CD19-Regulated Signaling Thresholds Control Peripheral Tolerance and Autoantibody Production in B Lymphocytes

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Summary

The CD19 cell surface molecule regulates signal transduction events critical for B lymphocyte development and humoral immunity. Increasing the density of CD19 expression renders B lymphocytes hyper-responsive to transmembrane signals, and transgenic mice that overexpress CD19 have increased levels of autoantibodies. The role of CD19 in tolerance regulation and autoantibody generation was therefore examined by crossing mice that overexpress a human CD19 transgene with transgenic mice expressing a model autoantigen (soluble hen egg lysozyme, sHEL) and high-affinity HEL-specific IgMα and IgDα (IgHEL) antigen receptors. In this model of peripheral tolerance, B cells in sHEL/IgHEL double-transgenic mice are functionally anergic and do not produce autoantibodies. However, it was found that overexpression of CD19 in sHEL/IgHEL double-transgenic mice resulted in a breakdown of peripheral tolerance and the production of anti-HEL antibodies at levels similar to those observed in IgHEL mice lacking the sHEL autoantigen. Therefore, altered signaling thresholds due to CD19 overexpression resulted in the breakdown of peripheral tolerance. Thus, CD19 overexpression shifts the balance between tolerance and immunity to autoimmunity by augmenting antigen receptor signaling.

B lymphocyte tolerance to self antigens is achieved by the negative selection and elimination of immature B cells that express high-affinity IgM receptors for autoantigens (1–4). Negative selection is antigen receptor-dependent but also relies on established triggering thresholds for intracellular signals (1, 5). If antigen receptor ligation generates inadequate intracellular signals because of a low affinity for autoantigens, or the valency or concentration of autoantigen is low, autoreactive B cells mature and leave the bone marrow but are rendered functionally anergic (1, 4–7). Intracellular signaling thresholds are likely to also play a major role in the regulation and maintenance of peripheral tolerance.

The CD19 cell surface molecule regulates intracellular signaling thresholds critical for B cell development and humoral immunity (8–13). B lymphocytes from mice that overexpress CD19 are hyper-responsive to antigen receptor crosslinking, which results in serum Ig levels that are increased by ~40% and humoral responses that are augmented several fold (12, 14, 15). Based on this, it was expected that CD19 overexpression by autoreactive B cells would either lead to their augmented negative selection in the bone marrow or result in a more profound state of peripheral anergy. Unexpectedly however, C57BL/6 mice that overexpress CD19 have two- to fourfold higher levels of anti-DNA autoantibodies and rheumatoid factor (8, 16). Increased autoantibody production in mice overexpressing CD19 correlates with dramatic increases in the number of B1 lineage cells. However, since IgG anti-DNA autoantibodies are preferentially increased in mice that overexpress CD19, the CD19-induced autoantibodies may alternatively result from alterations in conventional B cell tolerance.

Transgenic mouse models for autoreactive B cells (4, 7) provide a mechanism for determining the role of CD19 signaling in regulating peripheral tolerance and autoimmunity. B cells from transgenic mice expressing a model autoantigen (soluble hen egg lysozyme, sHEL1) and high-affinity HEL-specific IgMα and IgDα (IgHEL) antigen receptors enter the peripheral pool but are anergic to antigen receptor ligation and produce little, if any, spontaneous HEL-specific antibody (17). Mice that express a human CD19 (hCD19) transgene provide a model for examining augmented CD19 function in vivo (8, 14–16, 18, 19). Since hCD19 can re-

1. Abbreviations used in this paper: hCD19, human CD19; IgHEL, high-affinity HEL-specific IgMα and IgDα; sHEL, soluble hen egg lysozyme.
place the function of mouse CD19 in vivo, hemizygous hCD19+/+ transgenic mice express cell surface CD19 at a twofold higher density while hCD19++ transgenic mice express threefold higher densities of CD19 (16, 19). Therefore, shEL/IgEL double-transgenic mice were crossed with hCD19 transgenic mice to determine whether tolerance would be maintained in shEL/IgEL/hCD19 transgenic mice or autoantibodies would be generated. CD19 overexpression in shEL/IgEL double-transgenic mice resulted in the production of anti-HEL antibodies at levels similar to those observed in IgEL mice lacking this model antigen. Therefore, lowered signaling thresholds due to CD19 expression in some mice that were 1,000-fold higher than in shEL/Ig-1 mice. Thus, lowered signaling thresholds resulting from the overexpression of CD19 abrogated peripheral anergy in a significant proportion of 2-mo-old shEL/IgEL double-transgenic mice while CD19 mice were equivalent to those of Ig-1 transgenic mice.

Materials and Methods

Mice. hCD19 transgenic mice (h19-1 line, C57BL/6) were as described (12, 15). In the h19-1 line of mice, 9–14 copies of the hCD19 transgene are integrated into a single (or closely linked) site(s). These h19-1 mice used in this study were backcrossed onto a wild-type C57BL/6 background for 8 to 10 generations without a diminution of hCD19 expression and all mice express similar levels of cell-surface hCD19. Mice expressing shEL (M5 line) and IgEL (M4 line) were as described (17, 20). shEL/IgEL/hCD19 triple-transgenic mice were generated by appropriate backcrosses of shEL/IgEL double-transgenic mice with hCD19+/+ mice. Transgene expression was assessed as described (12, 15, 17, 20). Mice were housed in a specific pathogen-free barrier facility. All studies and procedures were approved by the Duke University Animal Care and Use Committee.

Immunization of Mice. 2-mo-old mice were immunized i.p. with 100 µg of HEL in complete Freund’s adjuvant (CFA; Sigma Chemical Co., St. Louis, MO) or PBS in CFA at day 0 and were boosted at day 21. Animals were bled just before the first immunization and 7, 14, and 28 d later.

Mouse Ig isotype-specific ELISAs. Serum levels of HEL-specific IgM allotype (IgM^a) antibody were measured by ELISA on HEL-coated plates as described (21). Absolute antibody concentrations were determined relative to a standard curve of HEL-specific IgM^a mAb (E1 clone) generated from an IgEL transgenic mouse immunized with HEL. The ELISA sensitivity limit was ~20 ng/ml of anti-HEL IgM^a antibody.

Immunofluorescence Analysis. Antibodies used in this study included: FITC-conjugated and biotin-coupled goat anti-mouse IgG isotype-specific antibodies (Southern Biotechnology Associates, Inc., Birmingham, AL); anti-B220 (CD45R A, RA3-6B2, provided by R. L. Coffman, DNAX Research Inst., Palo Alto, CA), anti-I-A (M5/114.15.2, clone TIB120; American Type Culture Collection [ATCC], Bethesda, MD), anti-HSA (M1/69; Pharmingen, San Diego, CA), anti-CD5 (53-7.313, clone TIB104; ATCC), anti-B7-2 (GL-1; Pharmingen) and anti-mouse IgM^a (DS-1; Pharmingen) monoclonal antibodies. Phycoerythrin-conjugated streptavidin (Fisher Scientific, Fair Lawn, NJ) was used to reveal biotin-coupled mAb staining. Phycoerythrin-conjugated goat anti-rat IgG antibodies (Caltag, Burlingame, CA) were used to visualize anti-CD5 mAb staining. Cells reacting with biotin-coupled HEL were stained with phycoerythrin-conjugated streptavidin. Isolated lymphocytes were analyzed on a FACScan flow cytometer (Becton-Dickinson, San Jose, CA) as described (16).

Measurement of Intraacellular Calcium. Splenocytes were isolated, loaded with indo-1 and stained with FITC-labeled anti-B220 antibodies as described (22). Relative intracellular Ca^++ levels ([Ca^++]i) were assessed by flow cytometry after gating on the B220^+ population of cells. Baseline fluorescence ratios were collected for 1 min before HEL and/or specific mAb were added at final concentrations of: HEL, 100 ng/ml; anti-mouse CD19, 40 µg/ml (M B19-1, IgA) (16); and anti-human CD19, 40 µg/ml (H B12b, IgG1) (23). An increase in the ratio of indo-1 fluorescence indicates an increase in [Ca^++]i.

Statistical Analysis. All data are shown as mean values ± SEM. Analysis of variance (ANOVA) was used to analyze the data, and the Student’s t test was used to compare population sample means. The Mann-Whitney test was also used to compare population frequency distributions. The 95% confidence interval for anti-HEL antibody levels observed in shEL/IgEL mice was determined using the log normal distribution (mean ± 2 SD) of antibody values with undetectable levels (~20 ng/ml) assigned the value of 10 ng/ml.

Results

Autoantibodies in shEL/IgEL/hCD19 Transgenic Mice. Serum anti-HEL IgM^a autoantibody levels in IgEL transgenic, shEL/IgEL double-transgenic, and shEL/IgEL/hCD19 triple-transgenic mice were determined to assess the status of B cell tolerance in each set of mice. Serum antibody levels for each individual mouse are shown in Fig. 1 and mean autoantibody levels for each set of mice are provided to simplify discussion of the results.

2-mo-old shEL/IgEL double-transgenic mice produced very low or undetectable levels of anti-HEL IgM^a antibodies (mean levels 31 ng/ml) when compared with IgEL transgenic mice (mean 16,700 ng/ml, Fig. 1). However, 45% (14 of 33) of shEL/IgEL/hCD19 19+/+ mice had anti-HEL IgM^a autoantibody levels (mean 2,430 ng/ml) that were significantly greater than those found in shEL/IgEL mice (P < 0.001, Fig. 1).

Anti-HEL IgM^a antibody levels were also elevated in 38% (14 of 36) of shEL/IgEL/hCD19 19+/+ mice (mean 10,500 ng/ml, P < 0.01, Fig. 1). Autoantibody levels in some shEL/IgEL/hCD19 mice were equivalent to those of IgEL-transgenic mice not expressing shEL. In fact, overexpression of CD19 resulted in anti-HEL autoantibody levels in some mice that were 1,000-fold higher than in shEL/IgEL mice. By comparison, overexpression of CD19 in IgEL/hCD19 19+/+ mice resulted in only a fourfold increase in anti-HEL antibody levels (mean 77,300 ng/ml, Fig. 1). Thus, lowered signaling thresholds resulting from the overexpression of CD19 abrogated peripheral anergy in a significant proportion of 2-mo-old shEL/IgEL mice.

The breakdown in peripheral tolerance and the development of autoantibodies in shEL/IgEL mice that overexpressed CD19 correlated with mouse age. By 5–10 mo of age, all shEL/IgEL/hCD19 19+/+ mice produced significantly higher levels of autoantibodies (mean 144,222 ng/ml) than shEL/IgEL mice (303 ng/ml, P < 0.01, Fig. 1). The lowest autoantibody level found in a 5-mo-old shEL/IgEL/hCD19 19+/+ mouse was 2,255 ng/ml. Therefore, the breakdown of tolerance in shEL/IgEL/hCD19 19+/+ mice had 100% penetrance by 5 mo of age.
Abrogation of Peripheral Tolerance in sHEL/Ig HEL/mice. Whether B cell anergy could be surmounted in young mice that overexpress CD19 was assessed by immunizing 2-mo-old mice with HEL in CFA. Mice without detectable levels of spontaneous anti-HEL antibody concentrations for each group are provided for reference. The dashed horizontal line (arrowhead) delimits the 95% confidence interval for the log normal distribution of anti-HEL antibody levels observed in unimmunized 2-mo-old sHEL/IgHEL mice as described in Materials and Methods.

A breakage of Peripheral Tolerance in sHEL/IgHEL/hCD19 mice. Whether B cell anergy could be surmounted in young mice that overexpress CD19 was assessed by immunizing 2-mo-old mice with HEL in CFA. Mice without detectable levels of spontaneous anti-HEL antibodies were also injected with CFA alone to mimic a nonspecific inflammatory stimulus. Immunization of sHEL/IgHEL/hCD19+/+ mice with HEL generated primary anti-HEL antibody responses in some mice, and a mean secondary antibody response that was 200-fold higher (P < 0.05) than that of sHEL/IgHEL mice (Fig. 2 A). A measurable antibody response was only detected in sHEL/IgHEL mice after second¬ary immunization (Fig. 2 A). A striking result was that the inflammation induced by CFA alone induced sHEL/IgHEL/hCD19+/− mice to produce anti-HEL antibodies in response to endogenous sHEL autoantigen (Fig. 2 A). In this case, the mean secondary antibody response was 4,000-fold higher than in sHEL/IgHEL mice (P < 0.05). In fact, the anti-HEL antibody levels induced in some sHEL/IgHEL/hCD19+/− mice were equivalent to those of IgHEL mice (Fig. 2 B). Similar results were obtained with sHEL/IgHEL/hCD19+/− mice although anti-HEL autoantibody levels were intermediate (data not shown). In sHEL/IgHEL/hCD19+/+ mice that already expressed detectable anti-HEL antibodies, autoantibody levels were also dramatically augmented after CFA administration (data not shown). Therefore, inflammatory responses induced by the administration of CFA revealed a breakdown in tolerance and resulted in autoantibody production in anergic mice that overexpressed CD19.

Effects of CD19 Overexpression on B Cell Development. The effects of CD19 overexpression on B cell development was assessed to elucidate the cellular basis for the breakdown of peripheral tolerance in sHEL/IgHEL mice. The breakdown in tolerance did not result from relaxed negative selection, since the number of mature IgM + B220+ or HSA10 B220+ B cells in the bone marrow, blood, and spleen of IgHEL/hCD19+/+ mice was significantly reduced in the absence or presence of sHEL (Fig. 3, Table I). A similar decrease in the generation of mature B cells occurs, presumably as a consequence of increased clonal elimination, in wild-type mice that overexpress CD19 (15). However, since all B cells bear the same receptor with the same affinity for antigen, the partial decrease in generation of mature B cells in the bone marrow of IgHEL/hCD19+/+ mice and sHEL/IgHEL/hCD19+/− mice suggests that this develop-
expression of CD19 may alter the generation of mature B cells through mechanisms in addition to increased negative selection. Nonetheless, the breakdown in tolerance did not result from relaxed negative selection.

Peripheral B cell numbers were significantly reduced in both IgHEL mice and shHEL/IgHEL mice overexpressing CD19 (Fig. 3, Table I). Overexpression of CD19 reduced circulating B cell numbers by 87% in IgHEL mice and 78% in shHEL/IgHEL mice. CD19 overexpression reduced spleen B cell numbers by 42% in IgHEL mice and 48% in shHEL/IgHEL mice. Conventional B cells within the peritoneum were also reduced by >90% in IgHEL/hCD19+/− and shHEL/IgHEL/hCD19+/− mice. Overexpression of CD19 did not induce the generation of B cells with the phenotypic characteristics of either B1a or B1b cells and only small numbers of CD5− B220lo B cells were observed in any of the 2-mo-old transgenic mouse lines (Fig. 3, Table I). In addition, all of the HEL-binding B cells in each line of mice were conventional B cells since they were CD5−, CD23+, IgDhi, and B220hi (data not shown). Thus, the dramatic increase in the levels of autoantibodies generated in shHEL/IgHEL/hCD19+/− transgenic mice were even more significant given the >50% reduction in numbers of peripheral B cells in these mice.

Chronic stimulation through the B cell antigen receptor in shHEL/IgHEL mice results in a unique IgMlo IgDhi phenotype with increased expression of class II (I-A) antigens (17, 21, 24). In comparison, the overexpression of hCD19+/− in these mice resulted in an even lower IgM expression and higher I-A expression (Fig. 3, Table I). Despite the decrease in surface IgM, all B cells from shHEL/IgHEL/hCD19+/− mice still bound shHEL in vitro in proportion to their IgM density (Fig. 3C). B cells from IgHEL/hCD19+/− transgenic mice had an intermediate IgMlo I-Ahi phenotype even in the absence of shHEL (Fig. 3, Table I). B cells from mice that overexpressed CD19 also expressed significantly elevated levels of cell surface CD86 (B7-2) (data not shown). Therefore, CD19 overexpression appeared to augment the phenotypic outcome of signaling through the B cell antigen receptor in the absence or presence of autoantigen. However, B cells from shHEL/IgHEL/hCD19+/− mice still exhibited a phenotype that is characteristic of anergic B cells.

Peripheral tolerance in shHEL/IgHEL mice results in the failure of anergic B cells to mobilize intracellular Ca++ in response to HEL-mediated antigen receptor crosslinking in vitro (25). B cells from shHEL/IgHEL/hCD19+/− mice were equivalent to anergic B cells from shHEL/IgHEL mice in their failure to mobilize Ca++ in response to HEL (Fig. 4A). B cells from shHEL/IgHEL/hCD19+/− mice that generated high levels of autoantibodies also failed to mobilize Ca++ in response to HEL (data not shown). B cells from IgHEL/hCD19+/− mice generated normal Ca++ responses (Fig. 4B). Therefore, the development of autoimmunity in shHEL/IgHEL/hCD19+/− mice does not result from a CD19-induced recovery of early signaling responses in the bulk of anergic B cells.

Although antigen receptor ligation did not induce Ca++ responses in anergic B cells, crosslinking human and mouse
CD19 induced a normal Ca\(^{2+}\) response in anergic B cells from shEL/Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice (Fig. 4 A). Crosslinking mouse CD19 in shEL/Ig\(^{H\text{EL}}\) mice also induced a normal Ca\(^{2+}\) response (data not shown). The presence of HEL during CD19 crosslinking resulted in a Ca\(^{2+}\) response that was significantly greater than that observed with CD19 crosslinking alone in shEL/Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice (P < 0.01, Fig. 4 A). However, the magnitude of the CD19/HEL-induced Ca\(^{2+}\) response in shEL/Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice was less than that observed in Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice (Fig. 4 B). Of interest was the observation that the magnitude of the CD19-induced Ca\(^{2+}\) response in shEL/Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice was always significantly higher than the Ca\(^{2+}\) response in Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice (n = 3, P < 0.05). The increased Ca\(^{2+}\) responses of B cells from shEL/Ig\(^{H\text{EL}}\)/hCD19\(^{+/+}\) mice presumably results from the endogenous ligation of antigen receptors by HEL encountered in vivo. These results indicate that CD19 ligation can induce a relatively normal Ca\(^{2+}\) response in anergic B cells. Moreover, CD19 ligation can also augment transmembrane signals generated through the B cell antigen receptor despite clonal anergy.

### Discussion

The striking induction of autoantibody production in shEL/Ig\(^{H\text{EL}}\) mice that are normally functionally anergic directly implicates CD19 signaling thresholds as a regulator of peripheral tolerance in B cells. CD19 overexpression by only twofold to threefold caused a breakdown of peripheral B cell tolerance in a clear and dramatic fashion, with autoantibody levels increased several thousandfold in some shEL/Ig\(^{H\text{EL}}\)/CD19\(^{+/+}\) mice (Figs. 1 and 2). These dramatic increases in autoantibody levels in shEL/Ig\(^{H\text{EL}}\)
hCD19+/+ mice are even more significant given the >50% reduction in numbers of peripheral B cells in mice that overexpress CD19 (Fig. 3, Table 1). Overexpression of CD19 alone did not induce anergic B cells to produce autoantibodies, as evidenced by the fact that some sHEL/IgHEL/hCD19 mice were anergic and did not produce spontaneous autoantibodies until 5 mo of age (Fig. 1). However, significant autoantibody production was induced in 2-mo-old anergic sHEL/IgHEL/hCD19 mice by inducing inflammation with CFA (Fig. 2). Autoantibodies in sHEL/IgHEL mice that overexpressed CD19 are likely to have originated from a breakdown in tolerance in conventional B cells, since HEL-specific B1a or B1b cells were not detected in IgHEL (26), sHEL/IgHEL or sHEL/IgHEL/hCD19 mice (Fig. 3, Table 1). These findings suggest that alterations in CD19-related signaling thresholds breaks peripheral tolerance, which predisposes B cells to the induction of autoantibodies.

The levels of autoantibody production observed in sHEL/IgHEL mice that overexpress CD19 (Figs. 1 and 2) clearly demonstrates that tolerance was abrogated in a significant portion of B cells. Since IgHEL B cells are constantly exposed to antigen in transgenic sHEL mice, autoantibody production in sHEL/IgHEL/CD19+/+ mice is most likely induced through an antigen receptor-dependent process. Autoantibody production in sHEL/IgHEL/CD19+/+ mice may relate to the observation that CD19 ligation can augment transmembrane signals generated through the antigen receptor despite clonal anergy (Fig. 4). We have recently demonstrated that genetic alterations in CD19 expression have significant effects on the signal transduction pathways activated after B cell antigen receptor engagement (27). Therefore, one pathway to autoantibody production in sHEL/IgHEL mice may be via concomitant CD19 overexpression, chronic antigen receptor ligation, and the influence of inflammatory mediators triggering the simultaneous breakdown of tolerance and autoantibody production in anergic IgHEL B cells. Alternatively, inflammatory mediators such as those generated by CFA administration may induce the expansion or differentiation of antigen-stimulated IgHEL B cell clones subsequent to a CD19-induced breakdown in tolerance. The latter possibility is supported by the finding that B cells from mice that overexpressed CD19 maintained a phenotype characteristic of anergic B cells (Fig. 3) and failed to generate Ca2+ responses after antigen receptor ligation (Fig. 4). The spontaneous development of autoantibodies in sHEL/IgHEL/hCD19 mice may also require a breakdown in T cell tolerance, since HEL-specific helper T cells are anergic due to chronic sHEL exposure (6). Thereby, soluble factors induced by CFA administration may replace the requirement for T cell help during autoantibody production in sHEL/IgHEL mice. Although the exact pathway to autoantibody production has yet to be determined, the current results suggest strongly that inappropriate CD19 expression or function contributes to autoimmunity by disrupting tolerance.

Variability in the timing and magnitude of autoantibody production in individual sHEL/IgHEL/CD19+/+ mice is similar to what has been observed in many mouse models of autoimmunity (28). Overexpression of CD19 in sHEL/IgHEL mice resulted in significant autoantibody production in a large portion of 2-mo-old mice, while all triple-transgenic mice produced significant autoantibodies by 5 mo of age (Fig. 1). The expansion and/or accumulation of B cell clones that have escaped tolerance may explain why all sHEL/IgHEL/hCD19+/+ mice produced high levels of spontaneous autoantibodies by 5 mo of age. This contrasts markedly with old sHEL/IgHEL mice which did not produce significant levels of anti-HEL autoantibodies (Fig. 1). Previous studies of sHEL/IgHEL mice have demonstrated that functional inactivation of autoreactive B cells is maintained throughout life (29). The variability and delayed onset of autoantibody production in sHEL/IgHEL/CD19+/+ mice may also result from their confinement to a specific-pathogen-free barrier facility. Autoantibodies first appear in some mouse models of lupus (NZB and MRL strains) around 2 mo of age, but are dramatically increased by 4–5 mo of age in all mice. Variability in onset and magnitude of autoantibody production also occurs in individual mice of these inbred mouse strains. Therefore, variability between the triple transgenic mice examined in this study is not surprising despite their identical genetic background and the fact that the hCD19 transgene is expressed to the same extent in all animals. Consistent with the current studies, an association between CD19 overexpression and autoimmunity in humans has been suggested (30). The etiology of autoimmunity in humans has also been historically linked with the accumulation of inflammatory episodes or infectious agents and often varies in degree and time of onset. Therefore, many of the current findings in sHEL/IgHEL mice mimic the...
evolution of autoreactive B cells and autoantibodies in humans.

The role of antigen receptor signaling strength in the development of autoreactive B cells has recently been examined in sHEL/IgHEL mice (26, 31). Mutations in the SHP1 protein tyrosine phosphatase of motheaten viable (me) mice abrogates the negative regulatory role of SHP1 in antigen receptor signaling, resulting in the generation of autoantibodies in nontransgenic mice. In IgHEL mice, the me mutation lowers signaling thresholds, which incites the negative selection of IgHEL B cells in the bone marrow of sHEL mice (26). The SHP1 deficiency thereby prevents autoantibody generation but facilitates the development of peritoneal B1 cells reactive with HEL (26). These results contrast markedly with the results of the current study, in which lowering B cell signaling thresholds by increased CD19 expression resulted in a breakdown in tolerance and autoantibody production rather than the total negative selection of IgHEL B cells in the bone marrow of sHEL mice (Figs. 1 and 3). Thus, SHP1 may play a key role in setting thresholds for negative selection in the bone marrow, while CD19 may preferentially regulate peripheral tolerance. Alternatively, tolerance may be finely tuned, with CD19 and SHP1 altering signaling strengths to differing extents. In contrast with CD19 overexpressing B cells, signaling in response to antigen receptor ligation is diminished in CD45-deficient IgHEL B cells (31). Diminished signaling in CD45-deficient B cells leads to reduced negative selection in the bone marrow and prolonged retention in peripheral lymphoid tissues of mice expressing sHEL. Since the in vivo functional capacity of peripheral CD45-deficient IgHEL B cells or their production of autoantibodies has not been examined, it is difficult to assess how diminished signaling in those studies relates to the current studies. Nonetheless, all of these studies demonstrate that antigen receptor signaling strength influences positive or negative selection, and the current studies demonstrate a direct and active role for CD19 in regulating peripheral tolerance and autoantibody generation.

It could be argued that the breakdown in tolerance observed in this study results from the inadvertent insertion of the CD19 transgene into a locus that controls B cell tolerance. However, this is unlikely for several reasons. First, the human CD19 transgenic mice used in these studies reconstitute normal B cell function when crossed with CD19-deficient mice (19). The h19-1 line of mice has also been backcrossed extensively onto a wild-type C57BL/6 background without a diminution of human CD19 expression. This suggests that only one transgene integration site exists and that the heterogeneity in autoantibody production observed between triple transgenic mice does not reflect the segregated inheritance of transgenes that have integrated into multiple sites. We have also generated and analyzed seven independent lines of hCD19 transgenic mice (15, 18). In all cases, B cells from each mouse line demonstrate identical functional abnormalities: hyper-responsive B cells and enhanced autoantibody production. The magnitude of these abnormalities correlates directly and linearly with the level of hCD19 overexpression (19). In addition, the abnormalities observed in hCD19 transgenic mice are reciprocal of what we have observed in CD19-deficient mice (12, 14, 16). Therefore, the effects observed in the current study most likely relate directly to augmented CD19 function rather than an interruption of other genes involved in tolerance regulation.

CD19 is a signaling component of a multimeric complex that includes CD21, the receptor for the C3d fragment of complement that covalently associates with antigens during complement activation (23, 32). C3d binding to CD21 can thereby act as a ligand for the CD19 complex that links complement activation with B cell function (33-35). Since CD21-deficient mice manifest developmental and functional defects similar to those of CD19-deficient mice (12, 13, 36, 37), overexpression of CD19 in vivo may mimic C3d ligation of CD21 by augmented signaling through the CD19 complex (8). Because the roadblock to B cell Ca++ responses in anergic B cells was transcended in vitro by simultaneous CD19 ligation and antigen receptor signaling (Fig. 4), C3 cleavage products binding to the CD19 complex may provide a molecular mechanism for bypassing peripheral B cell anergy in vivo. The inappropriate or prolonged generation of C3d during inflammatory or infectious episodes in vivo may increase the responsiveness of autoreactive B cells to weak self antigens through augmented CD19 function, resulting in a breakdown of tolerance and the clonal amplification of autoantibody-producing B cells. Because altering CD19 complex function provides a mechanism for breaking self-tolerance in vivo, CD19 function may be a molecular mechanism linking inflammation with the development of autoimmune disease.

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