Systemic lupus erythematosus patients produce a broad spectrum of autoantibodies (auto-Abs), including anti–double-stranded (ds)DNA (1, 2). In healthy individuals these auto-Abs are regulated at multiple checkpoints during B cell development (3, 4), and defects, such as loss of T cell tolerance (5), late-stage B cell deregulation (6), and clearance defects of nuclear and cytoplasmic antigens, can lead to autoimmunity.

A reproducible appearance of auto-Abs can be achieved by induction of a systemic lupus erythematosus–like syndrome by a chronic graft-versus-host (cGvH) reaction (7). To initiate cGvH, lymphocytes from a Bm-12 mouse are transferred into a C57BL/6 (B6) recipient (8, 9). The Bm-12 and the B6 mouse differ at the MHC class II protein by three amino acids; hence, the cGvH is thought to be mediated by Bm-12 CD4$^+$ T cells (10). cGvH leads to B cell hyperactivation followed by the production of a variety of auto-Abs directed to chromatin, Sm, DNA, and other nuclear antigens. In addition, the mice develop glomerulonephritis (8, 11). The cGvH is thought to require B cell receptor cross-linking (12). The spectrum of auto-Abs induced by cGvH is narrowed in an anti-DNA transgenic mouse, referred to as 56R/B6 (13–15). This 56R H chain site-directed knock-in mouse was generated using the VH from an anti-DNA Ab isolated from a diseased MRL/lpr mouse (16). The DNA reactivity of the 56R H chain is due to positive charges of arginines (Arg) in the complementarity-determining regions (CDRs) (17).

The 3H9 and 56R transgenic models have been useful for studying the mechanism of receptor editing (18, 19). L chain rearrangement in the 56R model seems to persist until the self-reactivity of the transgenic 56R H chain is altered. Such B cells bearing the 56R transgenic H chain are allowed to migrate into the periphery when paired with so-called “editor” L chains. Such editor L chains have low isoelectric points due to a high frequency of aspartates (Asp) in their CDRs, and we think they block the interaction between H chain Arg and DNA (17).

The 56R H chain on a BALB/c background pairs mainly with the $\kappa$ L chain V$\kappa$21D, an efficient editor of anti-DNA activity (17). On a B6 background, however, less efficient editors V$\kappa$38C and V$\kappa$20 L chains are used (17, 20, unpublished data). Among $\lambda$ chains, $\lambda$x has been shown to modify DNA binding of the 56R H chain (21, 22) and may result in altered self-reactivity. In 56R/BALB/c mice, a new
population of edited 56R H chain transgenic B cells has been identified. In this mouse, B cells express two receptors of which one is $\kappa$ and the other is $\lambda$ (17), and these “partially edited” B cells are thought to reside in the marginal zone (MZ) (23, 24).

Using cGvH, we asked whether incompletely and partially edited B cells were activated and secreted auto-Abs in cGvH. We find that the induction of anti-DNA Abs is accompanied by the induction of other auto-specificities. Hybridoma panels derived from cGvH-induced mice were analyzed and showed that auto-Abs arose from B cells that were polyreactive and bound to self-antigens and proteins, such as dsDNA, phosphatidylserine (PS), myelin basic protein (MBP), thyroglobulin, histone, insulin, cytochrome C, and $\beta$-galactosidase. We found that the observed polyreactivity was a feature of the 56R H chain in combination with the editor $\kappa_{20}$ and $\kappa_{38C}$. Consistent with the expression of both the $\kappa$ and $\lambda$ L chain in the serum after cGvH disease, hybridoma panels derived from cGvH-induced mice included partially edited B cells with $\lambda_1$ L chain that coexpressed $\kappa$ L chain. The B cells with the $\lambda$ and $\kappa$ L chain bound to dsDNA, MBP, and PS and could account for some of the auto-Ab expression in cGvH. In addition, those double positive B cells may be located in the MZ.

RESULTS

Induction of auto-Abs in cGvH

The cGvH reaction induces a lupus-like syndrome that includes activated lymphocytes and the production of auto-Abs (8, 9). In previous studies, cGvH was induced by the transfer of splenocytes (7–11, 14, 15, 25, 26) or partially purified CD4$^+$ T cells (27) from Bm-12 mice to B6 recipients. The cGvH induces a variety of auto-Abs in B6 mice (8, 10), but in the case of 56R/B6, the auto-Ab repertoire is enriched in anti-DNA specificities as has been observed in studies on 56R Abs from unmanipulated and cGvH 56R/B6 mice (13–15, 28). To rule out possible donor B cell response in the cGvH, we used purified Bm-12 CD4$^+$ T cells (CD4$^+$CD3$^+$ > 96%; not depicted). Under these conditions, we found that cGvH caused an increase of anti-dsDNA Abs (Fig. 1 A), accompanied by an increase of anti-PS Abs (Fig. 1 B), but we also find induction of anti-MBP Abs (Fig. 1 C).

The cGvH also activated peripheral B cells in the 56R/B6 recipient as seen by the transient expression of CD69 and CD86 as well as by the increased levels of MHC class II and Fas (CD95). A slight decrease of CD24 was also observed (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20060075/DC1). We conclude that cGvH induced by purified CD4$^+$ T cells leads to secretion of anti-DNA Ab and activation of B cells in the 56R/B6 recipient. The induction of different auto-Ab specificities in 56R transgenic mice raised the following questions: Do these specificities come from different nontransgenic B cell populations, or can these specificities be attributed to polyreactive Abs expressed by 56R H chain with editor L chains as has been found to be the case for $\kappa_{20}$ and $\kappa_{38C}$? The 56R transgene in B6 mice both accelerates the onset and increases the titers of anti-dsDNA Abs in cGvH as reported previously (14, 15). To study the role of 56R in auto-Ab induction, we tested the expression of 56R H chain by flow cytometry and in hybridoma panels.

Role of 56R H chain in auto-Ab induction

To test whether the anti-DNA, anti-PS, and anti-MBP Ab specificities (Fig. 1) were associated with the 56R H chain, we used allotype-specific Abs that distinguish the transgenic allele Igha (16, 29) and the nontransgenic endogenous allele at a serum dilution of 1:180. IgM and IgG Abs to dsDNA (n = 6 for control group and n = 7 for cGvH group) (A), to PS (n = 5 for control and cGvH group) (B), and to MBP (n = 3 for control group and n = 6 for cGvH group) (C) were tested by ELISA. MRL/lpr sera were tested at the same concentration for comparison (●, n = 4). Results represent means ± SEM.
Igh\textsuperscript{b} of the 56R/B6 mouse. In the control mice, 45–48% of the B220\textsuperscript{+} splenic B cells were IgM\textsuperscript{+} and IgD\textsuperscript{+} at day 60. An almost identical number (44–45%) was found in the cGvH group. Of the B220\textsuperscript{+} B cells, 8–10% expressed endogenous H chain according to allotype IgM\textsuperscript{b} and IgD\textsuperscript{b} expression. This may result from aberrant VH replacement followed by rearrangement of the untargeted allele. The IgM\textsuperscript{b} and IgD\textsuperscript{b} fraction did not change after cGvH (Fig. S2 A, available at http://www.jem.org/cgi/content/full/jem.20060075/DC1). From the correlation of allotype expression and auto-Ab production, we conclude that 56R H chain is mainly associated with the cGvH-induced auto-Ab.

**mAbs isolated after cGvH**

To test whether \kappa-edited 56R Abs are responsible for the different autoreactivities found in the serum (Fig. 1), we studied mAbs secreted by the hybridomas in two independent fusion panels generated 4 and 10 wk after cGvH. We tested the hybridomas for several characteristics: binding to multiple self-antigens, the H chain isotype, and the type of L chains.

**Specificity and isotype.** In the 4-wk fusion, 53 out of 78 hybridomas secreted IgM or IgG (Table I A and Table S1, which is available at http://www.jem.org/cgi/content/full/jem.20060075/DC1). Similar results were obtained in the 10-wk fusion, as 42 out of 139 hybridomas secreted IgM (Table S2). Both fusions yielded a large number of clones that were IgM IgG\textsuperscript{+}. These “nonsecreters” were not analyzed further. Supernatants from both fusions were tested for binding to dsDNA, PS, and MBP. Most of the mAbs in the 4-wk fusion bound dsDNA as well as PS. In addition, these mAbs bound MBP (Table I A). Polyreactive mAbs that bound to dsDNA, PS, and MBP were also found in the 10-wk fusion (see below; Table II).

H chain gene usage in polyreactive Abs. To determine whether these polyreactive mAbs were associated with 56R, we tested whether the hybridomas have an intact 56R gene. PCR products from primers complementary to the 3H9/56R (56R) H chain leader sequence and the CDR3 of the 56R transgene (16) indicate the presence of the complete 56R transgene. In the 4-wk fusion, we found 33 clones out of 53 secretors to be 56R\textsuperscript{+} (Table I A). The remaining 20 clones did not amplify the 56R transgene by this assay and are classified as 56R\textsuperscript{−}. 11 of these 20 56R\textsuperscript{−} clones did amplify a product with a 56R CDR3 reverse primer and the JHCH primer. This PCR product represents 56R VH genes that have been truncated or VH replaced (13). No PCR product was detected with either of the 56R PCRs in 9 out of 20 56R\textsuperscript{−} clones, and they were not analyzed further (Table S1).

Most of the clones that bound dsDNA, PS, and MBP were 56R\textsuperscript{+} (Table I A). A few 56R\textsuperscript{−} clones bound to PS, dsDNA, and MBP, but their affinity for these antigens relative to the 56R\textsuperscript{+} mAbs was low (Table S1).

**L chain gene usage.** We determined the types of L chains of these polyreactive mAbs. The V\kappa genes from both the 56R\textsuperscript{+} group and the 11 56R\textsuperscript{−} mAbs that have truncated or VH

Table I. B cell hybridomas generated from 56R/B6 splenocytes 4 wk after cGvH reaction

| Total clones (78) | 56R tg\textsuperscript{+} (33) |
|------------------|-----------------------------|
| IgM\textsuperscript{+} (29) | 24 |
| Reactivity to dsDNA | 23 |
| PS | 18 |
| MBP | 21 |
| IgG\textsuperscript{+} (24) | 9 |
| Reactivity to dsDNA | 3 |
| PS | 2 |
| MBP | 3 |

| B Total clones (36) | V\kappa12–13 | V\kappa20 | V\kappa21D | V\kappa38C | V\kappa(S) |
|---------------------|-------------|----------|-----------|-----------|-----------|
| 56R tg\textsuperscript{+} (25) | 1(J2)\textsuperscript{a} | 1(J2)\textsuperscript{a} | 4(J2)\textsuperscript{a} | 1(J2) | 1(J4) |
| 56R tg\textsuperscript{−} (11) | 2(J5)\textsuperscript{a} | 1(J2)\textsuperscript{a} | 1(J4) |

\textsuperscript{a}Groups that contain clones with two L chain rearrangements.

\textsuperscript{b}56R\textsuperscript{+} clones from fusion 4 wk after cGvH (Bm-12 into 56R/B6) are polyreactive (A). \kappa L chain usage in 56R\textsuperscript{+} and 56R\textsuperscript{−} clones from fusion 4 wk after cGvH (Bm-12 into 56R/B6). Only clones with H and L chain secretion are listed and were tested for rearrangement of \kappa L chains using specific primers for V\kappa12-13, V\kappa20, V\kappa21D, and V\kappa38C, and a degenerate primer V\kappa(S) for other \kappa rearrangements (B). For more details see Table S1.

Table II. Binding of MBP and PS of 56R\textsuperscript{+} IgM\textsuperscript{+} B cell hybridomas with different L chains

| V\kappa-gene | MBP/dsDNA binding | PS/dsDNA binding |
|-------------|--------------------|------------------|
| V\kappa20/12 | 0.77 ± 0.10 (n = 7) | 5.55 ± 1.78 (n = 3) |
| V\kappa20/14 | 0.58 ± 0.05 (n = 14) | 3.62 ± 1.02 (n = 6) |
| V\kappa38c/J2 | 0.61 ± 0.09 (n = 7) | 0.40 ± 0.12 (n = 3) |
| V\kappa38c/J4 | 0.92 ± 0.06 (n = 7) | 0.51 ± 0.27 (n = 3) |
| V\kappa38c/J5 | 0.35 ± 0.03 (n = 11) | 1.06 ± 0.24 (n = 4) |
| V\kappa(S)/J1 | 1.03 ± 0.07 (n = 7) | 0.25 ± 0.02 (n = 3) |

| V\kappa-gene | MBP/dsDNA binding | PS/dsDNA binding |
|-------------|--------------------|------------------|
| V\kappa20/J2 | 1.46 ± 0.05 (n = 3) | 14.26 ± 0.85 (n = 3) |
| V\kappa20/J4 | 0.85 ± 0.08 (n = 7) | 10.68 ± 2.41 (n = 7) |
| V\kappa38c/J5 | 1.36 ± 0.14 (n = 3) | 1.25 ± 0.06 (n = 3) |

56R\textsuperscript{+} IgM hybridomas are compared from two fusions. Clones were polyreactive to at least two antigens in EUSA assays testing for dsDNA, PS, and MBP. The mean of the MBP/dsDNA and PS/dsDNA was calculated from at least three clones, and ± indicates the error (SEM). The values show the binding to PS and MBP relative to DNA binding. Fusion panels are from 4 (A) and 10 wk (B) after GvH. For more details see Table S2.
We searched for the origin of polyreactivity by flow cytometry. After cGVH induction, there was an increase in B220⁺ IgG⁺ splenocytes expressing the plasma cell marker CD138 (2.57 and 4.05% of the B220⁺ cells by days 28 and 60 after cGVH, respectively). We considered these surface Ig⁺-expressing B cells to be pre-plasma cells (Fig. 3 A). We also found an increase of B220⁺ IgM⁺ B cells expressing CD138 (0.37 and 0.75% of the B220⁺ cells by days 28 and 60 after cGVH, respectively), indicating that B cells with the H chain transgenic allele IgH⁺ of the 56R/B6 mouse differentiated into plasma cells, and were giving rise to anti-DNA Abs. B220⁺ IgM⁺ B cells expressing CD138 (2.29 and 2.95% of the B220⁺ cells by days 28 and 60 after cGVH, respectively) were also increased and most likely to be of the IgG class (Fig. S2 B).

**Contribution of L chain to polyreactivity**

The 56R⁺ mAbs had distinguishable reactivities depending on the Vk and Vκ/Jκ combination. Vκ21D vetoed binding to dsDNA and the other antigens: PS, MBP, thyroglobulin, cytochrome C, histone, β-galactosidase, and insulin. Both Vκ20 and Vκ38C Abs were polyreactive, but their binding pattern was different (Fig. 2 A). 56R⁺ clones with Vκ20 or Vκ38C bound dsDNA with higher relative affinity than Vκ21D. In addition, Vκ20 and Vκ38C rearranged to J2 with dsDNA with a higher relative affinity than the same V regions rearranged to J4 or J5 (Fig. 2 A). We studied the binding to PS, MBP, and the other proteins relative to their binding to dsDNA. We showed that binding to PS was favored over binding to dsDNA in clones with the L chain Vκ20, whereas Vκ38C L chains bound to dsDNA and PS with similar affinity. Affinity for MBP, thyroglobulin, cytochrome C, histone, and β-galactosidase was higher in Vκ38C as compared with Vκ20, and no difference was found between Vκ20 and Vκ38C for insulin (Fig. 2 B).

Relative binding to PS and MBP was also assessed by calculating the ratios of PS to DNA and MBP to DNA binding of the mAbs that expressed the 56R transgene from clones in each group. The OD₄₀₅ of the binding assays was normalized based on the Ig concentration of 1 μg/ml for each supernatant. All clones were IgM⁺ and expressed the indicated editor L chain. The following groups were tested: Vκ20/J2: n = 2 (clones 50.1 and 50.2 from one original clone); Vκ20/J4: n = 4 (clones 71.1, 71.5, 73.2, and 73.4 from two original clones); Vκ21D/J2: n = 2 (clones 79.1 and 79.5 from one original clone); Vκ38C/J2: n = 2 (clones 160.1 and 160.5 from one original clone); Vκ38C/J4: n = 2 (clones 222.4 and 222.6 from one original clone); Vκ38C/J5: n = 3 (clones 89.4, 89.6 and 105.5 from two original clones); and Vκ(S)/J1: n = 2 (clones 206.1 and 206.5 from one original clone). All clones tested were sequenced using specific primers for Vκ20, Vκ21D, and Vκ38C and were unmutated. All tested binding specificities are shown in comparison for one representative result for each group. SEM from triple values in the ELISA assay (A). To analyze the binding of the subclones (to PS and the proteins MBP, thyroglobulin, histone, insulin, cytochrome C, and β-galactosidase) relative to dsDNA binding, dsDNA binding of each subclone was set to 1; therefore, only clones with considerable binding to dsDNA were included in this representation. Relative binding to the tested antigens is shown for clones expressing the indicated L chain. Numbers show p-values determined by the Anova Single Factor Test (B).

**Plasma cell differentiation in cGVH**

We searched for the origin of polyreactivity by flow cytometry. After cGVH induction, there was an increase in B220⁺ IgG⁺ splenocytes expressing the plasma cell marker CD138 (2.57 and 4.05% of the B220⁺ cells by days 28 and 60 after cGVH, respectively). We considered these surface Ig⁺-expressing B cells to be pre-plasma cells (Fig. 3 A). We also found an increase of B220⁺ IgM⁺ B cells expressing CD138 (0.37 and 0.75% of the B220⁺ cells by days 28 and 60 after cGVH, respectively), indicating that B cells with the H chain transgenic allele IgH⁺ of the 56R/B6 mouse differentiated into plasma
Potentially pathogenic auto-Abs are thought to home to the MZ (35, 36); therefore, we tested whether MZ B cells were affected by cGvH in the 56R/B6. This is relevant because it has been shown that 56R/BALB/c has an enlarged MZ B cell compartment. Here we show that in the 56R/B6, the MZ B cell population (CD21 high and CD23 low) comprised as much as 50% of the B220+ splenocytes. The percentage of the MZ B cell population decreased after cGvH, whereas the CD21–CD23– B cell population increased. The percentage of follicular B cells (CD21 intCD23 high) slightly increased (Fig. 3 B, dot plots). The majority of the CD138+ B cells was found in a subpopulation of CD21–CD23– B cells (Fig. 3 B, histograms). Yet, during cGvH, the number of lymphocytes in the spleen increased on average 2.5-fold and the spleen weight increased 3.2-fold (not depicted); therefore, we cannot conclude that the absolute numbers of MZ B cells in cGvH were decreasing. However, we observed a decrease of CD21 expression by the mean fluorescence intensity on MZ B cells (not depicted). This suggests that MZ B cells were activated during cGvH. The concommitant increase of plasma cell differentiation and the decrease of CD21 expression in MZ B cells suggest that potentially autoreactive B cells from the MZ were affected by the cGvH reaction and may have differentiated to become auto-Ab–producing B cells.
T cells in the spleen of 56R/B6 recipients were also activated as seen by the down-regulation of CD3 (not depicted) and CD4 as well as by the increased expression of CD69 (tested at day 14; Fig. 4). The expression of the T cell co-stimulatory molecules CD28 and ICOS also increased, and CD4+ T cells displayed a memory phenotype as shown by the expression of CD44_{high}, CD45R{low}, and CD62L_{low} (Fig. 4). We could not distinguish between donor and recipient CD4+ T cells in these studies, but the activation of T cells as late as day 60, by which time we would not expect Bm-12 donor T cells, indicates a role of endogenous T cells in antigen activation. We conclude from these data that cGvH induced by purified Bm-12 CD4+ T cells leads to plasma cell differentiation (Fig. 3) and a sustained activation profile of T cells in the 56R/B6 recipient (Fig. 4). The role of endogenous T cells in the cGvH has been established by studies showing that CD4 knockout mice are unresponsive to cGvH (37).

Expression of \( \lambda \) L chain in cGvH

We tested anti-DNA Abs for the usage of \( \kappa \) and \( \lambda \) L chain in the serum after cGvH and developed our anti-dsDNA assay with anti-\( \lambda \) and anti-\( \kappa \) reagents. We found \( \lambda \) - and \( \kappa \)-containing anti-dsDNA Abs (Fig. 5). The strong induction of \( \lambda \) anti-dsDNA Abs led us to ask whether these Abs come from isotypically included B cells that express \( \lambda 1 \) and \( \kappa \) L chain (18). We also asked whether these isotypically included B cells are activated by cGvH and differentiate into anti-dsDNA–producing plasma cells. This is relevant because it has been shown that MZ B cells in the 56R/BALB/c mouse can coexpress a \( \kappa \) and a \( \lambda \) L chain. The \( \lambda 1 \) L chain with 56R H chain binds DNA (38), but the \( \kappa \) L chain is an editor (17). These isotypically included B cells are thought to be sequestered in the MZ (23, 24). We analyzed the composition of B cells in the MZ of a 56R/B6 mouse using a single cell PCR technique and tested for B cells with both \( \lambda 1 \) and \( \kappa \) L chain (Table III, A and B). L chain rearrangements were identified....

Figure 5. \( \lambda \) and \( \kappa \) L chain usage in anti-dsDNA Abs in 56R/B6 after cGvH. For induction of cGvH, Bm-12 CD4+ T cells were injected into 56R/B6 (■). As a control, B6 CD4+ T cells were injected into 56R/B6 (△). Anti-DNA Abs were detected by ELISA with an anti-\( \lambda \) (n = 6 for cGvH group and n = 3 for control group) and anti-\( \kappa \) (n = 4 for cGvH group and n = 3 for control group) detection Ab by ELISA at a serum dilution of 1:180. Results represent means ± SEM. MRL/lpr sera were used for comparison (●; n = 4) (A). Detection of L chain expression during cGvH was analyzed by Western blot. Sera from mice after cGvH (Bm-12 CD4+ T cells injected into B6 or 56R/B6 recipients) or sera from control mice (B6 CD4+ T cells were injected into B6 or 56R/B6 recipients) were run under reducing conditions. \( \lambda \)x was detected using a polyclonal anti-V\( \lambda \)x Ab (reference 22). Serum samples were diluted at 1:100 for \( \lambda \) and at 1:2,000 for \( \kappa \) L chain detection. The expression of \( \lambda \)x after exposure for 5 min, \( \lambda 1,2,3 \) after exposure for 20 min, and \( \kappa \) after exposure for 30 s is shown. Results are representative of seven experiments. \( \lambda \)x, \( \lambda 1,2,3 \), and \( \kappa \) in the serum are shown from one mouse (B).
using nested PCRs on cDNA from single cells. Specific primers were used for VA1/2 and CA1/2, Vk21D and Ck, and Vk38C and Ck (20). Other κ L chains were amplified using the degenerate primer Vκ(S) (39). We found that all B cells with λ L chain also had a κ L chain rearrangement. Sequence analysis showed that B cells with in-frame λ L chain had rearranged the editor L chain Vk21D and Vk38C. Two examples with in-frame λ L chain had, in addition to the κ editor, another κ rearrangement. Other λ L chain PCR products had large deletions in the V or V/J regions. Rearrangement of λ L chain in-frame or out of frame was always accompanied by an in-frame κ L chain that was often the editor Vk38C or Vκ20 (Table III B). Thus, the MZ contains a population of B cells that seems to have rearranged excessively and introduced λ deletions.

We tested whether κ/λ 56R B cells were activated by cGvH by analysis of a fusion panel 10 wk after cGvH. We found isotypically “included” B cells that coexpressed a λ1

### Table III. Single cell analysis of B cells from the MZ of a 56R/B6 mouse

**A**

| Single cells (179) | 56R tg⁺ (124) | Vκ(S) (124) | Vκ38C (100) | Vκ21D | Vλ (26) |
|-------------------|----------------|-------------|-------------|--------|---------|
| Plate I           | 67             | 80          | 30          | 11     | 15      |
| Plate II          | 57             | 44          | 70          | n.d.   | 11      |
| 56R tg⁺ (124)     | Vκ(S) (87)     | Vκ38C (64)  | Vκ21D       | Vλ (17)|         |
| Plate I           | 58             | 29          | 10          | 9      |         |
| Plate II          | 29             | 35          | n.d.        | 8      |         |

**B**

| Cell no. | 56R | Vκ(S)/Ck | Vκ38C/Ck | Vκ21D/Ck | Vλ1/Cλ |
|----------|-----|----------|----------|----------|--------|
| Plate I  |     |          |          |          |        |
| 10       | +   | +        | +        | +        |        |
| 18       |     |          |          |          | +      |
| 20       | +   | (Vκ21-4 (98%) J2 (100%) IF) | + (Vκ38C/J5 IF) | + (Vλ1/Jλ1/Cα1 IF) |
| 22       |     |          |          | +        | a      |
| 40       | +   |          |          |          |        |
| 43       |     |          |          |          | +      |
| 53       | +   | + (Vκkk4 (91%) J5 (95%) IF) | + (Vκ38C/J5 IF) | + (Vλ1/Jλ1/Cα1 IF) |
| 59       |     |          |          | +        | a      |
| 62       | +   | +        | (Vκ38C/J4 IF) | + (Vλ1/Jλ3/Cα2 OF) |
| 63       | +   | (Vκ21-4 (98%) J2 (100%) IF) | + (Vκ38C/J5 IF) | + (Vλ1/Jλ1/Cα1 OF) |
| 74       |     |          |          |          |        |
| 75       | +   | (Vκ21-4 (98%) J1 (100%) IF) | + (Vκ38C/J5 IF) | + (n.d.) |
| 76       |     |          |          |          | +      |
| 95       |     |          |          |          | +      |
| Plate II |     |          |          |          |        |
| 13       | +   | + (Vκ21-4 (98%) J2 (100%) IF) | n.d. | + (Vλ1/Jλ1/Cα1 OF) |
| 14       |     | + (Vκ19-23 (98%) J5 (100%) IF) | n.d. | + (Vλ1/Jλ1/Cα1 OF) |
| 29       |     | + (Vκ21-4%) J2 (97%) IF) | n.d. | + (Vλ1/Jλ3/Cα2 OF) |
| 33       | +   | + (Vκ19-15 (92%) J5 (100%) OF) | + (Vκ38C/J4 IF) | n.d. | + (Vλ1/Jλ3/Cα2 OF) |
| 34       |     | + (Vκ38C/J4 IF) | n.d. | +        |
| 49       |     |          |          |          | +      |
| 52       | +   | + (Vκ19-23 (98%) J2 (100%) IF) | n.d. | + (Vλ1/Jλ1/Cα1 OF) |
| 59       | +   | + (Vκ21-4 (98%) J2 (96%) IF) | + (Vκ38C/J4 IF) | n.d. | +      |
| 60       |     |          |          |          | +      |
| 71       | +   | + (Vκ21-23 (97%) J2 (100%) IF) | + (Vκ38C/J4 IF) | n.d. | +      |

Single cells from the MZ (B220⁺ CD21m⁺ and CD23⁺) in a 56R/B6 mouse were analyzed by PCR for Igκ, Igλ, and 56R H chain. All PCR product-positive cells were included in the table (A). B cells contained Igκ⁺ cells. B cells with in-frame λ L chain show message for one or two in-frame κ L chain(s) (in bold) (B). For further details see Table S3. Numbers in parentheses indicate sequence similarity to the germline gene in percentages.

n.d., not done.
IF, in frame.
OF, out of frame.
⁺a, sequence similarity in percent, determined for κ L chains amplified with the degenerate primer Vκ(S).
and a \(\kappa\) L chain. 5 out of 139 clones were \(\lambda\)1, and 4 of these expressed the 56R H chain. One \(\lambda\)1 clone did not express the 56R H chain and was negative for IgM and IgG. All of the 56R/\(\lambda\)1-expressing clones coexpressed a \(\kappa\) L chain by ELISA, and in three of those examples, we found rearrangement of \(\lambda\)x by PCR. In addition, we found one clone that expressed the 56R H chain with \(\lambda\)2. All clones that expressed the 56R H chain with \(\lambda\)1/\(\lambda\)2 L chain bound to dsDNA, PS, and MBP. None of the clones expressed only \(\lambda\)1, indicating that B cells that express the 56R H chain with \(\lambda\)1 alone are eliminated from the repertoire (Table IV). Thus, partially edited B cells coexpressing \(\lambda\) and \(\kappa\) L chain were activated in cGvH and could be fused. We also found \(\kappa\)/\(\lambda\)-expressing B cells from cGvH-induced mice by flow cytometry (Fig. 3 C); however, the number of isotypically included B cells was higher as found by single cell PCR analysis (Table III). We are considering that this phenomenon is due to the slight cross-reactivity of anti-\(\lambda\) and anti-\(\kappa\) Abs used in flow cytometry analysis (Fig. 3 C).

The editor L chain, \(\lambda\)x, is also induced by cGvH. \(\lambda\)x is known to bind MBP both as free L chain and in association with H chains (40, 41), including 56R (unpublished data). We tested the possibility that cGvH might induce \(\lambda\)x-associated Abs by a Western blot of sera developed with \(\lambda\)x antisera (Fig. 5 B). We were unable to detect \(\lambda\)x in the serum of 56R/B6 mice or in nontransgenic B6 mice, but after initiation of the cGvH, we found that \(\lambda\)x levels in both 56R/B6 and nontransgenic B6 mice rose rapidly and were maintained until day 60 (Fig. 5 B; not depicted for day 60). Expression of \(\lambda\)x after cGvH was significantly higher in the 56R transgenic than in the nontransgenic B6, suggesting that this L chain is associated with the 56R H chain. Consistent with the ELISA data (Fig. 5 A), we also found increased levels of \(\lambda\) and \(\kappa\); however, in 56R/B6, expression of those L chains did not increase to the same extent as \(\lambda\)x (Fig. 5 B).

**DISCUSSION**

The 56R H chain transgene enhances the anti-DNA production in cGvH (14, 15) and in the quasi-autoimmune B6 (30–34 and unpublished data). This is unexpected because Abs with the 56R H chain in BALB/c are edited. DNA binding is either vetoed by L chains, such as V\(\kappa\)21D (17) and Ax (21, 22), or DNA binding on the cell surface is diluted because more than one receptor is expressed (17, 23, 24). In this study, we show that editing can also be incomplete (Fig. 2). Editors were originally defined in BALB/c transgenics, a strain that does not convert to autoimmunity even with an anti-DNA transgene. Most of the B cells from this mouse express the 56R H chain with V\(\kappa\)21D, an L chain that vetoes DNA binding completely (17). In the B6 mouse, the L chain repertoire is reversed, very few 56R/V\(\kappa\)21D combinations are found, and the majority of B cells express 56R H chain with incomplete editors, such as V\(\kappa\)20 and V\(\kappa\)38C (reference 20, unpublished data, and Table III). We think that the difference lies in the tolerance threshold of the strain: BALB/c, the prototypic “healthy” strain, tolerates only completely edited Abs, whereas B6, a lupus-susceptible strain (30–34), is more permissive. We show that the cGvH-induced autoimmunity in 56R/B6 mice activates B cells using the incomplete editors V\(\kappa\)20 and V\(\kappa\)38C (Fig. 2 and Table I B). Thus, incomplete editing contributes to autoimmunity by determining an available repertoire that includes autoreactive B cells. In support of this idea is the finding that 56R/B6 and the congenic 56R/Sle2/B6 (expressing the additional lupus susceptibility gene Sle2) spontaneously produce anti-DNAs that include 56R/V\(\kappa\)20 Abs (unpublished data). Evidence for such a “lupus repertoire” is also found in lupus-affected individuals (3, 4).

Incomplete editing is inevitable. The ability of certain L chains to edit is correlated with certain structural features of these L chains, in particular, a high frequency of Asp residues in or near the CDRs (17). Hence, we think editing may be the result of an interaction between the VH Arg residues and the VL Asp residues to interfere with the VH Arg interaction with DNA (38, 42, 43). However, editor Asps may not neutralize every Arg and might account for residual anti-DNA activity (44). Similarly, not all Asps need to interact with Args and might be available for interactions with basic proteins such as MBP, as in the 56R/\(\lambda\)x.

The incompletely edited Abs, such as 56R/V\(\kappa\)20, retain activity for DNA and cardiolipin. This is thought to be a cross-reaction based on binding to a phosphate-containing moiety shared by both (45). But a novel source of polyreactivity comes from binding activities that some editors confer from binding activities that some editors confer.
A relevant example has been described in which the 3H9 H chain gives different patterns of antinuclear antigens depending on the L chain with which it is associated (46). We conclude from these results that editor L chains are correlated with anti-self activity. J gene usage also seemed to play a role in binding some proteins, namely MBP, thyroglobulin, cytochrome C, and histone, as shown for its relevance for Vk20 and Vk38C in binding to dsDNA (Fig. 2, A and B). Although polyreactivity can come about by the creation of different combining sites, it can also be due to enhanced cross-reactivity, as in the case of Vk20.

Receptor editing acts in B cells with autoreactive receptors; hence, rearrangement persists and can produce isotypically included B cells as we have described in the 56R/BALB/c transgenic mouse. One of the two receptors (56R/α1) is antinuclear (38), and the other is a 56R/κ editor (24). Such isotypically included κ/λ double-expressing B cells in 56R/BALB/c are found in the MZ of the spleen (24) and in hybridsomas from LPS-activated fusions. Fusion panels generated after cGvH include clones that coexpressed α1 and κ L chain and secrete anti-DNA Abs. In this study, none of the clones (Table IV) expressed α1 L chain only. Therefore, we think the κ/λ B cell may represent a state of tolerance. In this sense, a B cell may have a self-reactive receptor and expression of two L chains may alter or reduce the affinity of the B cell to an antigen. Other examples of dual expression come from recent studies in 56R mice in which the κ locus has been knocked out. Hybridoma panels from these mice showed that the vast majority of the 56R/λ1 B cells coexpressed αx L chain, an efficient editor, and only very few have rearranged α1 alone (47 and unpublished data), but these 56R/α1 clones bound DNA even when αx L chain was coexpressed. cGvH appeared to activate these κ/λ double-expressing B cells because we find anti-DNA Abs with α1 in the serum in cGvH (Fig. 5 A). By single cell analysis we tested whether these κ/λ B cells may be in the MZ of 56R/B6 mice and confirmed the existence of κ/λ-included B cells (Table III). Single cell analysis of MZ B cells also revealed interesting properties that may act on the λ locus. All λ L chain rearrangements, whether out-of-frame or in-frame, were accompanied by a κ L chain, which was (except in one case) in-frame (Table III).

As discussed above, we think that DNA and/or MBP binding can result from incomplete editing. For example, the Asp-rich λx editor in association with H chains including 56R binds MBP (40, 41, and unpublished data). The acidic property of λx to which we attribute the MBP activity is also a feature of κ editors, and these L chains might also confer MBP activity to 56R. And, as we show here, 56R with Vk38C and Vk20 also bound MBP as well as DNA and PS (Fig. 2, A and B) (28). Therefore, an incompletely edited Ab may have both Arg(s) and Asp(s) available for binding both to DNA and proteins.

During the cGvH reaction there was a change in the percentages of MZ B cells, follicular B cells, and CD21+CD23− B cells (Fig. 3 B). MZ B cells may have activated by alloreactive Bm-12 T cells, which in turn may have activated endogenous T cells of the 56R/B6 recipient. These T cells could give help to the differentiation of plasma cells. This idea is consistent with recent findings that showed that MZ B cells, not follicular B cells, are potent activators of naive CD4+ T cells. MZ B cells migrate to T cell areas and differentiate upon interaction with T cells into plasma cells (48). This could be the case in the cGvH reaction, and plasma cells found in the CD21+CD23− population might be a source of auto-Abs to DNA (Fig. 3 B, histograms). Support for this idea comes from features of 56R/Sle2/B6. Sle2 further increases the tendency to autoimmunity of 56R/B6, and in 56R/Sle2/B6, the MZ and the CD21+CD23− B cell populations are increased as compared with 56R/B6. These CD21+CD23− B cells secreted auto-Abs of the IgM allotype, the transgene-encoded allele, in culture (unpublished data). We also find Abs reactive to DNA expressing the H chain of the Iγ allotype (Table I). Assuming that both Sle2 and cGvH enhance autoimmunity, we think that the increase of the CD21+CD23− cells in 56R/Sle2/B6 is analogous to the increase of CD21+CD23− B cells in cGvH-induced autoimmunity. The question remains whether partially edited κ and λ double positive B cells are able to move out of MZ to become plasma cells.

Conclusions

Autoreactive and polyreactive Abs. Here we present a detailed description of the auto-Abs induced by cGvH. The study reveals the structural basis for autoreactivity of certain Abs and the origin of these auto-Abs.

Origin. Natural auto-Abs are found in abundance in both healthy and autoimmune individuals, and there is a vast and confusing literature on their origin, rationale, and implications for autoimmunity and immune protection (49). Our study answers these questions by showing that at least some if not all “natural” auto-Abs result from receptor editing. Of course, the abundance of these auto-Abs in our model is in part because of the anti-DNA transgene. This produces an immature B cell population, most of which must be edited to survive. But even nontransgenics must have a substantial repertoire that is shaped by editing. Based on the mechanisms that generate the immature VH repertoire, we estimate that at least 50% of this repertoire is anti-DNA (50, 51). Thus, editing of this population will have a profound influence on the frequency of natural auto-Abs in the peripheral B cell repertoire.

Relevance. A question raised by natural auto-Abs is how to explain their presence in a self-tolerant individual. This dilemma is usually explained by invoking affinity differences; i.e., natural auto-Abs are low affinity anti-self and are ignored by tolerance mechanisms (52). But the question remains: Why is there a large abundance of these auto-Abs? The only explanation is positive selection (53) of this population, and several roles for natural auto-Abs have been proposed.
These roles range from housekeeping Abs to stimulators of the repertoire to anti-virus Abs (49). If indeed natural auto-Abs are important in normal immunity, their existence in the repertoire can be understood. It has been shown that the B cell repertoire that is induced by the cGvH is not random but a selective process and requires B cell receptor cross-linking (12) and is supported by the involvement of endogenous T cells (as seen in this study [Fig. 4] and by others [37]). In addition, certain edited auto-Abs appear to be selected as suggested by differences in the repertoire of editors between strains; for example, the distribution of editors in B6 as compared with BALB/c.

Implications. We show here that the peripheral repertoire is shaped by receptor editing and selection. Negative selection can continue to act on edited B cells if they retain self-reactivity after the initial editing event. In this sense, a healthy BALB/c mouse may only tolerate perfectly edited, nonautoreactive B cells. Other strains may be more permissive and allow incompletely edited B cells to enter the periphery. The ability to regulate this population may determine lupus susceptibility by supplying the peripheral B cell population with the precursors to pathogenic auto-Abs. That these B cells are involved in the onset of disease is demonstrated by these findings.

MATERIALS AND METHODS

Mice. The generation of site-directed 3H9/56R knock-in mice has been described previously (13, 17). The 3H9/56R transgene was determined by PCR amplification of tail DNA (16, 17). B6 and coisogenic Bm-12 (B6.C-H2bm12/KhEg) mice were obtained from The Jackson Laboratory. All mice were maintained in our mouse colony. All experiments were conducted in accordance with the Animal Welfare Act.

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cGvH. Mice were 2–8 mo old at time of cGvH initiation. All cGvH experiments for flow cytometry, serum data, and the fusion experiment at 10 wk after cGvH were performed with 107 Bm-12 (for cGvH) or B6 (for controls) splenocytes prepared using standard techniques, or with 107 CD4+ T cells (purity >96%) from The Jackson Laboratory. All mice were maintained in our mouse colony at the University of Chicago. All animal care and procedures were conducted in accordance with the Animal Welfare Act.

Flow cytometry. Before staining, cells were preincubated with excess mouse anti-IgG3/II/III (2.4G2; BD Biosciences) to block Fc receptors. Dead cells were excluded by propidium iodide staining (0.33 μg/ml PBS). The generation of site-directed 3H9/56R knock-in mice has been described previously (13, 17). The 3H9/56R transgene was determined by PCR amplification of tail DNA (16, 17). B6 and coisogenic Bm-12 (B6.C-H2bm12/KhEg) mice were obtained from The Jackson Laboratory. All mice were maintained in our mouse colony. All experiments were conducted in accordance with the Animal Welfare Act.

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PCR on hybridoma DNA. Genomic DNA was purified from individual hybrids. 150 pg DNA was used in each reaction. Primers and conditions for H and L chain PCR assays have been detailed previously (13, 47, 54). The presence of 3H9/56R H chain transgene was identified by PCR using primers complementary to the 3H9 H chain leader exon and the CDR3 sequence (16), or with a 56R CDR3 reverse primer (the complement of the 56R/CDR3 primer described above) and the JHCH primer. The JHCH primer binds to a site between JH4 and CH5 (55). This primer amplifies truncated or replaced 3H9 VH genes because both occur at the embedded heptamer 5’ of CDR3.

For typing of 𝜅 L chain rearrangements, 𝜅-specific forward primers (Vκ12-13, Vκ20, Vκ21D, and Vκ38C) and Vκ(S) (39) or L5 (50) forward primers were used under conditions described previously (17). The Vκ(S) PCR primer should amplify ~70% and the L5 primer should amplify ~60% of Vκ genes. Using jκ2 or jκ5 reverse jκ primers (57), the size of the PCR product corresponds to the jκ segment participating in the rearrangement event as described previously (54). To determine editor rearrangements, PCRs were performed in 1X buffer II (PerkinElmer) with a final concentration of 200 μM of each dNTP (Boehringer), 50 pmol of each primer, 1.5 mM MgCl₂, and 1 U of AmpliTaq Gold (PerkinElmer). Selected PCR products were sequenced using primers for Vκ20, Vκ21D, and Vκ38C as indicated in Results.

Single cell sorting, cDNA synthesis, and PCR. Single cell PCR was performed as described previously (3). MZ B cells B220⁺C21D⁺C21D’hif from one 56R/B6 mouse were sorted on a FACS Vantage (Becton Dickinson) into 96-well PCR plates containing 4 μl lysis solution (0.5X PBS containing 10 mM dithiothreitol, 8 U RNAsin [Promega], and 0.4 U U-5’-3’ Prime RNA Inhibitor [Eppendorf]) and were immediately frozen on dry ice. All samples were stored at −70°C. The cDNA was synthesized in a total volume of 14 μl in the original 96-well PCR plate. RNA from single cells was reverse transcribed at 37°C for 55 min with 150 ng random hexamer primer (pd[N6]; GE Healthcare), 0.5 μl dNTP mix (10 mM each), 1 μl of 0.1 M dithiothreitol, 0.05% (vol/vol) Nonidet P-40, RNase inhibitors (4 U RNAsin and 6 U Prime RNase inhibitor), and 50 U Superscript III reverse transcriptase (Invitrogen). Ig gene rearrangement was tested using the degenerate primer Vκ(S) (39) with nested Cκ reverse primers, specific nested primers for Vκ21D and Vκ38C with nested Cκ reverse primers (for primers see reference 29), primers for the 56R H chain transgene (20), and primers for Vλ designed as follows: Vλ12-13, vn 5’ cagtgttgtgatcgcagattc3’; Vλ20 vn 5’ cagcactaacctgtgctcaca3’; Cκvn 5’ gcaggggctatccttatc3’; Cκvn 5’ gcaggtctacaagggattc3’). Products were amplified by nested PCR (for Vκ(S)Cκ “semi-nested PCR”) in 40-μl reactions containing 20 μM primers and 1.2 U HotStar Taq DNA polymerase (QIAGEN). The PCR conditions were as follows: Denaturation for 4 min at 94°C, followed by 30 s at 94°C and 30 s at 55°C (for Vκ21D, Vκ38C, and 56R H chain), or at 50°C (for Vκ(S) and Vλvn) and 55 s (first PCR) or 45 s (nested PCR) at 72°C for 50 cycles. Elongation was allowed for 10 min at 72°C. As a template, 2 μl cDNA was used for the first PCR and 3 μl of the first PCR product was used for the nested PCR. PCR products were sequenced using primers for Vκ20, Vκ21D, Vκ38C, Vκ(S) and Cκ, and Vκ1/2 and Cκ as indicated in Results.

Western blot. Serum samples were diluted to 1:100 for detection of λ and 1:2,000 for detection of 𝜅 L chain in Tris- and SDS-containing buffer. Proteins were denatured with β-ME and boiled for 5 min. Proteins were resolved on a 12% SDS page and transferred to a PVDF membrane (Bio-Rad Laboratories) at 100 mV, 500 mA. After blocking for 2 h with 4% milk in PBS-T, the blot was probed at 4°C overnight with a rabbit anti-mouse VxX polyclonal Ab (22) or with a horseradish peroxidase-labeled goat anti-mouse Vx or Vλ12-23 polyclonal Abs (both Southern-Biotech) in 1% milk in PBS-T. Membranes were washed three times at room temperature with PBS-T. For λx, membrane was incubated with horseradish peroxidase-conjugated goat anti-rabbit IgG (Bio-Rad Laboratories) for 2 h in 1% milk in PBS-T. Membranes for λx, λ1, and k were developed using ECL reagent.

Statistics. Statistical significance was determined using Anova Single Factor Test for analysis of ELISA. Anova Single Factor Test is appropriate for multiple statistical comparisons on a single dataset. Numbers in figures give the actual p-value determined by this test (Fig. 2 B). For all other ELISA assays on single clones, the SEM was determined for triple repeats (Fig. 2 A) and on serum from the indicated number of mice in each group (Figs. 1 and 5 A).

Online supplemental material. Online supplemental material gives greater details about fusion 4 wk after cGVHD, fusion 10 wk after cGVHD, and single cell analysis of MZ B cells isolated from a 56R/B6 mouse. Figs. S1 and S2 and Tables S1–S3 are available at http://www.jem.org/cgi/content/full/jem.20060075/DC1.

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