Functional Heterogeneity of Marginal Zone B Cells Revealed by Their Ability to Generate Both Early Antibody-forming Cells and Germinal Centers with Hypermation and Memory in Response to a T-dependent Antigen

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Abstract

Marginal zone (MZ) B cells play a major role in the first-line responses against blood-born T-independent bacterial antigens (TI), but the full scope of their immune functions is not known. Here we compare the responses of MZ and follicular (FO) B cells to a T-dependent antigen (TD), hapten-(4-hydroxy-3-nitrophenyl)acetyl (NP) coupled to chicken γ-globulin, in a cell transfer system. Consistent with the conventional paradigm, MZ B cells but not FO B cells rapidly generated the early burst of NP-specific antibody-forming cells (AFC), high levels of IgM Ab, and early IgG with relatively high affinity to NP. However, MZ B cells were also capable of forming germinal centers (GCs), albeit with a delay, compared with FO B cells. The early AFCs and the GCs originated from different MZ precursors, but the MZ- and FO-derived GCs were similar in VH gene repertoire, somatic mutation, and production of late AFC and IgG Ab. Surprisingly, the MZ but not the FO memory response included IgM Ab. We conclude that MZ B cells are heterogeneous, comprising cells for both early AFC response and GC/memory pathway against TD antigens.

Key words: B cells • marginal zone • T-dependent Ag • antibody-forming cells • germinal center

Introduction

Newly formed B cells in adults migrate from BM to peripheral lymphoid tissues where they continue to differentiate into functionally and anatomically distinct subsets (1). Mature peripheral B cells include B1 cells, which reside mainly in peritoneal and pleural cavities (2), and the conventional B2 B cells. The mature B2 cell population is heterogeneous, consisting of the major follicular B cells (FO), in LN and splenic lymphoid follicles, and a subset residing in the marginal zone (MZ) of the spleen (3). MZ and FO B cells are distinguished by differential expression of several cell surface markers: MZ B cells are IgDlowCD21highCD23low, whereas FO B cells are IgDhighCD21interCD23high (4, 5). In addition, MZ B cells express various activation markers, such as high basal levels of CD80, CD86, CD40, and CD44, and low level of CD62L (5, 6). As a functional corollary, MZ B cells exhibit rapid and robust proliferation and Ig secretory responses to stimulation with LPS, anti-IgM, and CD40 ligands (5–7). The apparent hyperreactivity of MZ B cells and their unique anatomic localization at the red pulp junction strongly suggest that these B cells mediate rapid Ab responses to blood-born antigens (8).

There is increasing evidence for selection of B cells into the MZ or follicular pool through B cell receptor (BCR)-mediated signals (9, 10). Studies on transgenic mice demonstrated that B cells expressing BCR with specificity for different Ags had accumulated either in the follicles or in the MZ (11). In particular, B cells specific for the bacterial epitope, phosphorylcholine, homed to MZ and responded to the TI form of the Ag (3, 11). The ligands that control cellular selection and homing are not known (3). Interestingly, in humans (12, 13), and to a lesser extent in rats (14), some MZ B cells express somatically mutated IgV genes, suggesting

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Abbreviations used in this paper: AFC, antibody-forming cell; BCR, B cell receptor; CGG, chicken gammaglobulin; CSR, class-switch recombination; DN, double negative; FO, follicular; GC, germinal center; HRP, horseradish peroxidase; MZ, marginal zone; NIP, 4-hydroxy-3-nitrophenyl acetyl; NP, (4-hydroxy-3-nitrophenyl)acetyl; PALS, periarteriolar lymphoid sheath; PNA, peanut agglutinin; R/S, replacement to silent; SA-ALPH, Streptavidin conjugated to alkaline phosphatase; SHM, somatic hypermutation; TD, T-dependent Ag; TI, T-independent Ag; TI-I, TI type I Ag; TI-II, TI type II Ag.
that they already encountered Ags and became memory cells. Collectively, these findings suggest that the repertoire of MZ B cells is skewed by stimulation with environmental Ags.

Fagarasan and Honjo (15) proposed that MZ B cells respond to T-independent Ags (TI) and may not be regulated by T cells. This notion was based primarily on the phenotype of Pyk-2–deficient mice (16) that had a severely reduced MZ B cell population and diminished Ab response to TI type I Ag (TI-I) and TI type II Ag (TI-II). However, this defect was restricted only to the IgG2a and IgG3 isotypes, whereas IgG1 and IgG2b Ab responses to TI-I and TI-II Ags were unaffected. Moreover, the disruption of Pyk-2 also inhibited the IgM response to T-dependent Ags (TD). These conflicting observations suggest that the function of MZ B cells and their relationship with T cells are complex. Indeed, Tanigaki et al. (17) failed to find a relationship between MZ B cells and TI Ab responses. They generated mice lacking expression of RBP-J, a mediator of Notch signaling, in B cells using conditional mutagenesis. Such mice had no MZ B cells, but Ab responses to TI-I, TI-II, and TD Ag were unaffected.

The phenotypes of the Pyk-2 (16) and RBP-J (17) mutants have shown that MZ B cells are not dedicated only to TI Ab responses. MZ B cells present Ags and deliver costimulatory signals to T cells more efficiently than FO B cells in vitro (5), suggesting that MZ cells can mount rapid Ab responses requiring cognate T cell help (18). Indeed, using a hapten-carrier system, Liu et al. (19) demonstrated a rapid appearance of specific, hapten-binding MZ B cells in rats that had been primed previously to the carrier protein.

We hypothesized that the Ab response of MZ B cells to TD Ags is qualitatively different from that of FO B cells and that the two cell subsets differ in their potential to form antibody-forming cells (AFC), germinal centers (GCs), and memory cells. To test this prediction rigorously, we reconstituted scid mice with purified MZ and FO B cells from naive WT C57BL/6 donors, supplemented them with carrier-primed T cells, and then stimulated the chimeras with the hapten-[(4-hydroxy-3-nitrophenyl)acetyl (NP) coupled to chicken gammaglobulin (CGG). The NP-specific Ab response of IgG mice has been well characterized at the cellular and molecular level: NP-binding V_{H} regions are encoded by the group of V186.2/V3 genes of the J58 family; the dominant clonotype expresses the V186.2 segment rearranged to DFL16.1/2 and J_{H}2 segments in combination with the λ L chain (20–23). This response to NP thus provides a precise tool for comparing potential differences between MZ and FO B cells in repertoire and function. Our results show an unexpected functional heterogeneity of MZ B cells. Upon stimulation with TD Ag, MZ cells rapidly produce large numbers of AFC that have distinct clonotypic repertoire; however, these cells also give rise to GCs with characteristic somatic hypermutation and generate immunological memory.

Materials and Methods

Animals. Normal C57BL/6, B6.SJL-Ly5.1 (CD45.1) (both 8–12 wk), and C57BL/6 scid mice (8–10 wk) were purchased from The Jackson Laboratories and maintained in microisolator cages in the animal facility of the University of Maryland, Baltimore.

Antigens. NP and its analogue (4-hydroxy-5-iodo-3-nitrophenyl)acetyl (NIP) (Cambridge Research Biochemical) were conjugated at various substitution ratios to CGG (Sigma-Aldrich) or BSA (Amersham Biosciences) as described (24).

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**Figure 1.** Purification of splenic MZ and FO B cells. (a) T cell–depleted splenocytes were stained for CD21–FITC and CD23–PE, and the CD23^{hi} FO cells were separated by autoMACS with anti-PE beads (b). The CD23^{-} fraction (c) was stained with B220–APC, and the B220^{+}CD23^{-} MZ cells (d) were separated by FACS from the CD21^{−}CD23^{−} DN fraction (e). The FO and MZ B cell fractions were reanalyzed before the cell transfer (b and d).
Antibodies. Anti-Thy1.2 (HO13-4), anti-CD4 (GK1.5), and anti-CD8 (3.155) hybridomas (American Type Culture Collection) and anti-CD3 hybridoma (145-2C11, provided by Dr. Jeffrey A. Bluestone, University of California, San Francisco, CA) were grown in our laboratory, and the Abs were isolated from culture supernatants by salt precipitation. Anti-B220-APC (RA-6B2), anti-CD23-PE (B3J4), anti-CD21-FITC (7G6), anti-CD19-P (1D3), anti-CD11b-biotin (M1/70), anti-CD11c-biotin (HL3), anti-CD45.2-biotin (104), anti-CD45.1-biotin (A20), and GL-7-FITC were purchased from BD Biosciences. Horseradish peroxidase (HRP)-conjugated goat anti-mouse IgM, IgG, IgG1, IgG2a, IgG2b, IgG3, λ, and κ were obtained from Southern Biotechnology Associates, Inc.

Purification of MZ, FO B Cells. Single spleen cell suspensions were prepared by grinding spleens between two frosted glass slides in medium consisting of RPMI 1640 with 25 mM HEPES (Life Technologies) and 0.5% BSA (Sigma-Aldrich). B cell–enriched populations were prepared by depleting T cells using two treatments with an antibody cocktail consisting of anti-CD4 (GK1.5), anti-CD8 (3.155), anti-Thy1.2 (HO-13-4), and normal rabbit serum, at 37°C for 40 min. The enriched B cells were stained with anti-CD23-PE on ice for 15 min followed by incubating with anti-PE microbeads (Miltenyi Biotec), and the CD23+ B cells (FO B cells) were separated from the CD23− B cells by autoMACS (Miltenyi Biotec). The resulting CD4 T cells were enriched by passing the spleen cell suspension through a MACS LS column (Miltenyi Biotec). The resulting CD4 T cells contained 100% of CD8 T cells and CD20+ FO cells.

Adoptive Transfer and Immunization. ~2–2.5×10^6 of purified MZ or FO B cells, together with 4×10^6 CGG-primed CD4 T cells were injected i.v. into C57BL/6 mice, and the recipients were immunized i.p. with 40 μg of NP-CCG in alun. Blood and/or spleen samples were collected for analysis of the primary response at days 4, 8, 14, 36, 60, and 85. To measure memory responses, splenocytes from the recipient mice were collected at day 85 after primary immunization, and 5×10^6 cells were transferred into sed mice. The mice were challenged with 10 μg of soluble NP-CCG i.p., and memory responses were examined at days 6 and 9 after the boost. The sed recipients that were not challenged, and the recipients of naive splenocytes that received the boost dose of soluble Ag served as controls.

In experiments with mixed B cell subsets, MZ cells were prepared from C57BL/6 mice expressing the Ly5.2 allele of the common leukocyte antigen CD45, whereas the FO cells and the CD21CD23− labeled as “double negative”) B cells were from BL6.SJ-Ly5.1 (CD45.1) donors.

Antibody Measurement. Serum NP-specific Abs were measured by standard ELISA using 96-well plates coated with 100 μg/ml of NP18-BSA. The relative affinity of serum Ab was determined by titration on ELISA plates coated with BSA coupled to NP at different ratios, respectively, NP18-BSA and NP18-BSA. The plates were developed with HRP-conjugated goat Ab specific for mouse IgG, that is the major isotype of anti-NP Ab (>80% of IgG) in this system. Binding (OD450) to each plate at the linear portion of the titration curve was determined, and the relative affinity of Ab was expressed by binding to NP18-BSA as a percentage of binding to NP18-BSA (Eq. 1):

\[
\text{affinity} = \frac{\text{OD450}_{\text{w},\text{NP18-BSA}} - \text{OD450}_{\text{w},\text{BSA}}}{\text{OD450}_{\text{w},\text{NP18-BSA}}} \times 100.
\]

Phenotypic Analysis of GC B Cells by FACS©. Mice were killed at 14 d after the cell transfer and immunization, and the splenocyte suspension was stained with anti-CD19-PE, GL7–FITC, and biotinylated anti-CD45.2 (Ly5.2) or anti-CD45.1 (Ly5.1) followed by SA–APC. GC B cells were enumerated as GL7+/Ly5.1+ or GL7+/Ly5.2+ cells within the CD19 gate.

Immunohistochemical Staining of GC and APC and Microdissection of Target Cells. Frozen splenic sections were prepared and stained as described previously (25). The specific, NP-reactive GCs were identified by dual staining with peanut agglutinin (PNA) coupled to horseradish peroxidase (PNA-HRP) (EY Laboratories, Inc.) in combination with a biotinylated NIP-BSA followed by Streptavidin conjugated to alkaline phosphatase (SA-ALPH) (ImmunoPure). NP-specific AFCs were visualized as plasmacytes stained with biotinylated NIP-BSA. These cells formed typical foci close to the borders of B cell follicles. PNA+NP+ GC cells and NIP-positive AFCs were recovered using a sharpened micropipette controlled by micromanipulator (model MM188; Nikon). The recovered cells were lysed with PCR-lysing buffer (1× PCR buffer containing 100 μg/ml proteinase K, 0.05% Tween 20) at 56°C for 1 h followed by inactivation of proteinase at 95°C for 10 min. Resulting samples were used for PCR amplification.

In mice reconstituted with cells expressing different CD45 alleles, the GCs in splenic sections were visualized by dual staining with PNA-HRP and anti-Ly5.1-biotin plus SA-ALPH. The dual-stained PNA+Ly5.1+ GCs were readily distinguishable from single-stained PNA+Ly5.1− GCs in the same section (see Fig. 4 in Results), and the latter were then scored as Ly5.2+. The anti-Ly5.2 conjugate could not be used for identification of GCs in sections because of the high background staining.

PCR Amplification, VH Gene Repertoire Screening, and Mutation Analysis. Amplification of VDJ rearrangements using two pairs of primers for nested PCR was performed as described previously (25, 26). PCR products were cloned using pBluescriptII SK plasmid (Stratagene) and DH5α Escherichia coli and screened by hybridization with VH gene–specific, 32P-labeled oligonucleotides (26). All transformed colonies hybridized with the probe (5′-GTAGGCTGGCCTTGACAGGA-3′) corresponding to the region of genomic DNA (as position 21–27) that is shared by the V186.2/V3 group of 22 genes of the J558 family. A proportion of these colonies also hybridized with the probe (5′-TACCCAGCAGGACATCTTTCC-3′), which identifies a DNA region (as position 50–57) specific for the V186.2 gene sequence. The number of 21–27+/50–57− colonies has been shown to correlate with the number of B cells expressing one of the analogue (non-V186.2) genes of the V186.2/V3 group (26). Plasmid DNA from the double-positive (21–27+/50–57+) colonies was extracted and sequenced by the Biopolymer Laboratory of the University of Maryland, School of Medicine using an automatic DNA sequencing system (Applied Biosynthesis). All double-positive colonies invariably yielded the rearranged V186.2 sequence. Mutations were scored based on nucleotide substitutions in the VH gene only.
Shared mutations within a set of clonally related sequences (according to the CDR3) were counted as one mutational event.

**Results**

Distinct Cellular Responses of B Cell Subsets. Scid mice were reconstituted with either highly purified MZ cells (“MZ mice”) or FO cells (“FO mice”) plus carrier-primed T<sub>H</sub> cells as described in Materials and Methods and Fig. 1. These recipients demonstrated marked differences in their responses to immunization with NP-CGG (Fig. 2). MZ mice exhibited an early extrafollicular AFC response with the appearance of NP-binding plasma cells on day 4 and large AFC foci formation by day 8 (Fig. 2 a and b), whereas the formation of the GC was relatively slower. Only few follicles stained faintly with PNA on day 8 (Fig. 2 c); PNA<sup>+</sup> GCs were seen in less than half of splenic sections, numbering less than three GCs/section (Table I). However, by day 14 the GCs in MZ mice reached typical size (Fig. 2 d) and numbered up to six per section (Table I).

In contrast, FO mice had no AFC on day 4 (Fig. 2 e) and only few, scattered NP-binding plasmacytes were found on day 8 (Fig. 2 f); AFC foci were not seen until day 14 (not depicted). However, FO mice developed large PNA<sup>+</sup> GCs from day 8 (Fig. 2 g) that were apparent in every splenic section (Table I) and that continued on day 14 (Fig. 2 h).

Although the purity of MZ cells was >95% (Fig. 1 d), we considered the possibility that the GCs in MZ mice resulted from a small (≈5–10%) cell contamination. Since the CD23<sup>+</sup> FO cells were removed by autoMACS before FACS<sup>®</sup> sorting for MZ cells, we considered that the most likely contamination was from the CD21<sup>lo</sup>/CD23<sup>+</sup> double negative (DN) immature B cells that might have completed their maturation to FO cells and form GCs after the adoptive transfer. We addressed this concern by reconstituting scid mice with purified Ly5.2<sup>+</sup> MZ cells (2.7 × 10<sup>6</sup>) that were mixed with 3 × 10<sup>5</sup> cells from the CD21<sup>lo</sup>/CD23<sup>+</sup> B cells (Fig. 1 e) that were isolated from Ly5.1<sup>+</sup>

### Table I. Germinal Centers in the Spleen of Mice with MZ and FO B Cells

| Days postimmunization | B cells transferred | GC/section (range) | Positive/total<sup>a</sup> |
|-----------------------|--------------------|--------------------|-----------------------------|
| 8                     | MZ                 | 0–3                | 10/21                       |
|                       |                    | 0–1                | 1/4                         |
|                       | FO                 | 0–8                | 10/11                       |
|                       |                    | 1–4                | 14/14                       |
| 14                    | MZ                 | 0–6                | 5/10                        |
|                       |                    | 0–2                | 5/10                        |
|                       |                    | 0–1                | 3/9                         |
|                       |                    | 0–5                | 6/10                        |
|                       |                    | 0–3                | 9/14                        |

<sup>a</sup>Serial longitudinal sections were prepared from two to three spleens/group (20–50 sections through ≈50–60% of the splenic thickness). PNA<sup>+</sup> germinal centers were scored on several sections separated by ≈20–30 μm.

<sup>b</sup>Positive sections contained at least one GC.
donors (see Materials and Methods). It should be emphasized that CD21 expression in the DN fraction is variable, with a few cells staining nearly as bright as the MZ cells (Fig. 1 c). Therefore, substantial numbers of GC B cells in these chimeras should be Ly5.1\+/H11001 if the GCs originated from the immature cells rather than MZ cells. However, at 14 d after the transfer and immunization, the numbers of GC B cells were nearly proportional to the input: 90% of GL-7\+ GC B cells were Ly5.2\+/H11001 by FACS® (Fig. 3 A, Table II, Group A), and all but one GC in splenic section were PNA\+/Ly5.1\+/H11002 (Table II, Group A). Thus, GCs in the MZ mice do not arise from immature B cells.

MZ Cells Form GCs in Competition with FO Cells. It could be argued that in normal spleen the potential of MZ cells to form GCs is low, because the majority of GCs are generated by FO cells. However, in these chimeras, FO cells are scarce, and thus the GC potential of MZ cells is revealed. FO cells are likely not essential for GC formation, as MZ cells can form GCs without FO cells in the absence of FO cells. However, FO cells may play a role in the regulation of GC formation, as FO cells are known to secrete cytokines that are essential for GC formation. In conclusion, the data presented here suggest that MZ cells have the potential to form GCs, and that FO cells may play a role in the regulation of GC formation.

Table II. Phenotypes of GC in Mice Reconstituted with MZ Plus FO and/or DN B Cells at Various Ratios

| Number of injected B cells | GC B cells in the recipient spleens\(^a\) |
|---------------------------|------------------------------------------|
|                           | FACS\(^b\) analysis (%) | GC enumeration | Ratio |
|                           | (LY5.2\+/LY5.1\+) | (LY5.1\--GL-7\+) | (LY5.2\+/LY5.1\+) | (LY5.1\--PNA\+) | (LY5.1\--PNA\+) | (LY5.1\--/LY5.1\+) |
| Group | MZ | DN | A | 2.7 \times 10^6 | 3 \times 10^5 | 9:1 | 20 | 2 | 10 | 14.6 ± 5 | 1\(^d\) | 14 |
|       | MZ | FO | B | 2.7 \times 10^6 | 3 \times 10^5 | 9:1 | 16 | 10 | 1.6 | 17.4 ± 4.7 | 7.4 ± 3 | 2.4 |
|       | MZ | FO | C | 3 \times 10^5 | 2.7 \times 10^6 | 1:9 | 4 | 13 | 0.3 | 2.6 ± 1 | 12.6 ± 2.7 | 0.2 |

\(^a\)Day 14 after adoptive transfer and immunization; three recipients/group.

\(^b\)Pieces (~20%) of the spleens were pooled, stained, and gated on CD19 bright cells.

\(^\)Number of PNA\+ centers in 10 spleen sections; mean from three mice ± SD. See Materials and Methods and Fig. 4 for details on visualization of Ly5.1\+ and Ly5.1-GC.

\(^d\)One PNA\+ center was found in one out of three mice.
cells to form GCs is eclipsed by a much larger population of FO cells. We examined this point by cotransferring the purified Ly5.2+ MZ cells and Ly5.1+ FO cells at ratios of 9:1 (group B) and 1:9 (group C) together with T\textsubscript{H} cells. The numbers and the Ly5 phenotype of splenic GL-7\textsuperscript{+/H11001} input. Group B received nine times more Ly5.2 and their respective numbers vary according to the cellular than Ly5.1\textsuperscript{+/H11001}.Ly5.2\textsuperscript{+/H11001} contains three PNA\textsuperscript{+/H11001} staining is shown in a splenic section from group B that described in Materials and Methods. An example of the dual stainings were shown by costaining with anti-Ly5.1 as described in FACS® analysis (Table II). GC B cells in the mixed chimeras were formed by either the Ly5.2+ MZ cells or Ly5.1+ FO cells, and their respective numbers vary according to the cellular input. Group B received nine times more Ly5.2+ MZ cells than Ly5.1+ FO cells (ratio 9:1), but the numbers of Ly5.2+ GC B cells were only about twofold higher than Ly5.1+ GC B cells as determined by either FACS® (Fig. 3 B) or by section staining (Table II). This is consistent with the notion that GC precursors in the FO cell subset are more frequent and/or robust compared with MZ B cells. However, this trend was not apparent when the MZ:FO input ratio was reversed to 1:9 in group C (Table II). The numbers of Ly5.2+ GC B cells (Fig. 3 C) were nearly proportional to the input of Ly5.2+ MZ cells. It is conceivable that these results were influenced by the mechanisms of homeostasis and cell expansion that may be different for MZ and FO subsets. Nevertheless, the results argue strongly that the GCs detected in MZ mice are not derived from contaminating FO cells.

Serum Antibody Responses. Consistent with the pattern of cellular responses in single-reconstituted animals, the MZ mice mounted a robust NP-specific IgM serum Ab response during the first week after the immunization (Fig. 5 a) and switched to IgG Ab on day 8 (Fig. 5 b). FO mice did not produce any detectable serum Ab until day 8; the level of IgM was 10-fold lower than in the MZ mice, whereas the IgG titers had a similar range among individual mice in both groups. Thus, the massive, early AFC formation in the MZ mice correlates well with the anti-NP IgM response. The origin of serum Ab in FO mice can be either from the dispersed plasmacytes (Fig. 2 f) or perhaps from B cells in the follicles, which might have secreted Ig without differentiation into typical plasmacytes. Interestingly, the MZ mice maintained 10-fold higher IgM Ab titers throughout the entire 2-mo period of observation, and as described below, they also displayed a brisk, recall IgM Ab response to the Ag boost. Each of the two independent experiments shown in Fig. 5 included groups of control mice reconstituted either with whole unseparated splenic B cell population together with purified T\textsubscript{H}, or with T\textsubscript{H} alone, and immunized with NP-CGG. Notably, mice with unseparated B cells maintained intermediate levels of IgM Ab relative to the MZ and FO mice, confirming that MZ B cells, which represent 5–10% of spleen B cells, make the major contribution to IgM responses. None of the mice from the control group reconstituted with only T cells produced detectable anti-NP serum Ab, demonstrating that neither a contaminating donor B cell population nor “leaky” host B cells contributed to the anti-NP responses (not depicted).

VD Gene Repertoire of Anti-NP Responses. The rearranged V\textsubscript{H} genes in the NP-responding B cells in FO and MZ mice were examined by screening bacterial transformants with probes that discriminate between the V186.2 segment and the remaining “analogue genes” of the V186.2/V3 family. The V\textsubscript{H} gene repertoire was expressed as the ratio between V186.2 and non-V186.2 (analogue) genes, and the rearranged V186.2/D/J segments were sequenced. The most prominent finding in the MZ mice was the difference in the repertoire between the early AFC and the GC. Most AFCs (>70%) at day 8 used the V186.2 segment, but surprisingly few, if any, of those V\textsubscript{H} genes were
rearranged to the DFL16.1/2 segment (Fig. 6 a). Instead, the CDR3 was encoded by DSP 2–5, and other unidentified D genes and their length varied from 7 to 11 aa (Fig. 6 c). In contrast, the incipient GCs in MZ mice on day 8 were dominated by the non-V186.2 analogue genes; only 20% of GC B cells used the V186.2 segments, half of which were rearranged to DFL16.1/2 genes (Fig. 6 a). Unlike the early AFC repertoire, AFC recovered from MZ mice on day 14 expressed the canonical V186.2/DFL16.1/2 rearrangements (Fig. 6 a) with uniform CDR3 length (11–12 aa) (not depicted). Moreover, the repertoire of these late AFC was similar to that of the GC, and clones with identical CDR3 were found in both cell populations.

As expected, most of the V186.2 sequences recovered from early AFC foci in MZ mice were unmutated (27); only one focus contained somatically mutated AFC with an average of 1.2 mutations/VH (Table III). It is unlikely that the mutated AFC originated from the rare, nascent GCs in MZ mice because the focus was microdissected from a splenic region devoid of PNA⁺ follicles, and the VD gene repertoire of early AFC in these mice was very different from GC/late AFC repertoire. Thus, consistent with previous studies (21), the presence of mutations in early MZ AFCs suggests that some MZ B cells are immunologically experienced.

In FO mice, microdissection of the rare, early AFC for repertoire analysis was not technically feasible as they were not organized in foci (Fig. 2 f). However, 50% of GC B cells on day 8 used the V186.2 segment, and this proportion increased further on day 14, which is typical of the repertoire of NP-reactive GCs in intact BL/6 mice (20). Approximately half of the V186.2 segments in FO GCs were rearranged to DFL16.1/2 (Fig. 6 b). Day 14 AFCs in these mice expressed a similar repertoire as the GC B cells and often shared CDR3 of uniform length (11–12 aa).

**Somatic Hypermutation in GC and Late AFC.** Studies have shown that immunocompromised mice (28, 30) or mice immunized with the bacterial epitope phosphorylcholine (28, 29) may develop GCs that appear to be morphologically normal but are deficient in somatic mutation
and/or affinity maturation. Therefore, it was important to assess whether the GCs in both MZ and FO mice supported somatic hypermutation in the V186.2⁺ B cells. Other V₃₄ clonotypes, which were identified by hybridization with a crossoptive DNA probe (see Materials and Methods) were not sequenced. As shown in Table III, ~60–70% of V186.2 sequences from day 8 GCs were mutated, with an average of 1.7 mutations/V₃₄ in FO mice and 1.9 mutations/V₃₄ in MZ mice. These numbers increased to 2.5 mutations/V₃₄ and 2.8 mutations/V₃₄, respectively, by day 14, which is consistent with the mutation frequencies observed in the V186.2⁺ GC B cells in NP-immunized intact C57BL/6 mice (20). There were high ratios of replacement to silent (R/S) mutations in CDR1 and 2, indicative of positive selection in both FO (R/S = 8) and MZ (R/S = 11) GCs (Table III, day 14).

On day 14, the majority of AFCs in both FO and MZ mice was somatically mutated (Table III) with high R/S ratios in CDRs and the canonical NP-specific mutation in V₃₄ position 33 that replaces Trp(W) with Leu (L) (27), which are unmistakable signs of their origin from GC. Indeed, clonally related AFC and GC cells that shared somatic mutations were occasionally recovered from adjacent sites. In the case shown in Fig. 7, sequences from two foci, AFC-5 and AFC-20, shared four mutations with the adjacent GC-5, and shared five additional mutations with each other. The sequences from AFC-5 also contained seven unique mutations. Because of limited sampling, we cannot determine whether all AFC mutations occurred in the GC or whether the process of somatic hypermutation was perpetuated in the AFC focus.

**Antibody Affinity and L Chain Repertoire.** We next asked whether the distinctive repertoire of the early AFC response in MZ mice is reflected in the serum Ab quality. Unexpectedly, the NP-specific IgG in MZ mice had higher affinity on days 8 and 14, as indicated by the relative binding to NIP₄-BSA/NIP₁₈-BSA (Fig. 8). IgG from FO mice did not bind to NIP₄-BSA on day 8 and only matured a little by day 14.
### Table III. Somatic Mutations in VH186.2 Genes in FO- and MZ-derived AFC Foci and GC

| Cells sampled | Days after immunization | Recipients (mice/group) | Number of sites sampled | VH186.2 sequences | Mutations/VH (average) | W33L mutation | R | S^d | CDR1 + 2 | FW |
|---------------|-------------------------|-------------------------|-------------------------|-------------------|-----------------------|----------------|---|---|-------|-----|
| AFC foci      | 8                       | FO (2 mice)             | ND                      |                   |                       |                |    |   |       |     |
|               |                         | MZ (2 mice)             | 1                       | 12                | 0                     | 0              |    |   |       |     |
|               |                         | 2^e                     | 10                      | 0                 | 0                     | 12             | 10.0 | 1.7 |
|               |                         | 1^e                     | 8                       | 60                | 1.2                   | 7              | 8.0  | 1.7 |
|               | 14                      | FO (2 mice)             | 3                       | 20                | 85                    | 1.5            | 12             | 10.0 | 1.7 |
|               |                         | MZ (3 mice)             | 4                       | 18                | 95                    | 1.6            | 7              | 8.0  | 1.7 |
| GC            | 8                       | FO (2 mice)             | 4                       | 25                | 62                    | 1.7            | 0              |     |     |
|               |                         | MZ (2 mice)             | 2                       | 14                | 75                    | 1.9            | 0              |     |     |
|               | 14                      | FO (2 mice)             | 7                       | 32                | 90                    | 2.5            | 8              | 8.0  | 1.4 |
|               |                         | MZ (3 mice)             | 4                       | 13                | 100                   | 2.8            | 0              | 11.0 | 2.1 |

^aNumber of scid mice reconstituted with either FO B cells or MZ B cells plus T+ cells and sacrificed on the indicated day after immunization with NP-CGG.
^bAt least one focus (~20–50 AFC) and one GC was microdissected from each spleen.
^cNumber of sequences with tryptophan → leucine replacement in VH position 33.
^dReplacement/silent mutation ratio.
^eThese foci were dissected from the same spleen.

Figure 7. VH186.2/DFL16.2 nucleotide sequences recovered from adjacent GC #5 and AFC foci #5 and #20 in mice reconstituted with FO B cells on day 14 after the immunization. Colored bases indicate common point mutations that were found in sequences from all three sites (green) and those found in GC #5 and AFC #5 (blue), GG #5 and AFC #20 (purple), and AFC #5 and AFC #20 (orange).
The Abs from both groups reached nearly equivalent binding to NIP4/NIP18 after 1 mo. The anti-NP antibody in MZ and FO mice also differed in the L chain isotype. Although Abs with lambda chain were prevalent in both groups (Fig. 8 a), the MZ mice had ~2 log higher titers of kappa\(^+\) anti-NP Ab than the FO mice (Fig. 8 b); the two groups reached similar levels of kappa\(^+\) Ab after 2 mo. An increased proportion of kappa\(^+\) Ab is typical of the late primary or secondary response to NP (31). Together, the noncanonical CDR3 repertoire of AFC, the higher affinity of early Ab, and the higher proportion of kappa\(^+\) Ab suggest that the rapidly responding NP-specific MZ B cells had been immunologically preselected for their role in early responses.

Anamnestic IgM and IgG Responses. Lastly, we examined whether the mice reconstituted with either MZ or FO B cells and primed with NP-CGG/alum developed immunological memory. Both groups maintained high titers of anti-NP IgG even at 3 mo after the immunization, which could have masked the de novo response to Ag boost. Therefore, the splenocytes from the NP-CGG-primed MZ and FO mice were adoptively transferred into C57BL/6/scid recipients, and the recipients were challenged with 10 \(\mu\)g of soluble NP-CGG (\(\Delta\), \(\bigcirc\), and \(\bigtriangleup\)) or not (\(\Delta\) and \(\bigcirc\)).

Discussion
We have delineated the functional differences and similarities between MZ and FO B cells in response to a TD antigen in vivo. We demonstrate, for the first time, that
hapten-reactive MZ B cells are functionally and clonally heterogeneous and that they are capable of forming GCs. Although the bulk of the early primary response by MZ and FO is skewed to the AFC and GC formation, respectively, evidence is provided that the functions of these two cell subsets, in fact, overlap.

MZ B cells have been identified previously as a primary source of early AFC response to TI Ags (8). The potential role of these cells in response to TD hapten–protein conjugates was explored by Liu et al. (19) using immunohistological techniques. In their view, the Ag challenge induced a rapid movement of hapten-specific B cells into the MZ where they differentiated into hapten-binding cells, which were regarded as likely candidates for transport of immune complexes back to the follicles. However, the hapten-binding MZ B cells appeared to have no relationship to the plasmacytes that arose concurrently in the splenic red pulp. We now provide direct evidence that MZ B cells are the major source of the early primary AFC plasmacytes in response to a TD Ag.

The repertoire of the rapid MZ AFC response to NP had several unexpected features. The usage of V_{H} 186.2 gene was higher than that typically observed in the primary GC (20, 26; Fig. 6 a), however, this V_{H} 186.2 segment was rarely joined to the DFL16.1/2 gene, contrary to the canonical V186.2/DFL16.1 rearrangements that are found later in the NP-driven response (21–23; Fig. 6, a and b). Interestingly, the use of noncanonical D segments was shown to be a characteristic of secondary anti–NP repertoire based on analysis of hybridomas (21). Here we show that in addition to the unique VD repertoire, the early MZ-derived AFC contained occasional point mutations in the V_{H} gene and produced circulating IgG Ab that had higher antibody affinity for NP than the primary IgG Ab produced by FO B cells. These results suggest that the first wave of AFC arises from a subset of NP-reactive B cells that have been selected into MZ pool by stimulation with an NP-crossreactive epitope, either a self-ligand (3) or an environmental Ag. We hypothesize that this “natural priming” fortuitously stimulates an expansion of cell clones that react with NP more avidly than the naïve NP-specific B cells. The theory that the BCR of these cells use kappa light chains instead of lambda may be conjectured from the observation of higher levels of kappa^{+} anti-NP Ab in the MZ mice (Fig. 8 b).

Figure 9. Working model of splenic B cell responses to a TD antigen. Antigen-reactive (“naturally primed”) MZ B cells produce the large foci of early AFCs (1) similar to the model of TI response in reference 5. Clonotypically distinct MZ B cells become activated (2) and either enter GC (4) or differentiate into late AFC (3); the majority of late AFCs derive from GC (5). MZ-derived memory cells (6) produce anamnestic IgM and IgG Ab upon restimulation. FO B cells are the main source of GC (2a and 4a), producing late AFCs either directly (3a) or from GC (5a) and generating IgG-producing memory B cells (6a). Early AFC may be formed either by an FO B cell subset or by the GC-committed population (1a).
The adoptive cell transfer system revealed the potential of purified MZ B cells to generate conventional GC upon stimulation with TD Ag in the presence of Ag-primed Th cells. Up to 20% of B cells in the recipient spleen expressed the GL-7 marker of GC, which is within the upper ranges of GC reaction reported in other studies (27, 32). Using B cell subsets expressing different Ly5 alleles, we showed that the majority of GCs in MZ-reconstituted mice came from the MZ cells rather than from contaminating immature cells and that MZ B cells were able to generate the GL-7+ GC cells even in the presence of excess FO B cells. In chimeras that received Ly5.2+ MZ and Ly5.1+ FO cells, individual PNA+ splenic GC were entirely either Ly5.1 positive or negative. We have not observed a mosaic staining pattern with a mixture of PNA+Ly5.1- and PNA+Ly5.1+ cells within a GC (Fig. 4), suggesting that individual centers were founded either by MZ or FO B cell precursors, consistent with the theory of a pauciclonal origin of GC (20). Thus, our data clearly demonstrate that MZ B cells are functionally heterogeneous, containing distinct precursors for the rapid AFC formation and for GC formation (Fig. 9). When BCR on MZ cells binds Ag, they move rapidly to the splenic T cell zones (4, 33). We conjecture (Fig. 9) that some of these cells receive helper signals for migration to follicles and GC formation (Fig. 9).

It was surprising that MZ B cells, despite their rapid switch from IgM to IgG, continued to produce high levels of IgM Ab during a 3-mo period after the immunization and that they could also produce a robust rapid IgM anamnestic response. This implies that a significant proportion of Ag-stimulated MZ B cells does not undergo class-switch recombination (CSR) and that the cells remain at the VDJ/μ configuration even during their differentiation into memory B cells in GC. An induction of a robust IgM memory by priming with TD antigens has been observed previously (34, 35); our results now identify MZ B cells as the source of this IgM memory. It may be that MZ B cells respond less well to TD signals (cytokines or cognate stimuli) that are presumably required for the accessibility of DNA for CSR downstream of Cμ (36). An alternative mechanism for the long-term IgM-producing B cells is suggested from the recent demonstration by Dudley et al. (37) that increased activity of AID, an enzyme required for CSR, may lead to deletion of the 5’ internal switch μ sequence in some IgM-producing B cells, thus preventing any further isotype switch. We hypothesize that this mechanism occurs in the “naturally activated” MZ B cells but not in the FO B cells.

The working model in Fig. 9 proposes that the FO cells are committed mainly to GC formation and the late primary AFC. It is not clear whether the first anti-NP serum Ab in FO cell–reconstituted mice originated from the rapidly expanding GC or from an independent, small population of AFC precursors (Fig. 9, 1a). Still, the model of TD response in Fig. 9 emphasizes distinction between the earliest wave of AFC from MZ B cells and the later AFC that were produced by both MZ and FO cells, which had a VH gene repertoire similar to that of respective GC. Clonally related cells with shared point mutations were readily recovered from AFC foci and GC on day 14, indicating that the AFCs originated from those B cells within GCs that up-regulate Blimp-1 and differentiate into plasmacytes (38) (Fig. 9, 5 and 5a). However, a proportion of the late AFCs was unmutated; yet, some of them may also share CDR3 with GC B cells (39). This suggests that daughters of single, antigen-driven MZ and FO B cells may either form GCs and/or become plasmacytes (Fig. 9, 3, 3a, 4, and 4a).

Our results support the notion that the response of MZ B cells to a single epitope, NP, can be heterogeneous both functionally and clonotypically. Further studies are needed to test the hypothesis that MZ B cells include distinct clones, either which produce the rapid AFC response or those with different genealogy and genetic program that form the GC.

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