The 3′ end of the heavy chain constant region locus enhances germline transcription and switch recombination of the four γ genes

Wesley A. Dunnick, Jian Shi, Kevin A. Graves, and John T. Collins

Department of Microbiology and Immunology, University of Michigan Medical School, Ann Arbor, MI 48109

The switch in immunoglobulin (Ig) heavy chain class is preceded by germline transcription and then mediated by a DNA recombination event. To study germline transcription and class switch recombination we used transgenic mice with a 230-kilobase bacterial artificial chromosome that included a rearranged VDJ gene and the entire heavy chain constant region locus. In addition to several lines with intact transgenes, we identified two lines in which the heavy chain locus transgene lacked Cα and everything 3′ of it, including the regulatory elements HS3a, HS1-2, HS3b, and HS4. B cells from both lines with the truncated transgenes make abundant transgenic (Tg) VDJCμ transcripts and IgM protein. Deletion of the 3′ end of the locus results in dramatically reduced expression of both germline transcripts and switched VDJCH transcripts of the γ3, γ2b, γ2a, and ε genes. In addition, the transgenes lacking the 3′ end of the locus express reduced amounts of γ1 germline transcripts and 2–3% of the amount of Tg IgG1 in tissue culture compared with intact transgenes. Finally, switch recombination to γ1 is undetectable in the transgenes lacking the 3′ elements, as measured by digestion circularization–polymerase chain reaction or by the expression of VDJCγ1 transcripts.

Switch recombination between the μ and γ, ε, or α heavy chain genes, and the germline transcription that precedes it, are likely to be regulated by cis-acting elements within the heavy chain locus (1, 2). Many experiments implicate promoter region sequences, and other elements associated with individual heavy chain genes in this regulation (3). A group of enhancers 3′ of the Cα gene is known to control switch recombination to some isotypes (2). These elements are associated with DNase I hypersensitive (HS) sites (4–6), and hence, are termed HS3a, HS1-2, HS3b, and HS4 (in the order that they appear in the heavy chain constant region locus). These elements are strong B cell specific enhancers, especially in combination with one another (5, 7–17). A spontaneous deletion of all four elements has been associated with a loss of immunoglobulin expression by a cell line (18). Replacement of individual HS sites with a promoter/neomycin resistance gene cassette down regulates the expression of some isotypes in tissue culture or in vivo (19–21). Finally, deletion of HS3b and HS4 from the germline reduces dramatically the expression of γ3 and γ2b, and reduces somewhat the expression of γ2a, ε, and α. However, expression of μ and γ1 was reduced by <50% by the HS3b + HS4 deletion (22).

However, the “clean” deletion of HS3a or HS1-2, without replacement by the promoter/neomycin resistance cassette, does not alter expression of any heavy chain isotype, and, with the aforementioned exception (22), it has been difficult to delete more than one of the HS sites from the mouse genome. We have investigated the regulation of germline transcription and switch recombination by the 3′ HS sites using a transgene of the entire heavy chain constant region locus, including the four HS sites named above. We identified two independent lines of transgenic (Tg) mice in which the transgene was truncated before insertion, and therefore, the integrated construct lacks all four HS sites. We report here the effect of these spontaneous deletions on the expression of Tg μ and four γ genes.

RESULTS

Composition of transgenes

In independent microinjections of three constructs that differ slightly (see Materials and methods), we produced mice with a transgene of the entire constant region locus, and tested
The deletion of the 3' end of the heavy chain locus had a minimal effect on expression of Cα genes (19, 20, 22). However, a heavy chain locus lacking all four 3' HS sites has not been derived using the technology of homologous recombination in embryonic stem cells. We tested the expression of germline transcripts and postswitch transcripts of heavy chain genes in the transgenes that included the four 3' HS sites and other transgenes that had deleted the 3' end of the locus, including all four HS sites. To control for transgene expression from line 1001, we used lines 551, 7898, 580, 578, or 556, because these lines have a wild-type γ1 gene within their arsonate (ARS)/Iγ transgene. To control for line 822, we used lines 847, 884, or 858, which share with line 822 the STAT6 site mutation in the γ1 promoter region (see Materials and methods).

We tested the expression of the μ, γ3, and γ2b genes in RNA that was derived from T-depleted splenocytes that were cultured in LPS. The deletion of the 3' end of the Tg heavy chain locus had a minimal effect on expression of Cμ transcripts (Fig. 2 A). All Tg lines expressed approximately equal amounts of VDJCμ transcripts, whether the lines included an intact heavy chain locus (7898, 578, or 858) or lacked the 3' end of the locus (1001 and 822).

To detect Tg Cγ or Cε germline transcripts, we amplified Tg and End germline transcripts, using oligonucleotides that were exactly complementary to the transgene and the End genes. We then cut the PCR products with restriction enzymes that would distinguish the transgene and End genes. In contrast with Tg VDJCμ expression, we were unable to detect Tg germline γ3 transcripts in lines lacking the 3' end of the heavy chain locus (1001 and 822; Fig. 2 B). To estimate the amount of switch recombination to Tg γ3, we measured the expression of Tg VDJCγ3 transcripts. As a positive control for the expression of postswitch transcripts, we amplified cDNAs using primers in the Iμ sequence and in the Cγ3 gene (Fig. 2 B). These primers anneal perfectly to the End genes and the transgene, and amplify RNA products from End and Tg γ3 genes that have undergone switch recombination (23). Thus, the Iμ.Cγ3 RT-PCR ensured that...
switch recombination and expression of postswitch transcripts, at least from the End genes, had taken place. In addition, we used the amount of \( \mu_c\gamma_3 \) transcripts (and other \( \mu_c \) transcripts, see next paragraph) to normalize samples for variability in B cell preparations and response to cytokines and activators in tissue culture. To the extent possible, we equalized the amount of \( \mu_c\gamma_3 \) transcripts for each cDNA sample, and then tested the same concentrations of cDNA for Tg VDJ\( \gamma_3 \) transcripts (Fig. 2 B). Hence, the various RNA samples were approximately equalized for amount of switch recombination. Transgenes that included most of the 3′ end of the heavy chain locus also expressed postswitch VDJ\( \gamma_3 \) transcripts from the transgene, whether they had three copies of this gene (lines 884 and 858), two copies (line 7898), or one copy (lines 551, 847, and 580; Fig. 3 B). However, lines 1001 and 822 did not express Tg VDJ\( \gamma_3 \) transcripts. This was despite the fact that RNA samples from these lines included transcripts from the End locus that had switched to \( \gamma_3 \) (\( \mu_c\gamma_3 \)).

Parallel results were obtained when the \( \gamma_2b \) and \( \epsilon \) genes were analyzed. Although the transgenes with an intact 3′ end expressed germline transcripts and switched VDJ\( \gamma_3 \) transcripts well for the \( \gamma_2b \) and \( \epsilon \) genes, the transgenes in lines 1001 and 822—lacking most of the 3′ end—did not (Fig. 2, C and D). Line 822 does express a small amount of Tg \( \gamma_2b \) and \( \epsilon \) germline transcripts, although it expresses barely detectable or undetectable Tg VDJ\( \gamma_2b \) and VDJ\( \epsilon \) transcripts (Fig. 2, C and D).

Expression of Tg \( \gamma_2a \) transcripts

To investigate expression of \( \gamma_2a \) germline transcripts from the transgene, we cultured T depleted splenocytes from nonTg (NTg) and Tg mice in LPS or in LPS + IFN\( \gamma \). IFN\( \gamma \) induces the expression of \( \gamma_2a \) germline transcripts and the secretion of IgG2a (24, 25); we observed increased amounts of both germline transcripts (Fig. 3 A) and Tg VDJ\( \gamma_2a \) transcripts (Fig. 3 B) from B cells that were cultured in LPS + IFN\( \gamma \). As noted by the arrows in Fig. 3 A, \( \gamma_2a \) germline transcripts from the End gene migrate slightly faster than those from the transgene. In addition, there is greater expression of \( \gamma_2a \) germline transcripts from the “a” allele associated with the transgene than from the “b” allele of the End gene (compare, for example, the LPS + IFN\( \gamma \) samples for NTg and line 578, with two copies of the transgene). We do not know the basis for this increased expression, but we also have observed the same magnitude of increased ex-

Figure 2. Transgenic heavy chain expression as determined by RT-PCR.

(A) Transgenic VDJ\( \mu \) expression. Two amounts of cDNA, differing by fivefold, were tested. Line 858 has three copies of the Tg \( \mu \) gene; the other four lines have two copies. (B) Transgenic \( \gamma_3 \) expression. Lines 822, 580, 551, and 847 have one copy of the transgene; the other lines have two or three copies. The expression of Tg \( \gamma_3 \) germline transcripts by B cells from lines 847, 551, and 580 is documented in Fig. S3 (available at http://www.jem.org/cgi/content/full/jem.20041988/DC1). (C) Transgenic \( \gamma_2b \) expression. The HPRT expression for some of these LPS samples is presented in Fig. 3 C. (D) Transgenic \( \epsilon \) expression. For \( \gamma_3 \), \( \gamma_2b \), and \( \epsilon \), the same amount of cDNA was used for \( \mu_c \) and for Tg VDJ\( \mu \) amplifications.
B cells from lines 1001 and 822 express abundant IgCγ2a transcripts and demonstrate good switch recombination of the End γ2a genes (Fig. 3, B and D). However, in comparison with B cells with one (Fig. 3 D) or two copies (Fig. 3 B) of the Tg γ2a gene, B cells with a deletion of the 3' end of the transgene express reduced amounts of Tg VDJCγ2a transcripts. For example, the VDJCγ2a transcript signal (by PhosphorImager analysis) for line 1001 is 14% that of line 580.

Expression of Tg IgG1 and γ1 transcripts

In studies from other investigators, none of the disruptions or deletions of the 3' enhancers have reduced the expression of the γ1 gene by more than 50% (19, 21, 22). We tested B cells with the two transgenes lacking the 3' end of the locus for secretion of IgG1 with the Tg ("a") allotype. As reported (26), intact transgenes produce abundant Tg IgG1 when exposed to IL-4 and either LPS or CD40L (Fig. 4 A). The induction by IL-4 is 10- to 100-fold. However, IL-4 does not induce Tg IgG1 secretion by B cells from lines 1001 and 822; this results in much smaller absolute amounts of IgG1* (Fig. 3 A). The mutation in the Stat6 binding site in Tg lines 847 and 858 does not alter the ability of B cells from these lines to express Tg IgG1 in tissue culture (26). Hence, the reduced expression of Tg IgG1 by B cells from line 822 is unlikely to be a result of the Stat6 site mutation in the γ1 gene of the transgene. The cultures of B cells from lines 1001 and 822 do express abundant IgG1 from the End locus, and when the results are expressed as Tg IgG1 divided by total IgG1 in the culture, B cells from lines 1001 and 822 secrete 2–3% of the amount of Tg IgG1 secreted by B cells with an intact transgene (black bars in Fig. 4 B). We assayed Tg IgM expression from LPS cultures set up in parallel with the LPS + IL-4 and CD40L + IL-4 cultures, and expressed the data as Tg IgM divided by total IgM in the cultures (Fig. 4 B, gray bars). B cells from lines 1001 and 822 express abundant Tg IgM; on average, the same amount of Tg IgM as B cells with an intact transgene. B cells from lines 1001 and 822 that were cultured in LPS + IL-4 also secreted similar amounts of IgM compared with B cells from lines 7898, 578, and 884 that were cultured in LPS + IL-4 (unpublished data).

PCR-based tests for γ1 nucleic acids confirmed these results for IgG1 protein secretion (Fig. 4, C–E). Transgenic germline transcripts (Fig. 4 C) and VDJCγ1 transcripts (Fig. 4 D) were barely detectable in B cells from lines 1001 and 822; the level of expression was at or slightly above the background level that was detected in cDNA from NTg B cells. Switch recombination of the End γ1 genes occurred in the same cell samples, as revealed by abundant IgCγ1 transcripts. These results with RNA were confirmed at the DNA level by a variation of the digestion circularization (DC)-PCR assay (27). We performed DC-PCR on DNA samples from B cells that were treated with LPS + IL-4, and then digested the final products with MboI, which differentiates the transgene and End genes. We detected the 263-bp fragment ("Tg") in DNA from B cells with an intact

Figure 3. Transgenic γ2a expression. (A) RT-PCR for germline transcripts. The size difference between germline transcripts of the End gene and the transgene in lines 556, 578, and 1001 without the four-bp insertion in γ2a is noted by the lower Tg arrow. The size difference between germline transcripts of the End gene and the transgene with the four-bp insertion in γ2a in lines 858 and 822 with the four-bp insertion is noted by the upper Tg arrow. (B and D) RT-PCR for Tg VDJCγ2a and IgCγ2a transcripts. Lines 556, 578, and 858 have two or three copies of the ARS/Igh transgene. Lines 1001 and 822 have two Tg VH copies, but one Tg γ2a copy. Lines 847, 551, and 580 have one intact copy of the ARS/Igh transgene, but line 551 has two copies of Tg γ2a. (C) RT-PCR for HPRT transcripts. Although we normalized the LPS + IFNγ samples using IgCγ2a expression, we present HPRT transcripts to allow comparison of RNA levels between LPS and LPS + IFNγ samples. HPRT transcripts were amplified beginning with two cDNA concentrations, one fivefold greater than the other. The amount of cDNA that was used for each primer was identical for germline, Tg VDJCγ2a, and IgCγ2a transcripts. The more abundant HPRT transcripts were amplified from smaller amounts of cDNA, but the relative quantities of the cDNA samples were consistent among the HPRT amplifications and the other three reactions.

Expression of Tg γ2a transcripts

We assayed γ2a transcripts by more than 50% (19, 21, 22). We tested B cells with the two transgenes lacking the 3' end of the locus for secretion of IgG1 with the Tg ("a") allotype. As reported (26), intact transgenes produce abundant Tg IgG1 when exposed to IL-4 and either LPS or CD40L (Fig. 4 A). The induction by IL-4 is 10- to 100-fold. However, IL-4 does not induce Tg IgG1 secretion by B cells from lines 1001 and 822; this results in much smaller absolute amounts of IgG1* (Fig. 3 A). The mutation in the Stat6 binding site in Tg lines 847 and 858 does not alter the ability of B cells from these lines to express Tg IgG1 in tissue culture (26). Hence, the reduced expression of Tg IgG1 by B cells from line 822 is unlikely to be a result of the Stat6 site mutation in the γ1 gene of the transgene. The cultures of B cells from lines 1001 and 822 do express abundant IgG1 from the End locus, and when the results are expressed as Tg IgG1 divided by total IgG1 in the culture, B cells from lines 1001 and 822 secrete 2–3% of the amount of Tg IgG1 secreted by B cells with an intact transgene (black bars in Fig. 4 B). We assayed Tg IgM expression from LPS cultures set up in parallel with the LPS + IL-4 and CD40L + IL-4 cultures, and expressed the data as Tg IgM divided by total IgM in the cultures (Fig. 4 B, gray bars). B cells from lines 1001 and 822 express abundant Tg IgM; on average, the same amount of Tg IgM as B cells with an intact transgene. B cells from lines 1001 and 822 that were cultured in LPS + IL-4 also secreted similar amounts of IgM compared with B cells from lines 7898, 578, and 884 that were cultured in LPS + IL-4 (unpublished data).

PCR-based tests for γ1 nucleic acids confirmed these results for IgG1 protein secretion (Fig. 4, C–E). Transgenic germline transcripts (Fig. 4 C) and VDJCγ1 transcripts (Fig. 4 D) were barely detectable in B cells from lines 1001 and 822; the level of expression was at or slightly above the background level that was detected in cDNA from NTg B cells. Switch recombination of the End γ1 genes occurred in the same cell samples, as revealed by abundant IgCγ1 transcripts. These results with RNA were confirmed at the DNA level by a variation of the digestion circularization (DC)-PCR assay (27). We performed DC-PCR on DNA samples from B cells that were treated with LPS + IL-4, and then digested the final products with MboI, which differentiates the transgene and End genes. We detected the 263-bp fragment ("Tg") in DNA from B cells with an intact

Pression of germline transcripts by the "a" allele of the End γ2a gene of BALB/c mice compared with expression from the "b" allele of C57BL/6 mice (unpublished data). B cells from lines 1001 and 822 express a small amount of IFNγ-inducible Tg γ2a germline transcripts, at most 5% of the amount of Tg γ2a germline transcripts from lines 556, 578, or 858 with transgenes with an intact 3' end.
gene (lines 7898, 578, 847, and 858); this indicated switch recombination by the transgene (Fig. 4 E). The amount of the Tg recombination varied from ~10% of the total SpSy1 fragments (line 847 with one copy of the transgene) to greater than 50% of the total SpSy1 fragments (line 578 with two copies of the transgene). However, the Tg fragment was not detected when DNA from B cells whose transgenes lacked the 3’ end of the heavy chain locus (1001 and 822) was tested (Fig. 4 E).

DISCUSSION

Loss of the 3’ end of the heavy chain locus and heavy chain gene expression

We have found that, relative to transgenes with an intact heavy chain constant region locus, transgenes lacking the 3’ end of the locus express markedly reduced amounts of postswitched transcripts from the γ3, γ2b, γ2a, and e genes (Figs. 2 and 3). However, Tg μ expression was reduced marginally, if at all (Figs. 2 and 4 B). Qualitatively, these results are similar to experiments that use replacement of single 3’ enhancer elements (19–21) or the deletion of HS3b and HS4 (22). Although it is a little difficult to compare our results with those of the HS3b + HS4 deletion because the assays for switch recombination are different, it seems that the loss of the entire 3’ end of the locus in a transgene results in quantitatively more profound reduction in the expression of γ2a and e. Also similar to other studies of alterations of the

were tested for IgG1 with the Tg allotype by ELISA. The signals for Tg IgG1 in lines 1001 and 822, for all four culture conditions, are near the limit of detection, and therefore, can fluctuate between 0 and 20 ng/ml. In particular, the relative signals for line 822 B cells that were cultured in LPS compared with LPS + IL-4 are not representative of the average twofold (not statistically significant) reduction in LPS + IL-4 compared with LPS alone. Transgenic γ1 copy number for each line is given directly below the line number. (B) The portion of IgM was calculated by dividing the amount of AD8+ (Tg idiotype) IgM by total IgM for LPS cultures. Means, with SEM bars, are presented for two to four independent cultures. The portion of Tg IgG1 was calculated by dividing the amount of IgG1a by the amount of total IgG1 for LPS + IL-4 or CD40L + IL-4 cultures. Means, with SEM bars, are presented for 5 to 11 cultures. As a negative control, the mean IgG1a/total IgG1 value from LPS + IL-4 and CD40L + IL-4 cultures of NTg B cells is presented as an open box. (C) Endogenous and Tg germline transcripts were identified by TaqI-digestion of RT-PCR products of RNA from B cells induced with LPS + IL-4 (23). The line 578 sample was derived from an independent experiment. B cells from line 847 express smaller amounts of γ1 germline transcripts as a result of the mutation of the Stat6 site in the γ1 promoter (26). (D) Transgenic VDJCy1 and γ1Cγ1 transcripts were determined by RT-PCR, using the same amount of cDNA for the two reactions. RT-PCR for HPRT used less cDNA, but the relative amounts for each sample were held constant. (E) DC-PCR for Tg γ1 switch recombination. Transgenic and End recombination products were distinguished by MboI digestion. MboI sites are indicated by arrows at the bottom of the figure. Sequences flanking the 5’ portion of Spμ are indicated by a gray bar and those flanking the 3’ portion of Spγ1 are indicated by a black bar. These DNA segments are joined at the junction of the gray and black bars by ligation at EcoRII ends to form a circle. “Percent Tg” was calculated by dividing the cpm in the Tg band by the sum of the cpm in the Tg and the two End bands.

Figure 4. Transgenic IgG1 expression. (A) Means of quadruplicate samples, with SD error bars, for a single set of cultures. Supernatant fluids
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3’ enhancers (19, 21, 22), loss of the 3’ end of the heavy chain locus in a transgene resulted in dramatically reduced amounts of germline transcripts of the γ3, γ2b, γ2a, and ε genes (Figs. 2 and 3). One limitation of our study is that both truncated transgenes lacked Ce, so we could not study the effect of the truncations on α gene expression. Only one of the two truncated transgenes retained Ce, so our study of the expression of ε is limited to a single founder line lacking the 3’ end of the locus. It will be important to determine the effect of deletion of all four 3’ HS sites on the expression of the α heavy chain gene, and, in more founder lines, on the ε heavy chain gene.

In the study by Pinaud et al., expression of γ1 germline transcripts and switch recombination was not reduced by deletion of HS3b and HS4 from the germline (22). Likewise, insertions of a neo cassette into HS1-2 or HS3A did not alter γ1 expression (19, 21). In our Tg system, loss of the 3’ end of the locus is correlated with loss of γ1 expression that is as profound as the other isotypes that were studied (Fig. 4). It is a formal possibility that the IgG1 expression was unaffected in the study by Pinaud et al. (22), but was reduced to 2–3% of wild-type levels in our study (Fig. 4) because the former involved the End locus, whereas ours involved a transgene inserted into an ectopic locus. However, the copy number dependence and insertion site independence of Tg IgG1 expression in vitro argues against this possibility (26). It seems more likely that the larger deletion in our study leads to the loss in γ1 expression. From the study by Pinaud et al., the 6-kb deletion of HS3a + HS4 does not alter expression of IgG1 (22). The truncation of the transgenes in lines 822 and 1001 by ~60 kb results in a dramatic reduction in Tg IgG1 expression (Fig. 4). Therefore, deletion of the additional 54 kb reveals elements that are important for γ1 gene expression (HS3a, HS1-2, or other unidentified elements). Redundancy among the 3’ enhancer elements could allow γ1 gene expression in a heavy chain locus with much of the 3’ end deleted, if some critical elements were present (e.g., HS3b and HS4). A more directed series of deletions, which are feasible by modification of the ARS/IGH bacterial artificial chromosome in Escherichia coli, will be useful to define better the role of each element in the regulation of heavy chain germline transcription and switch recombination.

Enhancement of germline transcription

Our results support the notion that the 3’ HS elements are enhancers of a small amount of transcriptional activity. This enhancer activity has been observed readily in tissue culture cells and normal B cells by transfection experiments and in conventional transgenes (5, 7–17). In our study, it is best identified for the γ2a and ε genes, where a small amount of germline transcripts are observed in lines 1001 and 822. Addition of the 3’ end of the heavy chain locus (e.g., in lines 578 and 858) results in expression of γ2a and ε germline transcripts that are at least 10-fold more abundant. The amount of these germline transcripts in lines lacking the 3’ end of the locus is entirely consistent with the amount of cytokine-inducible germline transcripts from a 10-kb γ2a transgene (14) and from transient transfection of constructs with the ε germline promoter (16). Thus, the promoter regions for the individual heavy chain genes may direct a small amount of regulated germline transcription, which is enhanced dramatically by the 3’ HS sites. Synergy between two or more 3’ HS sites may be particularly important for this enhancement (11–15).

MATERIALS AND METHODS

Transgenes and Tg mice. The construction of the 230-kb ARS/IGH transgene with the entire constant region locus has been described previously (26). The transgene is derived from strain 129 DNA (Ighb). Injection of C57BL/6xSJL F2 eggs and backcrosses to C57BL/6 maintained the End loci as Ighb. Lines 551 and 7898 were derived from the first microinjection of ARS/IGH DNA. The transgene in these two lines has a wild-type γ1 gene and a four-bp insertion at the EcoRI site in Iγ2a. Lines 556, 578, 580, and 1001 were derived from a second microinjection. The transgene in these four lines includes a wild-type γ1 gene and lacks the four-bp insertion at the EcoRI site in Iγ2a. Lines 847, 884, 882, and 822 were derived from a third microinjection. These lines have mutations of four consecutive bp in the Stat6 binding site at −123 bp 5’ of the promoter for germline transcripts of the γ1 gene and a 4-bp insertion at the EcoRI site in Iγ2a (26). Lines 556, 578, 580, 1001, 551, and 7898 are designated “wild-type,” which refers to their γ1 genes. Lines 847, 884, 882, and 822 are designated “MSTAT,” which refers to their γ1 genes. All transgenes have a 4-bp insertion in the BgIII site in Iγ1.

The presence and copy number for the following elements in the transgene was determined as described previously (26): JH3-4/Ep, Sy3, Sy1, Sy2a, Sr, Hs3a, Hs1-2, Hs3b, and Hs4. Sy2b was tested using a HindIII digest probed with nick-translation psSy2b/Eb.6 (28). The Tg HindIII fragment is 10 kb long; the End HindIII fragment is 11.5 kb long (29). DNA samples from Tg mice were tested for the presence of Tg Ce by PCR of tail DNA in the presence of 3P-dATP, using the primers 5’-GAGCTGTCTACATCATGACC-3’ (5’ end) and 5’-GAGCATACTTGAATGTGGC-3’ (3’ end). Denaturation was at 95°C for 1 min, annealing at 60°C for 1 min, and extension at 72°C for 30 cycles, followed by a 10 min extension at 72°C. Digestion of the 169-bp product with Nsil yielded the uncut fragment (End), and 108- and 61-bp fragments (transgene). Copy number was determined by PhosphorImager quantification of the Tg and End bands. The End band represented an internal standard of two copies.

Assay of germline transcripts and switch recombination. B cells were prepared from various Tg mice and cultured with various combinations of LPS, CD40L, IFNy, and IL-4 as described previously (30). RNA and cDNA were prepared from those B cells. For all RT-PCR reactions (in the presence of 3P-dATP), denaturation was at 95°C for 1 min, annealing for 1 min, and extension at 72°C for 30 cycles, followed by a 10 min extension at 72°C. Digestion of the 169-bp product with Nsil yielded the uncut fragment (End), and 108- and 61-bp fragments (transgene). Copy number was determined by PhosphorImager quantification of the Tg and End bands. The End band represented an internal standard of two copies.
various cDNA dilution series (Fig. S2, available at http://www.jem.org/cgi/content/full/jem.20041988/DC1).

For the amplification of germine transcripts each C region primer was paired with a primer for its respective I exon. After amplification, products were digested with restriction enzymes that distinguished the End and transgenes by polymorphisms between the I and/or C exons. Iyr3 primer: 5′-TGATGGGATATATCAGGATCC-3′, 61° annealing. Digestion of the 537-bp product with PstI results in the 392-bp fragment (End), 100-bp and 83-bp fragments (transgene), and a 143-bp fragment (both genes). 

Iyr2 primer: 5′-GGCGTGCTTCTTCGAGAACTA-3′, 58° annealing. Digestion of the 578-bp product with TaqI results in a 317-bp fragment (End), 127-bp and 151-bp fragments (transgene), and a 110-bp fragment (both genes). 

Iyr2a primer: 5′-GCTGATGATCCTACTCTGAGAGAG-3′ fragment (transgene), 691-bp and 82-bp fragments (End), and a 108-bp fragment (both genes). Hypoxanthine phosphoribosyl transferase (HPRT) transcripts were amplified as described previously (14). 

DNA samples from Tg B cells cultured with LPS + IL-4 for 4 d were digested with EcoRI and ligated to form circles (27). DC-PCR products were amplified in the presence of 32P-dATP using the primers 5′-GCAGCTGCTTCTTCGAGAACTA-3′, 61° annealing. Digestion of the 537-bp product with PstI results in the 392-bp fragment (End), 100-bp and 83-bp fragments (transgene), and a 143-bp fragment (both genes). 

Online supplemental material. Fig. S1 documents the semi-quantitative nature of several RT-PCRs. Fig. S2 presents additional data that are used in determining the transcript content for various Tg lines. Fig. S3 presents additional data on germline transcription of the γ3 gene in specific Tg lines. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20041988/DC1.

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