

# Receptor Editing Occurs Frequently during Normal B Cell Development

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## Summary

Allelic exclusion is established in development through a feedback mechanism in which the assembled immunoglobulin (Ig) suppresses further V(D)J rearrangement. But Ig expression sometimes fails to prevent further rearrangement. In autoantibody transgenic mice, reactivity of immature B cells with autoantigen can induce receptor editing, in which allelic exclusion is transiently prevented or reversed through nested light chain gene rearrangement, often resulting in altered B cell receptor specificity. To determine the extent of receptor editing in a normal, non-Ig transgenic immune system, we took advantage of the fact that  $\lambda$  light chain genes usually rearrange after  $\kappa$  genes. This allowed us to analyze  $\kappa$  loci in IgM $\lambda^+$  cells to determine how frequently in-frame  $\kappa$  genes fail to suppress  $\lambda$  gene rearrangements. To do this, we analyzed recombined V $\kappa$ J $\kappa$  genes inactivated by subsequent recombining sequence (RS) rearrangement. RS rearrangements delete portions of the  $\kappa$  locus by a V(D)J recombinase-dependent mechanism, suggesting that they play a role in receptor editing. We show that RS recombination is frequently induced by, and inactivates, functionally rearranged  $\kappa$  loci, as nearly half (47%) of the RS-inactivated V $\kappa$ J $\kappa$  joins were in-frame. These findings suggest that receptor editing occurs at a surprisingly high frequency in normal B cells.

Key words: receptor editing • recombining sequence recombination • immune tolerance • B lymphocytes • V(D)J rearrangements

The fact that virtually all B cells express a single H and L chain prompted many studies to elucidate the underlying mechanism. One process that clearly contributes to allelic exclusion is the imprecision of V(D)J rearrangement that generates a maximum of one in-frame rearrangement per three attempts (1), but more active feedback processes are also involved. Classic studies showing the ability of a H chain transgene (2, 3) or an L chain transgene (4, and for review see reference 5) to mediate feedback suppression of H and L chain rearrangements, respectively, established important paradigms that have been widely accepted. But in the case of L chain allelic exclusion, this paradigm was weakened by an increasing number of “exceptions”, in which ongoing L chain rearrangement occurred despite expression of functional  $\kappa$  chain (6–9). Studies with autoantibody transgenic (Tg)<sup>1</sup> mice suggested that many of the exceptions to the feedback regulation model of L chain allelic exclusion could be explained by postulating self-tolerance-induced receptor editing (10–15). In addition, recent in

vitro studies (16–18) and analyses of autoantibody Ig knock-in mice (19, 20) have shown that L chain gene receptor editing can be an important mechanism of B cell tolerance. Despite these findings, it is unclear how frequently receptor editing is used for tolerance induction in normal, non-Ig Tg autoreactive B cells, in part because the extent of autoreactivity in the preselected B cell repertoire is unknown.

The organization of the  $\kappa$  locus, with arrangement of V $\kappa$  genes in both sense and antisense transcriptional orientations, the absence of D region gene segments, and the presence of several J $\kappa$  gene segments facilitates sequential, nested V $\kappa$ -to-J $\kappa$  rearrangement attempts (for review see reference 21). In developing B cells, these secondary rearrangements can both rescue receptor expression in cells that fail to assemble in-frame L chains (1, 22) and rescue autoreactive B cells from tolerance elimination by replacing rearranged  $\kappa$  genes with new ones that alter specificity (for review see reference 23). Another way that the organization of the  $\kappa$  locus promotes receptor editing is suggested by the existence of the conserved element known as recombining sequence (RS) in the mouse (or the homologous “ $\kappa$  deleting element” in humans; reference 24). RS is located ~25 kb

<sup>1</sup>Abbreviations used in this paper: FWR, framework region; J $\kappa$ D/+, heterozygous  $\kappa$  deficient germline genotype; RS, recombining sequence; Tg, transgenic.

downstream of the C $\kappa$  exon (25) and has no coding function (26), but undergoes V(D)J recombinase-dependent rearrangement that inactivates the  $\kappa$  locus by deletional rearrangements in *cis* (26–28) (see Fig. 1). In an autoantibody knock-in model system, RS rearrangements can inactivate functional  $\kappa$  genes (20), but the extent of RS-mediated receptor editing in normal B cells remains unknown.

One approach to estimate the extent of receptor editing in normal B cells is to analyze V(D)J recombinational remnants that are the predicted residue of editing. In mouse B cells, which contain both  $\kappa$  and  $\lambda$  L chain loci,  $\lambda$  gene rearrangement almost always occurs after  $\kappa$  rearrangement (for review see references 29, 30). Thus, if an appropriate  $\kappa$  gene is not assembled, rearrangement at the  $\lambda$  locus often follows. In  $\lambda^+$  B cells, RS rearrangements usually have deleted the C $\kappa$  loci (27, 28, 31) either by recombining to V $\kappa$ s, through the well characterized heptamer–nonamer recombination signal sequences (Fig. 1 B), or to heptamer sites in the J $\kappa$ –C $\kappa$  intron (Fig. 1 C) (27, 28, 32). Besides destroying the function of the  $\kappa$  locus, this latter mode of RS recombination has two important effects: first, unlike nested V $\kappa$ J $\kappa$  recombinations, it eliminates the C $\kappa$ -associated *cis*-acting enhancer elements that are critical for V $\kappa$ J $\kappa$  expression and rearrangement (33–36), and second, it retains any V $\kappa$ J $\kappa$  join that was previously adjacent to C $\kappa$ . This physiological knockout of regulatory sequences required for  $\kappa$  gene rearrangement thus “freezes” the locus, allowing an analysis of the V $\kappa$ J $\kappa$  gene that was assembled adjacent to the C $\kappa$  exon just before RS and  $\lambda$  gene rearrangement.

In this study, we have isolated such V $\kappa$ J $\kappa$  joins from a large number of individual IgM $^+$  $\lambda^+$  B cells and determined their nucleotide sequences in order to ascertain the extent to which RS inactivates functional  $\kappa$  genes in a normal, non-Ig Tg immune system. The results indicate that in normal IgM $^+$  B cells RS-mediated receptor editing is induced by and frequently inactivates functionally rearranged  $\kappa$  genes, probably because of immune tolerance.

## Materials and Methods

**Mice.** Mice homozygous for the targeted deletion of the J $\kappa$ –C $\kappa$  locus (J $\kappa$ C $\kappa$ D/J $\kappa$ C $\kappa$ D; a gift from D. Huszar, GenPharm International, San Jose, CA; reference 33) were maintained under specific pathogen-free conditions in the animal care facility at National Jewish Medical and Research Center. J $\kappa$ C $\kappa$ D/J $\kappa$ C $\kappa$ D mice were bred with B10.D2nSn/J mice to generate B10.D2nSn/J–J $\kappa$ C $\kappa$ D/+ mice (J $\kappa$ C $\kappa$ D/+), which were used at 6–8 wk of age.

**Cell Sorting and Genomic DNA Isolation.** Splenic cells from J $\kappa$ C $\kappa$ D/+ mice were isolated and stained with goat anti-mouse IgM-PE (Caltag Labs., San Francisco, CA) and goat anti-mouse  $\lambda$ -FITC (Fisher Scientific Co., Pittsburgh, PA) and sorted on an ELITE flow cytometer (Coulter Corp., Miami, FL) to collect IgM $^+$  $\lambda^+$  B cells. Genomic DNA was isolated from cells by overnight proteinase K digestion in lysis buffer (100 mM NaCl, 10 mM Tris-Cl, pH 8, 25 mM EDTA, 0.5% SDS) at 55°C, followed by phenol/chloroform extraction and EtOH precipitation.

**Analysis of Direct PCR Amplified Ig Rearrangements.** Genomic DNA from sorted cells was used as a template to amplify V $\kappa$ J $\kappa$ -intron-RS rearrangements. As shown in Fig. 1, primers A (de-

generate V $\kappa$  framework region [FWR]3; reference 37) and B (RS-101, 5' ACATGGAAGTTTTCCCGGGAGAATATG 3') amplified a product of ~1,450 bps (for J $\kappa$ 5) using an amplification profile of 1 min at 94°C, 1 min at 58°C and 1 min at 72°C for 30–35 cycles. The resulting V $\kappa$ J $\kappa$ 5-intron-RS products were gel isolated and cloned into the TA II vector (Invitrogen, Carlsbad, CA) and colonies were screened by hybridization using the IVS probe (1). PCR clones were sequenced by the dideoxy termination method (Sequenase; United States Biochemical Corp., Cleveland, OH) using vector-specific forward and reverse primers, as well as antisense J $\kappa$ 5-specific (5' CTAACATGAAAAC-CTGTGTCTTACACA 3') and RS-specific (5' AAAGCTA-CATTAGGGCTCAAATCTGA 3') primers. Both DNA strands from the PCR clones were sequenced over the V $\kappa$ J $\kappa$  joins to verify the reading frame.

**Production of  $\lambda^+$  Hybridomas.** Splenocytes from B10.D2nSn/J–J $\kappa$ C $\kappa$ D/+ mice were cultured in DMEM supplemented with 10% fetal bovine sera plus 50  $\mu$ g/ml LPS (*E. coli* LPS; Sigma Chemical Co., St. Louis, MO) for 3 d, then fused with NSO-bcl2 (38) myeloma cells for fusion 1 or SP2/0 (39) myeloma cells for fusions 2 through 5. Hybridoma supernatants were screened by ELISA for secretion of micrometer/liter Ig and the lack of  $\kappa$  Ig secretion.

**Analysis of Hybridoma Ig Rearrangements.** Hybridoma genomic DNA was digested with EcoRI or BamHI, then fractionated on 0.8% agarose gels, blotted to nylon membrane, and hybridized with the RS 0.8 (27) or IVS probes (Fig. 1 A). V $\kappa$ -RS rearrangements were identified by genomic Southern blot analysis using the RS probe and/or by PCR amplification using primer A and primer B to yield a PCR product of ~255 bps. V $\kappa$ J $\kappa$ -intron-RS rearrangements were identified by genomic Southern blot analysis using both the RS and IVS probes and/or by PCR amplification using primer C (J $\kappa$  intron, 5' CTGACTGCAGGTAGCGTG-GTCTTCTAG 3') and primer B, and amplified for isolation and sequencing using primers A and B. The resulting products were isolated from 1.8% low-melt agarose gels, cycle sequenced directly (Dye Terminator Cycle Sequencing Ready Reaction kit; PE Applied Biosystems, Norwalk, CT) and analyzed using an ABI 377 DNA Sequencer (PE Applied Biosystems). To obtain near full-length sequences of hybridoma 1-2A7, 1-2E11, 2-2H11, 3-15D6, 3-15C4 and 3-17B10 V $\kappa$  genes, a consensus FWR1 oligo (amino acids –1 through 8; 5' GGTGACATTGTGCT-GACCCAGTCTCCA 3') was used with antisense J $\kappa$ -intron oligos for PCR amplification, followed by cycle sequencing of the products. For hybrids 1-3E8, 2-15E11, 3-7G5, 4-1D2, 1-10A11 and 1-11A4, V $\kappa$  leader specific oligos (Ig Prime kit; Novagen, Madison, WI) were used for amplification.

**Cloning and Expression of V(D)J Rearrangements for Analysis of H/ $\kappa$ L Chain Pairing.** The H chain V(D)J and the L chain V $\kappa$ J $\kappa$ -RS rearrangements from hybridoma 2H11 were genomically cloned as previously described (40), with the modifications that  $\lambda$ Zap (Stratagene, La Jolla, CA) was the cloning vector and the RS and IVS probes were used to screen clones for the V $\kappa$ J $\kappa$ -RS rearrangement. The 6.5-kb EcoRI fragment containing the V(D)J rearrangement and the 4.0-kb EcoRI–XbaI fragment containing the V $\kappa$ J $\kappa$  rearrangement were gel isolated and ligated to pR $\mu$ Sal, a C $\mu$  expression vector (41) and pSV2-neo-C $\kappa$ , a C $\kappa$  expression vector (42) respectively. The H chain from hybridoma 15E11 was cloned by PCR amplification using a leader intron oligo (5' GAAGTGGCAGGACCTGAGGTGAAAATGACA 3') and an oligo that spanned the XbaI site downstream of J $\mu$ 4 (5' CAG-GCTCCACCAGACCTCTCTAGA 3'). The resulting product was digested with EcoO109I and XbaI and ligated into EcoO109I and XbaI digested 8-1C $\mu$  (40), a C $\mu$  expression vector. The

V $\kappa$ J $\kappa$ -RS rearrangement from hybridoma 15E11 was also cloned by PCR using a leader intron oligo (5' TGGAATTCAGGT-TCTACTGGAGACATTGT-3') and an oligo which spanned the XbaI site downstream of J $\kappa$ 5 (5' ACGAATTCGTCTA-GAAGACCACGCTACCT 3'). The resulting product was digested with EcoRI and subcloned into a shuttle vector containing V $\kappa$ 21C leader and promoter elements. An XbaI fragment that contained the promoter elements, leader, and the 15E11 V $\kappa$ J $\kappa$  rearrangement was isolated and cloned into the XbaI site in pSV2-neo-C $\kappa$  (42). The 2H11 and 15E11 H and L chain constructs were then cotransfected into SP2/0 myeloma cells and selected for expression of IgM $\kappa$  as previously described (40).

**IgM $\kappa$  ELISA for Analysis of H/ $\kappa$ L Pairing.** Supernatants from 2H11 and 15E11 H and L chain transfectoma clones were assayed for IgM $\kappa$  expression by ELISA. In brief, goat anti-mouse IgM (Southern Biotechnology Associates, Inc., Birmingham, AL) was diluted in PBS, coated onto 96-well Immulon 2HB plates (Dynex Technologies, Inc., Chantilly, VA) and incubated at room temperature for 3 h. Plates were washed five times with PBS/Tween 20, then incubated with blocking buffer (PBS, 0.5% BSA, 0.4% Tween 20) for 1 h at room temperature. Serial dilution of transfectoma and parental hybridoma supernatants were added and incubated at room temperature for 2 h. Plates were washed and horseradish peroxidase-conjugated goat anti-mouse  $\kappa$  (Southern Biotechnology Associates, Inc.) was added and incubated for 2 h. After a final wash, the chromogenic substrate 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) (Sigma Chemical Co.) was added in McIlvain's buffer (84 mM Na<sub>2</sub>PO<sub>4</sub>/48 mM citrate, pH 4.6) with 0.005% H<sub>2</sub>O<sub>2</sub> and OD 410 nm was read using an automated plate reader (Dynatech, Alexandria, VA). Transfectoma and hybridoma antibody concentrations were estimated by comparison to a TEPC 183 ( $\mu\kappa$ ) standard curve.

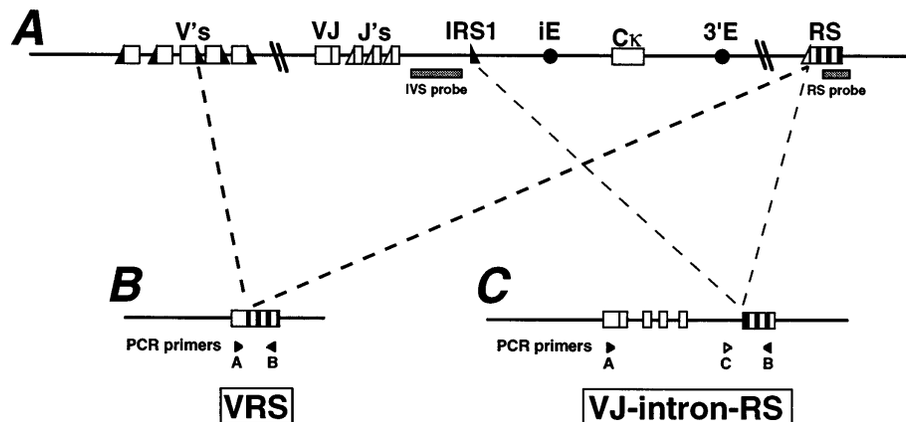
## Results

**Strategy for the Isolation of Editing Remnants.** To determine the extent to which RS recombination inactivates functional, in-frame V $\kappa$ J $\kappa$  joins in the preimmune B cell repertoire, IgM<sup>+</sup> $\lambda$ <sup>+</sup> splenic B cells were isolated by fluorescence activated cell sorting and their genomic DNA was analyzed by the PCR strategy outlined in Fig. 1 C. This cell sorting strategy should exclude from the template pool cells that are  $\kappa$ <sup>+</sup>, H chain isotype switched, surface (s)Ig<sup>-</sup>, or cells of a slg<sup>lo</sup>, germinal center phenotype. In a second series of ex-

periments, IgM $\lambda$  secreting hybridomas were isolated and their  $\kappa$  loci analyzed in detail. To simplify these analyses, all B cells analyzed were heterozygous for a targeted deletion of the J $\kappa$ -C $\kappa$  locus (J $\kappa$ D/+), in which only a single  $\kappa$  locus and RS allele could rearrange (33). The potential  $\kappa$  gene and RS element rearrangements are depicted in Fig. 1.

**Analysis of IgM $\lambda$  Cells Reveals Frequent Receptor Editing.** Genomic DNA from sorted IgM $\lambda$  cells was used as template for a PCR using a panspecific V $\kappa$  FWR 3 oligonucleotide primer, which recognizes ~80% of V $\kappa$  genes (37), together with an RS-specific primer to amplify V $\kappa$ J $\kappa$ -intron-RS rearrangements (Fig. 1 C, primers A and B). V $\kappa$ J $\kappa$ -intron-RS rearrangements containing V $\kappa$  genes rearranged to each of the four functional J $\kappa$  genes were detected by PCR amplification and Southern blotting (data not shown), but V $\kappa$ J $\kappa$ 5-intron-RS rearrangements were most abundant, in part because their smaller size promoted preferential amplification. Amplified V $\kappa$ J $\kappa$ 5-intron-RS rearrangements were gel-purified and cloned, and a total of 52 clones were sequenced across both the V $\kappa$ J $\kappa$  and the J $\kappa$ -intron-RS joins (Fig. 2). These two different recombination joins, present on each PCR product analyzed, provided markers for uniqueness. PCR products that were identical to one another, or that differed by just one nucleotide, were assumed to represent repeated isolates derived from the same initial template (i.e., derived from a single B cell clone). This represents an underestimate because the single base changes could have reflected real differences and because it was possible that some of the apparent repeats were independent events that happened to have identity in the portions of the genes studied, but not in upstream portions of the V genes. In this sample, at least 37 of the 52 clones represented independent events. Analysis of the V $\kappa$ J $\kappa$  join sequences allowed an assessment of the potential prior functionality of the V $\kappa$ J $\kappa$ 5 joins just upstream of intron-RS rearrangements. Surprisingly, 15 of the 37 clones (41%) contained V $\kappa$ J $\kappa$  joins that were in-frame (Fig. 2), and if the apparent repeats were not excluded 23 out of 52 (44%) were in-frame.

To verify the analysis of the PCR-amplified V $\kappa$ J $\kappa$ -intron-RS rearrangements and to increase the sample size, an independent sampling of V $\kappa$ J $\kappa$ -intron-RS rearrangements was derived from J $\kappa$ D/+ splenocytes in the form of B



**Figure 1.** RS rearrangements inactivate and preserve V $\kappa$ J $\kappa$  joins. A rearranged, potentially functional  $\kappa$  locus (A) can be silenced by two types of RS recombination: V $\kappa$ -RS (B) or V $\kappa$ J $\kappa$ -intron-RS (C). Type C retains the prior V $\kappa$ J $\kappa$  join, and the RS recombination event eliminates the known *cis* acting elements that are critical for efficient rearrangement and expression, thus freezing the locus from further V $\kappa$ J $\kappa$  recombination. Also shown are the intronic recombination sequence 1 (IRS1) (32), the intronic (iE) and 3' kappa (3'E) enhancers (35, 36), and the recombining sequence (RS) (27, 28) element. Probes IVS (1) and RS 0.8 (27, 28) are indicated by filled boxes.

**A**

CLONE	V $\kappa$	FWR3	CDR3	J $\kappa$ 5	IRS 1	RS	REPEATS
		70	88		TTTCCTGAGGA/ $\Delta$ AGGACCCCTAGT		
1	1	DFSLNIHPMEEDDTMYFC	QQSKEV	LTF	TTTCCT	CCCTAGT	2
2	4/5	SYSLIISMEAEADAATYYC	QQRSSYP	TF	TTTCCT	AGGACCCCTAGT	
3	4/5	SYSLTISRMEAEADAATYYC	QQRSSYP	LTF	TTT	GACCCCTAGT	
4	4/5	SYSLTISRMEAEADAATYYC	QQRSSYP	LTF	TTTCCT	GACCCCTAGT	
5	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP	LTF	TTTCCTGAG	AGT	3
6	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP	TF	TT	TAGT	
7	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP	LTF	TTTCCTG	GGACCCCTAGT	2
8	4/5	SYSLTISRMEAEADAATYYC	QQRSSSH	LTF	TTTCCTGAGG	CCCTAGT	
9	4/5	SYSLTISRMEAEADAATYYC	QQRSSYP	LTF	TTTCCTGAG	AGT	
10	4/5	SYSLTISRMEAEADAATYYC	QQRSSYP	LTF	TTTC	ACCCCTAGT	2
11	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP	LTF	TTTCCTGAG T	AGGACCCCTAGT	2
12	21	DFTLNHPVEEEDAATYYC	QQRSSNP	LTF	TTTCCT	GACCCCTAGT	
13	23	DFTLINSVETEDFGMYFC	QQRSSNP	LTF	TTTCCTGAGG	CCCTAGT	
14	23	DFTLINSVETEDFGMYFC	QQRSSNP	LTF	TTTCCTGAGG	CTAGT	
15	24/25	AFTLRISRVEAEDVGVYVC	MQHLEYF	TF	TTTCCTG	CCCTAGT	3

**B**

CLONE	V $\kappa$	FWR3	CDR3	IRS 1	RS	REPEATS
		70	88	TTTCCTGAGGA/ $\Delta$ AGGACCCCTAGT		
1	1	DFTLKRISRVEAEDLVYVC	FQGSVH*	TTTCCT	GACCCCTAGT	
2	1	DFTLKRISRVEAEDLVYVC	LQSTHP*	TTTCCT	AGGACCCCTAGT	
3	2	DFTLKRISRVEAEDLVYVC	MQGTHP*	TTTCCTGA	GACCCCTAGT	
4	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP*	TTTCCT	AGGACCCCTAGT	
5	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP*	TTTCCTGA	CT AGGACCCCTAGT	
6	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP*	TTTCCT	GACCCCTAGT	
7	4/5	SYSLTISRMEAEADAATYYC	QQRSSNP*	TTTCCT	GACCCCTAGT	
8	8	DFTLTISSVQAEADLVYVC	HQYLS*	TTTCCT	GACCCCTAGT	3
9	8	DFTLTISSVQAEADLVYVC	HQYLS*	TTTCCTG	AGGACCCCTAGT	
10	8	DFTLTISSVQAEADLVYVC	HQYLS*	TTTCCT	GACCCCTAGT	
11	8	DFTLTISSVQAEADLVYVC	HQYLS*	TTTCCT	AGGACCCCTAGT	
12	9	DYSLTISSESEDFADYYC	VQYAE*	TTTCCTG	AGT	
13	10	DYSLTISNLEQEDVYFC	QQRNTL*	TTTCCT	AGGACCCCTAGT	
14	10	DYSLTISNLEQEDVYFC	QQRNTL*	TTTCCT	ACCCCTAGT	
15	19/28	DFTLTISSVQAEADLVYFC	QQRSSNP*	TTTCCTGAG	AGGACCCCTAGT	
16	23	DYSLTISNLEQEDVYFC	LQYST*	TTTCCTGAG	A CCTAGT	2
17	23	DFTLINSVETEDFGMYFC	QQRSSNP*			
18	24/25	DFTLKRISRVEAEDVGVYVC	MQHLEY*	TTT	CCCTAGT	2
19	24/25	AFTLRISRVEAEDVGVYVC	MQHLEY*	TTTCCTGA	GACCCCTAGT	
20	24/25	AFTLRISRVEAEDVGVYVC	MQHLEY*	TTTCCTGAG	AGT	
21	24/25	AFTLRISRVEAEDVGVYVC	MQHLEY*	TTTCCT	GT	
22	24/25	DFTLKRISRVEAEDVGVYVC	MQHLEY*	TTTCCTG	AGGACCCCTAGT	4

**Figure 2.** Sequence analysis of (A) productive and (B) nonproductive V $\kappa$ J $\kappa$ -intron-RS rearrangements from FACS<sup>®</sup> sorted, IgM<sup>+</sup> $\lambda$ <sup>+</sup> splenic B cells. V $\kappa$  gene family and J $\kappa$  gene usage were assigned based on homologies to expressed V $\kappa$ J $\kappa$  genes (52) or homology searches of Genbank and the Kabat Ig database. Translated V $\kappa$  FWR3 (codons 70–88), CDR3, and J $\kappa$ 5 sequences to the conserved phenylalanine (F) are shown for productive rearrangements, whereas FWR3 and CDR3 sequences are shown for nonproductive rearrangements, with the asterisk (\*) adjacent to CDR3 in B denoting an out of frame V $\kappa$ J $\kappa$  join. The nucleotide sequences of the unrearranged J $\kappa$  intronic recombining sequence 1 (IRS1) and RS element (both of which contain a consensus heptamer sequence adjacent to the  $\Delta$  symbol) are shown above the sequences of the IRS1-RS joins present in each PCR clone. The RS join sequence for clone 17 was not determined. Underlined nucleotides could have been donated by either the IRS1 or the RS sequence, and N region addition (*bold*) and P-encoded nucleotides are shown between the joins. Repeats denote the number of times a particular sequence was observed.

cell hybridomas. A total of 133 IgM $\lambda$ -expressing hybrids were obtained from five separate fusions and their  $\kappa$  locus rearrangements were analyzed. Genomic Southern blot and PCR analysis revealed that at least 74% of the  $\lambda$ <sup>+</sup> hybrids (99 out of 133) had inactivated the wild-type  $\kappa$  locus by RS rearrangements (Table 1), a value in accord with previous estimates (31, 34). Two hybridomas apparently had undergone inversional V $\kappa$ -RS rearrangements, as they had unique restriction fragments that retained the C $\kappa$  locus as revealed by the intron (IVS) probe (data not shown), but scored positive in a V $\kappa$ -RS PCR (Fig. 1 B). Approximately 25% (26 out of 99) of the hybridomas with RS rearrangements had J $\kappa$ -intron-RS joins (Table 1), as detected with primers B and C (Fig. 1 C). Genomic Southern blot analy-

**Table 1.**  $\kappa$  Locus Rearrangement Status of IgM $\lambda$  Hybridomas

	$\kappa$ locus genotype				
	Hybridomas	RS <sup>-</sup>	V $\kappa$ -RS	V $\kappa$ J $\kappa$ -RS	Productive/ nonproductive
	<i>n</i>				
Fusion 1	37	12	16	9/9*	5/4
Fusion 2	44	10	30	4/4	2/2
Fusion 3	12	1	6	5/4	4/0
Fusion 4	25	9	11	5/2	1/1
Fusion 5	15	2	10	3/1	0/1
Totals	133	34	73	26/20	12/8

RS<sup>-</sup> denotes hybridomas that lacked a detectable RS rearrangement. V $\kappa$ -RS and V $\kappa$ J $\kappa$ -RS are defined in Fig. 1. Asterisk indicates total number of V $\kappa$ J $\kappa$ -intron-RS loci that were isolated followed by the number that we were able to PCR amplify with the consensus FWR3 oligo.

sis of 18 out of 20 hybrids scoring PCR positive for J $\kappa$ -intron-RS rearrangements demonstrated that the RS rearrangements colocalized with EcoRI restriction fragments hybridizing with the IVS probe (data not shown), thus independently confirming the V $\kappa$ J $\kappa$ -intron-RS rearrangement phenotype. V $\kappa$ J $\kappa$ -intron-RS rearrangements from individual hybridomas were PCR amplified and directly sequenced, rather than cloned, a procedure that diminishes potential Taq polymerase-generated mutations. Like the V $\kappa$ J $\kappa$ -intron-RS PCR clones, most of the V $\kappa$ J $\kappa$ -intron-RS loci from hybridomas used J $\kappa$ 5, although four hybridomas had rearrangements to upstream J $\kappa$ s, including one to J $\kappa$ 2 and three to J $\kappa$ 4, suggesting that developing B cells do not frequently undergo RS rearrangement until all of the J $\kappa$ s are rearranged. Sequence analysis over both the V $\kappa$ J $\kappa$  and intron-RS joins clearly showed that each cell line had a unique sequence at the V $\kappa$ J $\kappa$  join and that, remarkably, 12 of 20 (60%) of the V $\kappa$ J $\kappa$  joins were in-frame (Fig. 3 A).

**Diversity of V Gene Usage.** V $\kappa$ J $\kappa$ -intron-RS rearrangements were clearly diverse because at least 32 different V $\kappa$  genes representing 11 of the 19 V $\kappa$  families were identified among the 57 independent V $\kappa$ J $\kappa$ -intron-RS loci analyzed (data not shown). This value of V $\kappa$  gene representation in the V $\kappa$ J $\kappa$ -intron-RS loci analyzed is most likely an underestimate of the diversity because the 5' PCR primer lies in FWR3 and yields only a short stretch of V $\kappa$  gene sequence for interclonal comparison. Despite this limitation, multiple genes were observed within particular V $\kappa$  gene families. For example, within the V $\kappa$ 4/5 family, at least 11 different genes were represented among the 14 in-frame and 7 out-of-frame joins (data not shown). In addition, V $\kappa$  genes were sometimes found repeatedly in independent V $\kappa$ J $\kappa$ -intron-RS rearrangements and there was considerable overlap in usage among hybridoma and PCR clone sequences. 13 of the hybridomas used V $\kappa$  genes that were observed in

A				FWR/CDR SEQUENCES							
HYBRID	V <sub>K</sub>	J <sub>K</sub>	P/NP	FWR1	CDR1	FWR2	CDR2	FWR3	CDR3	FWR4	
1-2A7	4/5	5	P	DIVLTQSPALMSASPGKRVITMC	SARSSVSSSYLY	WYQQRPGSSPKLWLY	STSNLAS	GVPARFSGSGSGTYSYLTISSMEAEADAATFYC	QQYSGYPS	TF	
1-2E11	4/5	5	P	DIVLTQSPALMSASPGKRVITMC	SARSSVSSSYLY	WYQQRSGSSPKLWLY	STSNLAS	GVPARFSGSGSGTYSYLTISSMEAEADAATFYC	QQWSSNPP	TF	
2-2H11	4/5	5	P	DIVLTQSPALMSASPGKRVITMC	RASSSVSYMH	WYQQRPGSSPKLWLY	ATSNLAS	GVPARFSGSGSGTYSYLTISSVEAEADAATFYC	QQWSSNPPM	LTF	
3-15D6	4/5	5	P	DIVLTQSPALMSASPGKRVITMC	SARSSVSSSYLY	WYQQRSGASPKLWLY	RTSNLAS	GVPARFSGSGSGTYSYLTISSVEAEADAATFYC	QQWSSGYP	LTF	
1-3E8	19	2	P	DIVMTQSPKRFMTSIVGDRVSTIC	KASQWGTAVA	WYQQRPGSSPKLWLY	SASVRYT	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	QQYSGTPIH	TF	
2-15E11	19	5	P	DIVMTQSPKRFMTSIVGDRVSTIC	KASQWGTAVA	WYQQRPGSSPKLWLY	WASVRYT	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	QQYSGTPIH	F	
3-15C4	19	5	P	DIVLTQSPKRFMTSIVGDRVSTIC	KASQWGTAVA	WYQQRSGHSPKLLIK	YASQSLG	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	QQWSSYPL	LTF	
1-10A11	23	5	P	DIVLTQSPALMSASPGKRVITMC	RASQSLSNMLH	WYQQRSGHSPKLLIK	YASQSLG	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	QQWSSW	LTF	
1-11A4	23	5	P	DIVLTQSPALMSASPGKRVITMC	RASQSLSNMLH	WYQQRSGHSPKLLIK	YASQSLG	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	QQWSSW	LTF	
3-7G5	24/25	5	P	DIVMTQAAPSVPVTPGSEVSLSC	RSSKSLHSMGNTLYL	WELQRPQSPQLLY	RMSNLAS	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	MQHLEYF	TF	
4-1D2	24/25	4	P	DIVMTQAAPSVPVTPGSEVSLSC	RSSKSLHSMGNTLYL	WELQRPQSPQLLY	RMSNLAS	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	MQHLEYF	TF	
3-17B10	32	5	P	DIVLTQSPALMSASPGKRVITMC	ITSDIDDDDM	WYQQRPGSSPKLWLY	EGNLRP	GVDPDRFSGSGSGTDFPTLTISVQAEADLAVFYC	LQSDMPL	LTF	
B											
HYBRID	IR51	RS	HYBRID	IR51	RS						
TTTCCTGAGGA/ΔVAGGACCTAGT		TTTCCTGAGGA/ΔVAGGACCTAGT									
1-2A7	TTTCCT	AGGACCTAGT	4-12F2	TT	GT						
1-2E11	TTTCCT	AGGACCTAGT	1-2H6	TTTCCTGAGG	CCCTAGT						
2-2H11	TTT	CCCTAGT	2-7B5	TTTCCT	AGGACCTAGT						
3-15D6	TTT	CCCTAGT	5-6C2	TTTCCT	AGGACCTAGT						
1-3E8	TTTCCT	AGGACCTAGT	1-6C9	TTTC	CCCTAGT						
2-15E11	TTTCCTGAGG	CCCTAGT	1-5B6	TTTCCT	AGGACCTAGT						
3-15C4	TTTCCT	AGGACCTAGT	2-14A7	TTT	CCCTAGT						
1-10A11	TTTCCT	AGGACCTAGT	1-9E9	TTTCCTGA	AGGACCTAGT						
1-11A4	TTTC	CCCTAGT									
3-7G5	TTTCCT	AGGACCTAGT									
4-1D2	TTTCCT	AGGACCTAGT									
3-17B10	TTT	CCCTAGT									

**Figure 3.** Sequence analysis of V<sub>K</sub>J<sub>K</sub>-intron-RS rearrangements from IgM<sub>κ</sub> hybridomas. (A) Sequences of the V<sub>K</sub>J<sub>K</sub> rearrangements. The first digit in the hybridoma name indicates the fusion experiment number. Myeloma fusion partners were either NSO-bcl2 (fusion 1) or SP2/0 (fusions 2–5). V<sub>K</sub> gene family and J<sub>K</sub> gene usage were assigned as described in Fig. 2. P and NP denote productive and nonproductive V<sub>K</sub>J<sub>K</sub> rearrangements, respectively. Translated amino acid sequences of V<sub>K</sub> FWR, CDR, and J<sub>K</sub> sequences to the conserved phenylalanine (F) residue are shown for productive rearrangements, and V<sub>K</sub> FWR and CDR sequences, with \* denoting an out of frame V<sub>K</sub>J<sub>K</sub> join and # denoting an in-frame stop codon, are shown for nonproductive rearrangements. These sequence data are available from EMBL/Genbank/DBJ under accession numbers AF087023–AF087034 and AF087460–AF087467. (B) Sequences of the RS rearrangements (as described in Fig. 2).

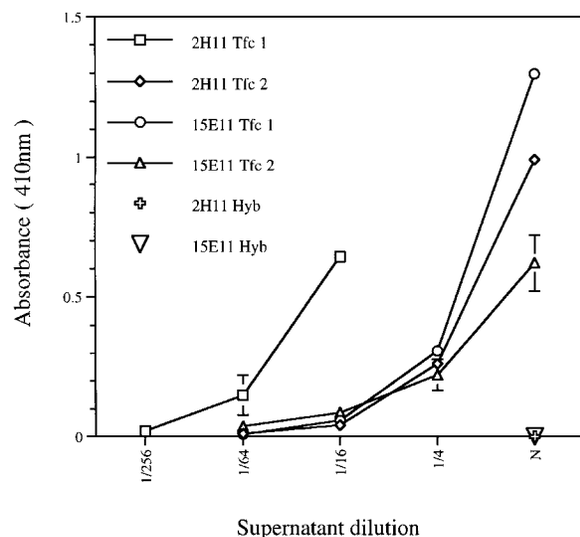
direct PCR-derived clones, whereas 6 hybridomas expressed distinct V<sub>K</sub> genes that were members of families observed in the PCR clone sample and 1 hybridoma expressed a V<sub>K</sub>32 gene, a V<sub>K</sub> family not seen in the PCR clone samples (Figs. 2 and 3 and data not shown).

**Intron/RS Joins.** The sequences of the J<sub>K</sub>-intron-RS joins in both the PCR clones (Fig. 2) and hybridomas (Fig. 3 B) were quite varied and were dominated by deletions at both sides of the joins, as up to nine nucleotides were missing from either the J<sub>K</sub>-intron or RS heptamer-flanking sequences. There did appear to be a bias for a particular join (e.g., clone 4, Fig. 2 A), which was observed to be associated with 13 independent V<sub>K</sub>J<sub>K</sub> rearrangements. Two of the intron-RS joins contained P nucleotides and one contained N-region addition nucleotides, consistent with findings described previously (7, 43).

**Rebuilding IgM<sub>κ</sub> Antibodies for Analysis of H/κL Pairing and Antigen Specificity.** To determine if the high frequency of in-frame V<sub>K</sub>J<sub>K</sub> rearrangements silenced by intron-RS recombination was due to the inability of H chains to pair with their κ L chains, the V(D)J and V<sub>K</sub>J<sub>K</sub> rearrangements from hybridomas 2H11 and 15E11 were cloned into C<sub>μ</sub> and C<sub>κ</sub> expression vectors, respectively. These H and L chain constructs were cotransfected into SP2/0 myeloma cells to generate transfectoma clones. Analysis of transfectoma supernatants by IgM<sub>κ</sub> sandwich ELISA revealed that the in-frame κL chains were able to pair with their hybridoma H chains (Fig. 4), suggesting that ongoing RS rearrangement was not due to the inability of H/L chain pairing. The specificity of the μκ transfectoma antibodies remains unknown, however. Attempts in flow cytometry assays to detect recombinant antibody binding to the surfaces of bone marrow cells were unsuccessful (data not shown).

## Discussion

In this report we examined the DNA sequences of V<sub>K</sub>J<sub>K</sub> joins located upstream of intronic-RS rearrangements in normal, non-Ig Tg B cells to determine the extent to



**Figure 4.** “Repair” of intron-RS recombination-silenced V<sub>K</sub>J<sub>K</sub> genes by restoration of C<sub>κ</sub> exon and surrounding elements reveals that silenced κL chains can pair with their original μ chain partner. The graph shows representative results from a μκ ELISA comparing several IgM<sub>κ</sub> transfectoma antibodies (Tfc) to their IgM<sub>κ</sub> parental hybridoma antibodies (Hyb). Antibodies in supernatants were captured on plastic using adsorbed anti-μ chain antibodies and revealed with anti-κ conjugates. Bars indicate the SD determined from antibodies assayed in triplicate. The concentrations of the hybridoma antibodies were at least 10-fold higher than those of the transfectoma antibodies based on comparison to a TEPC 183 (μ, κ) standard curve.

which RS-mediated recombination silences functionally rearranged  $\kappa$  genes. Nearly half of all  $V\kappa J\kappa$  joins inactivated by RS recombination were in-frame (27 out of 57). This high frequency is clearly incompatible with a strict feedback suppression model of L chain allelic exclusion, which predicts no in-frame  $V\kappa J\kappa$  joins upstream of the RS rearrangements. More strikingly, this high frequency is also significantly higher than 33%, the percentage of in-frame joins expected from random  $V\kappa J\kappa$  rearrangement, indicating that productive  $V\kappa J\kappa$  rearrangements actively induce intron-RS rearrangements. The data also demonstrate a physiological role for the RS element in normal B cell development—the inactivation of functionally rearranged  $\kappa$  genes.

To understand why we conclude that the RS rearrangements were actively induced by functional  $\kappa$  L chains, consider the extreme hypothetical cases of mice in which all  $\kappa$  gene rearrangements result in either autoreactive B cell receptors or nonproductive  $\kappa$  chains (Table 2). If  $V\kappa$ -to- $J\kappa$  and RS rearrangements proceed randomly, albeit with different relative frequencies, then in either case  $V\kappa J\kappa$  joins located upstream of intronic-RS rearrangements should be in-frame at a maximum frequency of one out of three. To significantly exceed this frequency, in-frame  $V\kappa J\kappa$  joins must stimulate the relative rate of (intronic) RS rearrangement. This argument applies to our data because the observed frequency of in-frame joins, 47.4%, is significantly higher than one out of three ( $P < 0.04$ , single sample test of a proportion based on a normal approximation). Since it is exceedingly unlikely that the stimulus for increased in-frame rearrangements is mediated by anything other than  $\kappa$  protein, and because  $\kappa$  chains can probably only be perceived by the signaling machinery of B cells through their association with H chains, we conclude that functional  $\kappa$  chains actively stimulate the rate of RS rearrangement based on B cell receptor antigenic specificity. These data also predict that in mice in which the  $C\kappa$  exon is inactivated, but surrounding *cis*-acting elements are left intact,  $V\kappa J\kappa$  rearrangement should be extensive, whereas RS rearrangement should be reduced. This is in fact the experimental observation (44).

The statistical argument also excludes the possibility that a high frequency of rearrangeable  $V\kappa$  pseudogenes, L chains that fail to pair with H chains, or a role for positive selection is responsible for our results. Furthermore, complete sequencing of the coding regions from all the in-frame  $V\kappa J\kappa$  rearrangements derived from  $\lambda^+$  hybridomas revealed no stop codons or other obvious defects that would have precluded function (Fig. 3 A). It is also unlikely that frequent aberrant H/L chain pairing is responsible for the high frequency of in-frame  $V\kappa J\kappa$  joins in the  $V\kappa J\kappa$ -intron-RS rearrangements, as demonstrated by the ability of H/ $\kappa$ L chains from two hybridomas to pair (Fig. 4). Moreover, there are few examples of L chains that fail to pair with H chains and most experiments suggest that virtually all random H/L pairs can associate (40, 45–48). Finally, if a lack of positive selection of surface Ig was responsible for the high frequency of in-frame joins, this would predict that B cells should frequently express two  $\kappa$  chains, a result that has not been observed.

The receptor editing events documented in this study probably do not represent renewed V(D)J recombination in mature B cells, such as has recently been described in the germinal center (49–51), because the cells analyzed expressed high levels of IgM and  $\lambda$  chain and because they were isolated and, in the case of the hybridomas, stimulated in a manner that should not have induced V(D)J recombination. Another indication that receptor editing in mature B cells is unlikely to explain our results is that the fraction of  $\lambda^+$  cells in newly formed and mature splenic B cells is nearly identical, suggesting that in unmanipulated mice mature  $\kappa^+$  cells rarely give rise to  $\lambda^+$  B cells (44). Overall, it would appear from our data that the RS rearrangements that we studied were actually stimulated, rather than inhibited, by productive  $\kappa$  gene rearrangements, probably as the result of immune tolerance-mediated receptor editing in immature B cells. To definitively test the prediction that the  $\kappa$  chains of the cells that we have analyzed generate autoantibodies in association with the same cell's heavy chain, it will be necessary to generate mice transgenic for these genes.

**Table 2.** Analysis of  $V\kappa J\kappa$  Joins in  $V\kappa J\kappa$ -intron-RS Sequences: Models and Predictions Compared with Experimental Data

Models	Predicted fraction in-frame	Experimental data	Observed percentage in frame
Perfect feedback regulation with functional $\kappa$ chain preventing $\lambda$ rearrangement	0%	27 out of 57 $V\kappa J\kappa$ -intron-RS loci	47.4% ( $P < 0.04$ )
Poor feedback regulation	$\leq 33\%$		
High frequency of $V\kappa$ pseudogenes	$\leq 33\%$		
High frequency of H/ $\kappa$ chain mispairing	$\leq 33\%$		
Any combination of the above	$\leq 33\%$		
Extreme model of editing with random RS rearrangements and all $\kappa$ s autoreactive	33%		

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