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Visualizing Antibody Affinity Maturation in Germinal Centers

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Abstract

Antibodies somatically mutate to attain high affinity in germinal centers (GCs). There, competition between B cell clones and among somatic mutants of each clone drives an increase in average affinity across the population. The extent to which higher-affinity cells eliminating competitors restricts clonal diversity is unknown. By combining multiphoton microscopy and sequencing, we show that tens to hundreds of distinct B cell clones seed each GC, and that GCs lose clonal diversity at widely disparate rates. Furthermore, efficient affinity maturation can occur in the absence of homogenizing selection, ensuring that many clones can mature in parallel within the same GC. Our findings have implications for development of vaccines in which antibodies with non-immunodominant specificities must be elicited, as is the case for HIV-1 and influenza.

The average affinity of specific antibodies increases dramatically over the course of an immune response (1, 2). This phenomenon is known as *affinity maturation*, and is the result of a Darwinian process that alternates stochastic somatic hypermutation (SHM) of

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immunoglobulin (Ig) genes with the selection and clonal expansion of B cells that have acquired affinity-enhancing mutations (2). Successive iterations of this cycle drive an increase in the overall affinity of antibodies over time, which is essential for their effectiveness in curtailing and preventing infection.

Affinity maturation takes place in germinal centers (GCs), defined microanatomical clusters containing up to a few thousand B cells that emerge in multiple copies within secondary lymphoid organs upon infection or immunization (3–7). Each GC is traditionally viewed as arising from a handful of independent "founder clones," many of which are lost as affinity-based competition leads to progressive focusing of the repertoire on the most successful progeny (which we refer to as "homogenizing selection") (8). Though it is inducible under experimental settings (9), the extent to which homogenizing selection takes place during normal immunization or infection has been difficult to determine, mostly due to technical limitations in the methods used to measure clonal diversity within individual GCs (10–14). Here, we report the development of two imaging-based approaches that expand our ability to study GC clonal composition in response to any antigen. Our findings highlight the heterogeneity of clonal evolution pathways in GCs.

Measuring clonal diversity in GCs using multiphoton microscopy

As an approach to estimating GC clonal diversity by imaging, we made use of a "brainbow" allele for multicolor fate mapping to permanently tag individual B cells and their progeny with different combinations of fluorescent proteins (15). This method, as implemented in the $Rosa26^{Confetti}$ allele (16), relies on stochastic Cre-mediated recombination to commit cells to expression of one of four possible fluorescent proteins, generating ten different color combinations when two alleles are recombined in homozygous mice (fig. S1A–B). Because recombination stops upon cessation of Cre activity, selective proliferation of an individual GC B cell would lead to the appearance of clusters of daughter cells of the same color.

We generated *Rosa26*^{Confetti/Confetti} mice carrying the Mx1-Cre transgene, which triggers Confetti recombination spontaneously during early life (17). Multiphoton imaging of popliteal lymph nodes (pLN) in naïve Mx1-Confetti mice showed that, in the absence of immunization, cells of different colors are intermixed, as expected from the polyclonal nature and migratory behavior of naïve lymphocytes (Fig. 1A, movie S1). In contrast, subcutaneous immunization with alum-adsorbed chicken gamma globulin (CGG) induced the appearance in the draining lymph node (LN) of clusters of cells of the same color, suggestive of monoclonal expansion (Fig. 1B, movies S2 and S3). Such clusters could also be found in mesenteric LNs (Fig. 1C) and even occasionally in Peyer's patches (fig. S1C) of unimmunized mice, where GCs form spontaneously in response to intestinal microbiota. Thus, clonal expansions within GCs can be readily detected by multicolor fate mapping, even in cases where the driving antigen is unknown.

Quantification of cell colors in GC dark zones at 15 days after immunization (Fig. 1D) revealed a wide range of color dominances (from 20.2% to 89.7%; median, 44.0%), and a relative paucity of predominantly single-colored GCs (6 of 40 (15%) GCs with color dominance > 70%; Fig. 1E). To validate these estimates, we used in-situ photoactivation

followed by FACS sorting (9, 18) to obtain single B cells from individual GCs, the clonal identity of which we then determined by *Igh* mRNA sequencing (Fig. 1F). This approach again showed varying degrees of clonal dominance among individual GCs (3/12 (25%) GCs with dominance >70%; range, 22.5% to 87.5%; median, 43.0%; Fig. 1G and fig. S2). It also revealed the frequent presence of clones that were shared between two individual GCs in the same LN, indicative of synchronous origin (colored slices in fig. S2). We conclude that GCs display variable degrees clonal dominance, even when induced synchronously by immunization. While predominantly monoclonal GCs do exist, these are relatively rare at the time point assayed.

Early GCs are highly diverse

The diversity of clonal dominance levels among mature GCs led us to question the generalizability of reports proposing that GCs are seeded pauciclonally (11, 12, 19). To address this, we generated mice expressing the photoactivatable-GFP transgene along with Cre recombinase driven by the endogenous Aicda locus (Aicda^{Cre}) (20), expressed in activated B cells, and a Rosa26 dox-stop-lox-tdTomato reporter. At 6 days after immunization with CGG alum, early GCs were identifiable in these mice as clusters of tdTomato⁺ cells within follicular dendritic cell (FDC) networks. We photoactivated two such clusters per LN, and sorted photoactivated GC B cells from each cluster for *Igh* sequencing (Fig. 2A and fig. S3A). Early GCs were highly and uniformly polyclonal, with 23 to 46 unique VDJ rearrangements detected when sequencing 34 to 77 single cells per GC (Fig. 2B). Extrapolation of these numbers using the Chao1 and ACE estimators of species richness (21) gave rough estimates of GC clonality that ranged from ~50 to ~200 clones per GC (median Chao1, 102 clones) for the 4 pairs of GCs analyzed (Fig. 2C). Overall mutation and class switching levels were low at this time point (fig. S3B), confirming that these were indeed early GCs. We were again able to identify B cell clones that were shared between two GCs in the same pLN (colored slices, Fig. 2B; mean 15.8% (SD 6.4) of clones found in one GC were also found in its neighbor). Similar experiments using other antigens showed that, although the extent of early GC diversity can vary depending on the immunizing antigen, high diversity is not a unique feature of the response to CGG (Fig. 2D and fig. S3C-E). Diversity was also not the result of non-specific recruitment of B cells to early GCs (fig. S3C and fig S4). We conclude that early GCs are highly diverse, containing tens to hundreds of distinct clones, depending on the antigen used for immunization. The progression from uniform diversity (Fig. 2) to variable degrees of clonal dominance (Fig. 1) in the CGG response implies that individual GCs display different rates of homogenizing selection acting subsequently to this coalescence.

Extent of homogenizing selection in individual GCs

Clonal dominance in GCs may arise by parallel selection of multiple members of a clone present in the early GC or by strong expansion of single SHM variants over cells of the same and of different clones. To investigate these dynamics, we generated mice in which recombination of $Rosa26^{Confetti}$ was driven by the tamoxifen-inducible $Aicda^{CreERT2}$ allele, specifically expressed in activated B cells (22). We thus delayed recombination until after GC formation, allowing different members of the same clone to express different color

combinations, each giving origin to an independent early GC "lineage." In this system, GCs dominated by the descendants of a single SHM variant would be identifiable as having "resolved" to a single dominant color (Fig. 3A).

Administration of tamoxifen to *Rosa26*^{Confetti/Confetti}. *Aicda*^{CreERT2/+} (AID-Confetti) mice at 5 days after CGG immunization triggered recombination, often of a single Confetti allele, in ~40–50% of GC B cells (fig. S5A–B) by the end of the 3 to 5-day period during which tamoxifen is active (23). Recombination thus ends near the onset of T cell-driven selection, at the beginning of the second week of the response (24) (Fig. 3A). GCs analyzed 3 days after tamoxifen treatment (8 days post-immunization) displayed roughly equal proportions of CFP⁺, YFP⁺ and RFP⁺ cells and a lower proportion of GFP⁺ and doubly-recombined cells (Fig. 3B), as expected from the original description of the Confetti allele (16). Only residual if any GCs containing recombined cells could be detected in unimmunized or alum-only control mice (fig. S5C).

Color distribution changed over time, with progressive emergence of GCs dominated by a single color/combination (Fig. 3B-G). To quantify color dominance, we calculated a normalized dominance score (NDS) for each GC, which approximates (25) the fraction of cells belonging to a single lineage originating at the time of recombination (fig. S5D). Kinetic analysis was extended to 15 days after tamoxifen treatment (20 days after immunization), near the end of the CGG-alum-induced GC response in most LNs. Median dominance increased monotonically, from 0.16 at day 3 to 0.39 at day 15 post-tamoxifen. GCs with NDS of 0.5 or more (corresponding to ~50% of all GC cells displaying the same color combination) appeared as early as day 9 post-tamoxifen, and a single color could account for close to 80% of GC B cells at day 11 and virtually all cells in a GC four days later (Fig. 3F). However, progression among GCs varied markedly, as dominance in the lowest-scoring GCs at days 11 and 15 after tamoxifen remained close to day 3 levels. Evaluation of color distribution in exceptional LNs in which GCs could still be found at very late time points showed that low-dominance GCs were present up to the very end of the response (fig. S6A-C). Therefore, GC selection does not inevitably lead to high color dominance within the lifetime of the response to CGG-alum.

To determine the extent to which progression of color dominance depends on affinity-based competition, we generated an adoptive transfer model in which wild-type hosts received B cells from *Aicda*^{CreERT2/CreERT2}. *Rosa26*^{Confetti/Confetti}. *Igh*^{B1-8hi/+} (AID-Confetti-B1-8) donors (in which Igλ⁺ B cells have uniformly high affinity to the NP hapten and do not undergo affinity maturation due to lack of a functional *Aicda* allele). NP-OVA-induced GCs in recipient mice approached high color dominance only rarely and with much delayed kinetics (Fig. 3E–F and fig. S6D). Therefore, differences in affinity between competing B cells likely contribute towards the rapid rise in dominance observed in a fraction of GCs. To quantify the effect of neutral competition—defined as the change in lineage abundances over time due to stochastic factors, in the absence of differences in affinity—in AID-Confetti-B1-8 GCs, we calculated a *divergence index*, which computes the difference between the expected and observed proportions of all 10 colors in each GC (fig. S5E), and is thus more sensitive to small changes in color abundance than the NDS. This metric showed marked divergence from baseline distribution in this population (Fig. 3G and fig. S6D), highlighting

the effect on GC selection of factors unrelated to affinity. Moreover, the least divergent GCs in "wild-type" AID-Confetti mice remained relatively close to baseline color distribution (at day 15 post-tamoxifen, 25.7% of AID-Confetti GCs scored below the median of the AID-Confetti-B1-8 distribution; Fig. 3G). We conclude that individual GCs are highly heterogeneous with respect to selection: while a fraction of these structures become heavily dominated by cells of one color in a matter of days—suggesting strong expansion of the descendants of a single SHM variant arising at or after the onset of GC selection—others deviate from baseline color distribution at levels that do not exceed those attained by neutral competition.

We extended our findings to a different antigenic system by measuring clonal dominance in GCs elicited by infection with Friend retrovirus (FV) (26). We treated infected AID-Confetti mice with tamoxifen at 20 days post-infection (an early time point in the delayed GC response to FV) and imaged spleen slices 10 days later (fig. S7A–B). A wide range of NDS and divergence scores were also observed in FV-induced GCs (Fig. 3H and fig. S7C). Therefore, heterogeneity in the outcome of selection is a common property of GCs induced by model antigens and by viral infection, and thus likely represents an intrinsic property of GC evolution.

Homogeneous GCs are the product of "clonal bursts"

To investigate the underlying clonal structure of the variation in homogenizing selection rates among GCs, we determined the *Igh* gene sequences of fluorescent B cells isolated from GCs with different degrees of color dominance. To achieve this, we dissected pLNs from immunized AID-Confetti mice into fragments containing single GCs using vibratome sectioning guided by multiphoton microscopy (Fig. 4A and fig. S8). From each LN, we sorted cells from one high-dominance GC and from a neighboring low-dominance GC for *Ig* sequencing.

We obtained Igh sequences from 52 to 74 single cells per GC from 3 pairs of pLN GCs harvested 10 days after tamoxifen treatment (15 days after immunization with CGG-alum) (Fig. 4B and fig. S9A). Comparison of SHM levels between high- and low-dominance GCs from the same LN (fig. S9B) and between Aicda-sufficient and haploinsufficient GCs (fig. S9C) indicated that heterogeneity was not the consequence of asynchronous GC formation or of substantially impaired AID activity. Most cells of the dominant color in the 3 highdominance GCs (Fig. 4B, and fig. S9A, left panels) derived from a single expanded clone. In all 3 GCs, dominance could be readily attributed to the selective expansion of a single SHM variant, 3 to 5 V_H mutations (4 to 9 total mutations) distanced from the unmutated ancestor (UA) (dashed boxes in Fig. 4B and fig. S9A). Thus, high-dominance GCs are predominantly monoclonal, and appear to result from "clonal bursts," in which a single SHM variant is heavily expanded over a short period of time, leading to extensive loss of clonal diversity concomitant with broad diversification of the expanded variant. On the other hand, lowdominance GCs (Fig. 4B and fig. S9A, right panels) were more clonally diverse than their high-dominance neighbors. Low color dominance is unlikely to be the result of failure to identify a dominant nonfluorescent lineage, since fluorescent and nonfluorescent B cells obtained from two independently-sequenced low-dominance GCs were largely clonally

related (fig. S10). Although low-dominance GCs consisted predominantly of small independent expansions, one of three GCs sequenced (LN2/GC2) contained a larger clone (accounting for 59% of all recombined cells), within which multiple colors were represented. This expanded clone carried heavy and light chain V(D)J rearrangements identical to the dominant clone in the single-colored GC sequenced from the same LN (LN2/GC1), indicative of a common cell of origin. In contrast to the clonal bursts of single-colored GCs, however, this multicolored expansion branched off from the *Igh* UA itself, and developed in parallel along several distinct lineages. Thus, expansion of this clone began in the pre/early GC period, prior to the end of the tamoxifen pulse, and subsequent competition in the mature GC failed to focus on a single dominant SHM variant. Together, these data indicate that, while clonal dominance in GCs can arise by parallel expansion of members of the same clone, loss of diversity is greatest in GCs in which strong clonal bursts rapidly expand and diversify single SHM variants.

Affinity maturation in the absence of homogenizing selection

To investigate the relationship between clonal bursting and affinity maturation, we measured the affinity of recombinant Fab fragments derived from B cells originating from high- and low-dominance GCs (Fig. 4B), as well as from their deduced or sequenced UAs. Fab affinities varied markedly between clones, from undetectable to low nanomolar K_D (Fig. 5). As expected, the two Fabs obtained from SHM variants that underwent clonal bursts had higher affinity for the immunizing antigen than their respective UAs (5.3-fold increase $(1.0 \times 10^{-7} \text{ to } 1.9 \times 10^{-8})$ for clone 1.1 and 14.5-fold increase $(4.1 \times 10^{-8} \text{ to } 2.9 \times 10^{-9})$ for clone 2.1, Fig. 5). However, affinity maturation was also evident in F_{ab}s cloned from lowdominance GCs. Of the three clones obtained from LN1/GC2, SHM variants of two clones (1.3 and 1.4) showed increased affinity over their UAs, whereas the affinity of the third clone (1.2) remained below detection level (Fig. 5). Clone 1.4 showed a gain in affinity of 12-fold (2.4×10^{-7}) to (2.0×10^{-8}) , notably larger than the 5.4-fold change observed for clone 1.1 (associated with a clonal burst in the neighboring single-colored GC). We also determined the affinity of six additional variants of clone 2.1 from LN2/GC2 (which shared its UA with the dominant clone of LN2/GC1, and are thus directly comparable between GCs). Gains in affinity were evident in all of these variants, ranging from 3.5 to 13.8 fold ($K_D 1.2 \times 10^{-8}$ and 3.0×10⁻⁹, respectively) (Fig. 5C–E). Although the highest of these was close in magnitude to the 14.5-fold change seen in variant 2.1M1 (associated with the clonal burst in LN2/GC1), the increase in affinity among GC2 Fabs was in general lower (median increase of 6.8-fold). Thus, affinity maturation can occur in the absence of clonal bursts, although limited data from the two LN2 GCs suggests that selection of high-affinity mutants in these cases may not be as efficient.

Finally, detailed analysis of the SHM trajectory of variant 2.1M1 showed that this burst was associated with a somatic mutation in *Igk* (C119>G, Ala>Gly) that alone increased the F_{ab} affinity by 7.6-fold (Eig. 5E-G). This mutation was also found to have occurred independently at least twice in LN2/GC2, but without the accompanying burst. Thus, identical mutations occurring within the same clone may still have distinct outcomes when in different GCs, again suggesting a role for stochastic factors in GC selection.

Conclusions

Using a combination of imaging methods and single-cell sequencing, we show that (i) early GCs can be highly diverse, containing up to hundreds of distinct B cell clones; (ii) individual GCs approach homogeneity at different rates, leading to variable loss of diversity; (iii) homogeneous GCs are the product of "clonal bursts" that lead to rapid expansion and further diversification of SHM variants with improved affinity; and (iv) affinity maturation can take place in the presence or absence of such bursts, and thus does not necessarily require radical loss of clonal diversity.

We speculate that affinity maturation arises through a balance between slow but steady elimination of lower affinity clones and variants, punctuated by sporadic clonal bursts of various magnitudes that generate a wealth of SHM variants of a clone, at the expense of clonal diversity. This balance would ensure that the polyclonality of the GC response is maintained, at the same time as very successful clones can be heavily diversified and exported to effector or memory fates, generating the range of GC outcomes observed by us and others (14, 27). Although it cannot be ruled out that differences in outcomes between GCs may be related to factors such as initial clonal composition or timing of the appearance of key mutations, the finding that GCs can deviate substantially from baseline color distribution even when all cells have equal affinity and cannot undergo SHM (Fig. 3F–G and fig. S6D) suggests that stochastic factors unrelated to affinity can strongly influence GC selection. A system in which B cell selection is driven by dynamic encounters with a limiting number of T helper cells would be especially sensitive to such effects (9, 27). The explosive B cell proliferation that can be driven by such encounters would provide a mechanistic basis for the clonal burst phenomenon (28, 29).

The finding that early GCs can contain a large diversity of clones is in contrast to the widely held view that these structures form pauciclonally (10–12, 30). Although this discrepancy may be partly attributable to methodological issues (31), Kuppers et al. (13), reported higher clonal diversity in one of two GCs originating from different reactive human lymph nodes, and speculated that pauciclonal GCs were likely the product of extensive selection acting on originally polyclonal early GCs. Our current data strongly support this view.

Our findings have implications for the design of vaccines against highly variable pathogens such as influenza and HIV, where broadly-protective antibodies targeting conserved, non-immunodominant epitopes must be elicited (32, 33). Understanding whether GC competition restricts the appearance of such clones, and how clonal bursts may promote the somatic diversification required for broad neutralization to emerge may prove key to the successful generation of broadly-protective antibodies by vaccination.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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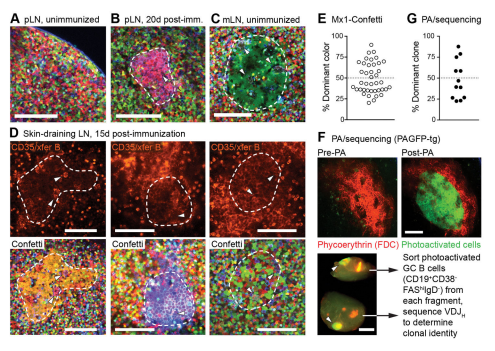


Figure 1. Visualizing clonal expansions in GCs using a brainbow allele (A) Popliteal lymph node (pLN) of an unimmunized Rosa26^{Confetti/Confetti}.Mx1-Cre mouse, imaged by multiphoton microscopy. (B) pLN of a mouse immunized 20 days previously with 10 µg CGG in alum in the hind footpad. The image shows a cluster of similar-colored cells corresponding to a GC (dashed line), as evidenced by the presence of tingible body macrophages (arrowheads). (C) Single-colored GC (dashed line) in a mesenteric lymph node (mLN) from an unimmunized mouse. (D) GCs in draining LN of mice immunized subcutaneously with CGG in alum 15 days prior to imaging. The location of the GC dark zone (dashed line) was determined by injection of labeled antibody to CD35 and surfacelabeled naïve B cells (top panels, fluorescence is from Alexa 633 label). For each GC, Confetti colors (bottom panels) were imaged independently and used for quantification. Confetti colors are as shown in fig. S1B. GC identity is confirmed by presence of tingiblebody macrophages (arrowheads). Scale bars, 100 µm. Second harmonic generation from collagen fibers is shown in blue. (E) Quantification of data as in (D). Each symbol represents one GC. Graph shows percentage of cells expressing the most abundant color combination. Data pooled from four mice, two independent experiments. (F) Quantifying GC clonality by photoactivation (PA). Photoactivatable-GFP-transgenic mice were immunized in the footpad with 10 µg CGG in alum and imaged 15 days later. FDC networks were labeled with phycoerytrhin immune complexes. Top, images of a single GC within a pLN, prior to and after photoactivation (Scale bar, 100 µm). Bottom, single pLN containing two photoactivated GCs (arrowheads) dissected into two fragments, each of which is separately processed for sorting of PA⁺ GC B cells and *Igh* sequencing (Scale bar, 500 μm). (G) Quantification of clonal dominance in multiple GCs. Data obtained as in (F), with clonal identity assigned based on Igh sequence. Each symbol represents one GC, with 2 GCs sequenced per LN (full clonality charts in fig. S2). Given is the percentage of cells belonging to the most abundant clone. Data are from 5 mice, 3 independent experiments.

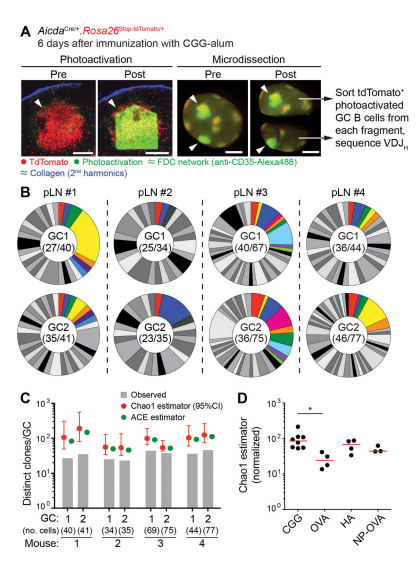


Figure 2. Clonal diversity in early GCs

(A) Photoactivation of single early GC clusters. photo activatable-GFP-transgenic. Aicda^{Cre}. Rosa26^{lox-stop-lox-tdTomato} mice were immunized in the footpad with 10 μg CGG in alum and imaged 6 days later. FDC networks were marked by injection of labeled antibody to CD35. Left panels show images of a single tdTomato⁺ cluster (arrowheads) within a pLN prior to and after photoactivation (Scale bar, 200 μm). Right panels show dissection of a single pLN with two photoactivated GCs (arrowheads) into two fragments, each of which is separately processed for cell sorting (Scale bar, 500 μm). (B) Pie charts showing clonal diversity in early GCs. Each slice represents one clone. Colored slices represent clones that were found in both GCs (upper and lower pie charts) from the same pLN. Numbers in the center of each chart are (number of clones observed/total number of cells sequenced). Clonal identity was assigned based on *Igh* sequence. Pairs are from 4 different mice in 3 independent experiments. (C) Estimation of total clonal richness in individual GCs using the Chao1 and ACE estimators. Graphs show observed clonal richness (from panel (B)), and total richness according to the indicated estimator. (D) Estimated clonal richness (Chao1) in GCs elicited by various antigens. Mice were immunized with 10 μg of the indicated antigen, and imaged/

photoactivated as in (A). Each symbol represents one GC, bar indicates median. For comparison purposes, estimates are normalized by interpolation to the size of the smallest sample (34 cells). Note that numbers for CGG GCs in this panel differ from those in panel (C) due to normalization. Further details in Fig. S3. * p < 0.05, Kruskall-Wallis test with Dunn's post-test. All other comparisons were not significant.

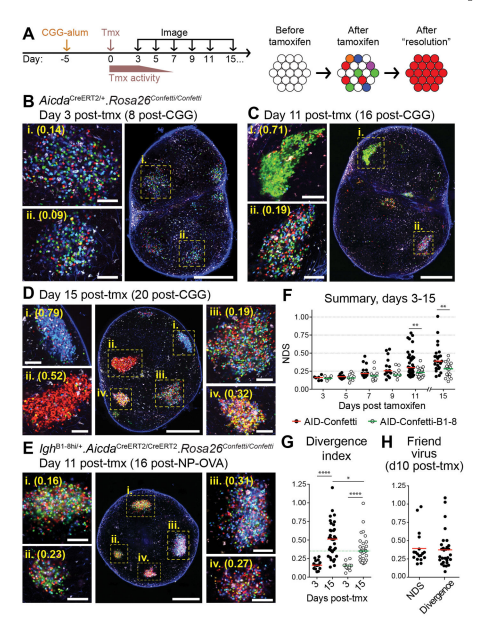


Figure 3. Kinetics of color dominance in individual GCs

(A) Graphic representation of the experimental protocol. AID-Confetti mice were immunized in the footpad with 10 µg CGG in alum, treated with tamoxifen (tmx) 5 days later, and imaged at the indicated time points. Tamoxifen triggers recombination of one or both Confetti alleles in individual GC B cells, independently of clonal origin. (B-E) Whole lymph node (large panel; scale bar, 500 µm) and higher-magnification images (side panels; scale bar, 100 µm) showing GCs at different times after tmx administration. Cell colors as in fig. S1B. Second harmonic generation from collagen fibers is shown in blue. (B-D), AID-Confetti mice imaged at the time points indicated. (E) WT recipients of $1-2 \times 10^4$ adoptively-transferred AID-Confetti-B1-8 Ig λ^+ B cells, immunized with NP-OVA as in (A) and imaged 11 days post-tmx. Higher-magnification panels show independently-acquired images of the GCs indicated in the overview panel. (F) Quantification of data as shown in

(B-E). NDS, normalized dominance score. Bars represent the median. (G) Divergence index for AID-Confetti and AID-Confetti-B1-8 mice at days 3 and 15 post-tmx. Bars represent the median. Green dotted line placed at the median of the day 15 AID-Confetti-B1-8 data for reference. (H) Quantification of GC selection in AID-Confetti mice infected with Friend Virus as detailed in fig. S7. Graph shows NDS and divergence index at day 10 post-tmx (day 30 post-infection). Bars represent the median. For panels (F-H), each symbol represents one GC. Data are pooled from 2–6 replicate experiments. For NDS quantification, we exclude GCs with density of fluorescent cells below 0.4 cells/100 μm^2 (equivalent to approximately 40% of cells having recombined a Confetti allele; see Supplementary Text). * p<0.02; ** p<0.01; **** p<0.0001, Mann-Whitney U test.

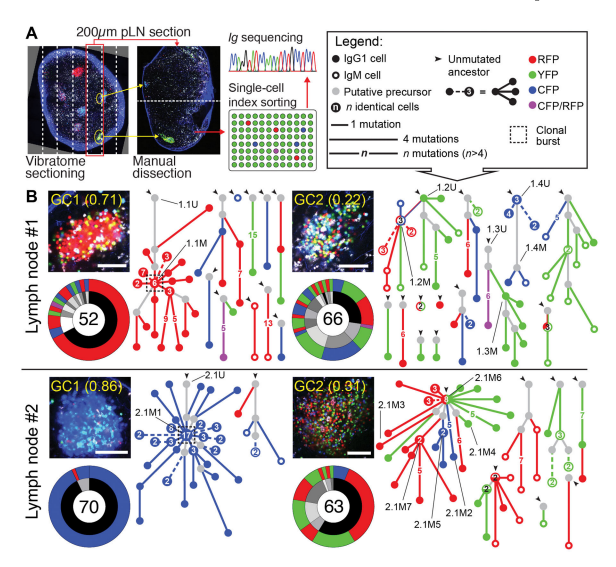


Figure 4. Clonal relationships among cells obtained from GCs with high or low color dominance (**A**) Method used to obtain *Ig* sequences. (**B**) *Igh* sequence relationship among B cells from 2 pairs of individual GCs from 2 pLNs of different mice, obtained 10 days after tamoxifen administration (15 days post-immunization), as described in Fig. 3. Each panel contains: *top left*, multiphoton image (scale bar, 100 μm, cell colors as in fig. S1B, second harmonic generation from collagen fibers is shown in blue); *bottom left*, clonal distribution pie-chart (with clones represented in grayscale in the inner ring and Confetti colors in the outer ring; number of cells sequenced is indicated in the center); and *right*, trees representing the phylogeny of *Ig* heavy-chain V-segment sequences within each clone (symbols according to the legend in the top right corner). Dashed lines within phylogenies indicate multiple variants distanced the same number of mutations from the originating node. IDs of variants for which affinity was measured in Fig. 5 are indicated by black lines. For each LN, GC1 and GC2 were considered as displaying high and low color dominance, respectively

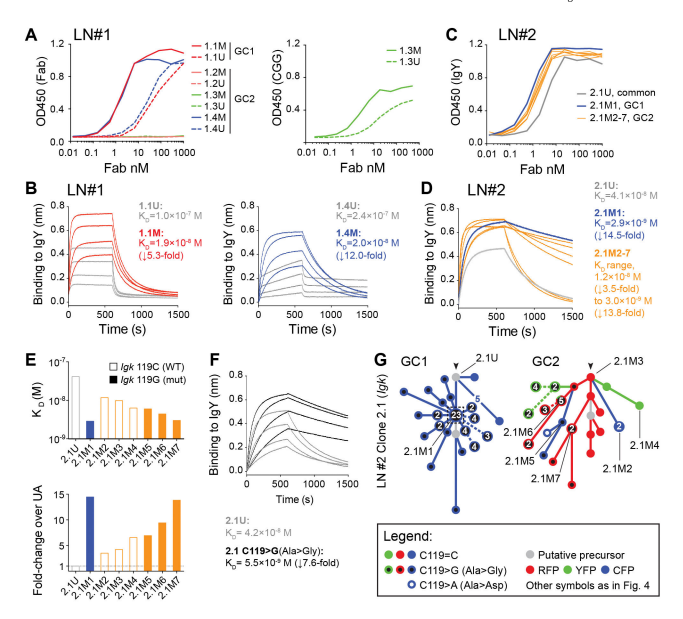


Figure 5. Affinity maturation in GCs with high or low color dominance

Affinity measurements for reconstructed F_{ab}s derived from B cell clones/variants indicated in Fig. 4B. (A) Binding of F_{ab}s cloned from LN1 to IgY (right) or CGG (left), measured by ELISA. (B) Bio-layer interferometry for F_{ab}s cloned from LN1 binding to IgY. (C) and (**D**) as in (A) and (B), respectively, but using F_{ab}s cloned from LN#2. (E) Affinity for IgY among variants of clone 2.1 (Fig. 4B) from LN2/GC1 (blue) and LN2/GC2 (orange), shown as K_D (top) and fold-change over UA (bottom). The unmutated ancestor (2.1U) is shown in gray. Open bars have the WT nucleotide (C) in the 119 position, closed bars have the C119>G (Ala>Gly) mutation. (**F**) Affinity of F_{ab}s reconstructed from clone 2.1, either unmutated (gray lines) or with replacement of a single *Igk* nucleotide (C119>G (Ala>Gly), black lines). (**C**) *Igk* sequence relationships among B cells from clone 2.1 from LN2/GC1 (left) or LN2/GC2 (right). Symbols according to the legend below the figure and in Fig. 4B. Cloned F_{ab} IDs are indicated by black lines. Note that, although not all cells yielded both *Igk*

and Igh sequences, clonal relationships are drawn from all available data, and therefore exact correspondence between the trees in panel (G) and in Fig. 4B is not expected. Bio-layer interferometry was performed with $F_{ab}s$ at 20, 40, 80, and 160 nM. Panel (D) shows only the 160 nM measurement. Reported affinities are the average of two measurements fitted globally at the 20–160 nM range.