box the first exon of the other (D2). The two different splice donor sites are indicated. Approximate primer locations for both real-time RT-PCR (B) and gel-based RT-PCR (C) of Jk transcripts are shown as small arrows. (B) Quantitative real-time PCR analysis of spliced Jk germline transcripts in sorted B cell subsets from wild-type mice. Distal promoter transcripts were quantified using primer #3 (transcript D1) or primer #9 (transcript D2) paired with the reverse primer in Ck. Proximal promoter derived transcripts (transcript P) were quantified using the same primer pair as for the alternatively spliced distal transcript D2. Harvested bone marrow from C57/BL6 mice was labeled as in (Fig. 1B) and sorted by flow cytometry into the indicated fractions before

result by amplifying this novel splice product from a bone marrow pre-B cell cDNA library and subjecting it to DNA sequence analysis (data not shown). In addition, Northern blot analysis revealed transcripts corresponding in size to the unspliced distal promoter transcript and a variant spliced from J κ 1 to C κ (data not shown). Hence the distal J κ transcript is alternatively spliced, producing at least two distinct mature transcripts (Fig. 3 λ 4 and Fig. S3).

Because of the presence of this alternatively spliced distal transcript, previous analyses of distal and proximal germline transcript abundance and promoter activity may be incorrect. To better analyze the relative amounts of transcription initiation from the distal and proximal promoters, we designed assays based on the 5' RNA ligase mediated rapid amplification of cDNA ends (5' RLM-RACE) protocol (Fig. S3) (19). RNA ligase is used to attach an adapter oligonucleotide to the decapped 5' end of mRNA which is then analyzed using a Taqman-based RT-PCR assay using start-site specific and adapter oligonucleotides, allowing the discrimination of transcripts based on their start site. As shown in Fig. 3D, the vast majority of transcription at each developmental stage is triggered by the distal promoter. Comparing this data with data obtained by conventional RT-PCR (Fig. 3B) allows us to conclude that most of the transcripts generally ascribed to the proximal promoter are actually due to an alternatively spliced form of the distal J κ germline transcript. Northern blot analysis confirmed that bone marrow pre-B cells express relatively little proximal promoter driven germline transcript (data not shown).

The Proximal $J_{\mathcal{K}}$ Promoter Is Only Weakly Active in Small Pre-B Cells.

Although the distal germline J_K promoter is active in all pre-B cells, the proximal promoter is active in only a small fraction. Given the low abundance of proximal germline transcript detected by the RLM-RACE assay in pre-B cells, we re-evaluated the expression of GFP in heterozygous κ^0 GFP knock-in mice. Using a stricter gating scheme we found, in contradiction to our previous report, fewer than 1% GFP positive cells amongst small pre-B cells (data not shown). This low percentage is in line with the RLM-RACE assay. Expression of GFP in all other B cell developmental subsets was similar to what we found previously (5).

Discussion

Our previous analysis of germline $J\kappa$ transcription, based on the $\kappa^0 GFP$ knock-in mouse, concluded that transcription was rare and stochastic in the pre-B cell population and that those rare transcriptionally active alleles were preferentially rearranged (5). However, the GFP insertion disrupted recombination of the locus and expression of the marker was not consistent with another report of biallelic germline transcription (15). We

harvesting cells for RNA isolation. Cntl represents a positive control RNA sample from AMuLV transformed pro-B cells. Values were normalized to Actin transcript abundance (± SD). Data are representative of two independent experiments. (C) RT-PCR analysis of Jk germline transcripts in primary pre-B cells. Pre-B cells from wild-type mice were sorted and processed for RNA isolation and subsequent RT-PCR. Various forward primers (numbered arrows) were paired with a common reverse primer in the C_K exon. An ethidium bromide-stained agarose gel analysis of the resultant products is shown. Lane marked L is a standard DNA size ladder. Numbers above the lanes indicate the forward primer used according to the diagram in (A). Data are representative of two independent experiments. (D) 5'RLM-RACE analysis of germline Igk transcripts in developing primary B cells. Primary cells were harvested and sorted from wild-type C57/BL6 mice and processed for RNA isolation and subsequent 5'RLM-RACE (strategy shown in Fig. S3). The graph represents the ratio of transcripts initiating from the distal and the proximal $\lg \kappa$ germline transcript promoters. No significant quantities of either transcript were detected in thymocytes or water controls. Data are representative of two independent experiments, with each PCR done in triplicate.

created the dκGT-hCD4 and Cκ-iYFP germline reporter mice in an attempt to reconcile these observations. Unexpectedly, the new mice revealed that germline J κ transcription is biallelic in all pre-B cells. In reconciling the new results with our previous report, we discovered that the two J κ germline promoters are not equivalent and that the vast majority of germline transcription in this locus derives from the distal germline promoter, 3.5 kb upstream of J κ 1. Analysis of the reporter mice in conjunction with 5' RLM-RACE revealed that, whereas the distal germline promoter becomes highly active in pre-B cells (fractions C' and D), the proximal germline promoter is only weakly active in pre-B cells and becomes somewhat more active as cells progress in development. The difference in activity between the two promoters has not been appreciated until now because the distal promoter generates two alternatively spliced transcripts, one of which completely overlaps with the proximal germline transcript (Fig. 3A). Hence transcripts from the distal and proximal promoters cannot be distinguished from one another by conventional RT-PCR.

The relative activity of the distal and proximal J κ germline transcript promoters (as determined by the 5' RLM-RACE assay and RT-PCR) matched well with the observed pattern of marker expression in both the d κ GT-hCD4 and κ ⁰GFP mice, and the overall amount of both germline transcripts is reflected in the $C\kappa$ -iYFP strain. Although the alternatively spliced distal transcript encompasses the GFP cDNA in κ^0 GFP mice, we believe that the unusually long 5' untranslated region (UTR) in the resulting mRNA prevents efficient translation of GFP, and, as a result, GFP is only translated from the proximal promoterderived transcript. These three $Ig\kappa$ knock-in mice are thus faithful reporters of the abundance of germline transcript originating from either promoter or from the locus as a whole.

Successful heavy chain locus rearrangement results in pre-B cell receptor (pre-BCR) signaling that induces several rounds of mitotic division and retargeting of recombinase activity to the light chain loci (20). Our new mice reveal that pre-BCR signaling results in the rapid onset of germline transcription of both $Ig\kappa$ alleles in all pre-B cells. It is unclear how allelic exclusion is maintained under such circumstances. It is possible that germline transcription is a requisite for recombination, but not sufficient, such that other modifications in addition to transcription are necessary for efficient rearrangement. Previous studies have demonstrated that locus accessibility increases in conjunction with increases in histone acetylation and methylation of histone H3K4 as cells transition from the pro-B to pre-B stage (20-23). However, it is not clear if any of these locus modifications occurs in an allele-specific manner to dictate preferential recombination. Similarly, although the Igk locus undergoes monoallelic DNA demethylation during B-cell development, this process occurs as a late step in the progressive modification of chromatin structure of the locus and it is unclear if the demethylation occurs before the onset of recombination (both $Ig\kappa$ alleles are largely methylated in pre-B cells) (22–24). The two $Ig\kappa$ alleles do re-localize in pre-B cells such that one allele tends to associate with heterochromatin (23, 25). However, given that germline transcription is biallelic and ubiquitous among pre-B cells, association with heterochromatin appears to have little effect on transcriptional competence. Although heterochromatin association is thought to be repressive, instances of transcriptional activity within this compartment have been noted previously (26, 27). Rather than serving to repress transcription, such localization may lead to preferential allele recombination by sequestering one allele away from the recombinase, chromatin modifying enzymes, or other nuclear compartments. Maintenance of transcriptional competence of the heterochromatin-associated allele may play a role in allowing this allele to recombine if the first fails to encode a suitable light chain. Another possibility is that germline transcription and accessibility of the $V\kappa$ gene segments is regulated on an allelic basis, perhaps by heterochromatin association. If the J κ cluster is in an "always on" state in pre-B cells, the regulated portion of Igk recombination may be accessibility of the $V\kappa$ segments. Whether $V\kappa$ germline transcription is allelically biased has not been explored.

Our results demonstrate that the distal $J\kappa$ promoter gives rise to the majority of germline transcript in this region. This promoter is approximately 3.5 kb upstream of the J κ cluster. This arrangement is very different from the germline promoters associated with other recombining loci that are within several hundred base pairs of the relevant gene segment (7). Recent experiments using the $TCR\alpha$ locus as a model found that blocking transcription from a germline promoter prevented rearrangement to downstream $J\alpha$ segments (9). However, the effect was local and only involved segments in close proximity to the transcriptional blockade; recombination to segments further downstream was unaffected. The distance from the distal germline $Ig\kappa$ promoter to the J κ cluster suggests that it might only weakly promote recombination at the J κ cluster. We propose that biallelic transcription from the distal promoter results in a modest increase in accessibility at the J κ cluster, resulting in infrequent, stochastic recombination. Once recombination occurs on either allele, the incoming V_{κ} promoter greatly increases accessibility to remaining J κ segments downstream, promoting further replacement rearrangements on the already rearranged allele until the recombinase is inactivated. In favor of this model, we found that the dκGT-hCD4 knock-in resulted in a slight recombination defect. We suggest that this defect is a result of the polyA transcription termination signal positioned between the distal promoter and the J κ gene segments (9). The polyA sequence prevents polymerase transit across the locus and further reduces the modest level of accessibility at the J κ cluster. However, because we cannot remove the polyA sequence from our targeted allele, we cannot test this formally. After a certain time period in the absence of a productive rearrangement (perhaps as levels of the pre-BCR decay or certain transcription factors increase), the proximal promoter might become increasingly active, enhancing local accessibility and rearrangement. Alternatively, proximal promoter activity, and thus $J\kappa$ accessibility, may increase in the setting of a receptor editing signal, a possibility we are currently exploring.

Methods

Generation of Knock-In Mice. Jκ-hCD4: A 400 bp fragment surrounding the reported distal Jk transcription start site and upstream of the 5' distal transcript splice site, was found to have greater than 60% homology between human, mouse, and dog and had robust promoter activity in luciferase assays. A QuickSite Mutagenesis kit (Stratagene) was used to introduce an AvrII restriction site 250 bp downstream of the transcription start site and mutate all ATG codons to alternate codons between the start site and AvrII site. A human CD4 cDNA lacking the 33 C-terminal amino acids (intracellular domain) and possessing a Y43I mutation (thought to abolish interaction with MHC class II) along with a SV40 intron/poly A and a floxed neomycin resistance gene were cloned into the AvrII site. This region was extended by 2 kb and 5 kb 5' and 3' homology arms and a diphtheria toxin subunit A (DTA) cassette to generate the final targeting construct (Fig. S1).

Cκ-iYFP. A floxed neomycin resistance gene followed by an IRES-YFP cassette (gift of Richard Locksley, University of California, San Francisco, CA) was cloned into the BsmBI site located 9 bp downstream of the stop codon of $C\kappa$. The fragment containing the $C\kappa$ exon was extended by 2 kb and 5.5 kb 5' and 3' homology arms and a DTA cassette to generate the final targeting

Targeting constructs were linearized and electroporated into Prm1-Cre ES cells followed by two weeks of neomycin selection. Neomycin resistant ES cell colonies were picked and initially screened for either knock-in by long distance PCR, PCR positive clones were further verified by Southern blot using probes outside of the homology arms. Two successfully targeted ES clones per construct were used for subsequent blastocyst injection. Both clones for each construct successfully gave rise to germline transmitting chimeric males. Males

were subsequently mated to 129/SvImJ females and the pups assayed for deletion of the neomycin resistance gene by PCR. Further breeding was used to remove the Prm1-Cre transgene from the background.

Flow Cytometry and Cell Sorting. Flow cytometric analysis of bone marrow and splenic B cell development in either strain was done as previously described (28, 29). For hCD4 knock-in mice, cells were additionally stained with human CD4 antibody (clone RPA-T4, eBiosciences). Cell sorting was done by flow cytometry or magnetic beads (Miltenyi). The identity of all antibodies is available upon request.

Transcript Analysis. Reverse transcription was done using SuperScript II or MoMLV-RT (Invitrogen) according to manufacturers instructions. Subsequent real time or conventional PCR was done using JumpStart Taq polymerase (Sigma) and Taqman labeled probes (for real time only). PCR cycling conditions were 95 °C for 4 min followed by 25 to 45 cycles of 95 °C for 30 sec, 60 °C for 30 sec, and 72 °C for1 min. Primer sequences are provided in

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Table S1. RLM-RACE was performed using a kit from Ambion. Linkerligated RNA was converted to first-strand cDNA using reverse transcriptase. This cDNA was analyzed using a nested PCR strategy that involved 14 cycles of amplification with linker and promoter-specific primers followed by a second round of amplification on an aliquot of the primary product using internal primers and a taqman probe (Fig. S3C). RLM-RACE products were cloned and subjected to sequence analysis to verify their identities. Individual cloned products served as absolute standards for quantification of distal and proximal promoter initiated transcript abundance by real-time taqman PCR.

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