Visualization of the Genesis and Fate of Isotype-switched B Cells during a Primary Immune Response

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Abstract

The life history of isotype-switched B cells is unclear, in part, because of an inability to detect rare antigen-specific B cells at early times during the immune response. To address this issue, a small population of B cells carrying targeted antibody transgenes capable of class switching was monitored in immunized mice. After contacting helper T cells, the first switched B cells appeared in follicles rather than in the red pulp, as was expected. Later, some of the switched B cells transiently occupied the red pulp and marginal zone, whereas others persisted in germinal centers (GCs). Antigen-experienced IgM B cells were rarely found in GCs, indicating that these cells switched rapidly after entering GCs or did not persist in this environment.

Key words: lymphocyte activation • marginal zone • T cells • transgenic mice • germinal centers

Introduction

IgG, IgA, and IgE antibodies participate in elimination of microbes from the host by activating complement and promoting phagocytosis. These antibodies are created when naive IgM+, IgD+ B cells undergo H chain isotype switch recombination, which involves a DNA deletion between switch regions upstream of the γ, α, or ε H chain constant exons (1). The accessibility of the recombinase to a particular switch region is correlated with germline transcription and splicing (2–5), whereas the recombination process itself is thought to be initiated by the activity of activation-induced cytidine deaminase (6, 7). Signals promoting germline transcription are regulated in vivo by cell interactions and secreted cytokines (8). Thus, although some switch recombination can occur in response to certain polyvalent antigens in a T cell–independent fashion, switching is greatly facilitated by cognate interactions with antigen-specific CD4 helper T cells (9).

In situ detection of antigen-specific B cells has revealed several aspects of CD4 T cell–dependent B cell activation and isotype switching. Naive IgM+, IgD+ B2 cells circulate through the B cell–rich follicles of secondary lymphoid tissues (10). After exposure to antigen, antigen-binding IgM+, IgD+ B cells migrate from random positions within the follicles to the border between the follicles and T cell–rich areas (11–13), where cognate interactions with antigen-specific CD4 T cells can occur (14). The B cells proliferate and foci of antibody-secreting plasmablasts, some of which contain isotype-switched antibodies, appear at the border between the T cell areas and the red pulp of the spleen, 6–8 d after immunization (11, 15). Shortly after the foci appear, clusters of isotype-switched B cells that
stain with the PNA lectin appear in germinal centers (GCs; reference 15) within the areas of the follicles occupied by follicular dendritic cells (16). B cells expressing B cell receptors (BCRs) consisting of the same germline VH and VL segments are present in the foci and GCs (17). However, those within the GCs contain somatic mutations in their BCRs, whereas those within the foci do not (18, 19). Because long-lived memory B cells contain somatic mutations (20), and because somatic mutations are thought to occur primarily in GCs (18, 21), it is thought that all memory B cells are derived from GCs.

These results have led to the current model in which antigen-specific B cells receive signals from CD4 helper T cells at the T cell–area follicle border, proliferate there and either migrate to the outer T cell area near the red pulp, differentiate into plasmablasts, and undergo isotype switching; or migrate to the GCs and undergo somatic mutation and isotype switching there. One limitation of this model is that the aspects related to isotype switching are based on studies in which rare antigen-specific B cells could only be detected after clonal expansion. Thus, critical events in the isotype switching process that occur before antigen-specific B cells undergo extensive proliferation could have been missed. This possibility is addressed here with a novel adoptive transfer system in which isotype switching by initially naive antigen-specific B cells can be monitored in situ before B cell clonal expansion.

Materials and Methods

Mice. All transgenic mice were bred in a pathogen-free facility according to National Institutes of Health guidelines. The DO11.10 TCR transgenic B10.D2Sn/J mice (22) were obtained from Dr. T. Mitchell (National Jewish Medical Center, Denver, CO), 3-83 IgM/D transgenic B10.D2Sn/J mice (23) and 3-83 knockin homozygous (KIH) mice (24, 25) were produced at the National Jewish Medical Center. The OT-II TCR transgenic C57BL/6 mice (26) were provided by Dr. L. Lefrancois (University of Connecticut, Farmington, CT) and crossed at the National Jewish Medical Center. The OT-II TCR transgenic C57BL/6 mice (26) were provided by Dr. L. Lefrancois (University of Connecticut, Farmington, CT) and crossed at the National Jewish Medical Center.

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Immunization. An OVA-mim was produced by inserting three tandem OVA 323–339 epitopes and the 17-amino acid mimotope peptide in frame into the pIII protein of M13, and purified as described previously (29). Purified OVA-mim, containing 20 μg pIII protein, was injected intravenously. A chemical conjugate (OVA–DEL) between OVA (Sigma-Aldrich) and duck egg lysozyme (DEL; Worthington Biochemical Corporation) was prepared with glutaraldehyde and purified as described previously (14). 12.5 μg OVA–DEL was mixed with 25 μg LPS and injected intravenously.

Adoptive Transfer. Lymph node and spleen cell suspensions from DO11.10 or OT-II TCR transgenic donors and 3-83 IgM/D, 3-83 KIH, or anti–HEL KIH transgenic donors were prepared for adoptive transfer as described previously (30). In some experiments, 3-83 IgM/D or 3-83 KIH cell suspensions were labeled with 5 μM carboxyfluorescein diacetate succinimidyl ester (CFSE; Molecular Probes) as described previously (31, 32). Lymph node and/or spleen cells containing a mixture of 2.5 × 10^6 TCR transgenic T cells and 7.5 × 10^5 BCR transgenic B cells were injected into the tail veins of recipient mice. DO11.10 T cells and 3-83 KIH B cells were transferred into B10.D2 recipients. For analysis of early times after immunization (days 2 and 4), OT-II T cells and anti–HEL KIH B cells were transferred into C57BL/6 recipients. For analysis of later times (day 11), OT-II T cells and anti–HEL KIH B cells were transferred into C57BL/6 F1 recipients. Because the anti–HEL KIH mice were on a mixed C57BL/6 and 129 background, use of the F1 recipients prevented rejection of the anti–HEL KIH B cells (which becomes prominent after day 4) due to minor histocompatibility antigen differences.

CD4 T cells were depleted by an intraperitoneal injection of 1 mg anti–CD4 mAb (GK1.5; American Type Culture Collection) 2 d before immunization.

ELISA. Sera were titrated in 96-well plates (Costar) coated with 20 μg/ml S23 mAb specific for 3-83 Ig (unpublished data) and blocked with 1% BSA. Plate-bound Ig was revealed by incubating the wells sequentially with biotinylated anti-IgM or anti–IgG2a (both from BD Biosciences), horseradish peroxidase (HRP)–labeled streptavidin (Sigma-Aldrich), and the chromogen 2,2′-azido-bis(3-ethylbenz-thiazoline-6-sulfonic acid) (Sigma-Aldrich). The titer at half the maximal OD was calculated for each sample.

Flow Cytometry. Exploded spleens were injected with, and minced in, a solution containing 400 U/ml collagenase (Boehringer). After a 15-min incubation at 37°C, 1 ml 100 mM EDTA was added for 5 min, followed by 8 ml 5 mM EDTA in EHAA (Biosource International). The tissue fragments were washed, passed through a fine mesh, washed, and were resuspended in culture supernatant containing 24G2 mAb (American Type Culture Collection) plus 1% mouse serum (Sigma-Aldrich) and 1% rat serum (Sigma-Aldrich) to block Fc receptors.

The resulting cell suspensions were incubated with fluorochrome-labeled antibodies (from BD Biosciences unless otherwise indicated) to detect transferred TCR or BCR transgenic cells. DO11.10 or OT-II T cells were detected with PE-labeled anti–CD4 and FITC-labeled KJ1–26 (Caltag) or CyChrome-labeled anti–CD4, PE-labeled anti–Vα2, and FITC-labeled anti–Thy1.1, respectively. Suspensions containing 3-83 KIH B cells were incubated with FITC-labeled S23 or FITC-labeled anti–IgMα, and antibodies specific for endogenous cells (PE-labeled anti–Thy1.2, Gr–1, anti–IgMγ, and F4/80; Caltag). The cells were fixed with 2% formaldehyde, permeabilized with 0.5% saponin (Sigma-Aldrich), and were incubated with biotinylated anti–IgMγ or IgG2a followed by PerCP-labeled streptavidin. The 3-83 KIH B cells were identified as FITC+, PerCP+, and PE– events.
3-83 KIH B cells were phenotyped by staining with allophycocyanin-labeled anti-B220, FITC-labeled S23, and PE-labeled antisyndecan; or PE-labeled anti-B220 and FITC-labeled GL-7, FITC-labeled anti-CD38, or FITC-labeled anti-I-A\(^d\). The cells were fixed, permeabilized, and stained with PerCP-labeled anti-IgM\(^a\) or IgG2a\(^a\) as described in previous paragraphs.

The anti-HEL KIH B cells were detected with 50 \(\mu\)g/ml HEL (Sigma-Aldrich), allophycocyanin-labeled anti-B220, FITC-labeled GL7 or anti-CD38, and biotinylated HEL (33), followed by PE-labeled streptavidin (Caltag). The cells were fixed, permeabilized, and stained with PerCP-labeled anti-IgM\(^a\) or -IgG2a\(^a\) as described in previous paragraphs.

**Immunofluorescence.** 6-\(\mu\)m cryosections were dehydrated in acetone and blocked with 1% \(\text{H}_2\text{O}_2\), culture supernatant containing 24G2 mAb plus 1% mouse and 1% rat serum, and avidin–biotin–blocking reagents (Vector). When multiple antigens were detected with biotinylated Abs on the same section, the staining was done sequentially, with 1% \(\text{H}_2\text{O}_2\) and avidin–biotin–blocking performed between each antigen. The DO11.10 T cells were detected with digoxigenin-labeled KJ1–26, followed by HRP-conjugated anti-digoxigenin (Boehringer) and TSA-direct Cy3-tyramide (NEN Life Science Products) according to the manufacturer’s directions. OT-II T cells were detected with biotinylated anti-Thy1.1, followed by HRP-conjugated streptavidin (NEN Life Science Products) and TSA-direct Cy3-tyramide (NEN Life Science Products).

The 3-83 and anti-HEL KIH B cells were detected with biotinylated anti-IgM\(^a\) or -IgG2a\(^a\) followed by HRP-conjugated streptavidin and TSA-direct Cy3-tyramide or TSA- indirect biotin–tyramide (NEN Life Science Products) and Cy5-labeled streptavidin (Zymed Laboratories). M13 bacteriophage was detected with biotinylated anti-phage Ab (Sigma-Aldrich) followed by HRP-labeled streptavidin (NEN Life Science Products), TSA- indirect biotin–tyramide (NEN Life Science Products), and Cy5-conjugated streptavidin (Zymed Laboratories). PNA\(^a\) cells were detected with biotinylated PNA (Vector) followed by HRP-labeled streptavidin and TSA-direct Cy3-tyramide.

Endogenous B220\(^+\) B cells were detected with FITC-labeled anti-B220 mAb. Alternatively, endogenous B220\(^+\) B cells were revealed with PE-labeled anti-B220 and IgD\(^+\) cells were revealed with FITC-labeled anti-IgD mAb. Slides were analyzed with a confocal microscope.

**Results**

**Adoptive Transfer Model to Follow B Cell Activation and Isotype Switching In Vivo.** To monitor early in vivo events in the cell biology of class switching, small populations of CD4 T and B cells from Ag receptor transgenic mice were transferred into histocompatible normal recipients, which were immunized with cognate antigen. CD4 T cells specific for chicken OVA peptide 323–339–I–A\(^d\) complexes were obtained from DO11.10 TCR transgenic B10.D2 mice (22), and were identified with the anti-clonotypic mAb, KJ1–26 (34). B cells were obtained from 3-83 KIH mice that express an IgM\(^a\) BCR specific for the H-2K\(^d\) MHC class I molecule (35, 36) or a mimotope peptide identified from a random peptide library (29). The 3-83 KIH B cells were tracked with an anti-idiotypic mAb, S23, and mAbs specific for the IgM\(^a\) or IgG2a\(^a\) allotypic determinants present in 3-83 KIH Ig H chains but not in the Ig H chains of the B10.D2 recipient mice. A bacteriophage containing linked epitopes (OVA-mim) that could be recognized by both the DO11.10 T cells and 3-83 B cells was used as an immunogen. Purified OVA-mim was injected intravenously to mimic a blood-borne antigen, such as a virus, which would be composed of multiple copies of a coat protein containing a T and B cell epitope.

Flow cytometry revealed that the transferred B and T cells were activated after injection of OVA-mimotope bacteriophage. Small populations of DO11.10 T cells and 3-83 KIH IgM B cells, which were not present in untransferred animals (Fig. 1, a and e), were detected in the spleens of recipient mice (Fig. 1, b and f) after transfer. 3-83 KIH IgG2a B cells were not detected in unimmunized recipient mice (Fig. 1 j). Injection of the recipient mice with OVA-mim resulted in a striking increase in the number of DO11.10 T cells and 3-83 KIH IgM B cells in the spleen 4 d later (Fig. 1, c and g), and was accompanied by the appearance of 3-83 KIH IgG2a B cells (Fig. 1 k). The antigen–stimulated populations of 3-83 KIH IgM and IgG2a B cells contained cells expressing either low or high levels of intracellular Ig (Fig. 1, g and k). The cells expressing high levels of IgM (Fig. 2, a–e) or IgG2a (Fig. 2, f–j) were plasmablasts or plasma cells, as evidenced by their large size, low expression of B220, and high expression of syndecan.

It was important to assess the T cell dependence of the 3-83 response to the OVA-mim because viruses and bacteria that display repetitive antigens induce both T cell–independent and –dependent B cell responses (9). In fact, OVA-mimotope bacteriophage can activate 3-83 B cells in the absence of T cells, perhaps owing to their epitope multivalency (37). As shown in Fig. 1, elimination of CD4 T

![Figure 1](https://example.com/f1.png)

**Figure 1.** Detection of transgenic lymphocytes. Normal mice or recipients of DO11.10 T cell and 3-83 KIH B cell mice were left untreated or depleted of CD4 T cells. Spleen suspensions were analyzed by flow cytometry 4 d after immunization with OVA-mim. (a–d) CD4\(^+\), KJ1–26\(^+\), and DO11.10 T cells (circled region); (e and f) surface IgM\(^a\), intracellular IgM\(^a\) 3-83 KIH B cells (boxed region); or (i–l) surface idioype\(^a\) (Id) and intracellular IgG2a\(^a\) 3-83 KIH B cells (boxed) are shown in the dot plots.
cells, including the transferred DO11.10 T cells (Fig. 1 d), greatly diminished the expansion of 3-83 IgM B cells (Fig. 1 h), and completely inhibited the appearance of 3-83 IgG2a B cells (Fig. 1 l) in mice 4 d after injection with the OVA-mimotope bacteriophage, indicating that these responses were T cell–dependent.

The T cell–independent component of the B cell response to OVA-mim was revealed with the more sensitive CFSE dye dilution assay. Naive 3-83 IgM/D B cells did not divide in unimmunized mice (Fig. 3 a), whereas virtually all of the 3-83 IgM/D B cells in recipients that contained DO11.10 T cells divided many times by day 4 after OVA-mim injection, as evidenced by nearly complete loss of CFSE (Fig. 3 b). In addition, a subset of the 3-83 IgM/D B cells divided in CD4 T cell–depleted recipients that were injected with OVA-mim. This proliferation was Ag-dependent and specific because it was not observed in mice 4 d after injection of wild-type bacteriophage (unpublished data) or LPS alone (Fig. 3 d). Thus, OVA-mim induced T cell–dependent clonal expansion and isotype switching to IgG2a, and like many natural antigens, also induced some T cell–independent B cell proliferation.

The Kinetics of Early B Cell Activation and Isotype Switching. The transferred B cells underwent rapid isotype switching. As early as day 2 after injection of OVA-mim, the number of 3-83 IgM B cells in the spleen increased over the starting level, and 3-83 IgG2a B cells containing low levels of intracellular Ig were detected for the first time (Fig. 4 a). The expansion of 3-83 IgM and IgG2a B cells peaked on days 3 and 4 and declined thereafter. The 3-83 IgM<sub>high</sub> plasmablasts, which appeared on day 2 and peaked on day 3, became undetectable by day 6 (Fig. 4 b). The 3-83 IgG2a<sub>high</sub> plasmablasts did not begin to accumulate until day 3, and were undetectable by day 10 (Fig. 4 b). The appearance of these Ig<sub>high</sub> plasmablasts in the spleen correlated with the presence of mimotope-specific IgM and IgG2a in the serum (Fig. 4 c). The persistence of antigen-specific IgG2a antibodies in the serum after the disappearance of the IgG2a plasmablasts was probably related to the long serum half-life of IgG2a (38), or the migration of plasma cells to other sites, such as bone marrow or gut (39).

The majority of the 3-83 IgM B cell expansion and IgG2a switching was dependent on the DO11.10 T cells because 5–10-fold fewer 3-83 IgM and IgG2a B cells were generated in mice that received only 3-83 B cells (Fig. 4, a and b, open symbols). In addition, because 3-83 IgG2a B cells could not be detected in CD4 T cell–depleted mice injected with OVA-mim (Fig. 1 l), the appearance of small numbers of 3-83 IgG2a B cells in mice that received only 3-83 KIH B cells was likely due to the helper activity of endogenous bacteriophage–specific CD4 T cells.

The cell division history of the early isotype-switched B cells was investigated because in vitro studies have shown that isotype switching correlates with cell division (40–42). As expected, the majority of naive 3-83 IgM cells in uninmunized mice were undetectable by day 6 (Fig. 4 b). The 3-83 IgG2a B cells were left untreated or depleted of CD4 T cells. CFSE levels in the transferred B220<sup>+</sup>, S23<sup>+</sup> B cells in spleens were analyzed by flow cytometry 4 d after the indicated intravenous injection.

Figure 3. Proliferation of 3-83 IgM B cells in CD4 T cell–depleted mice. Recipients of DO11.10 T cell and 3-83 IgM/D CFSE-labeled B cell recipients (shaded symbols) or 3-83 KIH B cell recipients alone (open symbols) were immunized with OVA-mim. At the indicated times after immunization, 3-83 KIH IgM<sub>high</sub> (circles) and IgG2a<sup>a</sup> (triangles) B cells in the spleen that expressed low (a) or high (b) levels of intracellular Ig were measured by flow cytometry. The total number of each B cell population was obtained by multiplying the frequency of the 3-83 B cell population by the total number of lymphocytes in the spleen. Each point represents the mean ± SEM from two to four mice. (c) Serum was collected from immunized recipients of DO11.10 T cells at the indicated times and analyzed for 3-83 Ig of the IgM<sub>high</sub> (circles) or IgG2a<sup>a</sup> (triangles) isotype by ELISA. Each point represents an individual mouse. The graph is representative of three independent experiments.

Figure 4. Kinetics of 3-83 B cell clonal expansion. (a and b) DO11.10 T cell and 3-83 KIH B cell recipients were immunized with OVA-mim. Serum was collected from immunized recipients of DO11.10 T cells at the indicated times and analyzed for 3-83 Ig of the IgM<sub>high</sub> (circles) or IgG2a<sup>a</sup> (triangles) isotype by ELISA. Each point represents an individual mouse. The graph is representative of three independent experiments.
munized recipients retained large amounts of CFSE (Fig. 5 a), 48 h after injection of OVA-mim, 98% of the 3-83 IgG2a B cells showed decreased levels of CFSE, indicative of three or more cell divisions (Fig. 5 c). This profile is consistent with the cell division time of 4–8 h reported for B cell centroblasts (43). In contrast, many of the 3-83 IgM B cells present at this time had undergone zero, one, or two divisions (Fig. 5 b). Therefore, 3-83 IgG2a B cells proliferaed to a greater extent than the 3-83 IgM B cells, demonstrating an in vivo correlation between isotype switching and cell division.

**Antigen-specific IgG2a B Cells First Appear in Follicles.** In situ studies were performed to identify the location of isotype switching in the spleen using mAbs specific for the IgM* or IgG2a* allotypic determinants present in 3-83 KIH Ig H chains and a sensitive in situ immunofluorescence detection method (44). In naive recipients, scattered 3-83 IgM B cells and DO11.10 T cells were found in the B cell–rich follicles and periarteriolar lymphoid sheaths (PALS) of each splenic white pulp cord, respectively (Fig. 6 a). 1 d after injection of OVA-mim, the 3-83 B and DO11.10 T cells increased slightly in number, and interactions between some of these cells were observed (Fig. 6, d and inset p) at the border between the follicles and PALS, as expected (14). Consistent with the flow cytometry findings, 3-83 IgG2a B cells were not detected until 2 d after immunization (Fig. 6, b, e, and h, and inset q). These first switched cells were located in B cell follicles rather than PALS foci, as expected based on previous work (11, 15). The majority of the switched B cells were apparently not in contact with DO11.10 T cells. Thus, the earliest detected switched B cells, which had divided at least three times (Fig. 5), appeared to have already migrated away from DO11.10 T cells. Importantly, this T cell–dependent IgG2a switching appeared to occur before detectable GC formation in the follicles.

![Figure 5](image)

**Figure 5.** Proliferation of 3-83 B cells 2 d after immunization. Recipients of DO11.10 T cells and CFSE-dyed 3-83 KIH B cells were not treated or immunized with OVA-mim. Intracellular staining was used to detect 3-83 KIH IgM* and IgG2a* B cells in spleen suspensions prepared 48 h after immunization. The dot plots show the CFSE dye levels on indicated 3-83 KIH B cell populations. The data are representative of three independent experiments.

**Antigen-specific B Cells Form Anatomically Distinct Populations of Plasmablasts, Pre-GC Cells, and Marginal Zone Cells.** Immunohistochemistry identified further heterogeneity of B cell phenotype and location as the immune response progressed. At the peak of clonal expansion on day 4, the 3-83 IgG2a B cells, and to a lesser extent the 3-83 IgM B cells, occupied three locations (Fig. 6, j–l). One population, located in the outer edges of the PALS near the red pulp, contained large amounts of Ig (as evidenced by intense staining with anti-Ig), and, thus, was probably the intracellular Ig^high^ population detected by flow cytometry. These cells were likely the equivalents of the PALS foci described in previous work (11, 15). Many of these plasmablasts expressed surface Ig and high levels of I-A^d^ (Fig. 2, d, e, i, and j) and were in apparent contact with the DO11.10 T cells present in the outer edges of the PALS (Fig. 6, j and k). A second population, which contained a lower amount of Ig, colocalized with antigen (Fig. 6, l and inset s) in the center of B cell follicles (Fig. 6, j, k, and inset r). Although these areas did not stain with the GC marker PNA, they may have been the antecedents of GC because they contained cells expressing CR1/2 (unpublished data) but not IgD (Fig. 6 v). DO11.10 T cells were observed in the pre-GC interacting with the 3-83 B cells (Fig. 6 r). A third population having a lower level of Ig was located in the marginal zone (Fig. 6 v). Therefore, by day 4 of the response, the progeny of the initial follicular B cells had migrated to the red pulp, pre-GC, and marginal zones.

3-83 IgG2a but Not 3-83 IgM B Cells Are Detected in GCs 10 d after Antigen Injection. The presence of 3-83 KIH B cells in pre-GC on day 4 raised the possibility that 3-83 KIH B cells participated in the later GC reaction. This possibility was tested by immunohistochemistry and flow cytometry. By day 10, after injection of OVA-mim, GCs containing PNA^+^ 3-83 IgG2a B cells (Fig. 6, n, o, and u) were detected in the follicles. These B cells showed evidence of many cell divisions (Fig. 7 g) when compared with naive 3-83 IgM B cells (Fig. 7 a), and contained a population of CD38^{low}\text{ and GL-7^+} GC phenotype cells (Fig. 7, h and i). 3-83 IgM B cells that had not switched were also present at this time, but were absent from GCs and resided in the follicles and marginal zones (Fig. 6, m and t) along with some 3-83 IgG2a B cells (Fig. 6 n). These B cells were CD38^{high}\text{ and GL-7^-} (Fig. 7, e and f), and, thus, resembled naive B cells (Fig. 7, b and c). However, they were not naive because they had divided many times (Fig. 7 d), and were likely to be in contact with antigen, which could still be detected in the red pulp and GCs (Fig. 6 o).

One explanation for the absence of Ag-experienced 3-83 IgM cells from GC was that surface IgM does not produce the appropriate signals for B cell retention or survival in GCs (45–48). To test this possibility, recipients of DO11.10 T cells and 3-83 IgM/D transgenic B cells (35), which cannot undergo isotype switching because the 3-83 V DJH/C^\mu^/C^\delta^ gene cassette is not integrated in the H chain locus, were immunized with OVA-mim. As shown in Fig. 7 j, a population of GL-7^+, 3-83 IgM transgenic B
cells was detected 10 d after immunization. The 3-83 IgM transgenic B cells could also be found in PNA+ GCs (unpublished data), demonstrating that 3-83 IgM-expressing B cells can indeed survive in GCs. Thus, although switched B cells normally dominate the GCs, the expression of a switched Ig is not necessary for survival in GCs, although it may provide a competitive advantage.

Anti-HEL KIH B Cells Behave like 3-83 B Cells. To assess if our results would be of predictive value in an independent system, transfer and immunization experiments were repeated using naive CD4 T cells from OT-II TCR transgenic mice, specific for chicken OVA 323–339–I-Ab complexes, naive B cells from anti-HEL KIH mice, and intravenous immunization with an OVA–DEL conjugate. DEL was used instead of HEL because the affinity of the transgenic BCR for DEL is much lower than for HEL ($10^7$ M$^{-1}$ vs. $10^{10}$ M$^{-1}$; reference 49), and, thus, more closely approximates the affinity of a naive BCR for its antigen. LPS was included as an adjuvant to ensure optimal T cell priming (50).

Anti-HEL KIH IgM B cells were found in the follicles and OT-II T cells in the PALS (Fig. 8, a and b) of the spleens of recipient mice after adoptive transfer. As in the 3-83 model, interactions between OT-II T and anti-HEL KIH B cells were observed at the T-B border on day 1 (unpublished data), and anti-HEL KIH IgG2a B cells with low levels of intracellular Ig first appeared in the follicles of the spleen on day 2 after intravenous injection of OVA–DEL plus LPS (Fig. 8, c and d). 4 d after antigen injection, the anti-HEL KIH IgM and IgG2a B cells increased and

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**Figure 6.** In situ localization of 3-83 B cells. Adjacent spleen sections from DO11.10 T cell and 3-83 KIH B cell recipient mice before immunization (a–e), or 1 d (d–f, and inset p), 2 d (g–i, and inset q), 4 d (j–l, close-ups r and s), or 10 d (m–o, insets t and u) after immunization with OVA–mim were stained and analyzed by confocal microscopy. B220 staining (green) identifies the B cell regions in a–u. The DO11.10 T cells appear red in a, b, d, e, g, h, j, k, m, n, p–r, t, and u. 3-83 KIH IgM cells appear blue in a, d, g, j, m, p, and t. 3-83 KIH IgG2a cells appear blue in b, e, h, k, n, q, r, and u. PNA appears red and bacteriophage blue in c, f, i, l, o, and s. In v, 3-83 KIH IgG2a cells appear green, cells that coinexpress B220 and IgD appear purple, and cells that express B220 alone appear red. Arrow points to the marginal zone in v.
formed the same populations of Ig\textsuperscript{high} plasmablasts in the red pulp, Ig\textsuperscript{low} pre-GC cells in the center of follicles, and marginal zone cells (Fig. 8, e and f) as were observed for 3-83 KIH B cells. On day 11 after immunization, the anti-HEL KIH IgM B cells were GL7\textsuperscript{-} and CD38\textsuperscript{high} (Fig. 8, i and j) like naive anti-HEL KIH IgM B cells (Fig. 8, g and h), whereas the majority of the anti-HEL KIH IgG\textsubscript{2a} B cells in the day 11–immunized recipients were GL7\textsuperscript{-} and CD38\textsuperscript{low} (Fig. 8, k and l). Therefore, as in the 3-83 model, anti-HEL KIH B cells first underwent isotype switching in the follicles, formed three anatomically distinct populations, and persisted at later times as GC phenotype IgG cells and non–GC phenotype IgM cells.

**Discussion**

By elevating the frequency of naive antigen–specific B and T cells, we were able to track the genesis and fate of isotype-switched B cells early on during a primary T cell–dependent immune response. It could be argued that elevating the frequency of these cells introduces variables that are not representative of the physiological situation. Although adoptively transferred lymphocytes undergo the activation process more quickly than their endogenous counterparts, evidence suggests that the quality and sequence of the events in this process are similar. For example, endogenous (51) and adoptively transferred naive antigen–specific CD4 T cells (30) undergo the same process of clonal expansion, contraction, and memory cell generation, although these steps appear to occur more rapidly in the transferred population. Similarly, in both the endogenous anti-NP response described by Jacob et al. (15) and in the adoptive transfer model reported here, plasmablasts appeared before GC B cells. However, both events occurred a day or two more quickly in the adoptive transfer situation. This could be related to the elevated frequency of helper T cells because the tempo of the primary anti-DNP B cell response was increased in hosts that contained previously primed cognate T cells (11). Therefore, although the rate of B cell activation may be influenced by the frequency of helper T cells, there is no obvious reason to believe that the order in which events occur during this process would be changed.

The finding that antigen–specific isotype-switched B cells were first detected in the follicles departs from the prevailing view that isotype switching occurs first in foci of plasmablasts in the red pulp and medullary cords (15, 52),
and later in the GCs (53, 54). Our results are consistent with a model in which IgM B cells, after receiving signals from helper T cells at the PALS–follicle border, migrate into follicles where they proliferate and undergo isotype switching. Isotype-switched B cells and unswitched IgM B cells then migrate to the red pulp, pre-GC, and marginal zones. Isotype-switched B cells in follicles may have been missed in previous studies of endogenous antigen-specific B cell populations because the frequency of these cells before clonal expansion is extremely low.

The first switched B cells did not appear to be interacting with antigen-specific T cells in the follicles. Because very few switched B cells appeared in mice that were not transferred with transgenic T cells, it is reasonable to suspect that critical signals for isotype switching were delivered earlier when IgM B cells interacted with antigen-specific T cells at the PALS–follicle border. Previous work has shown that T cell–dependent switching to the IgG2a isotype is defective in mice lacking the CD40 ligand (55) or the IFN-γ receptor (56). It is possible that the antigen-specific B cells received both of these signals from the interacting T cells at the PALS–follicle border and moved away into the follicle, already programmed to switch. This is plausible for the CD40 ligand because this molecule is induced on naive CD4 T cells within hours of TCR signaling (57). It is less clear how naive T cells could have provided IFN-γ so quickly. Naïve DO11.10 and OT-II T cells, like other naïve CD4 T cells, are very inefficient IFN-γ producers unless stimulated to differentiate into Th1 cells (50, 58). Another possibility is that antigen-specific B cells received the CD40 ligand signal from the interaction with the antigen-specific T cells, but switched in response to IFN-γ made by the B cells themselves (29), or from another source; e.g., NK cells (59).

The location of the antigen-specific Ig<sup>high</sup> B cells in the outer PALS and red pulp suggests that they are the counterparts of the NP-specific Ab-secreting PALS foci described by Jacob et al. (15). Consistent with previous findings (60), all of the antigen-specific Ig<sup>high</sup> B cells expressed surface Ig and MHC class II molecules. Thus, these cells did not lose molecules involved in antigen presentation, as has been reported for terminally differentiated plasma cells and myeloma cells (61, 62). Our finding, that the antigen-specific Ig<sup>high</sup> B cells were often juxtaposed with antigen-specific T cells at the border between the PALS and red pulp, is consistent with the idea that they were presenting antigen to the T cells, and in return receiving signals to differentiate further.

The antigen-specific IgM and IgG2a B cells present in the marginal zone after immunization could have been derived from naive follicular B2 cells or marginal zone B cells because both populations are present in the spleens of unimmunized BCR transgenic donor mice (63). We favor the former possibility because 3-83 KIH B cells still accumulated in the marginal zones in immunized recipients of 3-83 KIH lymph node cells, which do not contain marginal zone cells (Fig. 6 v). Activated B2 cells may traffic to the marginal zone because antigen is retained there by marginal zone macrophages or because the B cells acquire expression of integrins that are required for retention in the marginal zone (64).

The antigen-specific IgM, and particularly, the antigen-specific IgG2a B cells present in the follicles on day 4 were located in areas where antigen was concentrated, presumably on follicular dendritic cells (16). The interactions between antigen-specific T cells and antigen-specific B cells that were observed in these regions may be evidence of the process whereby helper T cells provide sustaining signals to B cells in GC (16).

The total number of antigen-specific B cells present in the spleen fell dramatically by day 10 after immunization. The loss was partly explained by the disappearance of antigen-specific Ig<sup>high</sup> plasmablasts, perhaps as a result of apoptosis or migration to the bone marrow (60, 65). Many of the surviving antigen-specific IgG2a B cells expressed GC markers and were located in PNA<sup>+</sup> GCs, whereas the surviving antigen-specific IgM B cells lacked GC markers, and were located outside of GCs. This absence of IgM cells from the GCs was particularly striking because these B cells were 10 times more abundant than antigen-specific IgG2a B cells at this time. Because the antigen-specific IgM cells were capable of entering early GCs, as indicated by their presence on day 4 in the follicular dendritic cell–rich regions, their absence from PNA<sup>+</sup> GCs on day 10 suggests an inability to persist in GC. The lack of persistence of antigen-specific IgM B cells in GCs is inconsistent with a previous paper that found PNA<sup>+</sup> IgM B cells persisting throughout a primary response to sheep red blood cells (66). However, this work followed a polyclonal B cell response, and, thus, it is possible that a steady-state level of newly activated, transiently migrating B cell clones were continuously being recruited into the PNA<sup>+</sup> IgM<sup>+</sup> fraction.

Several explanations are possible for the failure of antigen-experienced IgM B cells to persist in PNA<sup>+</sup> GCs. Subsequent engagement of surface IgG2a by antigen, but not IgM, may produce intracellular signals required for survival or retention in GCs (45–48). This does not appear to be the case because adoptively transferred 3-83 transgenic IgM B cells, and transgenic B cells in other models (67–69), can persist in GCs. Alternatively, the antigen-specific IgM cells that entered the follicular dendritic cell–rich regions may not have persisted because they underwent rapid isotype switching in this location, perhaps due to the abundance of antigen-specific T cells that were also present.

Finally, it is likely that some of the antigen-specific B cells present in the spleen 10 d after immunization were destined to enter the memory pool (70). The antigen-specific IgG2a B cells in the GCs are good candidates because it is thought that memory B cells are derived from the GC reaction. However, the antigen-specific IgM B cells that remained in the follicles and marginal zone are also candidates. The latter cells may correspond to the IgM memory population that survives in the marginal zone of rats for long periods of time after immunization (11, 71). Similarly, IgM B cells with a memory phenotype have been described.
in humans (72–74) and mice (75). Thus, it is possible that two populations of memory cells survive the early primary response: (a) switched B cells that pass through the GCs and (b) unswitched B cells that do not.

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