Stochasticity enables BCR-independent germinal center initiation and antibody affinity maturation

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Two immunoglobulin (Ig) diversification mechanisms collaborate to provide protective humoral immunity. Combinatorial assembly of IgH and IgL V region exons from gene segments generates preimmune Ig repertoires, expressed as B cell receptors (BCRs). Secondary diversification occurs when Ig V regions undergo somatic hypermutation (SHM) and affinity-based selection toward antigen in activated germinal center (GC) B cells. Secondary diversification is thought to only ripen the antigen-binding affinity of Igs that already exist (i.e., cognate Igs) because of chance generation during preimmune Ig diversification. However, whether stochastic activation of noncognate B cells can generate new affinity to antigen in GCs is unclear. Using a mouse model whose knock-in BCR does not functionally engage with immunizing antigen, we found that chronic immunization induced antigen-specific serological responses with diverse SHM-mediated antibody affinity maturation pathways and divergent epitope targeting. Thus, intrinsic GC B cell flexibility allows for somatic, noncognate B cell evolution, permitting de novo antigen recognition and subsequent antibody affinity maturation without initial preimmune BCR engagement.

INTRODUCTION

Adaptive humoral immunity depends on two systems of selection-coupled diversification to provide protection from a vast diversity of pathogenic threats. The first involves combinatorial assembly of IgH and IgL V region exons during B cell development in bone marrow to form the antigen recognition piece of the B cell receptor (BCR), initially expressed as IgM (Jung et al., 2006). The second involves activation-induced somatic hypermutation (SHM) of V exons and IgH class switch recombination by activation-induced cytidine deaminase (AID; Hwang et al., 2015). SHM is coupled to affinity-based selection of BCR toward antigen in germinal centers (GCs). Clones with mutated V exons that encode higher-affinity Ig/BCR competitively secure limiting cognate T cell help, leading to antibody affinity maturation (Victora and Nussenzweig, 2012).

Burnet’s clonal selection theory posits that chance antigen recognition by the preimmune BCR repertoire is required for the initiation and development of antigen-specific antibody responses. Under this conceptual framework, current models of how GC reactions are initiated involve initial B cell activation by antigen engagement of the BCR, followed by interactions of these B cells with antigen-specific T cells, which provide further activation stimuli (Victora and Nussenzweig, 2012; De Silva and Klein, 2015).

The degree of antigen recognition by BCR that is required at this initial stage is not fully understood. Low-affinity BCRs can seed robust GC reactions in the absence of competition from higher-affinity clones (Dal Porto et al., 2002; Shih et al., 2002; Schwickert et al., 2011), suggesting that competition between B cells may play a larger role than the absolute value of BCR affinity to antigen. In addition, antibodies cloned from activated B cells in GCs do not always bind to immunizing antigen (Di Niro et al., 2015; Kuraoka et al., 2016; Tas et al., 2016). Those studies relied on assays measuring antigen binding to secreted antibodies, which is less sensitive than testing reactivity to membrane-bound Ig/BCRs (Lingwood et al., 2012). However, they raise the possibility that B cells with very low-affinity—or potentially, noncognate—B cells may be activated and allowed to enter into the GC reaction, nonspecifically, to receive activating T cell signals. Processes allowing potentially nonspecific B cells to participate in GC reactions may be caused by poorly understood parameters possibly unrelated to BCR engagement, recently described as stochastic noise (Mesin et al., 2016).

Such noise mechanisms may have physiological relevance. In this regard, some high-affinity antibodies may have evolved from BCRs that may have had no initial recognition of antigen, as may be the case with the VRC01 class of anti–HIV-1 broadly neutralizing antibodies (Zhou et al., 2010; Scheid et al., 2011; Wu et al., 2011; Hoot et al., 2013). In addition, in vitro analysis of endogenously mutating B cell lines has uncovered a surprising diversity from SHM alone (Cumbers et al., 2002). However, whether nonspecific B cell activation and SHM, supported by stochastic noise, can generate de novo antigen recognition in GCs is unclear. In addition, whether B cells activated in this way could support develop-

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ment of high-affinity antibodies is not well defined. The swift Darwinian nature of the GC SHM/selection process theoretically could enable high-affinity antibodies to be generated from any starting point regardless of initial preimmune BCR recognition. If so, this would reveal a thus-far-undefined flexibility of the GC system.

Here we use a strict monoclonal system in which BCR lacks the ability to physically and functionally engage with OVA in the setting of OVA-specific T cells to explore BCR recognition requirements for B cell entry into the secondary/GC diversification program and to uncover possible outcomes of B cell maturation that may have had access only to evolutionary mechanisms of stochastic noise initially upon GC entry.

RESULTS AND DISCUSSION

To examine the degree to which noncognate antigen can influence GC B cell development and antibody evolution, we used a model system in which T cells, but not B cells, recognize immunizing antigen. T/B monoclonal mice (which we refer to as HRO mice) have hemagglutinin (HA)-specific monoclonal B cells (via knock-in IgH and IgL V regions) and T cells with transgenic TCR specific for an OVA peptide (DO11.10) in the setting of Rag-1 deficiency to ensure monoclonal expression of both BCR and TCR (Curotto de Lafaille et al., 2001).

We cloned and expressed the mAb from HRO mice and measured its affinity against OVA by surface plasmon resonance (SPR). As shown in Fig. 1 A, no binding was found, even at 10 mM OVA (the solubility limit of OVA), indicating that the association constant (K_a) of the mAb and OVA is less than 10^5 M^-1 (Fig. 1 A, left). In contrast, the mAb binds to an HA-OVA protein conjugate at nanomolar level, as expected (Fig. 1 A, right). Because membrane-bound Ig in the form of a BCR, may be able to bind to antigen better than soluble Ig (Lingwood et al., 2012), we examined the degree to which OVA—in either monomeric form or multimerized—could functionally engage HRO BCR on the surface of B cells. Both anti–mouse Igκ antibody and HA-OVA conjugate could trigger Ca^{2+} flux in HRO B cells (Fig. 1 B), indicating ability to engage BCR, as expected. However, neither monomeric nor multimerized OVA could induce a Ca^{2+} flux, even at high dose (500 µg/ml; Fig. 1 B).

In addition to Ca^{2+} flux, BCR engagement is known to induce BCR internalization. Consistent with the Ca^{2+} data, we found that both anti–mouse Igκ antibody and HA-conjugated protein could trigger HRO B cell BCR internalization, as indicated by reduced surface IgM, whereas monomeric or multimerized OVA could not (Fig. 1 C). OVA concentrations of up to 50 µg/ml were unable to induce OVA-specific T cell proliferation in co-culture assays with HRO B cells and DO11.10 T cells (Fig. S1 A). However, consistent with previous reports of B cell ability to take up and present antigen nonspecifically at high concentrations (Rock et al., 1984), OVA concentrations of 500 µg/ml were able to stimulate B cell–dependent T cell proliferation (Fig. S1 A). Collectively, these data indicate that OVA is essentially a noncognate antigen for HRO BCR, incapable of binding soluble antibody and unable to engage membrane-associated BCR, but capable of being processed in MHCII-containing processing compartments through what is likely a BCR–independent pathway.

To examine the degree to which OVA immunization can induce GC formation in the HRO system, we immunized HRO mice with OVA intraperitoneally and stained for GC B cell markers peanut agglutinin (PNA) and GL7 in CD19+ splenic, mesenteric lymph node (mLN), and Peyer’s patch (PP) B cells at 4, 8, and 12 d. At day 4, we found that some mice had clear induction of GCs in splenic and mLN B cells with OVA/alum immunization compared with alum alone (control; Fig. 1, D and E). However, GC B cells in PP were largely at background levels at baseline and were not induced with OVA immunization (Fig. 1, D and E). Because T cell activation is required for GC induction, the latter finding is likely caused by the absence of T cell antigen in the intestinal luminal compartment, sampled by PP.

We sorted single GC B cells from HRO mice 8 and 12 d after OVA immunization and found a substantial number of sequences with a mutation frequency of 1–4 in both IgH and Igk V regions (Fig. 1 F). Mutation analysis showed sequence changes clustering in complementary-determining regions (CDRs; Fig. 1, G and H), which are known to harbor intrinsic AID-mediated mutability (Yeap et al., 2015). These results suggest that, if given sufficient T cell help, B cells may harbor a permissiveness to participate in the GC fate and can initiate AID-mediated Ig diversification in the absence of any evidence of BCR engagement or measurable affinity for antigen.

To determine whether nonspecific B cells can enter GCs in the context of a competitive B cell environment, we injected HA-reactive HRO B cells into wild-type mice and immunized them with OVA the next day (Fig. S1 B). On day 6 after immunization, we observed that 3% of naive (IgD^− PNA^− ) splenic B cells were HA reactive (Fig. S1 C). In this setting, ~0.1% of GC (IgD^− PNA^− ) B cells were HA positive, and of these, one quarter to one third of them identified with the HRO sequence, with most mutations within AID target motifs (Fig. S1, D and E). We designed a TaqMan quantitative PCR (qPCR) assay with primers and probe that leverage the unique IgH CDR3 of HRO sequence. Analysis of activated GC B cells with this assay after OVA immunization indicated that ~1/1,000 (0.008 ± 0.007) are HRO B cells (Fig. S1 F). This represents the lower limit, as other HRO B cells may have mutated away from recognition in these assays. In addition, it is very likely that much more than 3% of B cells are nonspecific for any particular antigen. These experiments suggest that noncognate B cells can enter GCs and diversify through SHM in a competitive setting.

To test whether chronic stimulation with OVA can induce HRO antibody maturation toward de novo OVA reactivity, we serially immunized HRO mice with OVA in
alum adjuvant every 2–3 wk for several months and tracked OVA reactivity in plasma. All immunized HRO mice generated anti-OVA antibody responses (Fig. 2 A). Responses in HRO mice were delayed and lacked consistency in the timing of their appearance (Fig. 2 A). However, this delay and inconsistency may be explained by a monoclonal T cell system (Leung et al., 2013; Preite et al., 2015), as Tcr transgenic OT-II mice have a similar serological response upon OVA immunization (Fig. 2 A).

Flow cytometric analysis of splenic B cells of chronically immunized mice identified CD19 + B220lo OVA+ B-lineage cells (Fig. 2 B). We sorted single OVA+ B cells from three
mice (referred to as 1, 2, and 3) and sequenced the variable regions of heavy and light chain (VH and VL) from individual B cells. We also cloned and expressed seven representative mAbs (1A3, 1A4, 2B11, 2C1, 3A6, 3D1, and 3D7) from the sorted OVA+ B-lineage cells and measured their binding affinities to OVA by SPR. Each of the seven antibodies cloned and tested showed clear reactivity, with an affinity constant (Ka) in the range of 10^7–10^8 M^-1 (Fig. 2 C).

We analyzed VH and VK mutation patterns from single OVA+ HRO B cells from mouse 1, 2, and 3. We found some amino acid changes that were shared between mice, such as the N53Y, G55V, and Y59F changes in mouse 1 and 2, seen in >50% of the VH sequences. Although the N53Y and Y59F changes were seen in approximately half of the OVA+ VH sequences from mouse 3, other enriched changes were not common among the other mice (Fig. 3 A), such as S84N, S30F, and S31Y (Fig. 3 A and Fig. S2 A). Antibody clone 3D7, from mouse 3, which has high-affinity OVA binding (Fig. 2 B), has neither the G55V or Y59F changes, and clone 3A6, also confirmed to be a high-affinity OVA binder, has the Y59F but not the G55V or N53Y changes (Fig. 3 A and Fig. S2 A). Enriched amino acid changes were also seen in VK sequences but were not shared between the different mice (Fig. 3 A and Fig. S2 A). These data suggest that, although there are some common shared changes, a diversity of antibody sequences can arrive at OVA reactivity from the same nonreactive starting point through the secondary Ig SHM and selection system. Consistent with this, analysis of the clonal relationships between VH and VK sequences from OVA+ single cells showed a multiclad organization that was concordant between VH and VK pairs derived from the same cell, as shown by colored boxes linking the VH and VK clades in Fig. 3 B.
In general, sequences derived from one mouse occupied a single clade, with the exception of one, boxed in orange in Fig. 3 B, which is shared by clones from mouse 1 and 3. The clones tested and shown to have high-affinity OVA reactivity by SPR described in Fig. 2 C are indicated by stars and are represented in each clade in Fig. 3 B.

We sorted OVA\(^+\) cells from six different mice chronically immunized with OVA and subjected them to deep sequencing. We found that mutations accumulate at similar \(V_H\) and \(V_L\) locations, largely at DGYW (\(D = A, G, \text{ or } T; Y = C \text{ or } T; W = A \text{ or } T\)) sequence motifs, which is a favored AID target (Rogozin and Díaz, 2004; Fig. 3 C). Consistent with the single-cell clade analysis, phylogenetic analysis from the deep-sequencing data showed that sequences from individual mice are largely clustered together, suggesting that mutation pathways to OVA reactivity largely differ between mice (Fig. 3 D). In addition, IgM, IgG, and IgE isotypes are represented in the OVA-reactive pool (Fig. 3 E). We also did deep sequencing on the bulk population of B cells from unimmunized mice to determine the background HRO BCR mutation frequency. We found very low levels of background mutation in the range of sequencing error rate (Schirmer et al., 2016; Fig. S2 B), indicating that random mutations in naive mice favorable for OVA binding are extremely unlikely to occur at baseline. However, we cannot fully rule out this possibility.

To determine the epitope diversity of anti-OVA antibody response in OVA-immunized HRO mice, we constructed a random mutagenesis library of OVA (one to five random amino acid mutations per sequence) and displayed the library on the surface of yeast. We first screened the library for mutants failing to bind to OBI mAb, whose epitope is already known (Dougan et al., 2012; Fig. 4, A and B). We also screened yeast clones that failed binding and sequenced them to identify putative epitopes. We defined the positions with at least three mutations as epitope candidates and checked their locations on the structure of OVA. We considered that epitope candidates that are not clustered on the 3D OVA surface, or that are not exposed on the surface, as potential background signals. Among all eight candidate residues of 1A3, seven are exposed and clustered on the surface of OVA (the exception is H329), suggesting that 1A3 recognizes a conformational epitope consisting of H23, N25, R200, E203, Q204, E205, and S385 (Fig. 4, B and C). For 2C1, all the candidate residues are exposed except I158, which is partially covered. In addition, the mutations are enriched on positions E203, Q204, and E205, indicating that these are the core epitopes of 2C1. Although 3D1 has more candidate residues than 1A3 and 2C1, only eight residues (N27, V201, T202, E203, Q204, E205, E257, and S385) remain after the nonexposed and nonclustered residues are excluded (Fig. 4, B and C). Compared with the other three mAbs, 3D7 has the most candidate residues, but only nine residues (F181, G183, L184, F189, D191, T194, E337, G339, and E341) are exposed and clustered together; the others are either buried or scattered on the surface (Fig. 4, B and C). Comparing these epitopes, we found that 1A3, 2C1, and 3D1 bind to a similar epitope, whereas 3D7, which was isolated from the same mouse as 3D1, targets a different epitope, demonstrating that the SHM/GC selection system permits multiple epitope targeting, despite initiating from a homogeneous sequence in the same animal.

We also serially immunized HRO mice with OVA conjugates, namely, OVA-fluorescein and OVA-TNP, simultaneously. Anti-fluorescein and anti-TNP responses could also be detected in chronically immunized HRO mice (Fig. 4, D and E). These results indicate that serological responses are not unique to OVA and are consistent with the findings that multiple haptens can be targeted.

By juxtaposing monoclonal noncognate B cells with T cells specific for immunizing antigen, our experiments were designed to identify how flexible the GC B cell system can be to respond to activating cues without regard to deterministic BCR signals. The current paradigm is that the secondary diversification/selection system depends on chance recognition of antigen by the preimmune Ig repertoire to engage GC processes that nurture affinities already present. The findings described here suggest that the GC system may be more flexible than previously anticipated.

It is possible that in vivo BCR engagement may somehow occur in a fashion not observed here experimentally, such as the association of OVA with other particles that can occur in a fashion not observed here experimentally, even at saturating concentrations of OVA. Second, BCR internalization and Ca\(^{2+}\) signals were not observed when treated with high-dose and multimerized OVA in context of naturally expressed, membrane BCR. Third, despite the monoclonal starting point, multiple distinct epitopes can be targeted, on both OVA itself and anti-hapten responses from OVA-hapten immunization. If OVA were able to trigger the HRO HA-specific BCR in vivo, affinity maturation around a single OVA-epitope would be expected, and responses to two different haptens, randomly chosen, would not be expected. Together, these findings strongly suggest that recognition can be generated during the SHM process within B cells nonspecifically activated by mechanisms unrelated to BCR engagement with antigen.

Mechanistically, for B cells to acquire OVA-specific T cell help, OVA would require access into B cells. Although BCR is the most efficient way to deliver antigen through the cell to surface MHCII (Kakiuchi et al., 1983; Rock et
Figure 3. **Sequence analysis of OVA⁺ HRO B cells.** (A) Amino acid mutation frequency in HRO V₃ and Vκ of OVA⁺ single cells from three HRO mice (#1, #2, #3) isolated 268 d after the first immunization. The number of sequences represented in each plot is indicated. (B) Maximum-likelihood phylogenetic trees for paired single-cell V₃ (left) and Vκ (right) sequences from three HRO mice. The different color markers represent sequences from three different mice (#1 red, #2 blue, and #3 green). The blocks mark different clades in V₃ (left) and Vκ (right). Dashed lines link the pairing clades. Black stars indicate
B cells can take up and process antigen through MHC-containing processing compartments to influence T cell interactions in a BCR-independent manner (Rock et al., 1984; Jellison et al., 2007; Dwyer et al., 2014). Low-level OVA, nonspecifically taken in and presented by HRO B cells, may be sufficient for OVA-specific T cells to provide B cell activation signals—inducing Ig diversity through induction of AID and SHM. B cell selection in this setting would initially be driven by BCR-independent stochastic factors. These factors could include (but are not limited to) parameters such as proximity to limiting T cells and differences of OVA uptake caused by uneven concentration gradients. Other pathways of B cell diversity unrelated to BCR affinity include asymmetric GC B cell division (Barnett et al., 2012; Thaunat et al., 2012) as well as other less well-defined intracellular stochastic factors intrinsic to B cells that influence behavior (Barnett et al., 2012). SHM, together with stochastic selection, can continue until BCR diversification is sufficient to generate de novo antigen recognition by the BCR within the GC itself. Because BCR involvement can increase the efficiency of B cell antigen presentation by at least ~1,000 fold (Kakiuchi et al., 1983; Rock et al., 1984), selection based on affinity maturation would dominate after acquisition of initial BCR recognition.

Although it remains to be determined whether conversion of noncognate B cells to antigen reactivity occurs in real physiological conditions, the BCR-independent ability of B cell antigen uptake and presentation may be physiologically relevant. In this regard, it may provide a safety net function to cover potential deficiencies (i.e., holes) in the primary Ig repertoire to immunologically important epitopes, although perhaps at the cost of timeliness and consistency because of the rare involvement of nonspecific B cells in the setting of a competitive environment (Fig. S1, B–F). This may have relevance to antibody responses that appear to occur despite observations that unmutated Ig ancestors fail to bind native antigen, as is the case with the VRC01 class of broadly neutralizing antibodies (Zhou et al., 2010; Scheid et al., 2011; Wu et al., 2011; Hoot et al., 2013), which appear inconsistently and after unusual delays (Mascola and Haynes, 2013). In addition, stochastic noise has also been observed in B cell responses despite the presence of BCRs specific to immunizing antigen. Visualization of a system designed to fluorescently label B cells of identical affinity to antigen with diverse colors revealed that GC B cells were able to deviate substantially from baseline color distributions (Tas et al., 2016).

Our data may also suggest functions of a diverse primary Ig repertoire beyond providing anticipatory antigen recognition. A diverse Ig repertoire also furnishes a diversity of Ig substrates. In this setting, despite BCR nonreactivity to begin with, SHM in diverse—but nonspecifically activated—B cells with diverse IgV region sequence substrates would be expected to generate de novo antigen recognition more efficiently. Because this system may be used to target epitopes not otherwise targeted in the setting of natural preimmune BCR repertoires, our findings have implications for antibody bioengineering and vaccinology. We conclude that the GC somatic evolution system is sufficiently flexible such that preimmune BCR engagement is not required to generate de novo B cell–mediated antigen recognition and subsequent Ig affinity maturation.

**MATERIALS AND METHODS**

**Mice**

All mice were housed at the Karp Research Building at Children’s Hospital Boston. The T/B mice, referred to as HRO, maintained on a BALB/c background, were described previously (Curotto de Lafaille et al., 2001) and were provided by J. Lafaille (Skirball Institute of Biomolecular Medicine, New York, NY). Some HRO mice were interbred with BALB/c Rag 1 knockout mice, mouse strain 003145 (The Jackson Laboratory), to produce HRO mice possessing one HA-specific allele and another unassembled allele. Rag-deficient monoallelic and biallelic anti-HA BCR mice were used in the experiments. Wild-type BALB/c controls were obtained from the Jackson Laboratory were bred and maintained in a specific pathogen–free mouse facility in the Karp Research Building. OT-II mice (B6.Cg-Tg(TcraTcrob)425CbiLyS/J) and C57BL/6J controls were obtained from the Jackson Laboratory. The Institutional Animal Care and Use Committee for Children’s Hospital Boston approved all mouse experiments for this study.

**Immunizations**

Mice 4–8 wk of age were immunized in cohorts up to 260 d, with recurrent immunization intervals 2–3 wk apart. Mice underwent intraperitoneal immunization with either a single immunogen, OVA (Sigma-Aldrich), or simultaneous administration of trinitrophenyl (TNP) (11)-OVA conjugate (Biosearch Technologies) plus fluorescein (0.8)-OVA (Biosearch Technologies). All immunogens were combined with Inj ect Alum (Thermo Fisher Scientific) in a 1:1 ratio and agitated.

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the clones tested by SPR. **(C)** Histograms showing stacked mutation frequencies at each position in VH and Vk as calculated from deep sequencing data of OVA B cells from six HRO mice (6TR, 6TLTR, 6TL, 5TL, 5BR, and 2BR). The sequences from each mouse are represented with different colors, with the number of sequences mentioned in brackets. The CDRs, DGYW AID target motifs, and AGCT motifs, which serve as an optimal version of DGYW for AID targeting, are shown. Only productive sequences are shown. Replacement and silent mutations are indicated. **(D and E)** Dendrograms of unique HRO VH and Vk from deep-sequencing data created by maximum-likelihood method. n, number of sequences. Trees are rooted at the unmutated common ancestors (indicated by black boxes). Colors represent different mice (D) or IgH isotype information (E), n = 5 (VH) and 6 (Vk). For clarity, sequences with more than 13 mutations are shown for Vk.
Figure 4. HRO B cells can evolve toward multiple epitopes and haptens. (A) FACS plots showing the screening of a yeast library with five anti-OVA mAbs. Yeast displaying unmutated OVA (left) or a library of OVA mutants (mut. lib., right) were stained with OBI, 1A3, 2C1, 3D1, and 3D7 together with an anti-c-myc antibody. The OBI mAb was used as a control, as it has a described epitope (Dougan et al., 2012). c-myc is a tag used to indicate the expression of OVA or OVA mutant proteins. Yeast displaying OVA mutants that fail to bind with mAb (c-myc single-positive population) were gated and sorted. Dozens of yeast clones sorted by each mAb were sequenced (n = 19–96). (B) Epitope analysis of anti-OVA mAbs. Sequences of sorted mutants were aligned with wild-type OVA, and the mutations on each amino acid position were calculated and plotted in the histograms shown. Positions with three or more mutations were defined as epitope candidates and labeled beside the corresponding column in different colors: black, buried residues; green, exposed but spatially scattered residues; magenta, epitope of OBI; red, epitope of 1A3, 2C1, and 3D1; blue, epitope of 3D7. The red dashed line indicates three times background mutation levels, which we used as a cutoff for candidate residues representing putative epitopes. The number of sequenced mutants for each mAb is noted in each histogram. (C) Spatial epitope mapping of anti-OVA mAbs. The residues in B are highlighted on the 3D structure model of OVA with the same color. (D and E) ELISA data showing anti-TNP (D) and anti-fluorescein (E) IgG1 from plasma collected from wild-type (green, n = 3) and HRO (red) mice serially immunized with OVA-TNP (n = 5) and OVA-fluorescein (n = 5) conjugate proteins, or PBS (blue, n = 4). Each line represents an individual mouse.
by vortex at room temperature for 30 min before injection. Controls for immunization included PBS 1× combined with Inj ect Alum in a 1:1 ratio. For administration of OVA, doses of 100 µg were used. For administration of OVA with either TNP (OVA–TNP) or fluorescein (OVA–fluorescein), doses of 30–45 µg of each conjugate were used.

Mouse plasma

Mouse blood was collected by facial vein bleeding at the time of immunization. Approximately 50–250 µl of whole blood was collected for each mouse per facial vein bleed in microtainer K3EDTA tubes (BD). The samples were centrifuged in a benchtop microcentrifuge (Eppendorf Model 5424 or 5427R) at 400 g for 6 min. Approximately 10–50 µl of plasma was collected for each mouse per facial vein bleed in microtainer K3EDTA tubes (BD). The samples were centrifuged (4°C), plates were washed six times with wash buffer before ELISA plates (C96 NUNC MaxiSorp; Thermo Fisher Scientific) were coated with OVA (50 µg/ml; Sigma–Aldrich), fluorescein (5)-BSA (10 µg/ml; Biosearch Technologies), or TNP (11)-BSA (10 µg/ml; Biosearch Technologies) diluted in bicarbonate coating buffer (3.03 g Na2CO3, 6.0 g NaHCO3, and 1× PBS, with a final pH of 9.6) and incubated overnight at 4°C. The next day, plates were washed with wash buffer (1× PBS with 0.05% Tween-20; Sigma–Aldrich) and blocked with 5% BSA (Sigma–Aldrich) in wash buffer for 1.5 h at 37°C. Blocking solution was removed by rinsing with wash buffer three times. Standards were applied in dilutions (1:2) from the initial starting concentrations as follows: anti–OVA IgG1 standard (Purified Anti-Chicken OVA; BioLegend) using 5 µg/ml, anti–fluorescein standard (IgG Fraction Monoclonal Mouse anti–fluorescein; Jackson ImmunoResearch) at 10 µg/ml, and anti–TNP standard (Purified Mouse IgG1 λ Isotype Control; BD Biosciences) at 10 µg/ml. Plasma was added at 1:50 to 1:500 dilutions in duplicate, followed by serial dilution of each sample (1:10) in three subsequent dilution wells for each sample. After incubation at room temperature for 2 h (or storage overnight at 4°C), plates were washed six times with wash buffer before being coated with their respective secondary antibodies as follows: 10 µg/ml anti–OVA IgG1 (AKP Rat Anti-Mouse IgG1 Clone: X5C; BD PharMingen), anti–fluorescein IgG1, and αTNP IgG1 ELISAs. Secondary antibodies were incubated for 1 h at room temperature. Plates were washed with wash buffer eight times, and 100 µl phosphate substrate buffer (2 × 10 mg substrate tablets; S0942; Sigma–Aldrich) diluted into 10 ml ELISA development solution (1 M glycine, 10 mM ZnCl2, and 10 mM MgCl2 in 1× PBS) was added to each well. Blank wells, coated with the target antigens and incubated with secondary antibodies without plasma or standards, were included for each plate in duplicate. Sample and standard dilutions were also performed in duplicate for each plate. Plate readings were taken at 5- to 10-min intervals using an ELISA Plate Reader set to 405 nm (tunable Versa Max microplate reader; Molecular Devices). SoftMax Pro software v.5.4.1, Microsoft Excel, and GraphPad Prism software were used to analyze plate data.

B cell staining and sorting

For GC B cell staining, total splenocytes from OVA-immunized HRO mice and control mice were stained with DAPI, PNA–FITC, anti–GL–7–PerCP–Cy5-5, anti–IgD–PE, anti–CD19–APC–Cy7, anti–IgM–PE–Cy7, anti–IgG1–APC, anti–CD4–BV510, anti–CD8–BV510, anti–CD11c–BV510, anti–F4/80–BV510, and anti–Gr1–BV510. GC B cells with the phenotype DAP1−, CD19+, Dumpt−(CD4, CD8, CD11c, F4/80, and Gr1), PNA−, GL−7−, and IgD− were gated and single-cell sorted into 96-well plates containing lysis buffer (0.5× PBS, 10 nM dithiothreitol, and 4 U RNaseOUT) on a BD FACSAria sorter and immediately stored at −80°C. For OVA-specific B cell staining, total splenocytes from OVA-immunized HRO mice and control mice were stained with DAPI, anti–CD19–APC–Cy7, anti–B220–PerCP–Cy5-5, OVA–Alexa Fluor 647, anti–IgM–PE–Cy7, anti–IgG1–APC, anti–CD4–BV510, anti–CD8–BV510, anti–CD11c–BV510, anti–F4/80–BV510, and anti–Gr1–BV510. OVA-specific B cells were selected for the phenotype DAP1−, CD19+, Dumpt−(CD4, CD8, CD11c, F4/80, and Gr1), B220−, and OVA−. OVA-specific B cells were single-cell sorted into 96-well plates or bulk sorted into TRlizol.

Single B cell RT-PCR, antibody cloning, and expression

Single B cell RT-PCR, antibody cloning, and expression were performed as previously described (Tiller et al., 2009). In brief, reverse transcription was performed with SuperScript III (18-080-044; Thermo Fisher Scientific) and random hexamer primer. Two rounds of PCR were performed to amplify the VH and VK regions. PCR products were performed using HotStar Taq DNA polymerase (Qiagen 203205) with the following primers: 5′-MsVHE-HRO, 5′-GGG AAT TCG AAT GGT GAG TCT GGT GAG ATC TTG G-3′; Cμ outer, 5′-AGGGGCTCTCCTGAGGAGACGAGG-3′; Cγ1 outer, 5′-GGAGAGTGTCACACCCTGGAGA-3′; and Cγ2 outer, 5′-GAGAGTGTCACGCCACCCCTGGAGA-3′. The second round of PCR reactions were performed with the following primers: MsVHE-HRO, GGGAATTCGAGGTGAGTCTGGAGAGATCCTGG-3′; Cμ outer, 5′-AGGGGCTCTCCTGAGGAGACGAGG-3′; Cγ1 outer, 5′-GGAGAGTGTCACACCCTGGAGA-3′; and Cγ2 outer, 5′-GGAGAGTGTCACGCCACCCCTGGAGA-3′. PCR products were then sequenced by Sanger sequencing. To express the mAbs from single B cells, the selected sequences were cloned into a human IgG1- or human Igκ-mammalian expression vector for antibody expression. Then heavy chain vector and light chain vector were cotransfected into 293T cells with polylethyleneimine, and supernatants were harvested after 48 h. The antibody expression vectors were provided by G.D. Victora (The Rockefeller University, New York, NY).
BCR internalization
HRO splenocytes were resuspended in RPMI 1640 cell culture medium and seeded in a 96-well plate (2 × 10⁵ cells/well). Anti-Igκ antibody, OVA conjugated with HA, OVA monomer, and OVA multimer were added, and the cells were incubated in CO₂ incubator for 4 h. Cells were harvested and stained with anti-CD19-APC-Cy7, anti-IgM-PE-Cy7, anti-CD4-APC, and DAPI. The cells were analyzed on a Canto II (BD), and FACS data were analyzed in FlowJo software (Tree Star). IgM fluorescence intensity shift was used to evaluate BCR internalization.

Calcium flux assay
Total HRO splenocytes at 10⁷ cells/ml were loaded with 5 μM Indo-1 acetoxymethyl (I-1203; Thermo Fisher Scientific) in HBSS with 2% FCS. DAPI, anti-CD19-APC-Cy7, and anti-CD22-PE were added for the last 15 min of incubation. Cells were washed twice and left at room temperature for 15 min in HBSS with 2% FCS. Cells were then equilibrated at 37°C, and signal at Indo blue and Indo violet of gated B cells was collected for 1 min to record baseline Ca²⁺ levels as the ratio of Indo violet/blue. Cells were then stimulated with anti-Igκ antibody, OVA conjugated with HA, OVA monomer, and OVA multimer, and Ca²⁺ levels were recorded for an additional 3 min. Ionomycin-stimulated cells were used as a positive control. Data were collected in a FACs LSR II Fortessa (BD) and analyzed using FlowJo.

Library preparation and sequencing
For Sanger sequencing, total RNA was extracted using the TRIzol method (Invitrogen), followed by cDNA first chain synthesis using Superscript III (Invitrogen) with random hexamers (Random Primer 6; New England Biolabs). Amplification for heavy (IgH) and light (IgL) chains was conducted in separate reactions. For heavy chains, all isotopes were run in one reaction. First-round PCR reactions included 2 μl cDNA, 8 μl of 5× High Fidelity (HF) buffer, dNTP mixture (final concentration 250 μM each), forward and reverse primers (final concentration 250 nM each, primer sequences follow), 0.5 μl Phusion DNA polymerase (Thermo Fisher Scientific), adjusted to a final volume with RNase- and DNase-free water into 40 μl. Second-round PCR primers were previously described (Tiller et al., 2009). We used the following first-round PCR program: 98°C for 30 s, then 18 cycles of 98°C for 10 s, 65°C for 30 s, 72°C for 30 s, and 72°C for 5 min. Second-round PCR reactions included 5 μl cDNA, 8 μl of 5× HF buffer, dNTP mixture (final concentration 250 μM each), forward and reverse primers (final concentration 250 nM each, primer sequences follow), and 0.5 μl Phusion DNA polymerase, followed by volume adjustment with RNase- and DNase-free water into 40 μl. Second-round PCR primers for IgH (Igμ, Igγ1, Igγ2b, Igγ2c, Igγ3) were MgHVE, 5′-GGGAATTCGAGGTGCAGCTGAGGTGATTGG-3′; Cμ inner, 5′-GGGAGAGAACCATTTTGGGAAGGAC-3′; Cγ1 inner, 5′-GCTCAGGGAAATAGGCCTTGAC-3′; Cγ2c inner, 5′-GCTCAGGGAAATACCCCTTGAC-3′; Cγ2b inner, 5′-AGTCAGGGAAATGACCCCTTGAC-3′; and Cγ3 inner, 5′-GCTCAGGGAGAATGCCCCCTTGAC-3′. Second-round PCR Primers for IgL (IgLk) were mVκ, 5′-GAYTATTGMTCGACMRWCMTCA-3′, and Biwi P-mJK04, 5′-GCCACCCCTACGTCCAGCTCGCTTCGTC-3′. The second-round PCR program was as follows: 98°C for 30 s, then 18 cycles of 98°C for 10 s, 65°C for 30 s, 72°C for 30 s, and 72°C for 5 min. Second-round PCR products were purified by gel purification kit (QiAquick Gel Extraction kit; Qiagen), cloned into pGEM-T vector (Promega), and sequenced by Sanger method (Eton Bioscience). For deep sequencing, total RNA was extracted using the TRIzol method (Invitrogen), followed by cDNA first-chain synthesis using SMAMRTScribe reverse transcription (Clontech) and a mixture of IgH (Igμ, Igγ1, Igγ2b, Igγ2c, Igγ3, Igκ) and IgL (IgLk) constant region specific primers (IDT) in which TdT is replaced by UdR as follows: Cmu outer, 5′-GGGAGGCGUCUCGCAGGAGACAGGAG-3′; Cμ1 outer, 5′-GGAAGUGUGAGCACCGCGACCACAGGAG-3′; Cγ2 outer, 5′-GGAAGGCTGGTCACCGCGACCACAGGAG-3′; Cγ2c outer, 5′-GGGAGGCTGGTCACCGCGACCACAGGAG-3′; Cκ outer, 5′-GGGAGGCTGGTCACCGCGACCACAGGAG-3′; Cκ1 outer, 5′-GGAAGGCTGGTCACCGCGACCACAGGAG-3′; and Cκ3 outer, 5′-GGAAGGCTGGTCACCGCGACCACAGGAG-3′. A total of 100 ng of PCR products were subjected to deep sequencing with Illumina HiSeq 4000 (Illumina) in single end mode (150 bp) using Illumina input kit. The sequencing data were analyzed using FastQC and Quality Control Analysis Tool (QCAT).
CTGGTGGAGTCTGG-3', for IgH, and UNI-8N-HROK forward, 5'-AAGCAGTGATCAACGCAGANNNNN NNNAGATTGATGACAGTCTCCATCC-3', for IgL. The barcoding reaction contained 5 µl cDNA products, 8 µl of 5× HF buffer, 1 µl dNTP mixture (final concentration 250 µM each), barcode primer (final concentration 400 nM), and 0.2 µl Phusion DNA polymerase, with adjusted volume using RNase- and Dnase-free water to 40 µl. The barcoding program was as follows: 98°C for 1 min, 55°C for 30 s, 72°C for 10 min. Extraneous barcode primers were removed with 1 µl Exonuclease VII (New England Biolabs) per 40-µl reaction, incubated at 37°C for 1 h, followed by heat-inactivation at 95°C for 15 min. First-round PCR was designed to add adaptors at the 5’ and 3’ ends of the amplicon. Amplifications for heavy and light chain were done in separate reactions. Specifically, for heavy chain, each isotype was run separately. The first-round PCR reaction contained 5 µl barcoded single-stranded DNA products, 8 µl of 5× HF dNTP mixture (final concentration 250 µM each), forward and reverse primers (final concentration 250 nM each), and 0.5 µl Phusion DNA polymerase, with adjusted volume of final volume with RNase- and Dnase-free water to 40 µl. First-round PCR primers for IgH (Igκ1, Igκ2, Igγ1, Igγ2, Igγ3, Igγ) were CS1-UNV forward, 5’-TACACTGACGACATGTTCTA CAAAGCAGTGTATCAACG-3′; mgLm reverse, 5’-TAC GGTAGCAGACCTTGACTGATGCTGGAAGAC ATTTGGGAAAGAC-3′; mgLgI reverse, 5’-TACGGTAGC AGAAGATGGCTCTCAAGCTCGGAAATGCCCC TTGG-3′; mgLgG2 reverse, 5’-TACGGTAGCAGAGAC TTTGGTTCTGAACGGGAAGTGGCCTTTGAC-3′; mgLg2C reverse, 5’-TACGGTAGCAGACCTTGICTC TACTGCTAGGGAATAACCTTTTGAG-3′; mgLgG3 reverse, 5’-TACGGTAGCAGACCTTGICTCTACGC TCAGGGGATACCGCTTTGAC-3′; and mgLgE reverse, 5’- TACGGTAGCAGACCTTGICTCTACGGGCAGGCC AGGGTCTAGGAA-3′. First-round PCR primers for IgL (Igκ) chain were CS1-UNV forward, 5’-TACACTGAC GACATGTTCTCAGAACGAGTACATCAACG-3′, and mgLκ-HRO reverse, 5’-TACGGTAGCAGACCTTG GTCTAGTATTCAGCTCGAGTGTGCTG-3′.

We used the following PCR program for first-round deep-sequencing PCR: 98°C for 30 s, then 18 cycles of 98°C for 10 s, 65°C for 30 s, 72°C for 30 s, and 72°C for 5 min. Second-round PCR reaction was designed to add the Illumina linkers and sample barcodes. Amplifications for IgH and IgL were done in separate reactions, with each IgH isotype run separately. First-round PCR products generated in the Wesemann Laboratory were submitted to the DNA Services Laboratory at the W.M. Keck Center at the University of Illinois at Urbana-Champaign. Second-round PCR reactions included 4 µl of first-round PCR product used for each sample. The following reagents were combined: 2.0 µl of 10× FastStart Reaction Buffer without MgCl2, 3.6 µl of 25 mM MgCl2, 1.0 µl DMSO, 0.4 µl of 10 mM PCR grade nucleotide mix, 0.2 µl of 5 U/µl FastStart High Fidelity Enzyme Blend, and 3.8 µl water. 11 µl reagent mix was combined with 4 µl diluted first-round PCR product and 4 µl Illumina linker barcodes with the primers P5-CS1, 5’-AATGATACGCCAGACGTATGATGCCTCACTGAGATGTGTT CTACA-3′, and P7-index-CS2, 5’-CAAGCAGACAG GCCATACGAGAT[unique molecular identifier]TACGCT AGCACAGCTCTGCTCTC-3′. The PCR cycling program consisted of 95°C for 10 min, 1 cycle at 95°C 15 s, 60°C for 30 s, and 72°C for 1 min. In total, 19 cycles were conducted followed by 72°C for a 3-min extension. A diagram of the deep-sequencing strategy is provided in Fig. S3. PCR products were quantified on a Qubit fluorimeter (Life Technologies) and stored at −20°C. All samples were run on a Fragment Analyzer (Advanced Analytics), and amplicon regions and expected sizes were confirmed. Samples were then pooled in equal amounts according to product concentration. The pooled products were size-selected on a 2% agarose E-gel (Life Technologies) and extracted from the isolated gel slice with a gel extraction kit (Qagen). Cleaned size-selected products were run on an Agilent Bioanalyzer to confirm appropriate profile and determine mean size. The final pools were quantitated using Qubit (Life Technologies) and diluted to 5 nM final concentration. The 5-nM dilutions were further quantitated by qPCR on a CFX Connect Real-Time System (Bio-Rad Laboratories) and pooled evenly. The pool was denatured and spiked with 15% nonindexed PhiX control library provided by Illumina and loaded onto the MiSeq V2 Nano flow cell at a concentration of 7 P/M for cluster formation and sequencing. The PhiX control library provides a balanced genome for calculation of matrix, phasing, and prephasing, which are essential for accurate base calling. The libraries were sequenced from both ends of the molecules to a total read length of 250 nt from each end. The run generated .bcl files, which were converted into demultiplexed compressed fastq files using bcl2fastq 2.17.1.14 (Illumina). A secondary deep-sequencing decompressed the fastq files, generated plots with quality scores using FastX Tool Kit, and generated a report with the number of reads per sample/library. The .bcl files were also processed in bcl2fastq 2.17.1.14 without demultiplexing and reported. Both sorted unsorted fastq files were .tgz compressed and posted to a password-secured FTP site.

**Deep-sequencing analysis**

The sequences obtained from Illumina MiSeq deep sequencing were merged using PEAR v.0.9.10. The paired-end reads were truncated at the ends if the residues had a Phred score of less than 22. The merge criteria included minimum overlap of 20 nt and a minimum merged length of 380 nt for the heavy chain and 350 nt for the light chain. For intermittent low score (<20), nucleotides were replace by “N”. All sequences with four consecutive N’s were truncated at that point. PCR repeats were excluded using the unique molecular identifiers. Sequences with the same unique molecular identifier and a mismatch of 2% (8 nt, considered PCR error) were considered PCR repeats of the same mRNA. A consensus was generated.
keeping the most frequently occurring nucleotides at each position. At this juncture, all sequences obtained by Sanger methodology were added to the analysis and subjected to the next steps alongside sequences derived from high-throughput methodology. MUSCLE v3.8.31 was used for multiple sequence alignment. Multiple sequence alignments were edited with Jalview v.2.10. The N's were either replaced during consensus formation or assumed to be unmutated. The initial 15 nt (heavy chain) and 3 nt (light chain) of the heavy and light chains were removed during editing, as they had higher probability of PCR error/sequencing error. Mutation frequencies for each position and the number of mutations per sequence were calculated taking unmutated HRO heavy and light chains as reference. All analysis was done on Bioconductor package v.3.4 (R v.3.3.1). The dendrograms were created using MEGA7 v.7.0.18.

SPR

All SPR experiments were performed using the Biacore 3000 Processing Unit (GE) in the Molecular Biology Core Facility for Proteomics at Dana Farber Cancer Institute, Boston, MA. All expression clone–derived antibodies were tested using individualized runs performed in triplicate. A protein A sensor chip (GE) was used for analysis of all expression clone–derived IgG binding to OVA or HA–OVA conjugate. BiA normalizing solution (70% wt/wt) glycerol; GE) was used for periodic normalization of flow cells. BIA regeneration buffer, pH 1.7 (GE Healthcare), was used to regenerate the sensor chip surface after each cycle for each experimental run. All SPR assays were conducted using buffer containing 0.01 M Hepes, pH 7.4, 0.15 M NaCl, 3 mM EDTA, and 0.005% vol/vol Surfactant P20 (HBS-EP; GE Healthcare). The HBS-EP running buffer was used as a sample diluent, as running buffer for all cycles including start-up cycles, and as a negative control for analyte affinity binding in the active and reference flow cells. Reference channels were subtracted. All SPR data shown reflects the use of expression clone–derived antibodies from 293 T cell tissue culture supernatants after transfection. Supernatants were collected from Falcon 175-cm² sterile tissue culture flasks (Thermo Fisher Scientific) 48 h after cotransfection of 293 T cells using 250 µl polyethylenimine (stock 1 mg/ml; Polysciences) along with 25 µg of both IgH and IgL expression vectors per flask. The 293 T cells were maintained for 48 h in DMEM containing 4.5 g/L glucose, 1-glutamine, sodium pyruvate, and 10% FCS (Corning Cellgro). Supernatants were centrifuged in a Beckman Coulter GPR large benchtop centrifuge at 1,100 g for 30 min to remove cell debris before SPR.

All supernatant–derived, expression clone IgGs were used either diluted or undiluted in HBS–EP buffer and bound to the surface of the protein A chip. Different levels of IgG (ligand) binding to the protein A surface were evaluated for multiple expression clones via resonance signal, although this was neither expected nor observed to significantly alter equilibrium binding constants. For unmutated expression clone IgGs, IgG binding to the protein A chip was evaluated for resonance ranging from 400 to 10,000 RU. For mutated expression clone IgGs, isolated from single–cell sorting and representing biological pairings of IgH and IgL, the range of resonance studied for IgG bound to the protein A chip was 300 to 2,000 RU. Three separate SPR runs were performed for expression clone IgGs, and Kₐ is reported as a mean of those runs. OVA (Sigma–Aldrich) and HA–OVA analytes, diluted in HBS–EP buffer, with concentrations ranging from nanomolar to millimolar, were injected over surface–bound expression clone IgG to perform kinetic binding affinity analyses. For any given SPR run, at least one analyte concentration was performed in duplicate, and HBS–EP buffer alone was used for initial start–up cycles and as a control cycle after each series of analytes. Additionally, for any given SPR run, five to seven different analyte concentrations were tested. All binding affinity analyses were conducted using Biacore 3000 software, with a particular focus on the kinetic analysis function using kinetic wizard templates. All SPR data were analyzed first using BIA evaluation software followed by analysis of text files in Microsoft Excel and then production of figures using Adobe Photoshop CS6.

Epitope mapping

Epitope mapping of the five mAbs was performed by screening the OVA random mutagenesis library displayed on the surface of yeast as previously reported (Zuo et al., 2015). OVA was originally amplified from pcDNA3–OVA and then cloned to yeast surface display vector pCTCON2. The pCTCON2–OVA construct was chemically transformed to yeast cell line EBY100 to display OVA on the surface of yeast. Two rounds of error–prone PCR generated the OVA random mutagenesis library. The PCR products, together with linearized pCTC ON2 plasmid, were transformed into yeast competent cells using electroporation. The induced yeast library was stained with each of the five mAbs, and mutants losing binding to the mAbs were gated and sorted. Yeast displaying OVA was used as a positive control to gate the target population. Anti–c–myc antibody was used to identify yeasts displaying the transformed constructs. The sorted yeasts were sequenced, and mutations in OVA were identified by sequence alignment. OBI mAb was provided by S.K. Dougan (Dana–Farber Cancer Institute, Boston, MA) and H.L. Ploegh (Boston Children's Hospital and Harvard Medical School, Boston, MA). Plasmid pCTC ON2 and yeast strain EBY100 were provided by D.K. Wittrup (Koch Institute for Integrative Cancer Research, Massachusetts Institute of Technology, Cambridge, MA). Plasmid pcDNA3–OVA was a gift from S. Diebold and M. Zenke (Max–Delbrück–Center for Molecular Medicine, MDC, Berlin, Germany; plasmid 64599; Addgene). The structure of OVA was reported previously (Stein et al., 1991), and the structure figures were made with PYMOL (http://pymol.org/).

T cell proliferation

RBC–lysed HRO splenocytes were separated from non–B cell antigen presenting cells by treating with a biotin–labeled
cocktail of anti-CD11b (eBioscience), anti-CD11c (BioLegend), and anti-Ly–6G/Ly–6C (GR–1; BioLegend) followed by anti-biotin–magnetic bead labeling (Miltenyi Biotec) and column separation. Unbound flow-through was subjected to further magnetic column separation with anti-CD4 (L3T4) and B220 microbeads (Miltenyi Biotec). T cells were stained with CFSE to assess proliferation. A 96-well flat-bottomed tissue culture plate was seeded with 2 × 10^5 B cells and 5 × 10^4 CFSE-stained CD4^+ T cells with HA-OVA (5 µg/ml) conjugate or OVA (50 or 500 µg/ml) alone. T cell CFSE was assessed cytometrically after 5 d of co-culture.

T cell transfer and qPCR to analyze the frequency of HRO-B IgH in GCs

1 d before immunization with 100 µg OVA, 30 million B220^+ purified HRO B cells were transferred intravenously into wild-type BALB/c mice, which were killed 7 d after transfer. HA-positive B cells were identified with a biotinylated HA peptide and streptavidin conjugated to BV711. To analyze the frequency of HRO-B cells in the GC pool, a standard curve was generated by spiking 0.01–1.0% HRO-B cells into the 10^5 wild-type B cells. Approximately 10^5 GC B cells were sorted from HRO-B cells transferred wild-type BALB/c mice. Total RNA was isolated from each of the samples using the Trizol method (Invitrogen). Preparation of cDNA was performed by random hexamer (New England Biolabs) priming and superscript III (Invitrogen) reverse transcription for 60 min at 50°C. To analyze the HRO-B IgH frequency, specific TaqMan probe and primers for HRO-B IgH CDR3 were used (forward primer, 5′-CACCCCTGTACCT GCAAATGA-3′; reverse primer, 5′-CCAGTAAAGCAAA CCCCCTTCT-3′; and TaqMan probe, 5′-ATTACTGTG CAAGACGGGAGGTA3′), and reactions were run on a Mx3005p Stratagene real-time PCR machine in triplicate. Frequency is presented as the difference in threshold cycle (Ct) values normalized with HPRT.

Data availability

The Illumina MiSeq raw sequence data for this study is accessible at the NCBI Sequence Read Archive (SRA) under BioProject accession no. PRJNA388102 with BioSample accession nos. SAMN07199532 through SAMN07199557. The Sanger sequences are available in GenBank under accession nos. MF320741 through MF321728. Source code for sequence analysis can be found at https://github.com/Wesemann-lab/Mutation-analysis-of-Immunoglobulin-Sequences.

Online supplemental material

Fig. S1 shows the involvement of nonspecific B cells to early GC reactions in a competitive environment. Fig. S2 displays multiple sequence alignment of V_H and V_K from paired single-cell sequences and HRO BCR background mutation frequency. Fig. S3 provides a schematic representation of the deep-sequencing strategy.

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Author contributions: J. Silver and T. Zuo designed experiments, performed experiments, and prepared the manuscript; N. Chaudhary performed computational analysis of sequencing data and prepared figures; R. Kumari, P. Tong, S. Giguere, A. Granato, R. Donthula, and C. Devereaux performed experiments; and D.R. Wesemann conceptualized the study, designed experiments, supervised research, and wrote the manuscript.

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SUPPLEMENTAL MATERIAL

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Figure S1. Involvement of nonspecific B cells to early GC reactions in a competitive environment. (A) Histograms of CFSE dilution in Rag-deficient OVA-specific DO11.10 T cells incubated alone (red) or with HRO B cells treated with media (green), HA-OVA conjugate protein (5 µg/ml, purple), or OVA at 50 µg/ml (orange) and 500 µg/ml (blue) for 5 d. Shown are results from two independent experiments. (B) Schematic representation of the adoptive transfer experiments relevant for panels C–F. Wild-type mice received 30 million HRO B cells. 24 h later, the mice were immunized intraperitoneally with 100 µg OVA in alum. The mice were killed, and splenic GCs were analyzed by flow cytometry 6 d later. (C) Flow cytometric analysis of the splenic naive and GC B cells from adoptive-transferred and immunized mice stained for HA reactivity. (D) Summary of flow cytometry and single-cell sequencing data of GC B cells isolated based on HA reactivity. Data are representative of experiments in three other mice. (E) Distribution of the mutations from the mutated HRO-VH sequences identified in sorted GC B cells. Black stars indicate mutations within DGYW motifs. Blue stars indicate mutations that are within four nucleotides of a DGYW motif. (F) TaqMan qPCR measurement of HRO B cell frequency in early GCs of OVA-immunized HRO B cell–transferred wild-type mice. Frequency was assessed against a standard generated by spiking known dilutions of naive HRO B cells into wild-type B cells, and HPRT expression was measured as a normalization control. The blue dashed line represents the mean, and the underlying pink indicates the SD from four mice.
Figure S2.  Multiple sequence alignment of $V_{\text{H}}$ and $V_{\kappa}$ from paired single-cell sequences and HRO BCR background mutation frequency. (A) The different colors and shapes represent three different mice corresponding to the phylogenetic tree of paired sequences. Stars mark the SPR-tested sequences. (B) Distribution of mutation frequencies for each nucleotide position in $V_{\text{H}}$ and $V_{\kappa}$ (VL) of unimmunized HRO mouse splenic B cells from deep-sequencing data are shown. Two different mouse data from independent experiments (Expt.) are shown. $n$, number of sequences.
Figure S3. **Schematic representation of deep-sequencing strategy.** Proceeding top to bottom on the diagram, extracted HRO B cell RNA was subjected to first-chain synthesis using reverse transcription and multiple IgH and IgL constant region specific primers (designated "Specific 1"). After removal of primers using uracil DNA glycosylase, heat inactivation of this enzyme, and reverse transcription, IgH and IgL underwent barcoding reactions to add a unique molecular identification, using barcoding primers designated "Specific 2," which also enable second-strand cDNA synthesis. The Specific 2 primer also contains a universal (UNV) linker sequence used for downstream addition of PCR adaptors. Residual Specific 2 primers were removed with exonuclease 1. Then, once second-strand cDNA synthesis was completed, deep-sequencing library preparation proceeded through two steps of PCR amplification. First-round PCR was designed to add adaptors at both 5′ and 3′ ends of the amplicon, designated "CS1" and "CS2." The CS1 adaptor uses a UNV linker sequence to anneal to the cDNA template and amplify. The CS2 adaptor uses a cDNA specific sequence, designated "Specific 3," to anneal to the cDNA template and amplify. First-round amplification for IgH and IgL, as well as different IgH isotypes, was conducted in separate reactions. Second-round PCR was designed to add the Illumina linkers and sample barcodes, designated "Illumina P5" and "Illumina P7," to the cDNA libraries. Second-round amplification for IgH and IgL, as well as different IgH isotypes, was also conducted in separate reactions. PCR amplicons subsequently underwent quality control measures and Illumina MiSeq analyses to produce the sequencing data.