UNIVERSITY OF BIRMINGHAM University of Birmingham Research at Birmingham

Variant proteins stimulate more IgM+ GC B-cells revealing a mechanism of cross-reactive recognition by antibody memory

Burton, Bronwen R; Tennant, Richard K; Love, John; Titball, Richard W; Wraith, David C; White, Harry N

DOI: 10.7554/eLife.26832 10.7554/eLife.26832

License: Creative Commons: Attribution (CC BY)

Document Version Peer reviewed version

Citation for published version (Harvard): Burton, BR, Tennant, RK, Love, J, Titball, RW, Wraith, DC & White, HN 2018, 'Variant proteins stimulate more IgM+ GC B-cells revealing a mechanism of cross-reactive recognition by antibody memory', eLife, vol. 7, e26832. https://doi.org/10.7554/eLife.26832, https://doi.org/10.7554/eLife.26832

Link to publication on Research at Birmingham portal

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

•Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.

•User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?) •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

1	
2	
3	
4	Variant proteins stimulate more IgM+ GC B-cells revealing a mechanism of
5	cross-reactive recognition by antibody memory
6	
7	
8	Bronwen R Burton ^{2,†} , Richard K Tennant ^{1,†} , John Love ¹ , Richard W Titball ¹ , David C
9	Wraith ³ and Harry N White ^{1*}
10	
11	¹ Department of Biosciences, University of Exeter, Exeter, EX4 4QD, UK.
12	
13	² Faculty of Biomedical Sciences, University of Bristol, Bristol, BS8 1TD, UK.
14	
15	³ Institute of Immunology and Immunotherapy, University of Birmingham, Birmingham, B15
16	2TT, U.K.
17	
18	*corresponding author, h.n.white@exeter.ac.uk
19	
20	[†] Equal contribution.
21	
22	
23	
24	
25	
26	
27	
28	
29	
30	
31	
32	
33	
34	
35	
36	
37	
38	
39	
40	
41	
42	
43	
43 44	
44 45	
45 46	
40	
47	
47	
40	
48	

49 Abstract

50	
50	Vaccines induce memory B-cells that provide high affinity secondary antibody
51	responses to identical antigens. Memory B-cells can also re-instigate affinity
52	maturation, but how this happens against antigenic variants is poorly understood
53	despite its potential impact on driving broadly protective immunity against pathogens
54	such as Influenza and Dengue. We immunised mice sequentially with identical or
55	variant Dengue-virus envelope proteins and analysed antibody and germinal-centre
56	(GC) responses. Variant protein boosts induced GC with higher proportions of IgM+
57	B-cells. The most variant protein re-stimulated GCs with the highest proportion of
58	IgM+ cells with the most diverse, least mutated V-genes and with a slower but
59	efficient serum antibody response. Recombinant antibodies from GC B-cells showed
60	a higher affinity for the variant antigen than antibodies from a primary response,
61	confirming a memory origin. This reveals a new process of antibody memory, that
62	IgM memory cells with fewer mutations participate in secondary responses to variant
63	antigens, demonstrating how the hierarchical structure of B-cell memory is used and
64	indicating the potential and limits of cross-reactive antibody based immunity.
65	
66	
67	
68	
69	
70	
71	
72	
73	

75 Introduction

76	Antibody-based immunity is underpinned by memory B-cells that have undergone
77	antibody somatic hyper-mutation (SHM) and selection for improved antigen binding
78	in germinal centres (GCs) (MacLennan et al., 1997). Re-challenge with the same
79	antigen stimulates a rapid, higher affinity, secondary antibody response.
80	Protective immunity to highly mutable viruses, like Dengue and Influenza, can
81	be induced by vaccination but the high level of variation often leads to immune
82	escape (Nabel & Fauci, 2010), leading to a focus on generating vaccine responses
83	against conserved antigenic regions (Wu et al., 2010; Corti et al., 2011; Wang et al.,
84	2015).
85	Memory B-cells of IgM and IgG isotypes can also re-instigate GCs after
86	secondary exposure (Dogan et al., 2009; Pape et al., 2011; McHeyzer-Williams et al.,
87	2015), but how this happens against variant antigens is poorly understood despite its
88	potential impact on driving the most broadly protective immunity.
89	Several studies suggest diversity in the memory B-cell population, showing
90	that cells can express IgM or IgG (Dogan et al., 2009; Pape et al., 2011), be mutated
91	or non-mutated (Kaji et al., 2012) and have low affinities (Smith et al., 1997), but still
92	persist in GCs (Kuraoka et al., 2016).
93	It has long been speculated that this diversity may facilitate the recognition of
94	antigenic variants (Herzenberg et al., 1980; Pape et al., 2011; Kaji et al., 2012) which
95	could stimulate secondary GCs derived from less mutated, naïve-like, memory B-cells
96	that still had an advantage over naive B-cells due to their increased numbers, pre-
97	selected V-genes and lower activation thresholds (Good et al., 2007; Good et al.,
98	2009).

99	By sequentially immunizing mice with the same or different Dengue-virus
100	envelope proteins, and analyzing serum antibodies and GC B-cells, we provide
101	evidence that supports the hypothesis that less developed memory B-cells are used in
102	secondary responses to variant antigens.
103	
104	
105	
106	
107	
108	
109	
110	
111	
112	
113	
114	
115	
116	
117	
118	
119	
120	
121	
122	
123	

124 Results

125 *E-protein variants elicit secondary serum antibody responses with different speed and*126 *cross-reactivity*

127 We chose Dengue-3 envelope protein (E3) for all priming immunisations. Boost 128 immunisations were performed 38 days later with identical E3 protein or variant E2 or 129 E4 proteins which have 68% and 63% overall sequence identity with E3, respectively. 130 The cross reactivity of E3-primed mouse serum IgG correlated with sequence 131 identity (Fig. 1A), and overall cross-reactivity also correlated (Fig 1B). 132 Boosting with homotypic E3 antigen induced a rapid antibody memory 133 response with anti-E3 titres rising rapidly to day 7, and not increasing further (Fig. 134 1D). E-protein boosted antisera was not reactive with an irrelevant His-tagged protein 135 (PR8 HA)(Fig 1C). 136 Heterotypic boosting with E2 induced a rapid and significant increase in anti-137 E3 titre, as might be expected if cross-reactive memory antibodies against the priming 138 E3 antigen were recalled (Figure 1D), that did not increase further by day 17. E4 139 boosting induced a modest but not statistically significant increase in the anti-E3 titre, 140 even by day 17, showing the E4 variant boost had not induced a significant anti-E3 141 antibody memory response, or the induced antibodies had a low affinity for E3 (see 142 discussion). 143 The anti-E2 titre induced by the E2 boost increased about 120-fold by day 7 144 (Figure 1E), and did not increase further by day 17, further indicating that E2 boosting 145 induced a rapid memory-like serum IgG response against E2 derived from cross-146 reactive E3 primed memory B-cells. Conversely the anti-E4 titre, induced by E4 147 boosting, rose significantly but to a lower level, about 20-fold, by day 7 (Figure 1F)

148 and showed a further rise by day 17. A boost alone did not induce a detectable

149	antibody titre however, ('BO', Figure 1D) suggesting a role for memory B-cells of
150	some type and/or cross-reactive T-cell memory, facilitating the E4 boost response.
151	
152	Increased levels of $IgM+GCB$ -cells with fewer mutations after variant protein
153	boosting
154	E3 and E2 boosting induced early GC B-cell levels similarly by day 7, to 4.5-5.5% of
155	total lymphocytes, which then reduced by two-thirds by day 17 (Figure 2B). E4
156	boosting induced GC B-cell levels about a third as high, which then reduced similarly
157	by about 60% at day 17, remaining 4-fold higher than controls.
158	Analysis of the proportion of IgM+ GC B-cells showed a highly significant
159	trend at day 7 after boosting, with the proportion of IgM+ GC B-cells correlating with
160	increasingly variant challenge (Figure 2C). This trend continued to day 17. The
161	proportion of IgM+ B-cells was also consistent between individuals in an
162	experimental group (Figure 2D).
163	Overall levels of VH mutations increased in all groups from day 7 to day 17
164	(Figure 2E), consistent with secondary affinity maturation. Sequences are available in
165	Supplementary File 1.
166	There were lower levels of SHM in IgM+ GC B-cells 7 days after the variant
167	boosts, particularly with the most variant protein E4, compared to the homotypic E3
168	boost (Figure 2F). Boosting with variant proteins, therefore, induced early GCs with
169	increased proportions of IgM B-cells that had fewer VH mutations.
170	Analysis of the VH clonality of GC B-cells after E-protein boosts showed that
171	almost every VH sequence was from a distinct B-cell clone (Figure 2G). These data
172	also showed that the two variant boosts elicited different repertoires of VH. 40% of
173	the VH sequences sampled at day 7 from E2 boosted mice were either VH14-3 or the

174	closely related VH14-4 (black dots, Figure 2G), suggestive of a secondary response
175	more focused on a particular epitope (see discussion) Some of these VH were also
176	present in the homotypic E3 boost day 7 samples. but neither were detected at day 7
177	after E4 boosting (Figure 2G).
178 179	Changes in serum affinity/avidity after variant antigen boosting
180	E2-variant boosting induced an immediate and significant increase in avidity by day 7
181	(Figure 3A) which did not detectably change until perhaps day 32, although data
182	variability is high. A modest but significant increase in serum affinity, however, was
183	detected by day 17, with a further increase detected by day 32 (Figure 3C). We
184	interpret this to mean that a relatively small portion of serum IgG underwent affinity
185	maturation by day 17 in response to the E2 boost and was not detectable by the Urea
186	avidity assay due to high variability and the high pre-existing IgG titres (Figure 1E),
187	or other limitations of the Urea assay (Alexander et al., 2015). Boosting with the E4
188	variant elicited slower increases in relative affinity and avidity, only detectable by day
189	32, but by then representing an equivalent, if not greater, increase compared to that
190	induced by E2 (Figure 3B & 3D)
191	
192	Similar memory T-cell stimulation by variant Dengue E-proteins
193	Memory T-cells are necessary for memory B-cell responses against haptens and viral
194	proteins (Aiba et al., 2010; Hebeis et al., 2004). We found no evidence that the
195	memory T-cell response to re-stimulation by variant E-proteins was any different
196	from re-stimulation by E3 (Figure 3E). These data imply that a deficiency in T-cell
197	recognition of these antigens cannot explain the differences in response to E2 and E4

- 198 challenge, and supports the idea that either T-cell receptors can recognize antigenic
- 199 peptides from regions with around 50% sequence difference (see discussion) or, more

200 likely, B-cells present peptides from different, more conserved regions than those201 their antibodies bind to.

202

203 The primary antibody and GC response to E4

204 For comparison with the E4 boost response, we performed primary immunisations

with E4 and analysed serum antibodies and GC B-cells at day 7 and day 17. Serum

206 levels of anti-E4 IgG rose to a moderate level by day 17 (mean EPT = 3.6, Figure

4A), being less than seen after E4 boosting (Figure 1F). GC B-cell levels rose to a

208 mean of 0.8% lymphocytes at day seven after E4 priming, half as much as after the E4

boost, then fell similarly to the post boost samples by around 60% by day 17 (Figure

4B). As with the E-boost GCs, the proportion of IgM+ GC B-cells fell over time

211 (Figure 4C) and levels of VH mutation in all B-cells and IgM+ B-cells increased

212 (Figure 4D & 4E). The median level of VH mutation in IgM+ GC B-cells at day 7

after E4 priming is less (=2) than after E4 boosting (=3) suggesting, not conclusively,

that GC Bells at day 7 after E4 boosting are memory derived. Antibody titres were

215 insufficient to do a relative affinity competition ELISA and no 7M Urea resistant IgG

216 was detected 7 or 17 days after E4 priming (data not shown).

217

218 IgM Antibodies from E4 boost GC show evidence of prior selection

If E4 boost induced B-cells are memory derived the antibodies should show evidence of pre-selection by the E3 prime. We made 48 recombinant antibodies (rAbs), 38 of which were IgM (supplementary file 2), 24 from E4 primed mice (day 7 and day 17) and 24 from E4 boosted mice (day 7). Figure 4F and Supplementary File 2, show the results from the initial screen of all rAbs against E4, indicating that the efficiency of detection of positive binding (deemed as O.D. > 0.1, useful for subsequent titration)

225	was quite low but consistent with the 30-50% binding frequency of GC rAbs
226	previously observed (Kuraoka et al., 2016), except for E4 prime day 7, which has only
227	2/13 rAbs binding strongly enough to be titrated. This might be expected of antibodies
228	from a day 7 primary response GC, and indicated they were overall of lower affinity.
229	Other rAbs from this group showed evidence of weak binding (supplementary file 2),
230	indicating that the rAb cloning efficiency for this group was not reduced and only the
231	two strongest binders were above the ELISA titration threshold. All but one of the
232	positive binding rAbs were IgM. Figure 4G shows the ELISA titration and Figure 4H
233	the derived endpoint titres, which we are using as a proxy of affinity. A more strongly
234	binding IgM rAb from E4 boost day 7, B5, and the only positive binding IgG1 rAb,
235	G6, are indicated on Figure 4H. The positive-binding rAbs from E4 prime day 17
236	show a higher affinity than those from prime day 7, consistent with affinity
237	maturation. Six of the seven positive-binding IgM rAbs from E4 boost day 7 show a
238	higher affinity than the two strongest binding IgM rAbs from E4 prime day 7. This is
239	consistent with pre-selection by the E3 prime immunization, and also considering the
240	higher proportion of rAbs with an anti-E4 O.D. > 0.1 , implies the GC B-cells
241	expressing these antibodies are memory derived. rAb affinities were generally low,
242	which might be expected of IgMs particularly in early GCs. We estimated the Kd of
243	rAbs B5 and G6 (an IgG1) as around 150nm and $1\mu m$ respectively (see Methods).
244	Other rAbs would be in the super-micromolar range. Figure 4I shows the cross
245	reactivity of rAbs with E3. Binding to E3 correlates with binding to E4, but because
246	of the generally low rAb affinities we suggest that the antibodies cannot discriminate
247	between similar epitopes. The higher affinity of E4 boost rAbs B5 and G6, and
248	binding to E3, suggest they may have genuine specificity for E3, thus consistent with

their derivation from anti-E3 memory. That rAb B5 is an IgM with only one VH (andone Vkappa) mutation, provides further support for the proposal of this study.

251

252 Discussion

253 The most variant protein we boosted with, E4, stimulated GCs with the highest 254 proportion of IgM+ cells and with the lowest levels of VH gene mutation, greater VH-255 gene diversity, and a slower, more specific, serum IgG response that resulted in 256 equivalent if not higher affinity, compared to the heterotypic E2 boost. This response 257 was higher than the primary response to E4. IgM rAbs cloned from E4 boost day 7 258 GC showed a higher affinity for E4 than those from E4 primed day 7 GC, implying 259 they were memory derived. This demonstrates that IgM memory cells with fewer 260 mutations, from 'lower' levels of the memory compartment, participate in secondary 261 responses to variant antigens, and further challenges the hypothesis that highly 262 mutated, class-switched cells elicited by homotypic antigen boosting are a 'mirror' of 263 the antibody memory compartment (Weiss & Rajewsky, 1990). The slower nature of 264 the E4 boost response also suggests a lower level of immediate differentiation of 265 memory cells into AFCs than seen with for example the homotypic or E2 response, 266 and is consistent with reduced numbers of high affinity class-switched memory cells 267 recognizing E4.

The serum antibody response to the closer variant, E2, was more rapid, more cross-reactive and evidenced some earlier affinity maturation. These observations are consistent with a response derived more from the 'higher' layers of the E3 specific memory compartment. The IgM+ cells induced by E2 boosting have more mutations than after E4 boosting, indicating they are memory derived. As there are higher proportions of these IgM+ GC B-cells, with fewer mutations relative to the homotypic

274	E3 boost this	nrovides fi	urther support	for the hypothesis	that IgM+ B-cells with
4/1	L_{2} 000st, this	provides re	under support	for the hypothesis	that ight D-cons with

275 fewer mutations furnish memory responses to variant antigens

276 Naïve B-cells may contribute to the IgM+ GC B-cells we observe after E4 277 boosting, although the higher affinities of the rAbs from this group suggest many are 278 memory derived. Also, the slightly higher median level of VH mutation and the 279 higher levels of IgM+ GC B-cells after E4 boosting (2x) compared to priming, 280 suggest IgM+memory B-cells are involved in the boost response consistent with the 281 well established presence of IgM+ memory cells with few or no mutations (Dogan et 282 al., 2009; Pape et al. 2011; Kaji et al., 2012) and the known lower activation threshold 283 of memory B-cells in response to antigen (Good et al., 2007 & 2009). 284 Whilst E3 specific memory cells may be expected to increase the anti-E3 titre 285 when stimulated by a cross-reactive E4 boost, the small but not significant effect we 286 observe (Figure 1D) is consistent with the lowest affinity, least mutated, E3-specific 287 memory cells being stimulated by an E4 boost. Antibodies from such cells may, 288 therefore not add much to the already high, affinity matured, anti-E3 titre induced by 289 E3 priming. The 14-fold higher anti-E4 titre at day 7 after boost (Figure 1F) versus 290 day 17 after prime (Figure 4A) also argues for a significant contribution from B-cell 291 memory.

The fusion-loop epitope in domain 2 of the dengue envelope protein is 100% conserved between strains and in humans, antibodies against this are prevalent in cross-reactive secondary responses (Lai et al, 2013, Chaudhury et al., 2017). The E2 boost response is consistent with this effect, especially considering the restricted clonality seen in VH sequences, but the low anti-E3 titre induced by E4 is not. A recent study (Chaudhury et al. 2017) showed that the mouse response to recombinant E-protein is predominantly focused on domain 3 of the protein, and so cross reactivity

299	with the fusion loop epitope (domain 2) should be less dominant. While E2 and E4 are
300	68% and 63% overall identical to E3, in domain 3, a focus of mouse antibodies, they
301	are 62% and 51% identical, a bigger difference in differences, helping explain the
302	responses we observe here.
303	
304	
305	
306	
307	
308	
309	
310	
311	
312	
313	
314	
315	
316	
317	
318	
319	
320	
321	
322	
323	

- 326 Materials and Methods
- 327 Animals, immunisations and antigens
- 328 Female 8-11-week old BALB/c mice were purchased from Charles River, U.K.
- 329 Primary immunisations were intra-peritoneal (IP) with 25µg recombinant Dengue
- envelope protein (Biorbyt) precipitated in alum with $2x10^7$ heat-killed *B.pertussis*.
- 331 Secondary immunisations were IP with 25µg recombinant Dengue envelope protein
- 332 (Biorbyt) dissolved in phosphate-buffered saline (PBS). At designated time points
- 333 mice were anaesthetized and bled for collection of serum and then humanely
- 334 sacrificed for collection of spleen cells. Dengue envelope (E) proteins were C-
- terminal His-tagged and expressed in *E-coli* prior to purification. Dengue proteins
- 336 were tested for endotoxin by LAL assay (Fisher Scientific) and contained it at a low
- level: E2, 5.4EU/µg; E3, 2.5EU/µg; E4, 3.1EU/µg. Endotoxin in this range does not
- 338 give a detectable physiological response in mice (Copeland et al., 2005).
- 339
- 340 ELISA for serum and rAbs
- ELISA plates (Nunc Maxisorp) were coated overnight at 4^oC with 1µg/ml protein in
- 342 0.1M bicarbonate buffer pH 9.3. Plates were washed three times in PBS/0.05%
- 343 Tween-20 (Sigma) (PBST) and blocked for 30mins at room temperature with
- 344 PBST/2% bovine serum albumin (BSA, Sigma). Plates were then washed three times
- and incubated with serum dilutions in PBST/1.0% BSA for two hours at room
- 346 temperature. After three washes plates were incubated with alkaline-phosphatase
- 347 conjugated goat anti-mouse IgG (Sigma) for one hour at room-temperature, washed
- 348 three times and developed with pNPP substrate (Sigma) for one hour. Absorbance

349	was measured at 405nm. For the initial rAb screen, rAbs were incubated at $100 \mu gml^{-1}$
350	in PBST/1.0% BSA for 2 hours at room temperature on plates coated with E4 and
351	blocked as above, and subsequently treated as above except with use of anti-human
352	IgG second layer (Sigma). Background binding to plates was determined using
353	binding of non-specific polyclonal human IgG at 100µgml ⁻¹ , because the rAbs were
354	expressed as chimeric constructs with human constant regions, and this was
355	subtracted from the rAb O.D. Positive binding rAbs were deemed to be those with
356	O.D. > 0.1 that could be subject to an ELISA endpoint titration. For the ELISA
357	titration and endpoint analysis, doubling dilutions of positive binding rAbs, and
358	polyclonal IgG background subtraction control, were used starting at $100 \mu gml^{-1}$.
359	Endpoint titre was set at $O.D. = 0.1$ and calculated using interpolation on Graphpad
360	Prism. The assay was repeated using E3 coated plates to determine the rAB cross
361	reactivity. The affinity (Kd) of rAbs B5 and G6 (the two strongest binding rAbs) was
362	estimated from the inflection point of the ELISA titration curve as indicating 50%
363	maximal binding, and on the assumption that at these higher antibody concentrations
364	binding of rAB to immobilized antigen will have a minor effect on concentration of
365	unbound rAb. We estimated the B5 inflection point to be at approximately 25ugml ⁻¹
366	(= approx. 150nM) and the G6 inflection point to be just above 100 ugml ⁻¹ (= approx.
367	1uM)

369 Competition ELISA

ELISA plates were coated as above with target protein, then washed, blocked and
washed as above except the blocking was done at 37^oC for one hour. Mouse serum
samples were diluted in PBST/1% BSA to twice the concentration of the maximum
dilution that gave an absorbance at 405nm =1.0 in ELISA to the target protein. Serial

374	six-fold dilutions of competitor protein were made in PBST/1% BSA, such that the
375	highest concentration of competitor was $2.4\mu g$ in $30\mu l$. $30\mu l$ of diluted serum was
376	mixed with $30\mu l$ of each competitor protein dilution and incubated in a polypropylene
377	96-well plate at 37° C for 1 hour. Serum/competing antigen mixture (50µl) was then
378	added to each well of the target antigen coated plate and incubated at 37° C for one
379	hour. Plates were washed as above and then 50µl of alkaline-phosphatase conjugated
380	anti-mouse IgG (Sigma) was added to each well followed by incubation at 37° C for
381	one hour. Plates were washed as above and incubated with $75\mu l$ per well of p-
382	nitrophenyl phosphate substrate (Sigma) for one hour at room temperature.
383	Absorbance was measured at 405nm. All individual serum dilutions were also reacted
384	in the absence of competitor, against BSA, following the same incubation protocol.
385	These background values were subtracted from the competition ELISA values
386	obtained above. The readings were then normalized so that the samples with the
387	maximum competitor dilution gave a value of 1.0

389 Urea Avidity ELISA

Adapted from Puschnik et al., 2013. Assay plates were coated with antigen and
blocked as for the ELISA protocol. 1/200 dilutions of serum in PBST/1% BSA were
incubated on plates for 2 hours at room temperature. Wells were washed once with
PBST, incubated for 10 minutes at room temperature with PBST or PBST/7M Urea,
washed a further two times with PBST and then treated as for standard ELISA. The
avidity index was calculated by dividing readings from 7M Urea treatment by
readings from PBST-only treatment, after subtraction of background absorbance.

398 *Flow cytometry*

- 399 Whole spleen cell-suspensions were red-cell depleted with Pharm-Lyse (BD
- 400 Biosciences) and incubated with anti-CD16/32 monoclonal antibody (Fc-block, BD
- 401 Biosciences) for 15minutes at 4^oC. Cells were then stained with APC anti-B220,
- 402 BV421 anti-CD38, PE anti-CD95/Fas (all BD) and FITC anti-IgM (eBioscience) for
- 403 45 minutes at 4^oC. After washing, cells were re-suspended in PBS 5% FCS (Gibco)
- 404 and analysed or single-cell sorted on a FACS Aria II (BD).
- 405

406 *GC B-cell antibody sequencing, cloning, expression and purification*

- 407 Single GC B-cells were sorted into half a 96 well PCR plate (less 3 control wells)
- 408 containing10µl of chilled 10mM Tris pH 8.0, 1U/µl RNAsin (Promega) and placed on
- 409 dry ice then at -80°C. One-Step RT-PCR (Qiagen) was performed according to
- 410 manufacturers instructions, by adding 15µl RT-PCR master mix, using first-round
- 411 primer sets described in Tiller et al., 2009, with heavy-chain and kappa-chain primers,
- 412 for 50 cycles, annealing at 53.6°C. Heavy-chain second-round PCRs were performed
- 413 using 2µl first-round product and the nested/semi-nested primer sets from Tiller et al.,
- 414 2009, with Hot Star Taq polymerase (Qiagen) for 50 cycles annealing at 56° C.
- 415 Second round PCR product (4µl) was analysed on a 1.2% agarose gel. Successful
- 416 PCRs were then Sanger sequenced. For this study the sequencing primer was the pan
- 417 VH primer 5'MsVHE (Tiller et al., 2009) which leaves part of the 5' of FR1
- 418 unsequenced. For this reason the FR1 sequence was not included in the analysis. VH
- 419 sequence identification and SHM analysis was done using the IMGT V-Quest online
- 420 platform. VH sequences are in Supplementary File 1. Further cloning, construction
- 421 and expression of antibodies as chimeric IgG1 rAbs was done according to Tiller et
- 422 al., 2009. Briefly, second round PCRs of in-frame VH and VK sequences were

423	repeated with V-gene specific primers that included a restriction site for sub cloning
424	(Tiller et al., 2009). These PCR products were purified (Qiagen), restriction digested,
425	purified (Qiagen) and ligated (instant sticky-end ligase, NEB) into the appropriate
426	expression vector containing either human IgG1 or Kappa constant regions, prior to
427	transformation into E. Coli NEB5-alpha (NEB). Expression constructs in transformed
428	colonies were verified by sequence analysis prior to preparation of plasmid mini-
429	preps (Qiagen). 293A cells were split and grown to 80% confluence in DMEM with
430	ultra-low IgG FCS (PAN Biotech) in 150mm plates prior to replacement of medium
431	with 20ml Panserin 293A serum free medium. 15ug each of matched VH and VKappa
432	constructs were added to 2ml saline with 90ug PEI, briefly vortexed and rested for
433	10mins. Transfection solution was added to plates and mixed gently. After 3 days
434	medium was collected, centrifuged at 800g for 10mins to clear debris, and further
435	medium added. After a further 3 days medium was collected, cleared of debris as
436	before and pooled. 100ul protein-G sepharose (GE Healthcare) was added to
437	supernatants and incubated with rocking overnight at 4 °C. Protein G sepharose was
438	collected by centrifugation at 800g for 10 mins and transferred in PBS to a PBS
439	equilibrated spin column (Bio-Rad). After 3 rounds of washing with 800ul of PBS,
440	rAbs were eluted in two 200ul passes of 0.1M Glycine (pH2.9) into a tube with 40ul
441	of 1M Tris pH 8.0, 0.5% Sodium Azide. Antibody concentrations were determined by
442	O.D. on a Nanodrop instrument (Thermo) and corrected for an extinction co-efficient
443	of 1.36.

445 *T-cell proliferation assay*

446 Spleens were harvested from female BALB/c AnCrl mice 39 days after challenge.

447 Splenocytes $(5x10^5)$ were cultured in triplicate with the indicated concentration of E-

448	protein in X-VIVO 15 medium. Cells were cultured for 96 hours and 0.5 μ Ci of [³ H]
449	thymidine was added to wells for 16 hours before measurement with a 1450
450	MicroBeta counter (Wallac).
451	
452	Statistics
453	For statistical analysis sample sizes were chosen to address group size reductions that
454	observe the ARRIVE guidelines. Cages of three mice were randomly allocated to
455	treatment groups. These group treatments were independently biologically replicated
456	to give a sample size of 6. Where statistical analysis was applied, data points were
457	analysed with Levene's test for equality of variance and where violated they were
458	subject to a two-tailed Students t-test for unequal variance, otherwise the two-tailed t-
459	test for equal variance.
460	
461	
462	
463	
464	Acknowledgements
465	We are especially grateful to the late Michael Neuberger for critical discussion and
466	early comment on the project. Thanks to Patrick Wilson and Christian Busse for
467	advice on single cell antibody PCR, Per Klasse for advice on antibody avidity assays,
468	James Cresswell for advice on statistics, Kai Toellner for discussions and Jamie
469	Gilman for extra FACS work.
470	
471	
472	References

473	Aiba, Y. et al. (2010). Preferential localization of IgG memory B cells adjacent to contracted germinal
474	centers. Proceedings of the National Academy of Sciences of the United States of America, 107(27),
475	12192–7.
476	
477	Alexander, M. R., Ringe, R., Sanders, R. W., Voss, J. E., Moore, J. P., & Klasse, J. (2015). What Do
478	Chaotrope-Based Avidity Assays for Antibodies to HIV-1 Envelope Glycoproteins Measure ?, J. Virol
479	<i>89</i> (11), 5981–5995.
480 481 482	Chaudhury, S., Gromowski, G. D., Ripoll, D. R., Khavrutskii, I. V, Desai, V., & Wallqvist, A. (2017). Dengue virus antibody database : Systematically linking serotype-specificity with epitope mapping in dengue virus, <i>PLoS Neg Trop Dis</i> , 11,1–17.
483	
484	Copeland, S., Warren, H. S., Lowry, S. F., Calvano, S. E., & Remick, D. (2005). Acute inflammatory
485	response to endotoxin in mice and humans. Clinical and Diagnostic Laboratory Immunology, 12(1),
486	60–67.
487	
488	Corti, D. et al. (2011). A neutralizing antibody selected from plasma cells that binds to group 1 and
489	group 2 influenza A hemagglutinins. Science (New York, N.Y.), 333(6044), 850-6.
490	
491	Dogan, I., Bertocci, B., Vilmont, V., Delbos, F., Mégret, J., Storck, S., Reynaud, C-A. & Weill, JC.
492	(2009). Multiple layers of B cell memory with different effector functions. Nature Immunology, 10(12),
493	1292–1299
494	
495	Gitlin, A. D., Boehmer, L. Von, Gazumyan, A., Shulman, Z., Oliveira, T. Y. & Nussenzweig, M. C.,
496	(2016). Independent Roles of Switching and Hypermutation in the Development and Persistence of B
497	Lymphocyte Memory Article Independent Roles of Switching and Hypermutation in the Development
498	and Persistence of B Lymphocyte Memory, 1–13.
499	
500	Good, K. L., Avery, D. T., & Tangye, S. G. (2009). Resting Human Memory B Cells Are Intrinsically
501	Programmed for Enhanced Survival and Responsiveness to Diverse Stimuli Compared to Naive B
502	Cells. The Journal of Immunology, 182(2), 890–901.
503	
504	Good, K. L., & Tangye, S. G. (2007). Decreased expression of Kruppel-like factors in memory B cells
505	induces the rapid response typical of secondary antibody responses. Proceedings of the National
506	Academy of Sciences of the United States of America, 104(33), 13420–13425.
507	
508	Hebeis, B. J., Klenovsek, K., Rohwer, P., Ritter, U., Schneider, A., Mach, M., & Winkler, T. H. (2004).
509	Activation of virus-specific memory B cells in the absence of T cell help. The Journal of Experimental
510	Medicine, 199(4), 593-602. http://doi.org/10.1084/jem.20030091
511	

512	Herzenberg, L. A, Black, S. J., & Tokuhisa, T. (1980). Memory B cells at successive stages of
513	differentiation. Affinity maturation and the role of IgD receptors. The Journal of Experimental
514	Medicine, 151(5), 1071–87.
515	
516	Kaji, T. et al. (2012). Distinct cellular pathways select germline-encoded and somatically mutated
517	antibodies into immunological memory. The Journal of Experimental Medicine, 209(11), 2079-97.
518	
519	Kometani, K. et al. (2013). Repression of the Transcription Factor Bach2 Contributes to Predisposition
520	of IgG1 Memory B Cells toward Plasma Cell Differentiation. Immunity, 39(1), 136-147.
521	
522	Kuraoka, M., Schmidt, A. G., Nojima, T., Feng, F., Watanabe, A., Kitamura, D., Harrison, S. C.,
523	Kepler, T. B. & Kelsoe, G. (2016). Complex Antigens Drive Permissive Clonal Selection in Germinal
524	Centers. Immunity, 1–11.
525	
526	Lai, C. Y., Williams, K. L., Wu, Y. C., Knight, S., Balmaseda, A., Harris, E., & Wang, W. K. (2013).
527	Analysis of Cross-Reactive Antibodies Recognizing the Fusion Loop of Envelope Protein and
528	Correlation with Neutralizing Antibody Titers in Nicaraguan Dengue Cases. PLoS Neglected Tropical
529	<i>Diseases</i> , 7(9), 1–11.
530	
531	MacLennan, I. C., Gulbranson-Judge, A., Toellner, K. M., Casamayor-Palleja, M., Chan, E., Sze, D.
532	M., Luther, S. A. & Orbea, H. A. (1997). The changing preference of T and B cells for partners as T-
533	dependent antibody responses develop. Immunological Reviews, 156, 53-66.
534	
535	
536	McHeyzer-Williams, L. J., Milpied, P. J., Okitsu, S. L., & McHeyzer-Williams, M. G. (2015). Class-
537	switched memory B cells remodel BCRs within secondary germinal centers. Nature Immunology,
538	16(3), 296–305.
539	
540	Nabel, G. J., & Fauci, A. S. (2010). Induction of unnatural immunity: prospects for a broadly protective
541	universal influenza vaccine. Nature Medicine, 16(12), 1389–1391.
542	
543	Pape, K. A., Taylor, J. J., Maul, R. W., Gearhart, P. J., & Jenkins, M. K. (2011). Different B cell
544	populations mediate early and late memory during an endogenous immune response. Science (New
545	York, N.Y.), 331(6021), 1203–7.
546 547	Puschnik, A., Lau, L., Cromwell, E. A., Balmaseda, A., Zompi, S., & Harris, E. (2013). Correlation between Dengue-Specific Neutralizing Antibodies and Serum Avidity in Primary and Secondary

548 Dengue Virus 3 Natural Infections in Humans, 7(6), 1–8.

550	Smith, K. G. C., Light, A., Nossal, G. J. V, & Tarlinton, D. M. (1997). The extent of affinity
551	maturation differs between the memory and antibody-forming cell compartments in the primary
552	immune response. EMBO Journal, 16(11), 2996–3006.
553	
554	Tiller, T., Busse, C. E., & Wardemann, H. (2009). Cloning and expression of murine Ig genes from
555	single B cells. Journal of Immunological Methods, 350(1-2), 183-193.
556	
557	Toellner, K. M., Gulbranson-Judge, a, Taylor, D. R., Sze, D. M., & MacLennan, I. C. (1996).
558	Immunoglobulin switch transcript production in vivo related to the site and time of antigen-specific B
559 560	cell activation. The Journal of Experimental Medicine, 183(5), 2303–2312.
561	Victora, G. D. (2014). SnapShot: The Germinal Center Reaction. Cell, 159(3), 700-700.
562	
563	Wang, S., Mata-Fink, J., Kriegsman, B., Hanson, M., Irvine, D. J., Eisen, H. N., Burton, D. R., Dane
564	Wittrup, K., Kardar, M. & Chakraborty, A. K. (2015). Manipulating the selection forces during affinity
565	maturation to generate cross-reactive HIV antibodies. Cell, 160(4), 785-797.
566	
567	Weiss, U., & Rajewsky, K. (1990). The repertoire of somatic antibody mutants accumulating in the
568	memory compartment after primary immunization is restricted through affinity maturation and mirrors
569	that expressed in the secondary response. J Exp Med, 172(6), 1681–1689.
570	Wu, X. et al (2010). Rational design of envelope identifies broadly neutralizing human monoclonal
571	antibodies to HIV-1. Science (New York, N.Y.), 329(5993), 856-861
572	
573	Figure Legends
574	Figure 1
575	Serum antibody responses after boosting with Dengue envelope protein variants
576	A, Cross-reactivity of E3 primed serum IgG with E-protein variants. Red bar shows
577	mean value. Serum used was from mice mock-boosted with PBS 37 days after E3
578	priming and obtained 7 days later; E3, Dengue-3 envelope protein; E2, Dengue-2
579	envelope protein; E4, Dengue-4 envelope protein; % identity, sequence identity
580	between E3 envelope protein and respective protein; end-point titre (EPT) values
581	plotted are log2 of 1/(end point dilution x 100), each unit increase represents a
582	doubling of titre. B, E3 primed mouse serum cross-reactivity with E2 versus E4. C,

583	Control. Anti-PR8 HA serum IgG titre of E3 day 7 boost serum. D, Anti-E3 serum
584	IgG titre after boosting with respective proteins. Red bar shows mean value. n=6 from
585	two independent experiments for each group except boost only, n=3; first set of data
586	points reproduced from panel A for comparison; numbers 3,2 and 4 refer to serotype
587	of Dengue-envelope protein used for boost; BO, adjuvant primed, E3 boosted,
588	analysed 7 days later; Day, days after boosting. p-values calculated using two-tailed
589	Students t-test after testing for equality of variance. E, Anti-E2 serum IgG titre after
590	E2 boost. Red bar shows mean value. n=6 from two independent experiments for each
591	group; labeling and statistics as for panel D. F, Anti-E4 serum IgG titre after E4
592	boost. Red bar shows mean value. n=6 from two independent experiments for each
593	group; labeling and statistics as for panel D.
594	
595	
596	
596	Figure 2
596 597	Figure 2 GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-
596 597 598	
596 597 598 599	GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-
596 597 598 599 600	GC B-cell levels, isotypes, VH mutation and clonality after boosting with E- protein variants
596 597 598 599 600 601	GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-protein variantsA, FACS gating strategy used to identify and sort GC B-cells and determine isotype.
596 597 598 599 600 601 602	 GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-protein variants A, FACS gating strategy used to identify and sort GC B-cells and determine isotype. B, GC B-cell levels after E-variant boosting, expressed as % total lymphocytes; Red
596 597 598 599 600 601 602 603	 GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-protein variants A, FACS gating strategy used to identify and sort GC B-cells and determine isotype. B, GC B-cell levels after E-variant boosting, expressed as % total lymphocytes; Red bar shows mean value; numbers 3,2 and 4 refer to serotype of Dengue-envelope
596 597 598 599 600 601 602 603 604	 GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-protein variants A, FACS gating strategy used to identify and sort GC B-cells and determine isotype. B, GC B-cell levels after E-variant boosting, expressed as % total lymphocytes; Red bar shows mean value; numbers 3,2 and 4 refer to serotype of Dengue-envelope protein used for boost; BO, boost only, adjuvant primed, E3 boosted day 37, analysed
 596 597 598 599 600 601 602 603 604 605 	 GC B-cell levels, isotypes, VH mutation and clonality after boosting with E-protein variants A, FACS gating strategy used to identify and sort GC B-cells and determine isotype. B, GC B-cell levels after E-variant boosting, expressed as % total lymphocytes; Red bar shows mean value; numbers 3,2 and 4 refer to serotype of Dengue-envelope protein used for boost; BO, boost only, adjuvant primed, E3 boosted day 37, analysed 7 days later; Day, days after boosting. C, % IgM+ GC B-cells, of total GC B-cells,

608	between E3 and other variants; p-values calculated using two-tailed Students t-test
609	after testing for equality of variance. D , Levels of IgM+ and IgM- GC B-cells in
610	individual boosted mice. E, Number of mutations detected in VH of all isotypes of
611	GC B-cells, from n=3 mice except E4 boost day 17, n=2. Red bar is median value.
612	VH region sequenced is CDR1 to FR3; labeling as panel B. F, Number of mutations
613	detected in VH of IgM+ GC B-cells, from n=3 mice except E4 boost day 17, n=2. Red
614	bar is median value. G, Clonality of sequences from single GC B-cells 7 days after
615	boosting; colours indicate different mice in each group; thin sectors, unique
616	sequences; thicker sectors two or three clonal sequences according to sector size;
617	black dots, VH 14-3 or VH14-4 sequences; numbers in circles, number of sequences
618	from that mouse; Identical VH clones had the same: V-gene, CDR3 length, J-gene, D-
619	gene if assigned, D-reading frame, three or fewer differences in CDR3 amino acid
620	sequence.
621	
622	
623	Figure 3
624	Relative serum affinity and avidity after boosting with E-protein variants, and
625	T-cell re-stimulation
626	A, Relative avidity of E2 boost serum for E2, measured by resistance to 7M Urea.
627	Red bar shows mean value; Day, days after E2 boosting; Day 0 sample was from mice
628	mock-boosted with PBS 37 days after priming with E3 and obtained 7 days later. B ,
629	Relative avidity of E4 boost serum for E4, measured by resistance to 7M Urea.
630	Labeling as for panel A; Day 0 sample was from mice mock-boosted with PBS 37
(04	
631	days after priming with E3 and obtained 7 days later C, Relative affinity of E2

632 boosted serum for E2. Inhibition by lower concentration of competitor implies higher

633	affinity of serum for competitor. Maximum competitor amount $2\mu g$ in 50µl followed
634	by six-fold dilutions of competitor; timepoint of samples and numbers of individuals
635	in group indicated. Open circles, E2 boost day 17 serum competed with irrelevant
636	His-tagged protein measured on E2 target D , Relative affinity of E4 boosted serum for
637	E4. Labeling as for panel A. E, T-cell proliferation measured by ³ H incorporation 96
638	hours after re-stimulation in vitro with indicated amounts of E-protein variants; error
639	bars indicate standard error of the mean; n=4 or 5 from two independent experiments
640	(see source data). Closed symbols, E3 primed mouse splenocytes re-stimulated with
641	indicated E-protein variant. Open symbols, adjuvant primed mouse splenocytes re-
642	stimulated with indicated E-protein variant.
643	
644	Figure 3 Source Data File
645	Source data for Figure 3 panels C, D and E
646	
647	
047	
648	Figure 4
	Figure 4 Primary response to E4 and rAb binding.
648	
648 649	Primary response to E4 and rAb binding.
648 649 650	Primary response to E4 and rAb binding.A, anti-E4 IgG titre after E4 priming; Red bars show mean titres; A, serum from
648 649 650 651	Primary response to E4 and rAb binding.A, anti-E4 IgG titre after E4 priming; Red bars show mean titres; A, serum from adjuvant-only primed mice at day 45; d7, 7 days after E4 priming; d17, 17 days after
648 649 650 651 652	 Primary response to E4 and rAb binding. A, anti-E4 IgG titre after E4 priming; Red bars show mean titres; A, serum from adjuvant-only primed mice at day 45; d7, 7 days after E4 priming; d17, 17 days after E4 priming; EPT, end-point titre calculated as for Figure 1. B, GC B-cell levels after
648 649 650 651 652 653	 Primary response to E4 and rAb binding. A, anti-E4 IgG titre after E4 priming; Red bars show mean titres; A, serum from adjuvant-only primed mice at day 45; d7, 7 days after E4 priming; d17, 17 days after E4 priming; EPT, end-point titre calculated as for Figure 1. B, GC B-cell levels after E4 priming; Red bars indicate mean levels; A, cells from adjuvant-only primed mice 7
 648 649 650 651 652 653 654 	 Primary response to E4 and rAb binding. A, anti-E4 IgG titre after E4 priming; Red bars show mean titres; A, serum from adjuvant-only primed mice at day 45; d7, 7 days after E4 priming; d17, 17 days after E4 priming; EPT, end-point titre calculated as for Figure 1. B, GC B-cell levels after E4 priming; Red bars indicate mean levels; A, cells from adjuvant-only primed mice 7 days after priming; other x-axis labels as for panel A. C, %IgM+ GC B-cells after E4
 648 649 650 651 652 653 654 655 	 Primary response to E4 and rAb binding. A, anti-E4 IgG titre after E4 priming; Red bars show mean titres; A, serum from adjuvant-only primed mice at day 45; d7, 7 days after E4 priming; d17, 17 days after E4 priming; EPT, end-point titre calculated as for Figure 1. B, GC B-cell levels after E4 priming; Red bars indicate mean levels; A, cells from adjuvant-only primed mice 7 days after priming; other x-axis labels as for panel A. C, %IgM+ GC B-cells after E4 priming; Red bars show mean values; x-axis labels as for panel A. D, Numbers of VH

658	Numbers of VH mutations in IgM+ GC B-cells after E4 priming and boosting; Red
659	bars show median values, from n=3 mice (d7), n=2 mice (d17) and n=3 mice E4Bd7;
660	x-axis labels as for panel A except E4Bd7, 7 days after E4 boosting which was 38
661	days after E3 priming. F, ELISA screen of binding of all 48 rAbs. rAbs incubated at
662	100µgml ⁻¹ . Number of rAbs in each group indicated. P7, 7days after E4 prime; P17,
663	17 days after E4 prime; B7, 7 days after E4 boost. As the antibodies were cloned as
664	chimeric human IgG1 antibodies the background from non-specific human polyclonal
665	IgG binding has been subtracted from O.D. readings. Values in supplementary file 2.
666	G , ELISA titration of rAbs that showed binding $O.D. > 0.1$ in panel F. All but one
667	were IgM. IgG1 rAb indicated. Background subtraction as for panel F, using
668	appropriate dilution of polyclonal IgG. H, Anti-E4 end point titre of positive-binding
669	rAbs, used as a proxy of rAB affinity. X-axis labels as for panel F. End-point titre
670	values plotted are log2 of 1/end point dilution (undiluted = $100\mu gml^{-1}$). Red bars show
671	median values (excluding any IgG1 data). Stronger binding IgM rAb 'B5', and IgG1
672	rAb 'G6' EPT readings indicated. I, anti-E3 versus anti-E4 endpoint titres. Star, E4
673	prime day 7 rAbs; Square, E4 prime day 17 rAbs; circle, E4 boost day 7 rAbs. IgG1
674	EPT reading indicated. End-point titre values plotted are log2 of 1/end point dilution
675	(undiluted = $100 \mu gml^{-1}$).

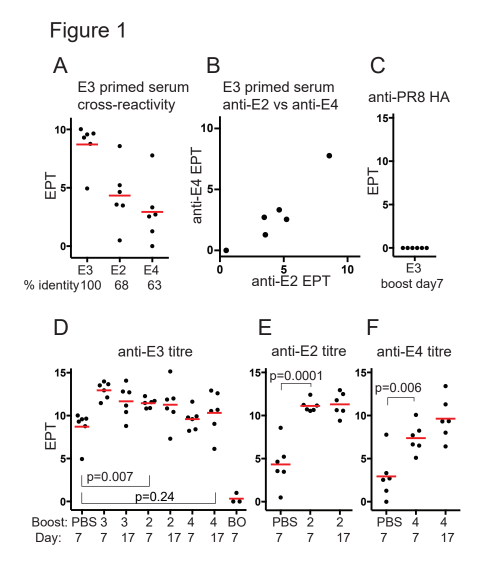
677 Supplementary File 1

678 GC B-cell VH Sequences

VH sequences from single sorted GC B-cells. Sequences are grouped into treatment
groups, and within this, arranged in blocks for sequences from individual mice. Raw
sequences were analysed by IMGT V-Quest. Due to cloning and sequencing primers
being at start of FR1 region, this region not included in mutation analysis. CDR1T,

- total mutations in CDR1; CDR1S, silent mutations in CDR1; CDR1R, replacement
- mutations in CDR1; likewise for FR2, CDR2 and FR3 regions; Tot Mut, total
- 685 mutations in CDR1 to FR3 regions.

- 687 Supplementary File 2
- 688 Data on recombinant antibodies



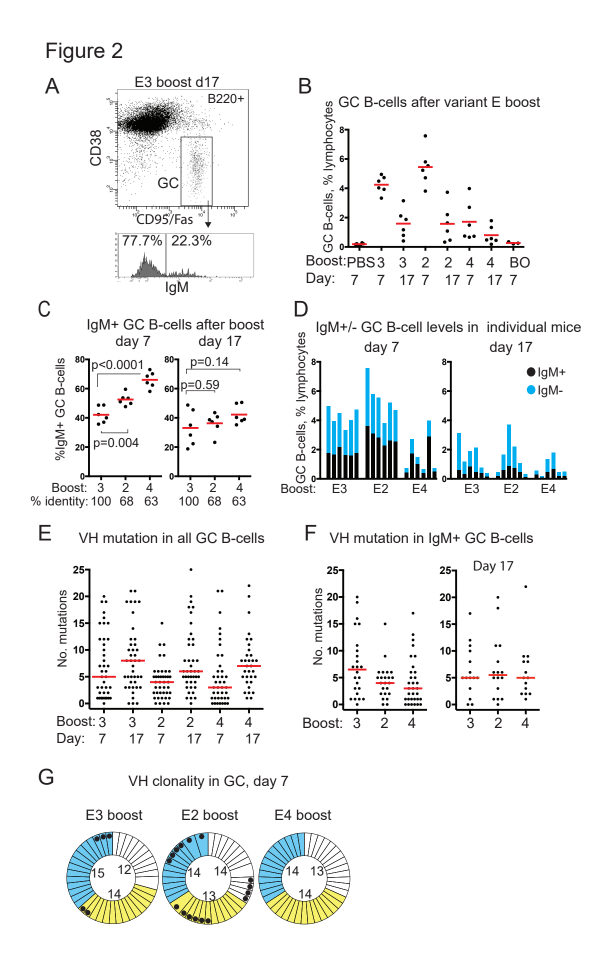


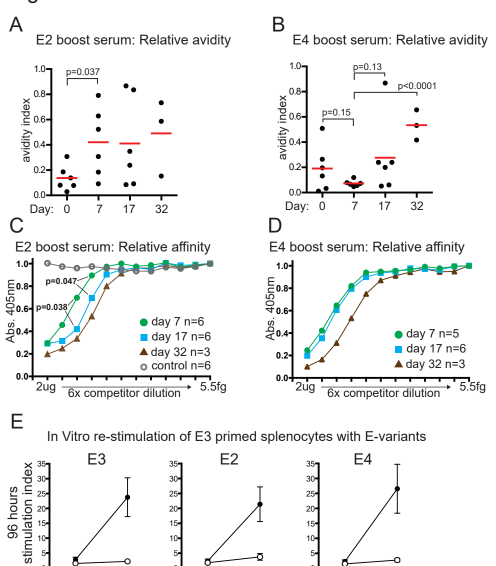
Figure 3

0

ug/ml

0.1

1.0



0

0.1

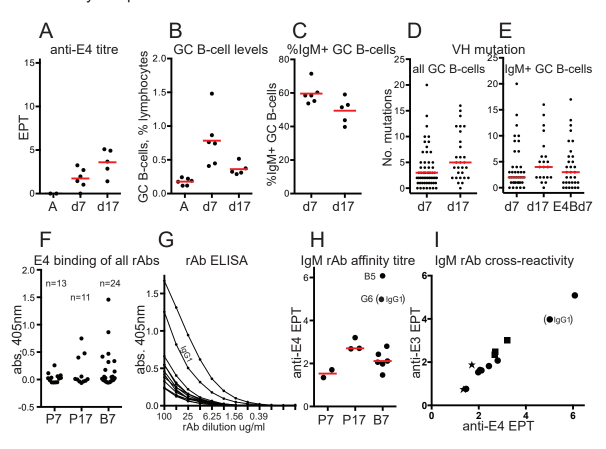
1.0

0-

0.1

1.0

Figure 4



Primary Response to E4