An immunoglobulin Cκ-reactive single chain antibody fusion protein induces tolerance through receptor editing in a normal polyclonal immune system

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Understanding immune tolerance mechanisms is a major goal of immunology research, but mechanistic studies have generally required the use of mouse models carrying untargeted or targeted antigen receptor transgenes, which distort lymphocyte development and therefore preclude analysis of a truly normal immune system. Here we demonstrate an advance in in vivo analysis of immune tolerance that overcomes these shortcomings. We show that custom superantigens generated by single chain antibody technology permit the study of tolerance in a normal, polyclonal immune system. In the present study we generated a membrane-tethered anti-Igκ/H9260-reactive single chain antibody chimeric gene and expressed it as a transgene in mice. B cell tolerance was directly characterized in the transgenic mice and in radiation bone marrow chimeras in which ligand-bearing mice served as recipients of nontransgenic cells. We find that the ubiquitously expressed, Igκ-reactive ligand induces efficient B cell tolerance primarily or exclusively by receptor editing. We also demonstrate the unique advantages of our model in the genetic and cellular analysis of immune tolerance.

Immunological tolerance is important in preventing autoimmunity and in promoting the efficiency of immune responses. Tolerance has been assessed experimentally in two general ways. First, tolerance in normal polyclonal individuals has been measured by evaluating responses revealed by subsequent immunization in vivo or in vitro (for review see references 1, 2). This first approach has the advantage of addressing a normal immune system, but because of the extreme heterogeneity of lymphocyte specificity and the low precursor frequency of antigen-specific lymphocytes, it makes analysis of tolerance mechanisms difficult or impossible. A second approach has involved the use of antigen receptor transgenic mice, which artificially increase the frequency of antigen-reactive cells by limiting or fixing expression of one or both receptor chains, permitting the visualization of cells with fully or partly defined specificities (3–6). Antigen receptor transgensics have become indispensable reagents in the study of lymphocyte biology, allowing analysis of development, selection, specificity, and memory. But transgenic models are designed to distort lymphocyte development, subverting the “allelic exclusion” mechanisms to generate quasi-monoclonal immune systems in defined lymphocyte subsets. Furthermore, antibody transgensics, or the more recent approach of targeted transgensics, have a number of inherent drawbacks, owing to nonphysiological changes, such as accelerated B cell development, a tendency to skewing in B cell subset, and nonphysiological D4/V to VDJH joining (7–10). Therefore, a variety of experimental conclusions based on the use of antigen receptor transgenic mice, in particular the relative contributions of different tolerance mechanisms to central and peripheral immune tolerance, require verification in normal, polyclonal immune systems.

One mechanism by which tolerance can occur in B cells is receptor editing, in which self-reactivity directly or indirectly induces ongoing recombinase activator gene (RAG) expression and secondary Ig light chain rearrangements that alter receptor specificity (6,
11–13). Receptor editing may be a major self-tolerance mechanism in immature B cells (14–20). Many studies indicate that the precursor frequency of autoreactive B cells is high and that autoreactivity may be corrected by light chain exchange (11, 12, 14, 21–26). Recent studies by Wardemann et al. show that reverse genetic analysis of human B cell antibody expression can be a fruitful approach to study repertoire and tolerance (23, 26). However, to date, studies demonstrating in vivo receptor editing have primarily relied upon analysis of receptor gene transgenic mice, the limitations of which have been discussed.

On the other hand, immune responses can sometimes also be analyzed in polyclonal models of the immune system, and with appropriate experimental tools this analysis can provide additional advantages over receptor gene transgenics because lymphocyte development is not radically altered. For example, lymphocyte responses can be modeled by stimulation with anti-Ig and anti-TCR reagents (27–34), or through the use of TCRβ-reactive superantigens, which among other things provided early evidence for clonal deletion and T cell anergy in lymphocytes (35, 36). B cell–reactive superantigens also exist, notably HIV gp120, Staphylococcal toxin protein A, and Peptostreptococcus protein L (37–39).

The premise of this study is that custom superantigens, generated by single chain Fv antibody engineering technology, can be expressed as transgenes to regulate or tolerate a fully normal, polyclonal immune system. To test this idea, we generated mice expressing a single chain antibody combining site reactive to the constant portion of mouse Ig L chain, which was engineered to be expressed as a membrane protein. We then used these mice to assess the mechanisms of B cell tolerance in vivo in a polyclonal immune system. This approach provides the advantages of a ligand that reacts identically with the antigen receptors of a high frequency of precursor cells, but does so without prior skewing of lymphocyte repertoire or subset, and in fact can be used with entirely normal cells. We call our synthetic superantigens “macroself” antigens to distinguish them from natural superantigens. We show here that a ubiquitously expressed Igκ-macroself antigen can promote central tolerance and receptor editing in a polyclonal immune system in vivo. Furthermore, we show that transgenic mice expressing κ-macroself antigen can facilitate the analysis of mutations that affect tolerance processes.

RESULTS

Generation of κ-macroself antigen constructs and transgenic mice

To express an Igκ-specific macroself antigen of the predicted structure indicated schematically in Fig. 1 A, we assembled the gene construct depicted in Fig. 1 B. The antigen specificity was generated by forming a single chain Fv from the variable genes of a rat anti–mouse Cκ hybridoma. We also chose to include hinge regions and Fc portion from rat IgG1 to promote protein stability, bivalency, and flexibility, and to project the binding sites away from the plasma membrane. The transmembrane and intracytoplasmic regions of the protein were derived from the H–2Kb gene, which we assumed were compatible with ubiquitous cell surface expression. A construct carrying the chimeric gene driven by a human cytomegalovirus promoter and zeocin resistance gene, generating plasmid pmsCA187ΔCH1. (C) Flow cytometry analysis of stably transfected cell lines. Two clones were analyzed for surface expression of the macroself Ag. (Left) Staining with an anti–rat IgG1 monoclonal antibody compared with empty vector-transfected control. (Right) Testing of binding specificity of the macroself antigen to mouse Ig light chain isotype. Two transfected cell lines were incubated with soluble mouse IgG2b,κ or IgG2b,λ and binding revealed with a secondary rat anti–mouse IgG2a/b reagent.

Figure 1. Design and in vitro testing of a synthetic B cell superantigen. (A) Schematic representation of the predicted protein structure of membrane bound anti–mouse Igκ-macroself Ag. A single chain Fv generated from the anti–κ hybridoma 187 is linked to the hinge and membrane proximal domains of rat IgG1 followed by transmembrane and cytoplasmic tail regions (Tm/Cy) of H–2Kb. (B) Gene construct encoding κ-macroself antigen showing intron/exon structure and selected features. Introns are depicted as thin lines. (G,G,S) refers to linker codons in one letter amino acid code: GGGGSGGGGGS. For stable transfection analysis, the gene shown was inserted into an expression vector providing a human cytomegalovirus promoter and zeocin resistance gene, generating plasmid pmsCA187ΔCH1. (C) Flow cytometry analysis of stably transfected cell lines. Two clones were analyzed for surface expression of the macroself Ag. (Left) Staining with an anti–rat IgG1 monoclonal antibody compared with empty vector–transfected control. (Right) Testing of binding specificity of the macroself antigen to mouse Ig light chain isotype. Two transfected cell lines were incubated with soluble mouse IgG2b,κ or IgG2b,λ and binding revealed with a secondary rat anti–mouse IgG2a/b reagent.
mothers that were Ig antibodies, microinjected zygotes were implanted in foster mouse MHC class I H-2K and the ubiquitin C promoter was chosen because it prosion in transient transfection by

Figure 2. Generation of κ-macroself antigen transgenics and in vivo transgene expression. (A) Schematic representation of DNA construct used for microinjection to generate transgenic mice. Elements shown are approximately to scale. The pUlik construct is a derivative of the gene shown in Fig. 1 B. It contains the κ-macroself Ag under the control of the human ubiquitin C promoter. The construct contains a Ig light chain gene leader exon and first intron. (B) Flow cytometry analysis of κ-macroself antigen expression in bone marrow, spleen, and lymph nodes of pUlik transgenic mice as detected using an anti–rat IgG1Fc antibody (dotted lines, nontransgenic cells; solid line, pUlik transgenic cells). Results from four different pUlik transgenic lines are shown.

ated using the construct shown in Fig. 2 A. A leader intron was included in this construct because it improved expression in transient transfection by approximately 10-fold (unpublished data) and the ubiquitin C promoter was chosen because it provided a more uniform and ubiquitous expression than a mouse MHC class I H-2Kb promoter tested (unpublished data). To avoid potential toxicity caused by maternal Igκ antibodies, microinjected zygotes were implanted in foster mothers that were Igκ deficient (40). Several transgenic lines were generated, four of which were selected for further study. As shown in Fig. 2 B, all lines expressed macroself antigen on virtually all tested cells: lines #2, #37, and #20 expressed at relatively high, uniform levels in lymphoid tissues, whereas line #26 was expressed more weakly, particularly in bone marrow.

Flow cytometry analyses were performed to determine if the κ-macroself antigen influenced the development of κ-expressing B cells and to see if there was an effect on λ+ B cells. All four transgenic lines had a similar, striking phenotype: in the κ-macroself transgenic mice, virtually all of the B220+ cells expressed λ L chains and failed to express κ L chains (shown in Fig. 3 for line #2 and #26 splenocytes, and summarized for all lines in Fig. 4, D and E). As shown in the lower right panels of Fig. 3, ~75% of B cells in the spleens of κ-macroself antigen mice were stained with a monoclonal antibody to λ1–3. The remaining cells likely expressed Vκx because these cells expressed slgM, but failed to react to anti-κ or anti-λ1–3 antibodies; moreover, κ-deficient mice (κ−/−) had a similar population of IgM+λ1–3 cells (Fig. 3). Similar losses of κ+ cells and increases in λ+ cells were observed in lymph nodes of the transgenic mice (Fig. 4, D and E). Peripheral B cell numbers were reduced ~50% in all κ-macroself antigen transgenic lines (total splenic cells and the B220+ fraction were reduced to, respectively, 80 and 60% of control levels; Fig. 4, A and C); in contrast, bone marrow B220+ cell numbers were unchanged in transgenic mice (Fig. 4 C). The percentages of λ+ cells in peripheral lymphoid tissues of κ-macroself transgenic mice were increased relative to nontransgenics by over sevenfold (Fig. 4 E), representing an overall numerical increase of three- to fourfold. These results indicate that in all κ-macroself transgenic lines κ+ cells were absent from the peripheral lymphoid organs and λ+ cell numbers were substantially increased.

Flow cytometry was also used to look for differences between normal and κ-macroself antigen-expressing mice in peripheral B cell subsets (Fig. 4, F and G). The κ-macroself antigen has the unique advantages of possessing comparable reactivity to receptors carried by most B cells present in all
target populations. Among B220\(^+\) splenic cells there appeared to be similar or somewhat increased percentages of newly formed splenic B cells in k-macroself transgenic lines compared with littermate controls, whereas the percentages of mature, follicular B cells were somewhat reduced and those of marginal zone B cells increased (Fig. 4 F). Taking into account the reduction of B220\(^+\) cells in the k-macroself antigen-expressing mice, we estimate that marginal zone B cell numbers were not significantly reduced in the k-macroself antigen transgenic mice, whereas follicular and newly formed B cells were reduced by ~50 and 20%, respectively. Peritoneal B-1 cells in the k-macroself antigen transgenic mice were similar in number to control littermates, but, like other peripheral B cells in these mice, expressed \(\lambda\), rather than K L chains (Fig. 4 G). Consistent with these observations, analysis of serum IgM and IgG2a/b revealed a loss of Ig\(\kappa\) and an increase in \(\lambda\)-carrying Igs in k-macroself transgenic mice (Table I). Thus, the immune system appears to readily adapt to the presence of an autoantigen reactive to 94% of normal B cell receptors.

### Analysis of receptor editing

To determine if the increase in \(\lambda\)\(^+\) B cells in k-macroself antigen-expressing mice was the result of receptor editing, rather than proliferation of preexisting \(\lambda\)\(^+\) cells, we first quantitated in the bone marrow the frequency of newly formed, B220\(\text{intermediate}/\lambda\)\(^+\) cells, a measurement that has been used as an indicator of the rate of new B cell production (41). As shown in Fig. 5, \(\lambda\) B cell production is increased three- to fourfold in k-macroself antigen-expressing mice compared with nontransgenic littermates. This increase was comparable to the increase seen in k\(^{-/-}\) mice, in which bone marrow output of \(\lambda\)\(^+\) cells is known to be elevated (41). To independently measure the rate of new \(\lambda\) B cell production, we quantitated Ig\(\lambda\) gene excision products (Fig. 6, A and C). These episomal DNAs generated by gene assembly fail to replicate upon cell division, and hence are a sensitive indicator of receptor editing (11). As shown in Fig. 6 C, the levels of VA\(^+\)-to-JA\(^+\) excision products were elevated approximately threefold in isolated B220\(^+\) bone marrow cells of k-macroself antigen-expressing mice, relative to littermate controls, consistent with the estimated increased bone marrow output of \(\lambda\)\(^+\) cells. An even greater increase in the relative levels of \(\lambda\) excision product was seen in splenic B220\(^+\) cells of transgenic mice (Fig. 6 C), suggesting that most splenic cells had undergone little or no proliferation since their generation. Once again, k-macroself antigen-expressing mice revealed a phenotype similar to k\(^{-/-}\) mice. We conclude that k-macroself transgenic mice have an increased rate of \(\lambda\) gene rearrangement and \(\lambda\)\(^+\) B cell production in the bone marrow.

Independent indices of receptor editing were measured to verify the additional predictions that recombine expression and destructive Ig\(\kappa\) rearrangements should be increased in k-macroself transgenic mice. RAG1 and RAG2 mRNA expression levels in k-macroself transgenic bone marrow B220\(^+\)

### Table I. Serum immunoglobulins of unimmunized k-macroself transgenic mice

<table>
<thead>
<tr>
<th>Mouse strain (n)</th>
<th>Total IgM</th>
<th>IgM, (\kappa)</th>
<th>IgM, (\lambda)</th>
<th>IgG2a/b, (\kappa)</th>
<th>IgG2a/b, (\lambda)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nontransgenic (4)</td>
<td>113 ± 78</td>
<td>75 ± 21</td>
<td>6 ± 4</td>
<td>211.0 ± 91.0</td>
<td>0.6 ± 0.1</td>
</tr>
<tr>
<td>k(^{-/-}) (4)</td>
<td>163 ± 60</td>
<td>&lt; 0.6</td>
<td>79 ± 39</td>
<td>0.6 ± 0.1</td>
<td>42.0 ± 8.0</td>
</tr>
<tr>
<td>Transgenic #2 (4)</td>
<td>188 ± 81</td>
<td>&lt; 0.6</td>
<td>76 ± 33</td>
<td>0.7 ± 0.2</td>
<td>101 ± 29.0</td>
</tr>
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cells were significantly elevated by 80 to 90% over nontransgenic controls (Fig. 6 B). Recombining sequence (RS; 42, 43) recombination to the acceptor sites in the J-Ck intron was also evaluated as a measure of destructive V(D)J recombination on the κ locus of B cells (Fig. 6 A). RS, which is found ~25-kb downstream of the Cκ exon, undergoes RAG-mediated recombination to Vκs and to two acceptor sites in the J-Cκ intron (IRS1 and IRS2) that inactivate the κ locus (44). RS recombinations are common in λ+ cells and are also seen in ~10% of normal κ+ cells (16, 45, 46). RS recombination to IRS1 and IRS2 was elevated in bone marrow and spleens of κ-macroself transgenic mice (Fig. 6 A, RS to κ intron rows). Taken together with showing similar increases in RAG expression and λ+ B cell production in κ-deficient and κ-macroself transgenic mice, the data suggested that κ-macroself antigen induces efficient central B cell tolerance in an otherwise normal immune system by a mechanism of developmental arrest and receptor editing.

Analysis of bone marrow B cell turnover and intracellular Ig expression

To further quantify the relative roles of receptor editing and clonal deletion in the tolerance induced by κ-macroself antigen, the intracellular Igκ expression and BrdU uptake of newly formed bone marrow B cells was assessed. Despite the data indicating excess RAG expression, RS recombination and λ gene excision product DNA in the bone marrow of κ-macroself mice, it remained possible that clonal deletion depleted developing κ+ cells, allowing excess survival and a nonproliferative buildup of λ+ cells. Such a hypothesis predicts an increased turnover of B cells in κ-macroself bone marrow and a drastic reduction of developing B cells carrying κ chains, whereas editing should not increase B cell turnover. Turnover was assessed by daily injection of BrdU and assessment of uptake in bone marrow B cells over time (47). As shown in Fig. 7 A (left), BrdU uptake in the B220intermediate bone marrow B cell population was similar or slightly slower in κ-macroself transgenic mice, compared with wild-type littermates, indicating that the κ-macroself antigen does not accelerate bone marrow B cell turnover. BrdU uptake was delayed in the slgM fraction of newly formed κ-macroself transgenic B cells compared with wild type, suggesting a longer average transit time (Fig. 7 A, right). These results were further supported by intracelluar immunoglobulin staining experiments as κ-macroself antigen mice had significant numbers of intracellular κ+ B cells in bone marrow, but not in the spleen (Fig. 7 B, lower rows). The frequency of cytoplasmic Igκ+ cells was only marginally reduced (by ~13%) among the newly formed, B220intermediate B cells in the presence of κ-macroself antigen (Fig. 7 D). Moreover, cytoplasmic κ+ cells cotained an IgM antibody (Fig. 7 C). Because both the anti-κ and anti-IgM antibodies used see only assembled immunoglobulins (48,

![Figure 5. Increased output of Igκ+ cells in κ-macroself antigen mice.](Image 1)

![Figure 6. Evaluation of molecular parameters of receptor editing in κ-macroself transgenic, littermate control, and κ−/− mice.](Image 2)
we infer that the bone marrow B cells of macroself antigen mice that carry slgM/B220<sup>int</sup> undergo receptor down modulation followed by receptor editing.

**Transplantation of mutant bone marrow in macroself-expressing hosts**

Because the macroself transgenics express antigen on all cells tested, including B cells, we assessed B cell tolerance in radiation bone marrow chimeras in which macroself transgenic line #2 mice served as hosts and donor bone marrow cells lacked macroself transgenes (Fig. 8 and Table II). Three types of bone marrow donors were used in these experiments: nonmutant controls (wt), mice carrying a hemizygous deficiency in the RAG1 gene (RAG<sup>−/−</sup>), and mice carrying a Bcl2 transgene expressed in the B cell compartment (51). Use of wild-type donors assessed the ability of macroself antigen to promote tolerance in a population of cells that was completely normal and lacking in any genetic

\[\text{Figure 7. Intracellular immunofluorescence analysis of BrdU uptake and immunoglobulin expression in macroself transgenic mice and littermates. (A) BrdU uptake with time of labeling in bone marrow B220<sup>int</sup> cells of transgenic mice (open circles) or littermates (diamonds). (Left) BrdU incorporation in total B220<sup>int</sup> cells; (right) BrdU incorporation in slgM<sup>+</sup>/B220<sup>int</sup> cells. (B) Comparison of surface lgk staining (top), with intracellular lgk staining (bottom). (C) Costaining for intracellular lgk and either slgM alone (left) or both surface and cytoplasmic lgM (right). (D) Statistical analysis of experiments shown in b and c. Left pair of bars shows the percentages of cells found in lower analysis boxes of B220/intracellular lgk stain (i.e., B, bottom). The right pair of bars shows percentages of cells in upper two quadrants of intracellular lgM/intracellular lgk stain with B220<sup>int</sup> gate (C, right). Means and standard deviations are indicated. Filled bars, transgenic; open bars, nontransgenic littermates.}

\[\text{Figure 8. Flow cytometry analysis of tolerance induction in radiation chimeras using macroself transgenic hosts. Mice were analyzed at 6 wk post reconstitution. (A and B) Analysis of bone marrow lymphocytes using anti-κ chain and B220 antibodies. Bone marrow donors are indicated to the left of the arrows, the recipient mouse genotypes are shown just below. Newly formed lymphocytes carrying Ig-κ or -λ were identified as falling in the lower analysis boxes, as indicated. (A) B6 (wt) or RAG<sup>−/−</sup> bone marrow was used to reconstitute lethally irradiated CD45.1<sup>−/−</sup>/CD45.2<sup>+</sup> macroself transgenic #2 (Tg #2) or littermate recipients. (B) Comparison of chimeras generated with bone marrow from Bcl-2 transgenic or littermate (wt) donors, using as irradiated recipients either transgenic #2 or littermate. (C) Analysis of spleen cells from the indicated radiation chimeras. Recipient mouse genotype is shown to the right of arrows above dot plots. Cells were stained with B220 and anti-κ antibodies (top) or anti-λ antibodies (bottom).}
Table II. Effect of Bcl2 overexpression and RAG1+/− mutation in donor cells on receptor editing and tolerance induction in κ-macroself radiation chimeras

<table>
<thead>
<tr>
<th>Donor → Host (n)</th>
<th>Cells /10⁶</th>
<th>B220</th>
<th>IgM</th>
<th>κ</th>
<th>λ</th>
<th>κ+/B220int</th>
<th>λ−B220int</th>
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<tbody>
<tr>
<td>Bone marrow</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Wild type</td>
<td>→ wt (5)</td>
<td>68 ± 6.6</td>
<td>78 ± 13</td>
<td>15 ± 2</td>
<td>21.0 ± 2.0</td>
<td>2.7 ± 0.5</td>
<td>12.8 ± 2.0</td>
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<tr>
<td>→ Tg#2 (4)</td>
<td></td>
<td>50 ± 13</td>
<td>74 ± 9</td>
<td>8 ± 3a</td>
<td>5.2 ± 1.8a</td>
<td>4.9 ± 1.4a</td>
<td>4.4 ± 1.3a</td>
</tr>
<tr>
<td>Bcl2</td>
<td>→ wt (4)</td>
<td>56 ± 13</td>
<td>84 ± 5</td>
<td>35 ± 2</td>
<td>33.0 ± 7.0</td>
<td>8.6 ± 1.2</td>
<td>22.5 ± 3.0</td>
</tr>
<tr>
<td>→ Tg#2 (3)</td>
<td></td>
<td>61 ± 17</td>
<td>74 ± 8</td>
<td>22 ± 2a</td>
<td>5.7 ± 0.4a</td>
<td>15.0 ± 2.5a</td>
<td>4.5 ± 0.4a</td>
</tr>
<tr>
<td>RAG1+/−</td>
<td>→ wt (4)</td>
<td>52 ± 7</td>
<td>67 ± 11</td>
<td>16 ± 2</td>
<td>21.0 ± 3.0</td>
<td>2.3 ± 0.7</td>
<td>9.7 ± 0.4</td>
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<tr>
<td>→ Tg#2 (3)</td>
<td></td>
<td>42 ± 2</td>
<td>60 ± 10</td>
<td>6 ± 2a</td>
<td>8.3 ± 2.0a</td>
<td>2.5 ± 0.7</td>
<td>7.4 ± 0.5a</td>
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<tr>
<td>Spleen</td>
<td></td>
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<tr>
<td>Wild type</td>
<td>→ wt (5)</td>
<td>67 ± 23</td>
<td>66 ± 3</td>
<td>50 ± 2</td>
<td>5.0 ± 6.0</td>
<td>4.0 ± 1.0</td>
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<tr>
<td>→ Tg#2 (4)</td>
<td></td>
<td>43 ± 7a</td>
<td>36 ± 7a</td>
<td>28 ± 5a</td>
<td>0.6 ± 0.3a</td>
<td>27.0 ± 7.0a</td>
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<tr>
<td>Bcl2</td>
<td>→ wt (4)</td>
<td>146 ± 33</td>
<td>84 ± 3</td>
<td>63 ± 14</td>
<td>65.0 ± 8.0</td>
<td>10.0 ± 2.0</td>
<td></td>
</tr>
<tr>
<td>→ Tg#2 (3)</td>
<td></td>
<td>83 ± 20a</td>
<td>73 ± 15</td>
<td>49 ± 10</td>
<td>1.0 ± 0.6a</td>
<td>41.0 ± 5.0a</td>
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</tr>
<tr>
<td>RAG1+/−</td>
<td>→ wt (4)</td>
<td>64 ± 6</td>
<td>68 ± 4</td>
<td>50 ± 3</td>
<td>61.0 ± 4.0</td>
<td>4.7 ± 0.5</td>
<td></td>
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<tr>
<td>→ Tg#2 (3)</td>
<td></td>
<td>39 ± 18a</td>
<td>29 ± 6a</td>
<td>26 ± 3a</td>
<td>0.2 ± 0.1a</td>
<td>19.0 ± 7.0a</td>
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Bone marrow sample data shows total cell counts, but percentages of B cell subsets are based on a lymphocyte gate. B220int refers to cells with intermediate levels of B220 and the indicated Ig light chain.

Items marked represent significant (P < 0.05) differences associated with the mutant donor type, compared to wild-type donor type in the comparable host. Items in bold represent significant (P < 0.05) differences between cells maturing in transgenic #2 compared to wild-type littermate hosts.

modifications. The results observed in line #2 hosts carrying wild-type donor cells were generally similar to those obtained in unmanipulated line #2 mice, except that cells in the bone marrow carrying low levels of Igκ were more apparent (Fig. 8 A, compare lower left pair of panels). Splenic B cells in transgenic #2 recipients were largely devoid of κ expression, but instead expressed λ chains (Fig. 8 C, right panels under “wt”). In addition, transgenic #2 recipients manifested an increased frequency of B220intmediate λ+ cells in the bone marrow (Fig. 8 A). Thus, nontransgenic B cells developing in κ-macroself line #2 appeared to undergo central tolerance and receptor editing.

The RAG+/− mutant and Bcl2 transgenic donors were tested to determine the feasibility of using κ-macroself transgenics to rapidly screen mouse mutant strains for defects in immune tolerance. We reasoned that enforced Bcl2 expression should suppress apoptosis, whereas heterozygous deficiency in RAG1 might hinder receptor editing. Bone marrow chimeras carrying RAG+/− or Bcl2 transgenic bone marrow manifested subtle, but reproducible, differences from chimeras reconstituted with wild-type bone marrow (shown in bold in Table II). Chimeras reconstituted with Bcl2 transgenic marrow had elevated frequencies of λ+ cells in the absence of κ-macroself antigen, and an even further enhanced λ frequency in κ-macroself recipients (Fig. 8 C); furthermore, these increases were paralleled by a rise in newly formed B220intmediate/λ+ cells in the bone marrow (Fig. 8 B). We conclude that the survival-enhancing effects of enforced Bcl2 expression do not block central B cell tolerance to κ-macroself antigen, but instead promote increased generation of λ+ cells. This increase in λ, along with the further-reduced frequency of κ+ cells in the bone marrows of κ-macroself recipients reconstituted with Bcl2 transgenic cells, likely occurs because of an extended lifetime for editing. In contrast, chimeras reconstituted with RAG+/− bone marrow had a subnormal peripheral B cell frequency that was exacerbated by the presence of κ-macroself antigen. Furthermore, RAG+/− cells had a significantly smaller κ-macroself antigen–induced increase in the rate of λ+ B cell production relative to wild type (Fig. 8 A). An additional anomaly observed in RAG+/− chimeras was a relative retention in the bone marrow of κ-macroself antigen recipients of cells carrying high levels of Igκ (Fig. 8 A, lower right). Thus, loss of surface Igκ expression in bone marrow B220+ cells of κ-macroself antigen transgenic mice is partly related to editing and is not wholly a result of receptor protein down modulation. Similar results were obtained in RAG+/− mice carrying the line #26 transgene (unpublished data). These results suggest that reduced RAG1 gene dosage suppresses or, more likely, slows tolerance-induced receptor editing, reducing the κ to λ isotype switch in developing B cells. This finding in turn may suggest that RAG1 levels are limiting for receptor editing.

**DISCUSSION**

The κ-macroself antigen approach that we introduce here is a strategy to study immunological tolerance in a large polyclonal cohort of cells. In the present study, we have used these mice to study immune tolerance in an otherwise unmanipulated immune system. We were able to revisit the
question of whether or not B cell tolerance to a ubiquitous, membrane-tethered self antigen occurs by receptor editing or deletion (or if it occurs at all in normal cells). Importantly, our approach allowed us to assess the relative contributions of editing and deletion in a normal immune system. The data indicate that both cell death and receptor editing contribute to tolerance because substantial increases in light chain gene recombination and new λ⁺ B cell production occur in κ⁺-macroself antigen mice, but their overall B cell numbers are reduced to ~50%. It is known that λ loci almost always recombine after κ loci (45, 52), typically ~24 h later (53, 54). However, we believe that apoptosis induced rapidly by autoreactivity (clonal deletion) appears to play a minimal role in our model, whereas the cellular time limit for editing to eliminate an autoreactive receptor exerts a more severe restriction. Because ~94% of B cells in non-transgenic mice express κ, the reduction in B cell numbers is less than predicted if generated κ⁺ B cells are rapidly eliminated by cell death and not replaced by other cells. Indeed, B cells in κ⁺-macroself transgenic mice resembled those of κ⁻/⁻ mice, which cannot be subject to negative selection of sIgκ (Fig. 3). Tolerance purely by rapid clonal deletion of developing B cells would be predicted to reduce κ⁺ B cell output while leaving λ⁺ B cell generation unchanged. Results of BrdU uptake studies and Igκ cytoplasmic staining were incompatible with significant rapid clonal deletion. Indeed, in the presence of κ⁺-macroself antigen the turnover of immature B cells was actually slowed, while the fraction of newly formed λ⁺ cells increased substantially. Rather, the results are most easily explained by a tolerance-induced developmental arrest, followed by significant, but incomplete, rescue by editing.

If editing occurs by developmental arrest without induced deletion in κ⁺-macroself transgenic mice these transgensics should resemble κ⁻/⁻ mice. This is in fact the case: their percentages of B cells in various lymphoid compartments are quite similar. We have assessed mice carrying κ⁺-macroself antigen on a background in which one of the κ constant alleles carried the human sequence (17) and found minimal B cell attrition compared with wild type (unpublished data). This would be consistent with findings indicating that the efficiency of rescue of autoreactive cells by editing is visibly affected only by severely reducing the number of L chain genes available for editing, such as by knocking out the κ locus, or limiting the available repertoire of Jκs (20, 25, 55–57). In contrast to these studies, however, our present results were obtained in a context in which B cell development was unaltered.

Anti-Ig suppression of B cell development in vivo and in vitro has been studied for many years (27–32, 34, 58–60). Work first done in chickens, then later in mice and rats, showing that antibodies introduced into developing embryos suppressed B cell development represented the first evidence for clonal abortion (for review see reference 61). In some studies, evidence of developmental arrest was seen, including “irreversible” receptor down-regulation (28, 59). Typically, mature B cells were absent from the periphery while B cell progenitors were retained in the bone marrow. However, in other studies, anergy was induced, rather than deletion (30, 32). Often, competitive advantage was had by B cells that did not react with antibody (31, 58, and for review see reference 61). Our results are similar in several ways to a study of anti-κ suppressed mice by Weiss et al., in which the loss of κ⁺ B cells and κ immunoglobulin in serum was compensated for by an increase of peripheral λ⁺ B cells and serum immunoglobulin λ (31). In any case, the prior studies were often consistent with our conclusions in that the immature B cells receiving anti-Ig constant region stimulus did not die immediately, though none of these studies proposed an editing type escape mechanism.

Our ability to probe immune tolerance using macroself transgenic mice as adoptive hosts for normal or mutant bone marrow is important for several reasons. First, it establishes that the macroself antigen need not be expressed by B cells to induce tolerance in wild-type B cells. Second, we establish that wild-type B cells are tolerance susceptible by editing in vivo, a finding that to our knowledge has never before been directly demonstrated in a non-Ig transgenic mouse. Third, we demonstrate that macroself transgenic mice allow one to screen mutant bone marrow for defects in immunological tolerance. We confirmed earlier results, obtained in 3–83 antibody transgenic mice carrying autoreactive receptors, indicating that enforced B cell expression of Bcl2 facilitated receptor editing, but did not rescue autoreactive B cells from central tolerance (62). This last is consistent with the notion that the incomplete editing to λ in κ⁺-macroself antigen transgenic mice is limited by the life span of editing competent cells, which can be artificially prolonged by enforced Bcl2 expression. In other words, we believe that autoreactive cells arrested in development and undergoing light chain rearrangements are subject to death by neglect, which can be slowed by Bcl2. Furthermore, we have made the novel additional finding that reducing RAG1 gene dosage impairs B cell production and receptor editing. (This result was independently confirmed using conventional Ig transgenic mice; unpublished data).

Although antigen receptor transgenic mice are a valuable resource for many studies, by design they perturb lymphocyte development, resulting in a number of anomalies. For example, targeted or randomly integrated transgenes are typically prematurely expressed, and are generally in an unusual antigen receptor gene context, such as in conventional IgH transgenics, which cannot undergo H chain class switch, or in targeted VDJh transgenics, where the introduced gene is downstream of germline D elements and can be eliminated by DJ joining. Furthermore, antigen receptor transgenes usually skew lymphocyte subset distributions, such as CD4/CD8 ratios in T cells or B-1/follicular ratios in B cells. Macroself antigen mice overcome many of these drawbacks. Importantly, the macroself approach can, in theory, be easily used for any antigen receptor for which there is a monoclonal antibody.
We note that B cells have the striking propensity to alter their Ig constant region usage during development, a feature that makes their analysis by challenge with macroself Ags potentially fruitful. Early in development, when bone marrow B cells express exclusively slgM, receptor editing can cause successive Ig-k allele usage, and can lead to a progression from usage of \( \kappa \) to \( \lambda \) L chain. Later in development, B cells up-regulate slgD, marking the beginning of a phase that includes migration to the spleen and final preimmune maturation. After encounter with antigen, reactive B cells often undergo a heavy chain class switch, leading to the loss of IgM and IgD, and their replacement by downstream H chain C regions. Hence, macroself Ags reactive to Ig-k, IgD, and IgG C-regions would be expected to promote immune tolerance in a polyclonal immune system at different developmental stages, and quite possibly by different mechanisms. As macroself antigens with specificity to other antigen receptors, such as TCRs, can be easily generated, they may provide useful tools for studies in other cell types. Future studies will focus on such possibilities.

With the advent of germline mouse knockout technology, autoimmune-prone congenic strains, and genome-wide ethyl nitrosourea mutagenesis studies, there is an increasing need to devise methods to rapidly screen mutant mice for immune tolerance phenotypes. At the present time, this is typically approached by crossing of the knockout in question to antigen receptor transgenic mice and then introducing the cognate (auto)antigen. Because it requires a minimum of two generations of breeding, and extensive mouse genotyping, this approach is time consuming and expensive, and therefore not ideal for the screening of large numbers of mutants. Generation of well-conceived macroself Ag mice could provide a means to greatly speed up this screening process. Our results show that mutant mouse bone marrow or fetal liver precursor cells could be used to reconstitute lethally irradiated macroself Ag-expressing mice, and the development and function of the transfected cells could be directly monitored. Because the macroself Ag mice carry specific ligands that react with a subset of normal lymphocytes, no special breeding is required, particularly if the macroself Ag transgene is maintained on the same genetic background. Macroself Ag transgenic mice with specificity for defined lymphocyte receptor elements can facilitate the analysis of mouse mutants, particularly those with known signaling defects.

MATERIALS AND METHODS

Generation of \( n \)-macroself gene constructs. RNA was isolated from the rat anti-\( \kappa \) hybridoma 187(49) (American Type Culture Collection), and expressed antibody genes were cloned by 5′ rapid amplification and cloning of ends (5′-RACE) using the RLJ-RACE PCR kit (Ambion) according to the manufacturer’s instructions. The variable genes were amplified using sense primer specific to the 5′ adaptor either with a rat Ck-specific antisense primer (5′-CTAACTGTTCCGTTGGATGGTG) to amplify the light chain variable region, or a rat IgG1 anti-sense specific primer (5′-GGTCTCCACGTTCCAGGTACGG) to amplify the heavy chain variable region, together with their respective leader sequences. PCR, products were then cloned in a plasmid vector using the TOPO TA cloning kit (Invitrogen) and several clones sequenced to obtain consensus sequences and to facilitate the identification of clones lacking mutations introduced by the amplification and cloning. Independently, rat IgG1 H chain constant regions were amplified from cDNA. To generate a single chain antibody gene and linked elements a PCR sewing approach was taken using the following oligonucleotide primers: primer 1 (5′V\( _{\text{H}} \)) 5′-CCCGCAATCGTGGACGATGGC-3′; primer 2 (3′V\( _{\text{H}} \)) 5′-CCTCCAGTCCCGCTCCGTCGTCCTCCGGT-3′; primer 3 (5′V\( _{\text{L}} \)) 5′-TCTGATGACTGAGCCGAGATTCACGCTGCAA-3′; primer 4 (3′V\( _{\text{L}} \)) 5′-CCCGGATCTTGGGCGTCG-3′; primer 5 (5′Ch) 5′-CATAGAAACCCGCGGGTATTGCAAGCC-3′; primer 6 (3′Ch) GGGAGTGAGGAGAGACTTCTCTCATGATTGTGCGG; 7 (5′F) 5′-GTCTGCTCCACTCCCAGGGTTTAAAGAGCCTCCTCCATCCAC; 8 (3′F) 5′-CCAACCT-TCACCTGAATCTGGCC. The “overlapping” sequences are underlined; those recognized by restriction endonucleases are in italics. PCR products corresponding to the LV\( _{\text{H}} \) and the V\( _{\text{H}} \) region were initially amplified with the primers 1 and 2, and 3 and 4, respectively. Rat Ck\( _{\text{H}} \) cDNA lacking the CH1 domain (C\( \text{y1CH1} \)) was amplified using primers 5 and 6. A segment of the genomic H-2Kb locus encoding the transmembrane (Tm) exon 5 was amplified using primers 7 and 8. The single chain Fv gene, including the intervening flexible peptide codons (Gly\( _n \)-Ser\( _n \)), was assembled by a second PCR step with the outer primers 1 and 4, and using as templates the LV\( _{\text{H}} \) and V\( _{\text{H}} \) PCR products at a 1:1 molar ratio (total 50 ng of DNA). Similarly, the hinge, Fc, and Tm coding regions were assembled together by recombinant PCR with the primers 5 and 8 using as DNA templates PCR products of primers 5 + 6 and 7 + 8. The Fv and the C\( \text{y1CH1} \) coding regions were then successively cloned upstream of the genomic cytoplasmic tail (exon 6–8) and the 3′ UTR of the H-2Kb gene in pBluescript II SK as Nnul/Xmal and Xmal/Bglll restriction fragments, respectively. The resulting chimeric gene was then cloned downstream the cytomegalovirus gene promoter (pCMV) as a Nnul /EcoRI restriction fragment into the pBluescript CE4.1 recombinant PCR using the following primers: primer 9 (5′-leader) 5′-TCTTCTCAGTATGGTGGATGACATGAGGGCCCATGCTC-3′; primer 10 (3′-leader) 5′-AGACATTCAGATGACCC; and primer 11 (see above). The leader intron and the scFv (5′-leader) 5′-TAACAGGTGCCTTGGCAGC-3′; primer 11 (3′-leader) 5′-CTCGTGACGACATCCAGACCCAGGTCGCC-3′; primer 10 (3′-leader) 5′-GTCGATCCTGATCTGCACAGCCGTCCCTTGGATGACGATGGC-3′; primer 9 (5′-leader) 5′-TGGCCGATCGCGGACAGTGCGATGAATCCAGACCCAGGTCGCCCTGTCCCACTCCAGCAC; and primer 4 (see above). The leader intron and the scFv 187.1 were first amplified by PCR using the primers 9 and 10, and 11 and 4, respectively. The leader intron and the scFv 187.1 were fused together by recombinant PCR using the outer primers 9 and 4, and then cloned as a Nnul/Xmal restriction fragment upstream of the mC\( \text{y1CH1} \) coding sequence. Finally, the 1.2-kb human ubiquitin C gene promoter (pCMV) was cloned upstream of the coding sequences as a Hindlll/Nnul restriction fragment in pBluescript II SK.

Stable transfection of L929 cells. L929 cells were transfected with pBudSCA187\( \Delta \)CH1 constructs using Lipofectamine/Plus reagent (Invitrogen) on six-well plates according to the manufacturer’s instructions. Stable transfectants were selected after 3 wk of growth in complete IMDM medium containing 100 \( \mu \)g/ml of Zeocin.

Mice. All mice were bred and maintained in the TSRI Animal Resources facility according to The Scripps Research Institute Institutional Animal Care and Use guidelines. C57BL6/J (B6), B6.RAG1\(/^{−/−}\) (50) and B6.CD4.1 mice were obtained from Jackson ImmunoResearch Laboratories. Germine Ig Jk-\( \kappa \)-Ck-deleted mice (40) were provided by D. Huzar (GenPharm Intl., San Jose, CA). EmuBcl-2-22 transgenic (Bcl2 Tg) mice (51), were provided by Drs. Strasser and Harris (The Walter and Eliza Hall Institute of Medical Research, Melbourne, Australia).
Production of transgenic mice. The 4-kb pU8-K transgene construct was separated from bacterial vector sequences by a digestion with HindIII/ EcoRI and agarose gel electrophoresis. The fragment was isolated and puriﬁed on Ethip–d columns (Schleicher & Schuell) according to the manufacturer’s recommendations and dialyzed overnight against zygote injection grade Tris-HCl-EDETA. Transgenic mice were produced by classical micro-injection techniques at the TSK J Mouse Genetics Core Facility. (B6 x DBA/2)F1 zygotes were microinjected with the pU8-K transgene and reimplanted into the oviducts of pseudo pregnant κ−/− foster mice. Analysis was then backcrossed three times to the B6.CD45.1 background.

Bone marrow chimeras. Recipient mice were κ-macrosel transgenics or littermate controls; all carried the CD45.1 allele. Recipients received 900 rad γ radiation from a Cs source on the day of transfer. Bone marrow donors were all of the CD45.2 allele. Bone marrow cell suspensions were depleted of T cells by treatment with biotinylated anti-Thy-1.2 antibodies and anti-biotin magnetic beads (Miltenyi Biotec) according to the suggested protocol. Five million cells were transferred i.v. per recipient. After 6 wk, recipient spleens were harvested to study their lymphocyte analyses by ﬂow cytometry. Only chimeras in which >98% of cells in bone marrow and spleen were donor-derived were included in the analysis.

Flow cytometry analysis. For the analysis of mouse cells ex vivo, nucleated cell suspensions were prepared from bone marrow, spleen, mesenteric lymph nodes, and peritoneal cavity. Erythrocytes were eliminated from the spleen and bone marrow preparations by ammonium chloride treatment. Cells were stained in staining buffer containing appropriately diluted dilutions of the following monoclonal antibodies: biotin-coupled mouse anti–rat IgG1 (BD Biosciences) developed with streptavidin–PE, PE and biotin rat anti–mouse IgG (187.1; BD Biosciences); biotin–anti-IgG3,1; (BD Biosciences) followed by PE or allophycocyanin–streptavidin; PerCP– or FITC-coupled anti–CD45SR/B220 (RA3-6B2; BD Biosciences); Cy5–coupled anti–IgM (331.12); PE–coupled anti–CD45.1 (eBioscience); FITC–coupled anti–CD45.2 (eBioscience). L929–transfected cells were harvested using PBS, 0.5 mM EDTA, washed twice, and resuspended in staining buffer (PBS, 1% BSA, 0.01% NaN3). Cells were incubated with biotin mouse anti–rat IgG1 (BD Biosciences), which reacts with the lunker region of the macrosel antigen, or mouse IgG2b mononuclear antibody carrying κ or λ L chains, followed by biotin–coupled anti–rat mouse IgG2b (BD Biosciences) to assess Fv binding site speciﬁcity. Biotin–coupled antibodies were revealed with streptavidin–phycocerythrin (PE; BD Biosciences). Cells were gated on the basis of forward and side scatter criteria to avoid contamination by dead cells or debris. For intracellular staining and BrdU uptake studies, surface stained cells were ﬁxed and permeabilized using a kit (Cyto-ﬁx/Cytoperm™, BD Biosciences) and stained according to the manufacturer’s instructions. FITC anti–BrdU antibody was used (BD Biosciences). Stained cells were analyzed on a FACSCaliber ﬂow cytometer (Becton Dickinson) using the FlowJo software package.

Excision product and RS-to-IRS PCR assays. B cells were isolated from spleen and bone marrow using an anti–B220 magnetic bead cell puriﬁcation system (Miltenyi Biotec). The purity of these preparations was >90% in all cases. Genomic DNA was extracted using the QIAamp DNA Mini Kit (QIAGEN) according to the manufacturer’s recommendations. PCR reaction were done in a ﬁnal volume of 50 μl containing 100, 25, 12, and 6 ng of B cell genomic DNA. VA1-to-Jα1 excision product DNA rearrangements were detected using the oligonucleotides and PCR conditions described (11). RS-to-IRS PCR assay was performed using primers B and C (16). Samples were ampliﬁed for 25 cycles: 1 min at 94°C, 1 min at 60°C, and 1 min at 72°C. PCR products were electrophoresed in 1.5% agarose gels, blotted on nylon membranes, and probed with intervening sequence probe as described previously (16).

Serum Ig determinations. Polyvinyl chloride plastic microplates (Falcon) were coated with a rat monoclonal antibodies speciﬁc for IgG2a,b (BD Biosciences), IgM (331.12), or IgA1 (R26–46; BD Biosciences). After washing and blocking, sera (diluted in PBS supplemented with 1% BSA) were incubated 3 h at room temperature. Bound Ig was detected using biotinylated anti–mouse IgG2a,b (R6–60.2; BD Biosciences) or horseradish peroxidase–conjugated anti–mouse IgG (187.1; BD Biosciences). Biotinylated antibodies were revealed using streptavidin–peroxidase (Sigma–Aldrich) followed by addition of the chromogenic substrate 2,2′-azino–bis(3-ethylbenzthiazoline–6-sulfonic acid) (Sigma–Aldrich) in McIlvain’s buffer (84 mM Na2PO4/48 mM citrate, pH 4.6). Absorbance was measured on a Spectra MAX250 plate reader ( Molecular Devices). Standard curves were obtained using a mouse IgM, (G 155–288; BD Biosciences) and a mouse IgMA (11E10; Southern Biotechnology Associates, Inc.) or a mouse IgG2b,κ (BD Biosciences).

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