B Lymphocyte Memory: Role of Stromal Cell Complement and FcγRIIB Receptors

Robert A. Barrington,1 Olga Pozdnyakova,1 Mohammad R. Zafari,2 Christopher D. Benjamin,2 and Michael C. Carroll1

1Center for Blood Research and Department of Pathology, Harvard University, Boston, MA 02115
2Biogen Inc., Cambridge, MA 02142

Abstract
To dissect the influence of CD21/CD35 and FcγRIIB in antigen retention and humoral memory, we used an adoptive transfer model in which antigen-primed B and T lymphocytes were given to sublethally irradiated wild-type mice or mice deficient in CD21/CD35 (Cr2−/−) or FcγRIIB receptors (FcγRIIB−/−). Cr2−/− chimeras showed impaired memory as characterized by a decrease in antibody titer, reduced frequency of antibody secreting cells, an absence of affinity maturation, and significantly reduced recall response. The impaired memory in Cr2−/− chimeras corresponded with the reduced frequency of antigen-specific memory B cells. Interestingly, FcγRIIB−/− chimeras showed a differential phenotype with impaired splenic but normal bone marrow responses. These data suggest that CD21/CD35 on stroma, including follicular dendritic cells, is critical to the maintenance of long-term B lymphocyte memory.

Key words: B lymphocytes • memory • complement receptors • FcγRIIB • follicular dendritic cells

Introduction
Long-term humoral memory is defined as the ability to generate a more effective antibody response after encounter with the original stimulating T-dependent antigen. In humans, humoral memory can persist for decades after initial antigen exposure as evidenced by the continued presence of antigen-specific immunoglobulin (1). Because serum immunoglobulins have half-lives for 1–3 wk (~4), the B lymphocyte compartment must continually produce specific immunoglobulin for long-term protection.

Effective recall responses to recurrent antigen exposure are postulated to require both preexisting, circulating antigen-specific antibody and memory B lymphocytes (5). Antibody-secreting cells (ASCs)* generate antigen-specific serum immunoglobulin that neutralizes harmful antigens and promotes efficient antigen trapping important for recall responses. Memory B lymphocytes quickly respond to cognate antigen, giving rise to new ASCs producing higher affinity immunoglobulins. Traditional models of humoral memory explain antibody persistence by continued differentiation of memory B lymphocytes into terminally differentiated ASCs (6). These models are called into question by the recent observation that some ASCs are long-lived, residing predominantly in the bone marrow (BM) and spleen (7, 8). Therefore, antigen-specific serum immunoglobulin titers could be maintained by these long-lived ASCs in the absence of continued differentiation of memory B lymphocytes. However, several classic studies suggest that continued differentiation of memory B lymphocytes is critical for optimal memory. Not only do serum immunoglobulin levels decrease more dramatically in the absence of antigen (9, 10), but the average affinity of serum immunoglobulin increases markedly with time after initial antigen encounter (i.e., affinity maturation; 11, 12), suggesting that the selection of high affinity B lymphocytes is an ongoing process. Therefore, despite the long-lived nature of some ASCs, continued differentiation of memory B lymphocytes likely plays a major role in humoral memory. Also implicit in these classic observations is that antigen is required for affinity maturation and for maintaining antibody titers. Therefore, understanding how antigen is retained long-term is paramount to understanding how affinity maturation and antibody titers continue after antigen exposure.

Memory B lymphocytes and long-lived ASCs are generated as end products of germinal center (GC) reactions (1, 13). Antigen sequestered on follicular dendritic cells (FDCs) within GC light zones provides the substrate for se-
lection of high affinity B lymphocytes into the memory compartment. FDCs harbor antigen over a long term, suggesting that these cells have a critical role in humoral memory. Consistent with this idea is the recent observation that LTβR-null mice, whose principally defined mutation is a lack of detectable FDCs and GCs, are incapable of maintaining high titer serum immunoglobulin levels (14). Despite the implied role for FDCs in sustaining humoral memory by making antigen available long-term, the mechanism of antigen retention is unknown.

Two classes of receptor molecules capable of trapping antigen on FDCs are complement receptors, i.e., CD21/CD35 and FcγRIIB receptors. CD21/CD35 have a major role in generating the memory B lymphocyte compartment. The depletion of complement component C3 before immunization led to diminished immune complexes on FDC surfaces, suggesting that C3-tagged antigen and/or immune complexes are trapped via CD21/CD35 (15). In addition, mice deficient in complement components C3 and C4 or receptors CD21/CD35 have reduced abilities to trap a wide array of both T-independent and T-dependent antigens, leading to markedly impaired short-term humoral responses (16–19). Short-term humoral responses appear less dependent on FDC-derived CD21/CD35 because chimeric mice (i.e., WT BM into Cr2−/− recipient mice) have normal to slightly reduced antibody responses (20). The role of CD21/CD35 in antigen deposition for long-term, i.e., memory responses, is less defined, although chimeric mice lacking CD21/CD35 on FDCs, but not on B lymphocytes, generate reduced antibody titers after secondary challenge (20, 21). Unlike CD21/CD35, FcγRIIB is not constitutive on the surface of FDCs and instead is inducibly up-regulated after immunization (22). FcγRIIB-deficient mice are hyperresponsive (23), although this likely results from diminished regulatory signaling on B lymphocytes rather than any perturbation in antigen deposition (24). Because FcγRIIB is up-regulated on FDCs after immunization and because persisting antibody titers are considered important for immune complex deposition, it was suggested that FcγRIIB might be critical for trapping IgG–antigen immune complexes required for maintaining humoral memory as well as for recall responses.

In this study, we investigate the influence of antigen retention on antibody titers and affinity maturation. We find that complement receptors CD21/CD35 on recipient stroma are critical in ensuring normal ASC and memory B lymphocyte frequency, high antibody titers, and continued affinity maturation. Moreover, FDC expression of CD21/CD35 is critical for effective recall responses. In contrast, antibody persistence and the recall response are less dependent on stromal cell expression of FcγRIIB receptors, although frequencies of splenic ASCs and memory B lymphocytes are reduced relative to WT chimeras.

Materials and Methods

Mice. Mice were housed at Harvard Medical School in a specific pathogen-free facility. C57BL/6 mice (Jackson ImmunoResearch Laboratories) and C57BL/6–CD45.1 congenic mice (National Cancer Institute) were used as sources of memory T and B lymphocytes. Recipient mice included the following genotypes: C57BL/6, Cr2−/− backcrossed 5 and 10 generations to C57BL/6, and FcγRIIB−/− mice backcrossed 3 generations on a C57BL/6 background (Taconic). Notably, no experimental differences were found in recipient mice with different degrees of C57BL/6 backcrossing.

Adoptive Transfer Protocol and Immunization Protocol. 4-hydroxy-3-nitrophenyl (NP)–specific memory B lymphocytes were generated using C57BL/6–CD45.1 congenic mice. First immunized with 50 μg 10% alum-precipitated (Sigma-Aldrich) NP–BSA (Biosearch Technologies) intraperitoneally, these mice were then boosted with the same antigen dose after 3 wk. Splenic mononuclear cells were isolated by ficoll-hypaque density centrifugation. To enrich for B lymphocytes, splenic mononuclear cells from NP–BSA–primed CD45.1 congenic mice were incubated with an antibody cocktail consisting of rat IgG anti–mouse CD4, CD8, CD11b/CD18, and IgD (Caltag and Amersham Biosciences) at concentrations of 10^7 cells/ml and 1 μg/ml/antibody. Anti–mouse IgD was added to the cocktail to restrict the recovery of memory B lymphocytes. Cells were then panned twice on Petri dishes coated with goat anti–rat IgG (mouse absorbed; Southern Biotechnologies) for 1 h at 4°C. B lymphocyte purity ranged from 70–80% with <10% T lymphocyte and <10% macrophage contamination (unpublished data). ~40% of adoptively transferred B lymphocytes were NP specific as determined by FACS®. The NP-specific memory response was IgG1/κ as determined by ELISA and BlAcore.

Keyhole limpet hemocyanin (KLH)–specific T lymphocytes were generated from C57BL/6 mice immunized intraperitoneally with 50 μg alum-precipitated KLH as described above. Similarly, splenic mononuclear cells were enriched for T lymphocytes by incubation with an antibody cocktail consisting of rat IgG anti–mouse B220, CD19, CD11b/CD18, and IgD. Splenocytes were then panned as described above. T lymphocyte purity was 50–70% in the transferred population with 10–30% B lymphocyte contamination and <10% macrophage contamination.

Single cell suspensions of 10^7 B and 5 × 10^6 T lymphocytes were adoptively transferred with 50 μg soluble NP–KLH intravenously into sublethally irradiated (650 RAD, Cs source) MHC-identical recipient mice (CD45.2). Irradiated control mice were given antigen but no memory B or T lymphocytes after irradiation. 3 or 16 wk after adoptive transfer, recipient and control mice were challenged intravenously with 50 μg soluble NP–KLH and 1 wk after mice were killed, tissues were isolated to measure recall responses. Recipient and control mice were bled periodically throughout the 16–wk rest period. Serum was analyzed by ELISA for specific antibody production to NP (see below).

For secondary adoptive transfers, single cell suspensions of 10^7 B lymphocytes from the primary chimeras were given intravenously with 5 × 10^6 T lymphocytes from KLH–primed mice and 50 μg NP–KLH were given to sublethally irradiated WT or Cr2−/− recipients. These mice were analyzed 3 wk later.

Anti-NP Response and Affinity Measurements by ELISA. Serum was collected from individual mice and NP–specific antibody titers were determined by sandwich ELISA. 96-well plates (Immulon 1B) were coated with 5 μg/well NP–BSA or NP3–BSA. Plates were blocked with the addition of 5% dry milk (Carnation®) in PBS (Blotto). NP–specific IgG serum antibody in serially diluted samples was detected by alkaline phosphatase–conjugated goat anti–mouse IgG (Sigma-Aldrich). Between incubations,
plates were washed with PBS containing 0.1% Tween. Color was developed using p-nitrophenyl phosphate (Sigma–Aldrich) as substrate and absorbance was measured at 405 nm using Softmax software package. Antibody titer was determined as the reciprocal of the greatest dilution whose absorbance remained above background. For relative affinity measurements, the ratio of titers to NP−BSA and NP15−BSA was calculated for individual mice using OD405 in the linear ranges of the assays as previously described (25).

Relative Serum Antibody Affinity to NP Determined by BLAcore. A BLAcore 2000™ was used to study the kinetics of anti-NP serum IgG binding to NP. Standard amine coupling was used to immobilize goat anti–mouse IgG Fcγ (Jackson ImmunoResearch Laboratories) diluted to 30 µg/ml in 10 mM acetic acid buffer, pH 4.0, to a CMS sensor chip (Biosensor AB; Amersham Biosciences). A second flow cell on the chip was prepared using the same procedure, without goat anti–mouse IgG Fcγ, to subtract out significant nonspecific responses. Experiments were performed at 25°C with a 10–μl/minute flow rate. Serum samples diluted 1:1,000 in HBS (BLAcore) containing 1% BSA were injected over the chip. Captured serum IgG was tested for antigen-binding kinetics by injection with NP−BSA. Apparent kinetic rate constants (kₐ and kᵣ) for NP−binding to serum IgG antibodies were estimated using nonlinear curve fitting with BLAevaluation 3.1 software (BLAcore). Equilibrium dissociation constants (Kₐ) were calculated as kᵣ/kₐ. All values were determined by first normalizing to the amount of IgG bound to the chip.

Enzyme-linked Immunospot Assay for NP-specific ASCs. Frequencies of NP-specific ASCs were quantitated as previously described (26). In brief, 24-well polystyrene plates (Costar) were coated with NP−BSA. After extensive washing, plates were blocked with 1% BSA in PBS for 2 h. Serially diluted splenic mononuclear cells or BM cells (10⁵ to 10⁶ cells/well) were added in DMEM media with 2% fetal bovine serum and incubated overnight at 37°C. Each dilution of cells was assayed in duplicate. Plates were then washed with PBS containing 0.1% Tween and incubated with alkaline phosphatase–conjugated goat anti–mouse IgG antibody (Sigma–Aldrich). Plates were developed using 5-bromo-4-chloro-3-indolyl phosphate at 1 mg/ml in 0.6% agarose to produce blue-colored spots identifying NP-specific ASCs. Spots were counted to determine ASC frequency. As controls, each sample was also plated in wells coated with BSA or KLH. Few BSA-specific ASCs were detected in the BM or spleen and <100 KLH-specific ASCs were observed per recipient spleen (unpublished data).

Immunohistology and Antigen Deposition. Chimeric mice were given 50 µg soluble NP−BSA intravenously 16 wk after adoptive transfer. Spleens were isolated 16–18 h after injection and prepared for immunohistology as previously described (27). 5-µm thick sections were examined for antigen retention within splenic compartments using streptavidin–coupled TRITC (Sigma–Aldrich), Peanut agglutinin (PNA; E-Y Laboratories) coupled to FITC–conjugated goat anti–mouse IgG (Southern Biotechnology Associates, Inc.), PE–conjugated rat anti–mouse CD19 (1D3; BD Biosciences), and biotinylated rat anti–mouse CD45.1 (A20; BD Biosciences). After washing, cells were incubated on ice with streptavidin–PerCP (BD Biosciences). Cells were analyzed with a FACSCalibur® (Becton Dickinson). CD19−CD45.1⁺NP⁺ cells within the lymphocyte gate were considered memory B lymphocytes. These cells were negative for cell surface expression of CD138, IgM, CD3, and CD11b.

Statistical Analysis. All statistical comparisons reported used the standard two-tailed t test assuming unequal variance.

Results

The mechanism of long-term antigen retention participating in humoral memory was examined using a hapten carrier adoptive transfer model. A similar adoptive transfer system was used previously in rats to demonstrate that persisting antibody titers and recall responses depended on the presence of antigen (9). This approach was used to create chimeric mice with differential expression of complement and FcyRIIB receptors on FDC stroma and B lymphocytes. To test the importance of CD21/CD35 in harboring antigen for long-term memory, NP-specific memory B lymphocytes, KLH–primed T lymphocytes, and antigen (NP–KLH) were transferred into sublethally irradiated recipient mice deficient in CD21/CD35 (Cr2−/−) or FcyRIIB (FcyRIIB−/−), as well as WT controls. Thus, chimeric mice have normal complement and FcyRIIB–sufficient B lymphocytes but their stromal cells and radioresistant myeloid cells are receptor deficient. To control for endogenous responses, parallel sets of recipient mice were treated identically except that they did not receive memory lymphocytes. Finally, to identify transferred memory B lymphocytes C57BL/6 mice congenic for the CD45.1 allotypic marker were used as donors, whereas recipient mice expressed CD45.2 exclusively.

Short-Term Responses. Short-term antibody responses were examined in the presence or absence of CD21/CD35 or FcyRIIB. All groups of chimeric mice had comparable anti-NP titers 3 wk after lymphocyte transfer with mean titers ranging from 16.7 × 10⁴ to 22.7 × 10⁴ (Table I). NP titers in WT chimeras deprived of antigen were substantially reduced. When WT and Cr2−/− chimeric mice were challenged with NP−KLH 3 wk after the initial transfer of antigen and memory lymphocytes, specific IgG titers were again comparable between WT and Cr2−/− recipient mice (unpublished data). These data suggest that the adoptively transferred memory B lymphocytes were generating equivalent short-term antibody responses irrespective of stromal expression of CD21/CD35.

Long-Term Antibody Persistence. To measure long-term antibody responses, changes in serum anti-NP titers were monitored over 16 wk for each chimeric mouse (Table I). WT mice receiving adoptively transferred cells in the absence of antigen generated two to three times less antibody compared with WT chimeras receiving antigen, demonstrating that optimal responses are antigen dependent. 6–8 wk after adoptive transfer, antibody titers in WT and FcyRIIB−/− chimeras dropped by ~25% of the initial titer,
Irradiated controls (WT, no antigen) found in splenic NP-specific ASCs of mice lacking CD21/CD35. WT and Cr2/ mice had similar frequencies of NP-specific ASCs by ELISPOT. The BM of WT and FcγRIIB/ mice 16 wk after transfer were analyzed for NP-specific ASCs. The presence of CD21/CD35 on recipient stroma is required for optimal production and/or sustenance of plasma cells.

One explanation for reduced ASCs in Cr2−/− chimeras is that ASCs are short-lived and require replacement by antigen-dependent precursors (28). To determine if there was a reduction in memory B lymphocytes in the Cr2−/− chimeras, frequencies of CD45.1+ (donor-derived) NP-binding λ+ (CD138−) B lymphocytes were assessed 16 wk after transfer. CD138− staining was used to distinguish memory B lymphocytes from ASCs. Based on FACS® analysis, a significant reduction in memory B lymphocytes was observed in the spleens of receptor-deficient chimeras, i.e., 310 ± 55 vs. 125 ± 25, 129 ± 29, and 63 ± 18 per 10⁶ splenocytes for WT, Cr2−/−, FcγRIIB−/−, and WT no antigen, respectively (Fig. 2 b). Similar reductions in donor-derived NP-specific memory B lymphocytes were found in BM, although only chimeras lacking CD21/CD35 stroma and WT no antigen chimeras had significant reductions relative to WT chimeras (Fig. 2 b).

Therefore, reduced ASC frequency and serum antibody titers in Cr2−/− chimeras might be caused by reduced frequencies of memory B lymphocytes.

Another potential contributing factor for reduced frequency of NP-specific ASCs and memory B lymphocytes

Table I. Persistence of NP Titers after Adoptive Transfer of Memory B Lymphocytes and During the Recall Response

| Chimera | Time after transfer of NP-specific memory B lymphocytes (NP-specific IgG titers, ×10⁴) |
|---------|---------------------------------------------------------------------------------|
|         | 2–3 wk | 6–8 wk | 10–12 wk | 16 wk | 17 wk |
| WT (n = 34) | 22.7 ± 3.8 | 19.7 ± 3.8 | 10.4 ± 1.9 | 6.4 ± 1.2 | 67.2 ± 19.1 |
| Cr2−/− (n = 31) | 16.7 ± 2.7 | 9.5 ± 1.7* | 5.8 ± 1.1* | 2.9 ± 0.7* | 6.4 ± 1.8* |
| FcγRIIB−/− (n = 32) | 22.3 ± 5.2 | 15.4 ± 4.4 | 7.3 ± 1.6 | 4.6 ± 1.5 | 38.4 ± 14.9 |
| WT, no antigen (n = 8) | 8.9 ± 3.4* | 7.7 ± 3.0* | 5.1 ± 2.3* | 2.1 ± 0.7* | 1.6 ± 2.1* |
| Irradiated controls (n = 41) | <1.6* | <1.6* | <1.6* | <1.6* | <1.6* |

Numbers represent mean anti-NP IgG titers (×10⁴) ± SEM at the indicated time points after adoptive transfer of NP-specific memory B lymphocytes. *, statistically significant differences upon comparison to WT. Results are pooled from six independent experiments.
in the absence of CD21/CD35 on recipient stroma is a general reduction in chimeric relative to WT recipients. The degree of chimerism was determined by comparing the ratio of donor B lymphocytes (CD45.1+CD19+) versus total B lymphocytes (total CD19+). On average, the frequency of donor B lymphocytes was slightly greater in WT mice compared with Cr2-/- mice, although the difference was not significant (35.8 ± 4.75%, 32.6 ± 4.1%, and 23.8 ± 3.3% in the spleens of WT, Cr2-/-, and FcγRIIB-/- chimeras, respectively). Therefore, the degree of chimerism does not explain the differences in frequencies of NP-specific ASCs and memory B lymphocytes between recipient groups.

**Affinity Maturation.** Given the importance of antigen in affinity maturation and the implication that CD21/CD35-bearing stroma is important for long-term antigen retention, it was proposed that affinity maturation might also be impaired in the absence of CD21/CD35. To estimate relative serum affinity, two approaches were taken. The first approach used ELISA to determine the ratio of antibody titers binding low (NP5) to highly (NP15) haptenated BSA. No significant differences in the ratio of NP5/NP15 titers were detected between the recipient groups 3 wk after the transfer of memory B lymphocytes (Fig. 3). In WT chimeras, the relative serum affinity increased modestly with time (from 52 ± 5% at 3 wk after transfer to 70 ± 7% at 16 wk, P < 0.03). Relative serum affinities did not change significantly with time in Cr2-/- chimeras (53 ± 4% to 48 ± 6%, from 3 to 16 wk, respectively). The most significant changes in relative serum affinities were observed in FcγRIIB-/- chimeras (60 ± 6% to 84 ± 6%, P < 0.006).

To confirm that relative serum affinities in FcγRIIB-/- chimeras increase markedly with time after the transfer of memory B lymphocytes, surface plasmon resonance was used to examine the real-time kinetics of anti-NP serum IgG binding to NP-BSA. Real-time measurements obtained by this technique allow an estimate of affinity based on relative on/off rates (K_D = k_on/k_off). WT chimeras demonstrated modest yet incremental increases in affinity whereas Cr2-/- chimeras failed to show statistically significant changes (for WT, ΔK_D of 1.5 × 10^-8 over 16 wk after cell transfer, P < 0.03; Table II). In contrast to Cr2-/-, FcγRIIB-/- chimeras demonstrated significant increases in affinity constants with time after cell transfer (ΔK_D of

### Table II. Changes in Relative Serum Affinity after Adoptive Transfer of Memory B Lymphocytes

| Chimera       | Mean ΔK_D (BIAcore) |
|---------------|---------------------|
|               | 1–12 wk             | 14–16 wk           |
| WT            | 1.6 × 10^-8*        | 1.5 × 10^-8*       |
| Cr2-/-        | 0.8 × 10^-8         | 1.4 × 10^-8        |
| FcγRIIB-/-    | 0.2 × 10^-8         | 4.8 × 10^-8**      |

Mean changes in relative affinity (ΔK_D; left side of table) were determined by calculating equilibrium dissociation constants (K_D) for each time point serum sample, followed by subtraction of starting K_D values 2–3 wk after cell transfer (n = 20 for WT, 11 each for Cr2-/- and FcγRIIB-/- chimeras). Results are summarized from two independent experiments. *, statistically significant differences upon comparison to mean 2–3-wk time point values (P < 0.03 for WT chimeras and P < 0.003 for FcγRIIB-/- chimeras).
4.8 × 10⁻⁸ over 16 wk after transfer, P < 0.003). Therefore, the absence of FcγRIIB-bearing stroma potentiates affinity maturation. Recall Responses. By definition, memory responses are faster and more robust than primary responses. The contribution of CD21/CD35 and FcγRIIB receptors on recall responses was tested in the current model by challenging chimeric mice intravenously with soluble NP₃-KLH 4 mo after adoptive transfer. 1 wk later, mice were analyzed for antibody titers and frequency of ASCs and memory B lymphocytes. Anti-NP titers increased approximately 10-fold in WT and FcγRIIB⁻⁻ chimeric mice after challenge (mean prechallenge to postchallenge titers = 6.4 × 10³ to 67.2 × 10³ for WT and 4.6 × 10³ to 38.4 × 10³ for FcγRIIB⁻⁻ chimeras; Table I). By contrast, the mean anti-NP titer increased approximately twofold in Cr2⁻⁻ chimeras after antigen challenge and was approximately 10-fold less than that observed in WT chimeras (from 2.9 × 10³ to 6.4 × 10³; Table I). Therefore, the recall response of Cr2⁻⁻ chimeric mice was substantially reduced compared with WT and FcγRIIB⁻⁻ chimeras.

To determine whether antibody titers correlated with the frequency of effector cells, NP-specific ASCs from the BM and spleen from all groups of mice were measured by ELISPOT after antigen challenge. The frequency of NP-specific ASCs increased ~10-fold in the BM and 100-fold in the spleen in all groups (compare Fig. 1, a to c and b to d). WT and FcγRIIB⁻⁻ chimeras produced similar frequencies of NP-specific BM ASCs, consistent with similar antibody titers generated as part of the memory response (Fig. 1 c). In contrast, the relative frequency of NP-specific ASCs was reduced two- to threefold in the spleen and BM of Cr2⁻⁻ chimeras compared with WT chimeras. This suggested that reductions in memory B lymphocyte frequency and/or reduced antigen localization via CD21/CD35 contributed to reduced recall responsiveness. Despite a near 10-fold increase in antibody titer, FcγRIIB⁻⁻ chimeras had significantly reduced frequency of splenic ASCs compared with WT chimeras.

To assess whether the renewed production of NP-specific memory B lymphocytes was affected in the absence of CD21/CD35 or FcγRIIB-containing stroma, splenic B lymphocytes isolated from chimeric mice were analyzed by FACS®. Again, as predicted for memory responses, an increase in the frequency of NP-specific memory B lymphocytes was observed both in the BM and spleen after antigen challenge in all chimeras (compare Fig. 2, b to d and c to e). For example, the frequency of NP-specific memory B lymphocytes in WT chimeras increased from a mean of 82 ± 12 to 420 ± 92 per 10⁶ cells in the BM and from 310 ± 55 to 3,280 ± 601 per 10⁶ splenocytes. The relative frequency of memory B lymphocytes was significantly reduced in the BM (160 ± 40/10⁶ cells) and spleen (1,720 ± 526/10⁶ cells) of CD21/CD35-deficient chimeric mice. Consistent with the reduced frequency of splenic ASCs, the relative frequency of memory B lymphocytes was reduced in spleens (1,560 ± 526/10⁶ cells) of FcγRIIB-deficient chimeric mice.

The finding of 10-fold differences in antibody production between WT and Cr2⁻⁻ chimeras, but only two- to threefold differences in memory B lymphocytes and ASC frequencies, suggested that donor-derived memory B lymphocytes in Cr2⁻⁻ mice might be functionally unresponsive. To address this question, serial adoptive transfers were performed. Splenic B lymphocytes from primary recipient mice were adoptively transferred along with antigen and T cells into sublethally irradiated WT mice. 3 wk after transfer, anti-NP titers and splenic ASC frequencies were determined. Consistent with earlier results, the mean anti-NP titer was approximately two- to threefold less in secondary chimeric mice receiving primed B lymphocytes from Cr2⁻⁻ primary chimeras compared with WT primary chimeras (2,267 ± 581 vs. 5,867 ± 533, P < 0.01; Fig. 4 a). Similarly, analysis of the frequency of splenic ASCs revealed an approximate twofold reduction in mice receiving primed B lymphocytes from Cr2⁻⁻ versus WT primary chimeras (55 ± 18.2 vs. 94 ± 7.4, P < 0.07; Fig. 4 c). Thus, the frequency of functional memory B lymphocytes was reduced in Cr2⁻⁻ chimeras as evidenced by their failure to respond optimally in WT recipients. The NP titer of recipients of primed B lymphocytes from primary FcγRIIB⁻⁻ donors was reduced by over fourfold relative to primary WT donors (1,200 ± 0 vs. 5,867 ± 533). Moreover, even greater reduction was observed in the frequency of NP-specific ASCs in secondary chimeras receiving B lymphocytes from FcγRIIB⁻⁻ primary chimeras (10.2 ± 1.8, P < 0.008; Fig. 4 c). Therefore, as observed in the spleens of primary FcγRIIB⁻⁻ chimeras, there was an impaired response after secondary transfer into WT recipients.
An additional factor in the diminished recall response of Cr2−/− chimeras is impaired antigen localization on challenge. To test this possibility, serial transfers were also performed using Cr2−/− mice as secondary recipients. An approximate twofold reduction in antibody titer was observed in Cr2−/− versus WT recipients (1,067 ± 133 vs. 2,533 ± 353, P < 0.03; Fig. 4 b). The difference in antibody titters between Cr2−/− secondary chimeras receiving either splenocytes from WT or Cr2−/− primary chimeras was again reflected at the level of ASC frequency (17 ± 4.9 for Cr2−/− vs. 25.8 ± 1.7 for WT; Fig. 4 d). Therefore, the 10-fold reduction in recall response of Cr2−/− chimeras was due in large part to both a reduction in memory B lymphocytes and a lack of complement receptors on recipient stroma.

**Antigen Deposition in the Absence of CD21/CD35.** To examine whether the ability to trap antigen efficiently contributes to differences in recall responsiveness between chimeric mice, the groups of recipient mice were injected with biotinylated NP-BSA 16 h before harvesting. Immunohistochemical analyses of splenic sections from WT and FcγRIIB−/− chimeras identified antigen within PNA+ GCs (Fig. 5, a and b). By contrast, antigen deposition was negligible within the GC of Cr2−/− chimeras (Fig. 5, c and d). The lack of localized biotinylated antigen in Cr2−/− mice was observed at multiple time points, ranging from 12–20 h after administration, suggesting that kinetic differences in antigen trapping are not responsible for the absence of detectable antigen in Cr2−/− chimeras (unpublished data). Importantly, antigen detected in WT and FcγRIIB−/− chimeras colocalized with FDCs within the light zones of GCs.

**Discussion**

The role of antigen in the maintenance of long-term B lymphocyte memory is subject to debate. In this study, the mechanism of antigen retention leading to long-term antibody production, persistence of memory B lymphocytes, and affinity maturation was addressed using an adoptive transfer model similar to that pioneered by Gray and Skarevall (9). Because memory responses using this model depend on introduced antigen, it is amenable to examining the contributions of receptors believed critical for capturing and retaining antigen by adoptively transferring memory B lymphocytes into mice deficient in either CD21/CD35 or FcγRIIB. We demonstrated that CD21/CD35 expression on radioresistant stomal cells (including FDCs) is important for maintaining serum titers (Table I), high frequencies of ASCs (Fig. 1) and memory B lymphocytes (Fig. 2), as well as for optimal affinity maturation (Fig. 3, Table II). Moreover, recall responses were severely compromised in the absence of CD21/CD35-bearing stroma (Figs. 1, c and d, 2, d and e, 4; Table I). In contrast, although splenic responses were impaired in FcγRIIB−/− chimeras, the absence of FcγRIIB on recipient cells did not affect overall antibody persistence or frequency of BM ASCs and surprisingly led to significantly increased serum affinity. These observations suggest that although both CD21/CD35 and FcγRIIB participate in trapping antigen, overall memory B lymphocyte responses are more dependent on stromal cell–associated complement receptors.

**Antigen-dependent maintenance of NP titers.** The experimental model used depends on antigen because WT mice receiving memory B and T lymphocytes without antigen at the time of cell transfer fail to sustain significant anti-NP titers. In addition, they are unable to mount significant recall responses 16 wk after the transfer of memory B lymphocytes (Table I). These data suggest that memory B lymphocytes are not maintained (in the absence of antigen) to the extent that they efficiently differentiate into antibody-producing plasma cells after antigen challenge. Interestingly, antibody titers in Cr2−/− chimeras were similar to WT chimeras not receiving antigen after 16 wk, suggesting that antigen retention via CD21/CD35 is critical for antibody persistence. However, chimeras lacking CD21/CD35+ stroma contained limited NP-specific ASCs. CD21/CD35 involvement in antibody persistence was examined previously, where it was reported that Cr2−/− mice immunized with T-dependent antigens generated reduced antibody responses (29, 30). In one report, it was noted that the decay of serum antibody after antigen challenge was faster in Cr2−/− mice compared with WT mice (29), consistent with the thesis that the ASC pool is not efficiently replenished in the absence of CD21/CD35. However, because these studies used Cr2−/− mice whose B lymphocytes and

![Figure 5](https://example.com/image.png)
folicular stroma lacked CD21/CD35, they were unable to distinguish CD21/CD35-dependent antigen trapping from its role as a B lymphocyte coreceptor. In this study, the responding memory B lymphocyte pool had intact B lymphocyte receptor signaling complexes and therefore the contribution of stroma-derived CD21/CD35 for antigen sequestration was isolated from coreceptor-associated B cell receptor signaling. The implication of reduced antibody titers in Cr2−/− chimeras in this study is that CD21/CD35 on FDCs is critical in long-term antigen retention, although other radioresistant CD21/CD35-expressing myeloid cells may also participate.

Differences in serum antibody levels in the various chimeric groups directly correlated with the frequency of NP-specific ASCs in BM. ASC frequency is reduced in Cr2−/− chimeras relative to WT chimeras, supporting the idea that active ASC replacement contributes to effective long-term B lymphocyte memory. Several recent reports addressed the possibility of long-lived ASCs. One report using bromodeoxyuridine labeling concluded that 60% of BM ASCs can survive in excess of 90 d without turnover (8). Another study demonstrated that a fraction of adoptively transferred ASCs continue to produce antibody for longer than 1 yr (7). The interpretations of the latter study were questioned, however, by another study using the same lymphocytic choriomeningitis virus model antigen system that failed to observe “antigen-independent long-lived persisting antibody” (28). Although experiments in this study did not address ASC lifespan directly, the observation that WT chimeras not receiving antigen still produced measurable, albeit significantly reduced, anti-NP titers is consistent with the presence of low numbers of long-lived ASCs (negligible ASCs were detected in the BM).

Interestingly, differences in antibody titers between WT chimeras and Cr2−/− chimeras was significant after 6–8 wk, which suggests that either fewer long-lived ASCs are generated initially after transfer and/or that continued production of ASCs is less efficient in mice without CD21/CD35 stroma. Long-lived ASCs, as well as memory B lymphocytes, are products of GC responses. Based on results using a Bcl-2 transgenic model, Smith et al. (31) proposed that predominantly high affinity ASCs produced in GCs are recruited to the BM. In the current model, the observation of similar antibody titers after 2–3 wk suggests that the initial GC responses between the chimeric groups are comparable. The subsequent differences in antibody titers observed between WT and Cr2−/− chimeras after 6 wk (i.e., after GC wane) could then be explained by less efficient replenishment of the BM ASC pool.

Affinity Maturation. To delineate whether post-GC selection contributed to altered antibody titers in WT and Cr2−/− chimeras, relative serum affinity changes were monitored. Early evaluation of affinity maturation demonstrated that affinity increases with time after initial antigen contact and that the degree of increase depends on the amount of antigen encountered (11, 32, 33). Because affinity increases with time, one presumes that either high affinity ASCs are selectively surviving or that the production and/or selection of new ASCs is ongoing (33, 34). Long-lived ASCs possess little to no surface Ig and are not responsive to antigen. Therefore, they are unlikely subject to selection (8, 24, 35, 36). In light of the data presented here, it appears that post-GC selection establishes new ASCs important for increases in relative serum affinity. Relative serum affinity was measured in this study using two independent assays. Previous work using the ELISA-based assay demonstrated that affinity measurements correlated with mutations within VDJ genes conferring high affinity binding (37, 38). Both techniques showed that over the course of 16 wk relative serum affinity increased modestly in WT chimeras whereas no statistically significant increase was observed in Cr2−/− chimeras (Fig. 3, Table II). Therefore, our results imply that antigen deposition via CD21/CD35 is important for post-GC selection, a thesis supported by the observation that Cr2−/− chimeras were less able to trap injected antigen (Fig. 5 c). Previous work using LTα−/− deficient mice concluded that FDCs are not required for affinity maturation (39). However, differences between these results versus those of this study are likely explained by the immunization protocols. Matsumoto et al. (39) immunized with much larger and more frequent doses of antigen in addition to using adjuvant, likely diminishing the importance of antigen deposition on FDCs for affinity maturation.

Memory B Lymphocyte Persistence. The longevity of memory B lymphocytes is proposed as antigen independent based on an experimental system in which the antigen specificity was switched (40). Additional support for antigen-independent persistence of memory B lymphocytes comes from reports that memory B lymphocytes are generally resting cells without measurable cell proliferation (41). Moreover, they persist in the absence of T cell help (42, 43) or chemokines and chemokine receptors required for FDC development (10, 39). In this study, the frequency of memory B lymphocytes is reduced significantly in WT chimeras not receiving antigen, although low frequencies were detectable predominantly in the spleen (Fig. 2 c). These memory B lymphocytes were donor derived because they were identified using an allo-specific marker and in addition, because irradiated control mice contained a negligible number of memory B lymphocytes. These data are consistent with reports suggesting that at least a fraction of the memory B lymphocyte pool is maintained in the absence of antigen. However, significantly more memory B lymphocytes were detected in WT chimeras receiving antigen, suggesting that a substantial fraction of the memory B lymphocyte pool is replenished with time. These frequencies of memory B lymphocytes detected by FACS® are similar to frequencies reported by Takahashi et al. (44). Similar to differences found with ASCs, optimal memory B lymphocyte frequency depended on stroma-derived CD21/CD35, a finding that was supported by three independent observations (Figs. 2 and 4). The consistent reductions in both ASC and memory B lymphocyte frequency observed between WT and Cr2−/− chimeras with and without antigen, in concert with changes in relative serum affinities, suggest that clonal selection continues...
after GC wane. Thus, it is suggested that both antigen-dependent and antigen-independent mechanisms drive B lymphocyte memory.

Role of Stroma-derived FcγRIIB. The importance of FcγRIIB expression on stroma was investigated by comparing memory responses in FcγRIIB−/− chimeras to WT chimeras. It was previously demonstrated that secondary responses in FcγRIIB−/− mice, as well as in FcγRI−/− mice (45), were normal (23). However, because these mice lack FcγRIIB on B lymphocytes and FDCs, these studies were unable to distinguish their relative contributions to the immune response. FcγRIIB is inducible on FDCs (22), making it a candidate for long-term retention of antigen via IgG immune complexes. Myeloid cells (including macrophages, monocytes, neutrophils, and mast cells) also express FcγRIIB along with FcγRI and RIII, and it is therefore possible that other radioresistant cells contribute to long-term antigen retention in the current system (46). Antibody persistence appears independent of FcγRIIB on FDCs because WT and FcγRIIB−/− chimeras had similar antibody titers and BM ASCs. However, both splenic ASC and memory B lymphocyte frequency were reduced in FcγRIIB−/− chimeras, suggesting either that antigen deposition via FcγRIIB is required for post-GC production of both splenic populations or that selection is more stringent in the absence of FcγRIIB (Figs. 1 and 2). Measurements of relative serum affinity seem to favor the latter hypothesis because affinity increases dramatically with time in FcγRIIB−/− chimeras (Fig. 3, Table II). One explanation is that selective pressure might be greater in FcγRIIB−/− chimeras due to increased negative signaling on B lymphocytes by unoccupied Fcγ portions of immune complexes not bound by FcγRIIB on FDCs. This possibility is based on the assumption that FcγRIIB on FDCs compete with FcγRIIB on B lymphocytes for binding IgG. Interestingly, FcγRIIB−/− chimeras are also similar to WT chimeras in their ability to trap injected antigen (Fig. 5 b). This might be due to increased trapping of antigen via CD21/CD35, although we did not directly address this possibility. An alternative hypothesis is that FcγRIIB has a signaling role on splenic FDCs that enhances the release of chemokines for retention of memory B lymphocytes and ASCs. FDCs are a source of at least one such chemokine, B lymphocyte chemokine-1, which promotes the localization of B lymphocytes within the splenic follicles (47, 48).

Recall Responses. B lymphocyte memory is best assessed functionally by measuring the recall response. Similar to findings of Gray and Skarvall (9), efficient recall responses in this study require antigen at the time of adoptive transfer. Although detectable donor-derived memory B lymphocytes in WT chimeras deprived of antigen, these mice did not generate a significant recall antibody response (Table I). Significantly, Cr2−/− chimeras demonstrated only a two-fold increase in antibody titer after challenge whereas titers in WT and FcγRIIB−/− chimeras increased 8–10-fold. Therefore, the recall response was impaired in the absence of CD21/CD35-bearing stroma. This result was additionally substantiated by secondary transfer of splenocytes from WT and Cr2−/− chimeras into WT secondary chimeras (Fig. 4, a and c). It is likely that several factors contributed to diminished recall responses in Cr2−/− chimeras. They had reduced serum antibody titers that potentially contributed to less efficient antigen trapping, as well as lower frequencies of memory B lymphocytes, necessary to mount a recall response. Interestingly, only two- to threefold reductions in ASCs and memory B lymphocytes were observed in Cr2−/− chimeras compared with WT chimeras, yet 10-fold differences in antibody titers were observed. This suggested that memory B lymphocytes in Cr2−/− chimeras were less responsive compared with those in WT chimeras due to less efficient trapping of antigen. This possibility seems likely because memory B lymphocytes from WT mice respond less efficiently after transfer into Cr2−/− recipients. Thus, the combined effects of fewer memory B lymphocytes and less efficient response in the absence of CD21/CD35 would explain the overall 10-fold reduction in antibody titers in Cr2−/− chimeras. Also intriguing is that despite a near normal antibody response in the FcγRIIB−/− chimeras, the frequency of splenic ASCs was significantly reduced both in the primary (Fig. 1 b) and secondary chimeras (Fig. 4 b). Thus, as discussed above, FcγRIIB seems to be required for the maintenance of memory B lymphocytes in the spleen independent of the amount of antigen trapping (Fig. 5 b).

In conclusion, the data presented here support a model whereby humoral memory is maintained by continuous antigen-dependent selection and/or differentiation of memory B lymphocytes and ASCs. FcγRIIB on splenic FDCs (or on radioresistant myeloid cells) also appears important for the maintenance of splenic memory B lymphocytes but is not required for overall memory recall responses. The implication from Cr2−/− chimeras is that ongoing clonal selection depends on antigen retained long-term on CD21/CD35-bearing FDCs. Lastly, the availability of CD21/CD35-bearing stroma provides the appropriate microenvironment for generating substantial recall responses.

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