The B Cell Receptor Itself Can Activate Complement to Provide the Complement Receptor 1/2 Ligand Required to Enhance B Cell Immune Responses In Vivo

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Abstract

B cells express complement receptors (CRs) that bind activated fragments of C3 and C4. Immunized CR knockout (KO) mice have lower antibody titers and smaller germinal centers (GCs), demonstrating the importance of CR signals for the humoral immune response. CR ligands were thought to be generated via complement fixation mediated by preexisting “natural” IgM or early Ab from inefficiently activated B cells. This concept was recently challenged by a transgenic (Tg) mouse model that lacks circulating antibody but still retains membrane IgM (mlgM) and mounts normal immune responses. To test whether CR ligands could be generated by the B cell receptor (BCR) itself, we generated similar mice carrying a mutated mlgM that was defective in C1q binding. We found that B cells from such mutant mice do not deposit C3 on B cells upon BCR ligation, in contrast to B cells from mlgM mice. This has implications for the immune response: the mutant mice have smaller GCs than mlgM mice, and they are particularly deficient in the maintenance of the GC response. These results demonstrate a new BCR-dependent pathway that is sufficient and perhaps necessary to provide a CR1/2 ligand that promotes efficient B cell activation.

Key words: C1q • IgM • germinal center • CD19 • CD35

Introduction

It has become clear that various signals from the innate immune system activate the adaptive immune system. Specifically, signals generated by the complement system can play an important role in B cell immune responses. B cells express complement receptors (CR1 [CD35] and CR2 [CD21]), which bind fragments of C3 and C4 (1–4) that can be deposited on Ags upon complement system activation. Mice lacking C3, C4, or CR1/2 had reduced Ab titers in serum and reduced size and number of germinal centers (GCs), particularly when the immunogenic stimulus was relatively weak (1–8). CR1/2 is widely expressed, notably on B cells and follicular dendritic cells (FDCs). Chimeric studies showed CR1/2 needs to be expressed on the B cell per se for optimal immune responses (6, 9). CR1/2 on B cells also enhances their entrance and survival in GCs, as shown by comparison of CR1/2 deficient and wild-type B cells in adoptive transfer experiments (10), thus implicating CR1/2 signaling in GC maintenance and presumably in memory B cell development.

Signaling via CR1/2 in B cells requires the formation of a complex with CD19/TAPA-1 (11–14), which transduces the intracellular signals. Coligation of this receptor complex with the B cell receptor (BCR) was shown to enhance BCR-mediated signaling and reduce the amount of Ag required for B cell activation (15–17). Moreover, coligation of BCR and CR1/2 prolongs BCR signaling via lipid rafts and enhances presentation of BCR bound Ag by class II MHC (18–24). These results provide a biochemical basis for the role of CR1/2 in B cell immune responses in vivo.

The fact that C3, C4, and C1q deficient mice all have defects in primary B cell responses suggests that complement fixation during the initial steps of an immune response is required to generate the CR1/2 ligand (25–29). In particular, the requirement for C4 and C1q implicates the classical pathway, which is primarily if not exclusively activated by immune complexes in vivo. Moreover, factor B–deficient mice (30), which are defective in the alternative pathway, appear to have normal immune responses.
It has been assumed that the requirement for early complement fixation to provide CR1/2 ligation during a physiologic B cell immune response was met by either preexisting (“natural”) IgM Ab or very early Ab that was secreted from a few inefficiently activated B cells (6, 31, 32). These preexisting Abs are thought to form immune complexes (ICs) with Ag, which could in principle activate complement via the classical pathway, leading to C3d deposition on ICs. A priori, a problem for this model is that it would require natural Ab with affinity and concentration high enough to form immune complexes. It seems unlikely this would be generally available for all important Ags. Additionally any ICs formed with natural Ab would be subjected to very efficient clearance mechanism via CRs on phagocytes, B cells, and red blood cells, which will at the least attenuate their ability to stimulate an Ag-specific B cell.

We have generated a Tg mouse model that has provided an additional challenge to this classical model (33). This mouse expresses a mutant Vh IgM Tg that lacks the secreted exon of IgM, and thus lacks circulating Ab. Surface Ig is expressed normally, and B cell development is rescued when the Tg is crossed on the Jh knockout background (JhD). The Tg carries a Vh186.2 VDJ rearrangement that confers NP specificity when paired with λ light chain. Though these mice lack detectable Ab, they had normal immune responses to NP-human serum albumin (NP-HSA) and NP-chicken gamma globulin (NP-CGG), with normal numbers and sizes of GCs and normal kinetics (33). This was unexpected if soluble preformed Abs were important to create ICs that in turn coligate CR1/2 and the BCR. To reconcile our findings with the observations that CR1/2 coligation is critical for optimal GC formation, we hypothesized that membrane IgM fixation of complement is both sufficient and necessary to provide local C3d deposition on mlgM and/or IgM-bound Ag, which in turn can coligate the CR1/2/CD19/TAPA-1 complex.

We have now tested this hypothesis by generating a mouse model similar to the mlgM mice that has an additional mutation in the CH3 domain which inhibits C1q binding to the BCR. We report here that these latter mice, which have no circulating Ig and are deficient in complement activation via the classical pathway by mlgM, show decreased responses to immunizations with NP-HSA and NP-CGG, in contrast to mlgM mice. By demonstrating the phenotype of mice that cannot activate complement via the surface BCR, we show the importance of this novel pathway for providing a ligand that can cocroslink CR1/2 and the BCR. Since the previously reported mlgM mice make normal responses and have this pathway intact, yet lack soluble Ab to perform these functions, it follows that the fixation of complement by mlgM must be sufficient, and possibly necessary, to start optimal immune responses.

Materials and Methods

Construct and Tg Mouse Generation. The Vh186.2-mlgM Tg mice have been described (33, 34). They lack the secreted exon of the IgM region and thus have 1,000-fold less than normal standing IgM levels. Their B cells neither secrete nor shed IgM in response to LPS in vitro and in spite of having 4% of B cells specific for the NP hapten, have undetectable levels of serum anti-NP. To generate mice with disabled C1q binding but the same properties as Vh186.2-mlgM mice we altered the Vh186.2-mlgM construct using a restriction fragment of the IgM constant region containing a point mutation encoding a Pro-Ser exchange at residue 436 [Vh186.2-mlgM(P436S)]. This mutation was shown to lower the Ka for C1 over 50-fold and to be >200-fold less active in hemolysis assays (35, 36). The altered construct was tested for expression by transfection of CH1, a λ/IgM-expressing cell line (37). CH1 transfectants expressed surface IgM anti-NP. The tested construct was then used to generate Tg mice by microinjection of purified DNA. Tg positive founders were identified by PCR and bred directly to the CB.17 mice carrying a homozygous deletion at the Jh locus (JhD/JhD.CB17) strain to eliminate endogenous H chain production. F1 generation mice that matched the previously generated mlgM Tg mice (33) in terms of mlgM expression and nitro-iodo-phenyl (NIP)-binding specificity as assessed by FACS® were selected for further backcrossing with JhD/JhD.CB17 mice. The constant part of the Vh186.2 heavy chain construct was used to perform Southern blots on tail DNA from mating pairs and their litters. The blots showed the stability of the unique integration sites over several generations and confirmed that each line has a single Tg locus. The mice were housed under specific pathogen free conditions and received Sulfatrim in drinking water every second week. Mice were used at backcross four for preliminary immunizations and the majority of experiments were done on backcross five and higher.

Antigens and Immunization. Chicken gamma globulin (CGG; Sigma-Aldrich), human serum albumin (HSA; Armour Pharmaceutical), and SRBCs (Colorado Serum) were haptenated with NP-hydroxysuccinimide ester (Cambridge Research Biochemicals). 6–12-wk-old mice were immunized intraperitoneally with NP-hydroxysuccinimide ester (Cambridge Research Biochemicals). 6–12-wk-old mice were immunized intraperitoneally with 50 μg or 5 μg alum precipitated NP-CGG, NP-CGG, or NP-CGG, or NP-CGG, as described. Controls were immunized with the same amount of alum used for precipitation of the Ags.

Flow Cytometry. FACS® was performed as described (38). In brief, spleens were mechanically disrupted and cells harvested. Red blood cells were lysed by 5 min incubation in ACT at room temperature. Purified splenocytes were counted and 10⁶ cells were stained in 96-well plates on ice for 20 min. Because of the low frequency of Ag+ cells we collected 10⁵ events per sample. PE haptenated with NIP-hydroxysuccinimide ester (Cambridge Research Biochemicals) was used to identify NP-specific B cells, as Vh186.2/λ recognizes NIP with high affinity. In addition splenocytes were stained with anti-λ-fluorescein (B374-PS84D; Southern Biotechnology Associates, Inc.) and anti-CD86-biotin/Cy5 (GL-1; BD Biosciences). In complement deposition assays, goat anti-C3-FITC (Cappel) was used in combination with anti B220-PE (RA3–6B2; BD Biosciences) and anti-λ-Alexa 647.

Histology. 5 μm frozen spleen sections were cut in a cryostat microtome, thaw mounted onto poly-L-lysine (Sigma-Aldrich)-coated slides, fixed in cold acetone for 10 min, and stored at −80°C (39, 40). Rehydrated sections were blocked with 3% BSA (Gemini Bioproducts) in PBS/0.1% Tween 20 and stained with anti-λ-alkaline phosphatase (Southern Biotechnology Associates, Inc.) and peanut agglutinin-biotin (PNA-bi; Vector Laboratories). Overlapping pictures of the whole spleen were captured and the area (in square μm) of GCs measured using.
Image Pro software (Media Cybernetics). The GC areas were added up for the whole section and then calculated as GC area per picture (each of the same unit area), thus normalizing for spleen and section size.

C3 Deposition Assays. Splenocytes were isolated from mlgM and mlgM(P436S) mice. 3 × 10^6 cells were incubated in 200 μl PBS/2% BSA on ice with 8 μg NP_{25}-BSA or BSA alone for 30 min. After washing off excess NP, the cells were injected into the tail vein of B cell–deficient mice. Half an hour after the injection, blood was collected directly into Heparin (5U)/EDTA (10 mM) to prevent blood clotting and nonspecific complement activation. Lymphocytes were separated from the anticoagulated blood by gradient centrifugation (Lympocyte Separation Medium; Cellgro) and stained with anti-B220, anti-λ, and polyclonal goat anti-C3 Abs (Cappel) and analyzed via flow cytometry. The anti-C3 detects intact C3 and its fragments, such as C3d.

Reconstitution of Soluble Antibody in mlgM(P436) Mice. Blood was taken via eye bleeding or cardiac puncture, as indicated in the figure legend. Antibody-free serum was obtained from JhD mice and NP anti-serum was obtained from (m+/+)lgM mice that had been immunized with NP_{25}-CGG 2 wk before. CGG-specific anti-serum came from BALB/c mice immunized with 100 μg CGG 2 wk before. Serum from naïve BALB/c mice was considered as “natural” antibody. The serum was tested for presence and absence of the expected specific antibodies by ELISA (not shown) and 200 μl were injected i.v. into the tail vein of mlgM(P436S) mice. 3 h later the mice were immunized i.p. with NP_{25}-CGG in alum. At day 12 after immunization the immune response was analyzed as described above.

Results

Complement Is Activated and C3 Fragments Are Deposited on B Cells in mlgM Mice but Not mlgM(P436S) Mice. In most species, complement is activated and leads to C3d deposition when whole blood or serum is kept ex vivo. Unfortunately, mouse complement is particularly susceptible to this: blood clotting at room temperature leads to substantial cleavage of C3 (41, 42). In preliminary experiments we found overwhelming deposition of C3d on B cells whether or not they were incubated with Ag, using either freshly prepared serum or plasma in a variety of tested buffers. This was predicted by results of several other groups using B cell lines and has been attributed to the ability of the CR itself to trigger C3d deposition on B cells in vitro but not in vivo (43–48). Therefore, to prevent too much nonspecific complement activation we performed a combined in vitro and in vivo experiment by separating the Ag binding step (in vitro) from the C3d deposition step (in vivo, by injecting the coated cells into B cell–deficient mice, see Materials and Methods). Note that we used a highly multivalent Ag (NP_{25}-BSA) that should effectively aggregate BCRs on the surface of B cells. Furthermore, by using B cell–deficient recipients as the in vivo source of complement, we avoided any potential influence of natural IgM in the recipient.

After injection, donor B cells are only a small fraction of PBL, and only ~3% of these are lambda positive, requiring us to collect 10^6 events per sample. mlgM cells showed complement deposition on their surface when incubated with NP-BSA, but no deposition was observed after incubation with BSA alone (Fig. 1 A). In contrast, B cells from mlgM(P436S) mice did not show complement deposition, no matter if they were treated with NP-BSA or BSA alone (Fig. 1 B). Although only λ^+ cells are specific for the Ag, we saw deposition of C3d on λ^+ cells as well, but only in the samples where we could find deposition on λ^+ cells (i.e., only in the mlgM but not P436S mutant mice and only when the B cells had been coated with Ag). This surprising but consistent finding indicated that injection of the Ag-coated cells had induced widespread, strong complement activation intravascularly in the recipients. Indeed, it is well known that injections of immune complexes can even cause anaphylactic death of mice presumably due to massive complement activation. In any case, these data show that mlgM, but not P436S mutant B cells can activate complement deposition directly on the B cell surface when the BCR is cross-linked.

Proliferative Immune Responses. Primary immune responses were initially evaluated at day 12, the known peak of the response in mlgM mice (33). To evaluate a range of different “strengths” of stimulation, two different carriers were chosen and haptenated at two different ratios. Each carrier and haptenation ratio was used for immunization at two different doses, 50 μg as used in all previous immunizations of mlgM mice, and 5 μg as a low Ag dose, for a total of 6 conditions. The B cell immune response was measured via the expansion of λ^+/NIP^+ cells as determined by FACS® analysis (Fig. 2). As will be detailed below, in all experiments performed we observed that the immune response of mlgM(P436S) mice was significantly lower than in mlgM mice with especially noticeable differences from the mlgM mice at the low dose of Ag (Table I). Nevertheless, the mlgM(P436S) mice responded to all immunizations regimes.

NP_{25}-CGG Immunization. The strongest response in both types of mice was seen to immunization with NP_{25}-CGG (Fig. 3 A). At 50 μg, both strains showed significant expansion of λ^+/NIP^+ cells. The λ^+/NIP^+ cell population
of splenocytes in mlgM mice increased nearly eightfold whereas this population expanded only fivefold in mlgM (P436S) mice. We performed this experiment two times each with two different mlgM(P436S) strains obtained from different founders, always comparing to mlgM mice immunized at the same time with the same batch of Ag. No differences were found in comparing the two mlgM(P436S) strains to each other and only small variations were seen between immunizations as indicated by the error bars representing the pooled data of these four experiments. Importantly, both of the mlgM(P436S) strains showed lower responses than the wild type mlgM mice in all experiments, arguing against an effect of the Tg integration site. At the lower Ag dose (5 µg) the same pattern (threefold increase of λ+/NIP+ cells in mlgM mice in comparison to twofold in mlgM(P436S) strains) can be observed during a weaker but still significant response to the Ag. It is interesting that there are nearly identical percentages after immunization with 5 µg in mlgM mice and with 50 µg in mlgM(P436S) mice at day 12 (the peak response, see time course below). In this sense the inability of mlgM(P436S) to bind C1q results in a 10-fold decreased sensitivity in terms of Ag dose.

Immunofluorescence responses were also analyzed by immunohistology of cryosections of spleens. Staining with PNA to detect GCs was of special interest, as ligation of CR1/2 on B cells can enhance survival in GCs (10). To confirm Ag specificity of GCs we included an Ab to the λ light chain. At day 12 after a 50 µg NP22-CGG immunization, mlgM(P436S) spleens had GCs of only ~2/3 the size of mlgM GCs (Figs. 4 and 6) and λ+ cells were visibly more abundant in mlgM spleens, confirming the flow cytometry data (Fig. 3 A). Nonetheless, spleens of both strains exhibited GCs homogeneously filled with λ+ cells. Distribution of λ+ cells in the white pulp was similar among the strains with most of the cells populating GCs and a smaller portion residing in the marginal zone. GC generation was also dose-dependent, as 50 µg immunizations always elicited bigger GCs than 5 µg immunizations (Fig. 4).

NP3-CGG Immunization. Again mlgM(P436S) mice did not respond as well to NP3-CGG as mlgM mice when analyzed by flow cytometry (Fig. 3 B). In these immunizations all of the responses were weaker than in NP22-CGG experiments, as expected from the lower haptenation ratio. Still, the results paralleled the NP22-CGG immunizations, with the mutant mice having attenuated responses. 12 d after immunization with 50 µg NP3-CGG, mlgM mice show 5.5-fold and mlgM(P436S) mice 3.4-fold increase of λ+/NIP+ cells. The responses to 5 µg of NP3-CGG on the other hand resulted only in 2-fold increase for mlgM and 1.4-fold in the mutant mice.

Immunohistologic examination of spleens from mice immunized with NP3-CGG showed that GCs form in response to this sparsely-haptenated Ag at the higher Ag dose, but are much smaller than in NP22-CGG immunizations. In immunizations with 5 µg of NP3-CGG we could observe λ+ GCs only in mlgM but not in mlgM(P436S) spleens. Thus, a more absolute difference between the mutant and wild-type strains emerged at low doses of a lightly haptenated Ag.

NP15-HSA Immunization. When HSA was used as a carrier for NP, B cell expansion was less robust than with CCG (Fig. 3 C), as we had observed before (33). Regardless, immunization with 50 µg NP15-HSA resulted in a weaker immune response in mutant mice (2.9% λ+/NIP+ cells) compared to mlgM mice (5.2% λ+/NIP+ cells). But after immunization with 5 µg NP15-HSA both mouse strains responded with only a ~50% increase in the λ+/NIP+ cell population.

### Table I. Summary of the Percentages of Total λ+/NIP+ Splenic B Cells in Response to Various Ags and Doses at Day 12 After Immunization

| Immunization | Ag Dose (µg) | mlgM % | Fold Increase |
|--------------|-------------|--------|--------------|
| NP22-CGG     | 5           | 2.3    | 3.2          |
|              | 50          | 7.3    | 6.5          |
|              | 150         | 17.5   | 1.9          |
|              | 600         | 22.2   | 1.4          |
|              | 1200        | 42.1   | 5.5          |
|              | 2400        | 120.1  | 2.6          |
| NP3-CGG      | 5           | 1.9    | 2.6          |
|              | 50          | 7.6    | 2.6          |
|              | 150         | 8.3    | 1.5          |
|              | 600         | 19.9   | 1.5          |
|              | 1200        | 44.1   | 2.1          |
|              | 2400        | 21.9   | 2.9          |
| NP15-HSA     | 5           | 1.4    | 1.4          |
|              | 50          | 5.3    | 3.4          |
|              | 150         | 1.3    | 4.2          |
|              | 600         | 1.9    | 1.2          |
|              | 1200        | 4.4    | 2.9          |
|              | 2400        | 2.1    | 2.1          |

a Percentage of live, splenic cells that are λ+/NIP+, as gated in Fig. 2.

b Fold increase in Ag-immunized spleens compared to alum-immunized spleens.
Stained sections of spleens immunized with NP$_{15}$-HSA had smaller GCs in comparison with CCG as carrier. GCs were also not homogeneously packed with $\lambda^+$ cells as seen at day 12 after NP$_{3}$-CGG immunization. Nevertheless, the differences between the mouse strains remained, with smaller GCs containing few $\lambda^+$ cells at the higher Ag dose for the mutant mice compared with larger GCs with more $\lambda^+$ cells in the mlgM mice. Results for the lower Ag dose showed parallel differences between the strains, with small GCs in both strains, but very few $\lambda^+$ cells only in the mlgM(P436S) mice.

**Primary Immune Response Kinetics.** To better define differences in the primary immune response of the mice, we measured the increase of $\lambda^+$/NIP$^+$ cells and the size of GCs at days 0, 4, 8, 12, and 20 after immunization, studying responses to both doses of NP$_{22}$-CGG.

After immunization with 50 $\mu$g NP$_{22}$-CGG, mlgM mice doubled the frequency of $\lambda^+$/NIP$^+$ cells at day four in comparison to nearly no response in mutant mice (Fig. 5). 8 d after immunization, mlgM mice displayed an even stronger increase of Ag specific cells. Interestingly, after overcoming the initial lag phase, the mutant mice show a growth rate between days 4 and 8 that is similar to mlgM mice. Given the selected time points for this time course experiment we observed the peak of the immune response of both mouse strains at day 12, as observed for mlgM mice previously. However, the shape of the growth curves of Ag-specific cells was quite different between the two strains. mlgM mice demonstrated a continuous increase in the fraction of $\lambda^+$/NIP$^+$ cells at each time point up to day 12 whereas in mutant mice, $\lambda^+$/NIP$^+$ cells increase little between days 8 and 12, a time when the GC response should be dominant (5, 33). This lower increase in Ag-specific B cells resulted in a plateau-shaped response in mlgM(P436S) mice, in contrast to the strong increase up to the peak in mlgM mice. Finally, at day 20, $\lambda^+$/NIP$^+$ cell frequencies are lower than on day 12 in both mouse strains.

Immunizations with 5 $\mu$g NP$_{22}$-CGG resulted again in lower responses of mlgM(P436S) mice during the whole response than in mlgM mice (Fig. 5). 4 d after immunization we observed, similar to the higher Ag dose, a much stronger response in mlgM mice than in the mutant mice. Between days 4 and 8 the mutant and mlgM mice show parallel but still weaker increases in Ag specific cell numbers, again similar to the higher Ag dose. Between days 8 and 12, $\lambda^+$ cells in mlgM(P436S) mice continue to expand, in contrast to the high Ag dose, almost matching the mlgM response to the same Ag. As will be seen below, in this case the percentages do not match the GC size data, indicating that most of the proliferation measured by FACS$^\circledR$ in this interval must have come from non-GC cells in the mutant mice, in contrast to the mlgM mice. The decrease in cell numbers between day 12 and 20 is similar in both mouse strains at 5 $\mu$g NP$_{22}$-CGG. On days 12 and 20 the percentages of $\lambda^+$ cells from 5 $\mu$g NP$_{22}$-CGG immunization in mlgM mice resemble those elicited by 50 $\mu$g immunizations in mlgM(P436S) mice.

Overall the mutant mice show a later onset of the response and less expansion of $\lambda^+$/NIP$^+$ cells. At the higher Ag dose the mutant mice seem to catch up between day 4 and 8 but are not able to sustain the increase in their response like mlgM mice up to day 12. At the low Ag dose mutant mice respond in a similar fashion to mlgM mice but with lower cell numbers. The rate of disappearance of Ag-specific cells between day 12 and day 20 is similar in both mouse strains. The observation that the mutant mice cannot sustain their response could be explained by the importance of complement for survival of B cells in GCs (10). Faster clearance within GCs could result in a lower $\lambda^+$/NIP$^+$ cell number that flattens out the response as seen with the 50 $\mu$g dose. To obtain more information about the influence of this mutation on the number and size of GCs, we followed the generation and disappearance of GCs during this 20-d time course.

**GC Reactions.** Immunohistologic analysis was performed on spleens from mice used for the time course of the primary immune response to NP$_{22}$-CGG. Similar to the flow cytometry data, GCs of mlgM(P436S) mice were smaller than in mlgM mice in nearly all groups. The peak of GC size in both strains occurred around day 12 (Fig. 6), in accord with the FACS$^\circledR$ analysis above.

After immunization with 50 $\mu$g NP$_{22}$-CGG we saw small GCs in both strains at day 4 that were only partly filled with $\lambda^+$ cells (Figs. 4 and 6 A). By day 8, GCs had in-
increased in size in both strains and were more densely filled with λ⁺ cells. The size of GCs of both mouse strains was similar after 8 d, a point at which the number of λ⁺ cells by FACS® most closely matched between the two strains.

Most importantly, however, the size of GCs continued to increase through day 12 in mlgM mice whereas GCs in mlgM(P436S) mice showed only a minimal increase after day 8. The halt in the growth of the GCs in the mutant

Figure 4. mlgM(P436S) mice generate smaller and fewer GCs throughout most of the primary immune response. Spleens were frozen from each experiment and 5 μm frozen sections were cut and stained with anti λ-AP (blue) and PNA-bi/SA-HRP (red). mlgM spleens consistently show more GCs of larger size than mlgM(P436S) spleens.
mice also demonstrates that the differences between the strains overall could not just be due to the slightly different numbers of /H9261/H11001 cells before immunization, as the size of GCs in both strains is nearly the same at day 8 but differs substantially at day 12. At day 20, the sizes of GCs were decreased in both strains. Still, at day 20, mIgM mice retained substantially bigger GCs (Fig. 6 A).

In responses to 5 μg of Ag, only mIgM mice had measurable /H9261/H11001 GCs 4 d after immunization (Figs. 4 and 6 B). At day 8, mlgM(P436S) mice caught up with mIgM mice and both strains had about the same size of /H9261/H11001 GCs which were also more densely filled with /H9261/H11001 cells. After 12 d we observed not only a stagnation in GC size in mlgM(P436S) mice that received 5 μg of Ag, as with 50 μg of Ag, but actually a decrease in GC size compared with day 8. mlgM mice on the other hand still showed growth in GCs up to day 12, reaching an average area four times greater than in the mutant mice. At day 20 both strains had smaller GC sizes, as was observed in the 50 μg Ag dose. Nonetheless, the mlgM mice retained GCs with areas ~5 times greater than in mlgM(P436S) mice.

Reconstitution of Soluble Antibody Restores Responses in mlgM(P436S) Mice. Natural autoantibody is thought to play a role in generating the CR1/2 ligand under normal circumstances (6, 31, 32). In fact, there is little direct evidence for this, with the best support for the idea coming from the phenotype of mice that lack secreted IgM (but not IgG) and have delayed kinetics of primary immune responses (49, 50). In contrast to the natural situation in nonimmune mice, it is very clear that if soluble Ab, particularly of IgM and certain IgG isotypes is added to Ag, this will potentiate the primary immune response (8, 51–54). One would predict that such Ab, which is absent in the mlgM(P436S) mice, might restore the responses of these B cells by bypassing the defect in the mutant mlgM. We therefore did two types of experiments (Fig. 7). In one case, we used polyclonal anti-carrier Ab raised by immunizing normal mice. Infusion of 0.2 ml of serum from such immune mice i.v. 3 h before i.p. immunization with NP22-CGG did indeed result in the restoration of proliferative responses to levels typically seen in mlgM mice (Fig. 7, A and B). Serum from nonimmune mice or from JhD mice had no significant effect. In a second approach, we immunized (m/H11001)sIgM Tg mice with NP-CGG (33) to generate secreted IgM anti-NP and used this serum. This also resulted in restoration of the response of mlgM(P436S) mice to nearly the levels of mlgM mice (Fig. 7 C). Thus, in the case of NP-CGG, immune but not nonimmune sera restored responses of mlgM(P436S) mice.

Discussion

In this paper we show the activation of complement on the B cell surface in vitro and demonstrate its importance...
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way activation by both eliminating soluble Ab and pre-
type that we could expect to see in disabling classical path-
in a partial immune response defect. This limits the pheno-
cr1/2 with the BCR, as in the CR KO mice, only results
deficient CR1/2-mediated signaling (5).

doses of Ag could at least in part compensate the effect of
play a role on FDC's. More recent results indicate that high
important for this phenotype, though CR1/2 could also

evertheless it was shown that CR deficiency on the B cell is
impairment of GCs in these mice were controversial. Nev-
all studies, although details varied. Also the results on the

Several groups have investigated the primary immune re-
creased signaling by cross-linking CD19 and the BCR.
which is associated with CD19 and TAPA-1 (14). There-
leads to increased signaling (11, 15). Later, this process was
first found that cocrosslinking of CD19 with the BCR

Whereas, in the system we studied, soluble Ab was not

Costimulation of B cells for an optimal immune response.
leading to deposition of C3 fragments that in turn provide
sufficient to start the classical pathway of complement,

We did this by genetically eliminating two sources of C1q-dependent comple-
ment activation: soluble Ab and membrane-bound IgM. Only the inactivation of the BCR-intrinsic (mlgM) source
gave a phenotype similar to the CR1/2 KO mice. Thus, we found that activation of complement by the BCR is
sufficient to start the classical pathway of complement,


Figure 7. Specific antiserum rescues the primary response in mlgM(P436S) mice. Serum was obtained from cardiac puncture of JhD and BALB/c (na-
ive and 2 wk after immunization with CGG) mice. Tail vein injection of 200 µl serum into mlgM(P436S) mice was followed by i.p. immunization with
5 µg of NP22-CGG three hours later. At day 12 the percentage of λ⁺/NIP⁺ (A) and λ⁺/PNA⁺ (B) splenocytes were determined by FACS® analysis. Admin-
istration of immune BALB/c serum (black) led to reconstitution of the primary immune response in comparison to mlgM(P436S) mice that received
no serum (white, P < 0.005), JhD serum (horizontal stripes, P < 0.005), naive BALB/c serum (diagonal stripes, P < 0.005), or immune BALB/c serum
(black, P < 0.005). The influence of NP specific IgM on the response of mlgM(P436S) mice (gray) was compared with the response in mlgM (black)
and mlgM(P436S) (white) mice at 5 µg and 50 µg NP22-CGG (C) (P < 0.01 for 5 µg and P < 0.05 for 50 µg between mlgM or mlgM(P436S)+Ab and
mlgM(P436S)).

for the immune response in vivo. We did this by genetically eliminating two sources of C1q-dependent comple-
ment activation: soluble Ab and membrane-bound IgM. Only the inactivation of the BCR-intrinsic (mlgM) source
gave a phenotype similar to the CR1/2 KO mice. Thus, we found that activation of complement by the BCR is
sufficient to start the classical pathway of complement,

Though it is widely accepted that complement partici-

pates in B cell signaling and subsequent B cell responses, its
relative importance and exact role are still debatable. It was
first found that cocrosslinking of CD19 with the BCR
leads to increased signaling (11, 15). Later, this process was
found to be mediated by C3b that links the BCR to CR2,
which is associated with CD19 and TAPA-1 (14). There-
fore, in vivo-activated complement is necessary for in-
creased signaling by cross-linking CD19 and the BCR.
Several groups have investigated the primary immune re-

For the journal The Journal of Experimental Medicine, published by the Rockefeller University Press, New York, New York 10021. Discussing activation of complement by the BCR, with a focus on the role of CR1/2-mediated signaling and the effects of specific antiserum in rescuing the primary immune response in mlgM(P436S) mice. The study highlights the importance of complement activation in B cell signaling and the role of CR1/2 in this process. The data suggest that removing both soluble Ab and membrane-bound IgM can lead to a phenotype similar to the CR1/2 KO mice, supporting the role of CR1/2 in B cell signaling. The use of specific antiserum to rescue the primary immune response in mlgM(P436S) mice further underscores the importance of this pathway in vivo.
due to our assay system with the sudden injection of high numbers of opsonized lymphocytes \((3 \times 10^7)\) in the peripheral blood. This massive complement activation on the B cell surface likely overwhelms soluble regulators of complement present in the blood which usually prevent spreading of C3. A similar situation probably occurs when immune complexes are injected into mice i.v., which can cause enough complement fixation to kill the animal. The intention of this assay was not to mimic physiologic complement activation but purely to show the ability of mIgM to activate complement in the absence of serum Ig. Nevertheless this assay also shows the lack of complement activation of Ag loaded mIgM(P436S) B cells, demonstrating the inability of this mutant mIgM to fix complement even in this setting where evidently activation can be amplified. These results demonstrated the plausibility of our hypothesis, directly showing that there is Ag-dependent complement activation even in the absence of secreted Ab as seen on mIgM B cells. Additionally, this experiment showed that the complement cascade can be started by the BCR on the target cells themselves when they are stimulated with Ag. Presumably complement on the surface of the B cell is activated by C1q binding that is catalyzed by the close proximity of BCRs after ligation by a multivalent Ag, much as IgG fixes complement only when several molecules are bound to Ag in close proximity.

As there are differences in deposition of complement on mIgM versus P436S Tg B cells, our hypothesis predicts differences in immune responses between the two types of Tg mice, given the known effects of complete inability to signal via CR1/2. To test this, we characterized the immune responses to different Ag carrier conjugations and different haptenation ratios. In all our experiments we found lower responses in the mutant than in mIgM mice. In the 50 \(\mu g\) group on the peak of the response of B cell expansion (day 12), we found only about half the frequency of Ag-specific cells in the mutant mice as in mIgM’s, varying somewhat depending on the Ag-carrier combinations used (Fig. 2). Consistently, the mutant mice showed the same extent of B cell expansion in response to 50 \(\mu g\) as mIgM mice had to 5 \(\mu g\), leading to the conclusion that the mutation leads to a 10-fold decrease in sensitivity to the Ags at the peak of the response. This finding is similar, perhaps coincidentally, to the \(\sim\)10-fold decrease in Ab titers observed in CR KO mice in comparison to wild-type mice (5–7), although in our system we are not able to measure Ab titers and therefore only looked at the percentage of Ag-specific B cell expansion. Moreover, the response to 5 \(\mu g\) of sparsely haptenated Ag (see NP\(_\gamma\)-CGG) or early in the response (NP\(_{22}\)- CGG at day 4) was very weak in the mutant mice and not much different from controls that were immunized with alum alone. These observations are in concert with earlier observations that complement coactivation boosts and prolongs the immune response particularly to weak or lower dose immunization. Nonetheless, we did see responses to lower dose Ag in P436S mice, albeit smaller ones than in mIgM mice, which we believe are again consistent with the notion that signaling via CR1/2 is not absolutely necessary for B cell responses, as CR KO mice also show at least a weak response.

Perhaps the most interesting defect in the mIgM(P436S) mice was found during the analysis of GCs. Initially, studying day 12 after immunization, we saw a difference in GC size between the mutant mice and mIgM mice, paralleling the data on B cell expansion. Data on the GC area during the time course of the primary immune response showed two major differences between the strains. First, GC responses (and to some degree proliferative responses) were slow to start in P436S mice, which was seen at both doses of Ag. This could indicate a rather critical role in the initial generation of GC precursor cells that may depend on deposition of C3 fragments catalyzed at the B cell surface. Nonetheless, it seems likely that there is a bypass of this pathway at least for a minority of GC precursor cells. This could be due to alternative methods to activate complement or simply that it is not absolutely required. Once GC precursors are selected, evidently the early part of the GC response proceeds similarly in the two mouse strains. At this point, we would suggest that CR1/2 signals generated in this way may be less important; this would also be before Ag concentrations would become limiting, as indeed the GCs are manifestly growing at this stage.

Significantly, by day 12 the GC responses stagnate (50 \(\mu g\) dose) or involute (5 \(\mu g\) dose) in the mutant P436S mice, but continue expanding in the mIgM Tg mice. This could reflect an increased requirement for CR-mediated cosignals during the late GC response, when Ag concentrations are waning. This generally agrees with prior results in CR1/2 KO mice (5–7, 10). It also fits with a relatively greater dependence of the GC response and memory response on CR signaling that was previously reported (10). However, there are conflicting results regarding GCs. One group (6) found reduced numbers and size of GCs whereas a second group (7) did not find differences in GC morphology after immunization with SRBCs (though it was unclear how this was quantitated). In their discussion, Molina et al. (7) note that this is surprising because CD19 KO mice fail to generate GCs. A recent paper by Chen et al. (5) also shows lower numbers of GCs in CR1/2 KO mice at low doses but diminished differences to controls at a high Ag dose. The latter finding is due to both an increase of GC numbers in CR1/2 KO mice and also a decrease in GC numbers in C57BL/6 controls. These findings are similar to ours at the low Ag dose but differ at the high Ag dose as both mIgM and mIgM(P436S) mice display larger GCs at the peak of the response, yet a difference remains between them. It could be that the higher precursor frequency of Ag-specific B cells in our mice accounts for these minor differences between our observations and those of Chen (5).

One very interesting paper investigated more closely the GC response in CR1/2 KO B cells (10) by transferring them into mice with ongoing GC responses. It was found that with low affinity Ag, CR-deficient B cells were quickly cleared from follicles and they also did not enter GCs. After stimulation with high affinity Ag CR-deficient
B cells were retained in follicles but only \(~1/10\) of the GCs had specific B cells in comparison to \(~1/2\) in CR-intact cells. It was also shown that CR KO cells did not divide, leading to the conclusion that CR-deficient B cells were only poorly sustained in GCs and only when the BCR cross-linking was relatively strong. Remarkably, it seems that our system has reproduced a similar phenotype by blocking both soluble IgM-mediated as well as membrane IgM-mediated C1q activation. We saw the most severe deficiencies of mIgM(P436S) mice in GC formation and especially with the low Ag dose. Moreover, the kinetic analysis showed that the GC response could not be sustained at a time when Ag should become limiting.

Lastly, we were able to test the role of immune and nonimmune serum to restore the immune response of mIgM(P436S) mice to levels seen in the wild-type mIgM Tg mice. We found that either anti-carrier or anti-hapten Ab could do this, but nonimmune serum had no effect. The anti-carrier serum would have been predominantly IgG, with some IgM, as it was raised by hyperimmunization of normal BALB/c mice. The anti-hapten serum would have been virtually all IgM, having been raised in the (\(m^+s\))IgM Tg mice (33). Thus, both IgG and IgM can have this effect, as expected from the ability of both isotypes to promote complement fixation and in accord with previous results (8, 51–54). Interestingly, at the doses used, nonimmune serum had no effect. This could be because we did not add enough serum, but in view of the positive results with immune serum, raises the point that for some antigens preimmune serum may not contain sufficient natural Ab to potentiate immune responses, further underscoring the relevance of the membrane-bound BCR-mediated pathway.

In summary, using a genetic approach and in vitro/in vivo assays as well as in vivo immunization, we found that complement can be activated, most likely via the classical pathway, on the surface of B cells as a consequence of cross-linking the BCR itself. We call this the BCR-intrinsic pathway of complement activation. This complement activation is sufficient for costimulation of B cells via CR1/2/CD19/TAPA-1 leading to a relatively enhanced immune response. We observed that in mice mutant for this pathway, 10 times the amount of Ag was necessary to induce the same peak B cell expansion as seen in mIgM mice. Differences in the kinetics and in particular the ability to sustain GC responses proved that the effect was independent of any differences in precursor frequency between the strains. Thus the mutant mice have revealed a novel pathway that is important in the initiation and development of B cell immune responses and links the complement pathway to adaptive B cell immunity. These results suggest a revision in the thinking on how the CR ligand is generated at the start of the B cell immune response. Although natural Ab could play a role if plentiful, it is fundamentally dispensable, whereas the BCR-intrinsic pathway is required, at least when sufficient preformed Ab to fix complement is absent. Physiologically, it seems likely that appropriate natural Ab will often be lacking and that the BCR-intrinsic pathway will play an important role.

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References

1. Carter, R.H., M.O. Spycher, Y.C. Ng, R. Hoffman, and D.T. Fearon. 1988. Synergistic interaction between complement receptor type 2 and membrane IgM on B lymphocytes. J. Immunol. 141:457–463.
2. Eden, A., G.W. Miller, and V. Nussenzweig. 1973. Human lymphocytes bear membrane receptors for C3b and C3d. J. Clin. Invest. 52:3239–3242.
3. Ross, G.D., M.J. Polley, E.M. Rabellino, and H.M. Grey. 1973. Two different complement receptors on human lymphocytes. One specific for C3b and one specific for C3b inactivator-cleaved C3b. J. Exp. Med. 138:798–811.
4. Pepys, M.B. 1972. Role of complement in induction of the allergic response. Nat. New Biol. 237:157–159.
5. Chen, Z., S.B. Koralov, M. Gendelman, M.C. Carroll, and G. Kelsoe. 2000. Humoral immune responses in Cr2–/– mice: enhanced affinity maturation but impaired antibody persistence. J. Immunol. 164:4522–4532.
6. Ahearn, J.M., M.B. Fischer, D. Croix, S. Goerg, M. Ma, J. Xia, X. Zhou, R.G. Howard, T.L. Rothstein, and M.C. Carroll. 1996. Disruption of the Cr2 locus results in a reduction in B-1a cells and in an impaired B cell response to T-dependent antigen. Immunity. 4:251–262.
7. Molina, H., V.M. Holers, B. Li, Y.-F. Fang, S. Mariathasan, J. Goellner, J. Strauss-Schoenberger, R.W. Karr, and D.D. Chaplin. 1996. Markedly impaired humoral immune response in mice deficient in complement receptors 1 and 2. Proc. Natl. Acad. Sci. USA. 93:3357–3361.
8. Applequist, S.E., J. Dahlstrom, N. Jiang, H. Molina, and B. Heyman. 2000. Antibody production in mice deficient for complement receptors 1 and 2 can be induced by IgG/Ag and IgE/Ag, but not IgM/Ag complexes. J. Immunol. 165:2398–2403.
9. Croix, D.A., J.M. Ahearn, A.M. Rosengard, S. Han, G. Kelsoe, M. Ma, and M.C. Carroll. 1996. Antibody response to a T-dependent antigen requires B cell expression of complement receptors. J. Exp. Med. 183:1857–1864.
10. Fischer, M.B., S. Goerg, L. Shen, A.P. Prodeus, C.C. Goodnow, G. Kelsoe, and M.C. Carroll. 1998. Dependence of germinal center B cells on expression of CD21/CD35 for survival. Science. 280:582–585.
11. Carter, R.H., D.A. Tuveson, D.J. Park, S.G. Rhee, and D.T. Fearon. 1991. The CD19 complex of B lymphocytes. Activation of phospholipase C by a protein tyrosine kinase-dependent pathway that can be enhanced by the membrane IgM complex. J. Immunol. 147:3663–3671.
12. Bradbury, L.E., G.S. Kansas, S. Levy, R.L. Evans, and T.F. Tedder. 1992. The CD19/CD21 signal transducing complex of human B lymphocytes includes the target of antiproliferative antibody-1 and Leu-13 molecules. J. Immunol. 149:2841–2850.
13. Kansas, G.S., and T.F. Tedder. 1991. Transmembrane signals
generated through MHC class II, CD19, CD20, CD39, and CD40 antigens induce LFA-1-dependent and independent adhesion in human B cells through a tyrosine kinase-dependent pathway. *J. Immunol.* 147:4094–4102.

14. Fearon, D.T., and R.H. Carter. 1995. The CD19/CR2/TAPA-1 complex of B lymphocytes: Linking natural to acquired immunity. *Annu. Rev. Immunol.* 13:127–149.

15. Carter, R.H., and D.T. Fearon. 1992. CD19: lowering the threshold for antigen receptor stimulation of B lymphocytes. *Science.* 256:105–107.

16. Weiss, A., and D.R. Littman. 1994. Signal transduction by lymphocyte antigen receptors. *Cell.* 76:263–274.

17. Tedder, T.F., L.J. Zhou, and P. Engel. 1994. The CD19/CD21 signal transduction complex of B lymphocytes. *Immunol. Today.* 15:437–442.

18. Dykstra, M., A. Cherukuri, and S.K. Pierce. 2001. Rafts and synapses in the spatial organization of immune cell signaling receptors. *J. Leukoc. Biol.* 70:699–707.

19. Dykstra, M.L., R. Longnecker, and S.K. Pierce. 2001. Epstein-Barr virus coopts lipid rafts to block the signaling and antigen transport functions of the BCR. *Immunity.* 14:57–67.

20. Stoddart, A., M.L. Dykstra, B.K. Brown, W. Song, S.K. Pierce, and F.M. Brodsky. 2002. Lipid rafts unite signaling cascades with Clathrin to regulate BCR internalization. *Immunity.* 17:451–462.

21. Pierce, S.K. 2002. Lipid rafts and B-cell activation. *Nat. Rev. Immunol.* 2:96–105.

22. Cheng, P.C., M.L. Dykstra, R.N. Mitchell, and S.K. Pierce. 1999. A role for lipid rafts in B cell antigen receptor signaling and antigen targeting. *J. Exp. Med.* 190:1549–1560.

23. Cheng, P.C., B.K. Brown, W. Song, and S.K. Pierce. 2001. Translocation of the B cell antigen receptor into lipid rafts reveals a novel step in signaling. *J. Immunol.* 166:3693–3701.

24. Cherukuri, A., P.C. Cheng, H.W. Sohn, and S.K. Pierce. 2001. The CD19/CD21 complex functions to prolong B cell antigen receptor signaling from lipid rafts. *Immunity.* 14:169–179.

25. Bottger, E.C., T. Hoffmann, U. Hadding, and D. Bitter-Suermann. 1985. Influence of genetically inherited complement deficiencies on humoral immune response in guinea pigs. *J. Immunol.* 135:4100–4107.

26. Fischer, M.B., M. Ma, S. Goerg, X. Zhou, J. Xia, O. Finco, S. Han, G. Keboe, R.G. Howard, T.L. Rothstein, et al. 1996. Regulation of the B cell response to T-dependent antigens by classical pathway complement. *J. Immunol.* 157:549–556.

27. Petry, F., M. Botto, R. Holtappels, M.J. Walport, and M. Loos. 2001. Reconstitution of the complement function in C1q-deficient (C1q−/−) mice with wild-type bone marrow cells. *J. Immunol.* 167:4033–4037.

28. Botto, M., C. Dell’Agnola, A.E. Bygrave, E.M. Thompson, H.T. Cook, F. Petry, M. Loos, P.P. Pandolfi, and M.J. Walport. 1998. Homozygous C1q deficiency causes glomerulonephritis associated with multiple apoptotic bodies. *Nat. Genet.* 19:56–59.

29. Cutler, A.J., M. Botto, D. van Essen, R. Rivi, K.A. Davies, D. Gray, and M.J. Walport. 1998. T cell-dependent immune response in C1q-deficient mice: defective interferon gamma production by antigen-specific T cells. *J. Exp. Med.* 187:1789–1797.

30. Matsumoto, M., W. Fukuda, A. Circolo, J. Goellner, J. Strauss-Schoenberger, X. Wang, S. Fujita, T. Hidvgei, D.D. Chaplin, and H.R. Colten. 1997. Abrogation of the alternative complement pathway by targeted deletion of murine factor B. *Proc. Natl. Acad. Sci. USA.* 94:8720–8725.

31. Klaus, G.G.B., J.H. Humphrey, A. Kunkl, and D.W. Dongworth. 1980. The follicular dendritic cell: Its role in antigen presentation in the generation of immunological memory. *Immunol. Rev.* 53:3–28.

32. Thornton, B.P., V. Vetvicka, and G.D. Ross. 1994. Natural antibody and complement-mediated antigen processing and presentation by B lymphocytes. *J. Immunol.* 152:1727–1737.

33. Hannum, L.G., A.M. Haberman, S.M. Anderson, and M.J. Shlomchik. 2000. Germinal center initiation, variable gene region hypermutation, and mutant B cell selection without detectable immune complexes on follicular dendritic cells. *J. Exp. Med.* 192:931–942.

34. Chan, O.T., L.G. Hannum, A.M. Haberman, M.P. Madaio, and M.J. Shlomchik. 1999. A novel mouse with B cells but lacking serum antibody reveals an antibody-independent role for B cells in murine lupus. *J. Exp. Med.* 189:1639–1648.

35. Arya, S., F. Chen, S. Spycher, D.E. Isenman, M.J. Shulman, and R.H. Painter. 1994. Mapping of amino acid residues in the C mu 3 domain of mouse IgM important in macromolecular assembly and complement-dependent cytolsis. *J. Immunol.* 152:1206–1212.

36. Wright, J.F., M.J. Shulman, D.E. Isenman, and R.H. Painter. 1988. C1 binding by murine IgM. The effect of a Pro-to-Ser exchange at residue 436 of the mu-chain. *J. Biol. Chem.* 263:11221–11226.

37. Lynes, M.A., L.L. Lanier, G.F. Babcock, P.J. Wettstein, and G. Haughton. 1978. Antigen-induced murine B cell lymphomas. I. Induction and characterization of CH1 and CH2. *J. Immunol.* 121:2352–2357.

38. Shlomchik, M.J., D. Zharhary, T. Saunders, S.A. Camper, and M.G. Weigert. 1993. A rheumatoid factor transgenic mouse model of autoantibody regulation. *Int. Immunol.* 5:1329–1341.

39. Jacob, J., R. Kasir, and G. Kelsoe. 1991. In situ studies of the primary immune response to (4-hydroxy-3-nitrophenyl)acetyl. I. The architecture and dynamics of responding cell populations. *J. Exp. Med.* 173:1165–1175.

40. Jacob, J., and G. Kelsoe. 1992. In situ studies of the primary immune response to (4-hydroxy-3-nitrophenyl)acetyl. II. A common clonal origin for parietalcell lymphoid sheath-associated foci and germinal centers. *J. Exp. Med.* 176:679–687.

41. Borso, T., and M. Cooper. 1961. On the hemolytic activity of mouse complement. *Proc. Soc. Exp. Biol. Med.* 107:227–237.

42. Klaus, G.G., M.B. Pepys, K. Kitajima, and B.A. Askonas. 1979. Activation of mouse complement by different classes of mouse antibody. *Immunology.* 38:687–695.

43. Johnson, A.A., A. Mirowski Rosengard, K. Skjodt, J.M. Ahearn, and R.G. Leslie. 1999. The structural basis for complement receptor type 2 (CR2, CD21)-mediated alternative pathway activation of complement: studies with CR2 deletion mutants and vaccinia virus complement-control protein-CR2 chimeras. *Eur. J. Immunol.* 29:3837–3844.

44. Marquart, H.V., S.E. Svehag, and R.G. Leslie. 1994. CR2 is the primary acceptor site for C3 during alternative pathway activation of complement on human peripheral B lymphocytes. *J. Immunol.* 153:307–315.

45. Nielsen, C.H., H.V. Marquart, W.M. Prodinger, and R.G. Leslie. 2001. CR2-mediated activation of the complement alternative pathway results in formation of membrane attack complexes on human B lymphocytes. *Immunology.* 104:418–
46. Nielsen, C.H., M.L. Pedersen, H.V. Marquart, W.M. Prodinger, and R.G. Leslie. 2002. The role of complement receptors type 1 (CR1, CD35) and 2 (CR2, CD21) in promoting C3 fragment deposition and membrane attack complex formation on normal peripheral human B cells. *Eur. J. Immunol.* 32:1359–1367.

47. Nielsen, C.H., and R.G. Leslie. 2002. Complement’s participation in acquired immunity. *J. Leukoc. Biol.* 72:249–261.

48. Olesen, E.H., A.A. Johnson, G. Damgaard, and R.G. Leslie. 1998. The requirement of localized, CR2-mediated, alternative pathway activation of complement for covalent deposition of C3 fragments on normal B cells. *Immunology.* 93:177–183.

49. Ehrenstein, M.R., T.L. O’Keefe, S.L. Davies, and M.S. Neuberger. 1998. Targeted gene disruption reveals a role for natural secretory IgM in the maturation of the primary immune response. *Proc. Natl. Acad. Sci. USA.* 95:10089–10093.