FcγRIV is required for IgG2c-mediated enhancement of RBC alloimmunization

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Passive immunization with anti-D can prevent maternal alloimmunization to RhD thereby preventing hemolytic disease of the fetus and newborn. Unexpectedly, anti-D fails in some cases and some monoclonal anti-D preparations paradoxically enhances alloimmunization. The underlying mechanisms modulating humoral alloimmunization by anti-D are unknown. We previously reported that IgG antibody subclasses differentially regulate alloimmunity in response to red blood cell (RBC) transfusions in a mouse model; in particular, IgG2c significantly enhanced RBC alloantibody responses. Initial mechanistic studies revealed that IgG2c:RBC immune complexes were preferentially consumed by the splenic dendritic cell (DC) subsets that play a role in RBC alloimmunization. The deletion of activating Fc-gamma receptors (FcγRs) (i.e., FcγRI, FcγRII, and FcγRIV) on DCs abrogated IgG2c-mediated enhanced alloimmunization. Because DCs express high levels of FcγRI, which has high affinity for the IgG2c subclass, we hypothesized that FcγRIV was required for enhanced alloimmunization. To test this hypothesis, knockout mice and blocking antibodies were used to manipulate FcγRI expression. The data presented herein demonstrate that FcγRIV, but not FcγRI or FcγRII, is required for IgG2c-mediated enhancement of RBC alloantibody production. Additionally, FcγRI is alone sufficient for IgG2c-mediated RBC clearance but not for increased alloimmunization, demonstrating that RBC clearance can occur without inducing alloimmunization. Together, these data, combined with prior observations, support the hypothesis that passive immunization with an RBC-specific IgG2c antibody increases RBC alloantibody production through FcγRIV ligation on splenic conventional DCs (cDCs). This raises the question of whether standardizing antibody subclasses in immunophrophylaxis preparations is desirable and suggests which subclasses may be optimal for generating monoclonal anti-D therapeutics.

KEYWORDS
Fc receptor, antibody, red blood cell, alloantibody, antibody mediated enhancement

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Introduction

Prevention of maternal alloimmunization to RhD by passive immunization with polyclonal anti-D (i.e., immunoprophylaxis) dramatically decreased rates of hemolytic disease of the fetus and newborn. Nonetheless, some anti-D preparations, under certain conditions, paradoxically enhance alloimmunization (1, 2). The underlying mechanisms by which anti-D modulates humoral alloimmunization remain unknown, but different hypotheses involve antigen modulation, increased RhD+ red blood cell (RBC) clearance, preferential ligation of inhibitory Fc receptors on B cells, and/or steric hindrance/antigen masking (3–5). Although polyclonal anti-D preparations offer numerous advantages (e.g., polyvalent specificity), several limitations apply, including the number of human volunteers required, highly variable time-to-response conversion to produce anti-D alloantibodies, and batch-to-batch variations (5, 6). Thus, there is a considerable need to develop a monoclonal antibody approach, which would provide a cost-effective, scalable option that offers batch-to-batch consistency. Although many monoclonal anti-D antibodies have been tested, some enhanced RBC alloimmunization responses whereas others suppressed (7–9). Additionally, there was discordance between RhD+ RBC clearance and immunoprophylaxis among the anti-D monoclonal antibodies evaluated (8), challenging the explanation that anti-D works by clearing RhD+ RBCs. Recently, two new monoclonal anti-D antibodies demonstrated efficacy in preventing RhD alloimmunization in pregnant women (10, 11), although additional studies with a larger number of participants and re-exposure to RhD+ RBCs are required to fully evaluate the efficacy of these new therapeutics.

Human polyclonal anti-D consists of four IgG subclasses: IgG1, IgG2a, IgG2b, and IgG3, each with distinct effector functions, including complement fixation and ligation of specific Fc-gamma (γ) receptors (FcγRs) (12). To gain insight into how different subclasses modulate immune responses, we previously reported the generation of monoclonal antibodies specific for model RBC alloantigens in mice, and the generation of panels of IgG switch variants for each murine IgG subclass: IgG1, IgG2a/c, IgG2b, and IgG3 (13, 14). Across these models, passive immunization with the RBC-specific IgG2c subclass consistently enhanced RBC alloimmunization rates and alloantibody levels (13–15). In initial mechanistic studies, IgG2c/RBC immune complexes were preferentially consumed by splenic CD11c+CD11b+ and CD11c+CD8+ conventional dendritic cells (cDCs), which are known to be important in immune responses to RBCs (13, 16, 17). In addition, the IgG2c-mediated enhancement of alloimmunization depended on specific FcγR expression; that is, deleting the common γ chain required for expression of all activating FcγRs (i.e., FcγRI, FcγRIII, FcγRIV) on splenic cDCs abrogated alloantibody production (13). By cellular surface plasmon resonance (cSPR), FcγRI and FcγRIV have been shown to have strong affinity for IgG2c (14). Based on prior observations demonstrating the involvement of FcγRIV in immune responses mediated by IgG2a (i.e., a genetic variant of IgG2c) (18), in initiating autoimmune responses (19, 20), and in CD4+ T cell priming by cDCs (21), we hypothesized that FcγRIV is required for IgG2c-mediated enhanced alloimmunization to RBC antigens.

Herein, we report that FcγRIV is required for enhanced RBC alloimmunization following passive immunization with IgG2c; that is, inhibiting FcγRIV abrogated RBC alloantibody production. FcγRIII was not required and FcγRI was not sufficient for enhancing alloimmune responses. Unexpectedly, FcγRI was not sufficient for increased alloimmunization but was alone sufficient for IgG2c-mediated RBC clearance from the circulation, providing further evidence that RBC clearance need not be directly linked to alloantibody production. Together, these data provide additional information into which FcγRs are responsible for that IgG2c-mediated enhancement of RBC alloimmunization.

Materials and methods

Mice

B6 (C57BL/6NCrl, stock #027) were purchased from Charles River and FcγRII-/+ (B6.129P2-Fcgr3<sup>tm1lvf/J</sup>, stock #003171) were purchased from The Jackson Laboratory; these mice were generated on a 129/B6 mixed background and backcrossed to C57BL/6 for 7 generations. Thus, to control for genetic background differences, FcγRIII<sup>−/−</sup> mice were bred to generate FcγRIII<sup>−/+</sup> mice for control recipients. For experiments, FcγRII<sup>−/−</sup> mice were bred to generate FcγRII<sup>−/−</sup> mice for control recipients. For experiments, FcγRII<sup>−/−</sup> mice were bred to generate FcγRII<sup>−/−</sup> mice. Mice expressing an RBC-specific triple fusion protein consisting of hen egg lysozyme, ovalbumin, and the human blood group molecule Duffy (HOD) were generated, as previously described (22). All mice were maintained in a pathogen-free environment on standard rodent chow and water in a light and temperature-controlled environment. Unless otherwise stated, mice were 8–24 weeks old and both male and female animals were used. All protocols used were approved by the Columbia University Irving Medical Campus Institutional Animal Care and Use Committee (IACUC).

Treatment of mice

Anti-HOD mAb IgG2c antibody (1 μg/mouse, which recognizes an epitope common to Duffy (Fy<sup>a</sup> and Fy<sup>b</sup>) blood group molecule contained within the HOD antigen) was passively infused into recipients 2 hours before an RBC transfusion, as previously described (13). Transfusions consisted of 50μL of leukoreduced CellTrace-CFSE
were performed, as previously described (13). Allogeneic RBCs, and subsequent alloantibody production, was previously described (13). Post-transfusion survival of delineate leukocyte subsets and Fc receptor expression, as antibody, spleens were collected and stained with antibodies to delineate RBCs and, and subsequent alloantibody production, were performed, as previously described (13).

Detection of alloantibodies by flow crossmatch

Sera was collected from experimental mice, diluted 1:100 and added to HOD target RBCs or B6 control target RBCs in FACS buffer (phosphate-buffered saline + 0.2mg/mL bovine serum albumin + 0.9mg/mL ethylenediaminetetraacetic acid). RBCs and sera were incubated for 20 minutes at room temperature, washed three times with FACS buffer and then stained with 1:100 goat anti-mouse immunoglobulins (IgM + IgG + IgA) conjugated to APC as secondary detection reagent. To determine the isotype and subclass of alloantibodies, we used directly-conjugated antibodies against IgM, IgG1, IgG2b, IgG2c, and IgG3. Samples were incubated at 4°C for 30 minutes and then washed three times with FACS buffer prior to analysis with a flow cytometer. To calculate the adjusted MFI, the sera incubated with B6 RBCs (negative control/background signal) was subtracted from the signal from sera incubated with HOD RBCs. The only difference between HOD and B6 RBCs is the expression of the HOD antigen, which is confirmed with PUMA6 and goat anti-mouse immunoglobulin-APC staining.

RBC labeling

HOD and B6 whole blood was collected into 14% CPDA-1 from donor animals by retro-orbital exsanguination and leukoreduced with an Acrodisc PSF syringe filter (Pall Life Sciences). For CellTrace-CFSE labeling, 10μl of CellTrace-CFSE (5 μ M) was added for every 1ml of pRBCs. For CellTrace-Far Red, 2ml of CellTrace-Far Red (5 μ M) was added for every 1ml of pRBCs. pRBCs were added directly into either working solutions of CellTrace-CFSE or CellTrace-Far Red, swirled to mix, and incubated at 37°C in the dark for 20 minutes, inverting after 10 minutes. RBCs were washed twice with PBS and resuspended at 20% hematocrit in PBS. Labelled HOD and B6 RBCs were then mixed at a 1:1 ratio. To determine pre-transfusion ratio, an aliquot of mixed RBCs was analyzed by flow cytometry.

Leukocyte isolation and FcγR staining

Spleens from mice were collected into complete RPMI, collagenase digested, and filtered, as previously described (13, 23). Single cells were washed with FACS buffer, RBCs were lysed, and the remaining cells were stained with antibodies to delineate leukocyte subsets (Supplemental Table 1). To enable detection of FcγRs, cells were stained with antibodies against FcγRI, FcγRII, FcγRIII, and FcγRIIV; to prevent potential steric hindrance, cells were stained for an individual FcγR to evaluate expression levels. Antibodies specific for Thy1.2, Ter119, PDCA1, CD115, Ly6G, CD11c, F4/80, and CD8 were purchased from Thermofisher those recognizing CD19, CD11b, FcγRI, FcγRII, and FcγRIIV were purchased from BioLegend. An antibody to FcγRIII was purchased from Bio-Rad. All staining was performed in FACS buffer, and cells were interrogated on an Attune NxT flow cytometer (ThermoFisher), and data analyzed with FlowJo software.

Statistical analysis

Statistical analysis was performed with a repeated measures 2-way ANOVA or one-way ANOVA with Tukey’s multiple comparisons test; p<0.05 was considered significant. Analyses were performed using Prism, version X (GraphPad Software, Inc.).

Results

Inhibiting FcγRIIV abrogates enhanced RBC alloantibody production induced by passive immunization with IgG2c

Passive immunization with anti-HOD mAb IgG2c was previously shown to enhance alloantibody production following transfusion with RBCs expressing the HOD alloantigen (13). This enhancement required expression of activating FcγRs on CD11c‘CD8‘ and CD11c‘CD11b‘ CDC subsets (13), although which individual FcγRs(s) were required remained unknown. To elucidate specific FcγRs, knockout mice and blocking antibodies were used. FcγRIIV activity was inhibited by infusion of an anti-CD16.2 antibody (clone 9e9; herein referred to as "9e9"). Non-specific effects were controlled for using either an IgG isotype control or PBS. 9e9 treatment inhibited detection of FcγRIIV on all splenic antigen presenting cell subsets evaluated, effectively reducing the staining signal to background levels (Figure 1A). Unexpectedly, 9e9 infusion also
led to decreased expression of FcγRIII on red pulp macrophages, inflammatory monocytes, resident monocytes, and neutrophils (Figure 1B) and decreased expression of inhibitory FcγRIIb on inflammatory monocytes (Figure 1C). No significant differences were observed with FcγRI expression (Figure 1D). Absolute counts of each cell subset revealed 9e9 infusion blocked FcγRIV but did not promote cell death (Supplemental Figure 1). In contrast, levels of each individual FcγR were similar between the IgG isotype control-treated and PBS-treated animals.

To assess whether FcγRIV inhibition affected RBC alloimmunization, recipient B6 mice were infused with 9e9, an IgG isotype control antibody, or PBS 30 minutes before passive immunization with anti-HOD mAb IgG2c or PBS control. Each recipient was then transfused 2 hours later with a 1:1 mix of fluorescently-labeled allogeneic HOD and syngeneic B6 RBCs (Supplemental Figure 2). Consistent with prior observations (13), B6 mice passively immunized with anti-HOD mAb IgG2c had significantly higher anti-HOD alloantibody production throughout the 21-day time course, as compared to (PBS + PBS) control mice (Figure 2A). In contrast, 9e9 pre-treatment significantly reduced anti-HOD alloantibodies over the 21-day time course, as compared to (PBS + anti-HOD mAb IgG2c) treated animals. No significant differences were observed between the (PBS + PBS) and the (9e9 + anti-HOD mAb IgG2c) treated groups, demonstrating that inhibiting FcγRIV abrogated enhanced RBC alloantibody production. Additionally, levels of anti-HOD alloantibody production were similar between anti-HOD mAb IgG2c groups pre-treated with PBS or the IgG isotype control, providing evidence that reduction in alloantibody production was specific to 9e9 recognizing its epitope, and not due to nonspecific antibody effects. Analysis of IgM and IgG subclasses showed similar a similar trend as total immunoglobulins (Supplemental Figure 3).

We previously reported that passive immunization with IgG2c induces significant allogeneic RBC clearance, which is mitigated by decreased expression of activating FcγRI, FcγRIII, and FcγRI (13). To test whether inhibiting FcγRIV affected HOD RBC clearance, post-transfusion survival was measured over a 3-week time course. Consistent with prior observations (13),...
HOD RBC survival at 24-hours was significantly reduced in (PBS + anti-HOD mAb IgG2c) treated animals compared to (PBS + PBS) controls with no anti-HOD mAb IgG2c (Figure 2B). However, no differences in clearance were observed between groups that received 9e9 or an IgG isotype control prior to anti-HOD IgG2c mAb. To evaluate potential long-term differences between experimental groups, HOD RBC survival was normalized to the (PBS + PBS) treated control group. No significant differences in HOD RBC survival were noted between groups passively immunized with anti-HOD IgG2c mAb (Figure 2C). Together, these data demonstrate that, although FcγRIV is required for IgG2c-mediated alloantibody production, but it is not required for allogeneic RBC clearance.

**FcγRIII is not required and FcγRI is not sufficient for enhanced RBC alloantibody production**

To test the roles of FcγRIII and FcγRI in IgG2c-mediated enhanced alloantibody production, knockout mice and 9e9 were used together. Passive immunization with anti-HOD IgG2c mAb followed by RBC transfusion into FcγRIII-/- animals induced significantly increased anti-HOD alloantibody levels at all time points, as compared to (PBS + PBS) treated control animals (Figure 3A). Because FcγRIII(−/−) mice were generated on a 129/B6 mixed background, FcγRIII(+/+) replete mice were used as controls to isolate the contribution of FcγRIII in alloimmune responses. These data demonstrate that FcγRIII is not required for IgG2c-mediated enhanced RBC alloimmunization. To test whether FcγRI alone was sufficient for enhancing alloantibodies with IgG2c, FcγRIII-/- mice were pre-treated with 9e9 or IgG isotype control. Pre-treating FcγRIII-/- mice with 9e9 (thereby eliminating both FcγRIII and FcγRIV, and leaving FcγRI signaling intact) significantly reduced anti-HOD alloantibodies to levels comparable to (PBS + PBS) treated control mice (Figure 3A, black triangles). RBC alloantibodies were not reduced in the IgG isotype control group (Figure 3A, gray diamonds). Thus, FcγRI is not sufficient for increasing alloantibody production upon passive immunization with anti-HOD IgG2c mAb. To exclude background genetics as a potential confounder, experiments were performed in FcγRIII(+/−) mice, in parallel, and similar trends were observed as in FcγRIII(−/−) mice (Figure 3B).

**FcγRI is sufficient and FcγRIII is not required for mediating RBC clearance by passive immunization with IgG2c**

Passive immunization with anti-HOD IgG2c mAb leads to rapid HOD RBC clearance (Figures 2B, C). Similar levels and
kinetics of clearance were observed upon blocking FcγRIV with 9e9 (Figure 2B), suggesting clearance is mediated by FcγRI, FcγRIII, or both. To evaluate FcγRIII-mediated clearance of allogeneic HOD RBCs by anti-HOD IgG2c mAb, 24-hour survival was determined in FcγRIII−/− and FcγRIII+/+ mice. HOD RBC clearance by IgG2c was equivalent in FcγRIII+/+ and FcγRIII−/− mice (Figure 4A). Thus, FcγRIII is not required for IgG2c-mediated clearance of HOD RBCs. To evaluate FcγRI function alone, FcγRIV was inhibited in FcγRIII−/− animals; this approach leaves only FcγRI signaling intact.

No significant change was observed in HOD RBC clearance between 9e9 or IgG isotype control treated FcγRIII−/− animals (Figure 4B), demonstrating that, FcγRI expression is alone sufficient for HOD RBC clearance by anti-HOD IgG2c mAb; similar results were observed in FcγRIII−/− control mice (Figure 4C). No significant differences in HOD RBC survival were noted between groups passively immunized with anti-HOD IgG2c mAb in FcγRIII+/+ or FcγRIII−/− mice (Supplemental Figure 4). Together, these data demonstrate that FcγRI is sufficient for IgG2c-mediated RBC clearance, and that FcγRIII is not required.

**FIGURE 4**

FcγRI is sufficient to mediate RBC clearance with IgG2c. Recipient mice were passively immunized with 1μg of anti-HOD IgG2c followed by an RBC transfusion 2 hours later. Some animals received an infusion of 1μg of 9e9 or IgG isotype control 30 minutes prior to passive immunization. Sera was collected from (A) FcγRIII−/− and (B) FcγRIII+/+ recipient mice weekly and analyzed for anti-HOD alloantibodies by flow crossmatch. Data shown are cumulative of 2 independent experiments with 5 mice per group. Statistical analysis was performed with a repeated measures 2-way ANOVA with Tukey’s multiple comparisons test; ****p<0.0001, ***p<0.001, **p<0.05, *p<0.05, ns, not significant.
Discussion

A murine model of RBC alloimmunization was used to test the hypothesis that FcγRIV is required for enhanced alloantibody production upon passive immunization with IgG2c. Inhibiting FcγRIV with 9e9, an antibody that blocks its binding site, reduced RBC alloantibody production to background levels. Passive immunization with anti-HOD IgG2c mAb into FcγRII−/− mice, which still express FcγRI and FcγRIV, enhanced RBC alloantibody levels, demonstrating that FcγRIII was not required. Likewise, inhibiting FcγRIV in FcγRIII+ animals, but leaving FcγRI signaling intact, failed to induce RBC alloantibodies, indicating that FcγRI was not sufficient for enhanced alloantibody production. In parallel, FcγRI was shown to be sufficient for anti-HOD IgG2c mAb-mediated RBC clearance, demonstrating that the mechanisms of IgG2c-mediated clearance and alloantibody production involve distinct FcγRs. These data, together with our prior publication (13), provide evidence that enhanced alloantibody production following passive immunization with antigen-specific IgG2c and transfusion of the corresponding antigen-expressing RBCs, requires FcγRIV expression on splenic cDCs.

The reliance on FcγRIV for IgG2a-mediated effector functions has been demonstrated in several models including B cell depletion, phagocytosis of platelets, and autoimmune responses such as autoimmune hemolytic anemia, rheumatoid arthritis, and nephritis (18, 24–27). Mechanistically, ligation of FcγRIV promotes cellular activation (e.g., CD86 expression) and increases antigen presentation (28). We, and others, have shown that immune complexes consisting of antigen:lgG2a (or IgG2c) are preferentially taken up by splenic cDC subsets, which promotes increased T cell activation and proliferation (13, 21). Because blocking FcγRIV can ameliorate autoimmune and alloimmune IgG2a/c-mediated pathology, FcγRIV is an attractive therapeutic target.

Treatment with the anti-FcγRIV 9e9 antibody significantly reduced detectable FcγRIV on the surface of all splenic antigen presenting cell subsets that were analyzed. Unexpectedly, there was also reduced expression of FcγRII and FcγRI (but not FcγRII) on different subsets. Thus, one must consider that some of the effects of 9e9 may be due to altering levels of FcγRII and/or FcγRII. However, we show herein that FcγRII−/− mice have normal induction of alloimmunization by IgG2c. Also, we have shown previously that FcγRII is not sufficient for alloimmunization, as common γ chain knockout mice have not IgG2c-mediated enhancement of RBC alloimmunization (13). Thus, we reject the interpretation that decreased FcγRII and FcγRIII contributed to blockade of alloimmunization by 9e9.

Interestingly, passive immunization with IgG2c anti-HOD mAb induced higher levels of enhancement in FcγRII−/− than in FcγRIII−/− mice. The reason for this difference is unclear, as FcγRIII−/− is not typically considered inhibitory; however, it could be the result of decreased competition with FcγRIV for the common γ chain required for FcγR signaling or contributions of the background genetics (29), which may influence immune responses.

Blocking FcγRIV with 9e9 significantly reduced HOD RBC alloantibody levels following passive immunization with anti-HOD IgG2c mAb. However, throughout a 3-week time course, alloantibody levels did gradually increase. This may be attributed to the half-life of the 9e9 antibody, as only one infusion was given. Repeating these studies in FcγRI and FcγRIV knockout mice would address the limitations of using a blocking antibody; however, FcγRI deficiency modulates expression of other FcγRs (18, 30). Finally, it is worth noting that passive immunization with anti-HOD IgG2c mAb in 9e9 treated FcγRII−/− animals resulted in much lower levels of RBC alloantibodies, as compared to wild-type B6 recipients. Thus, although FcγRIII was not required for enhancing RBC alloantibody levels, it may still contribute to alloimmune responses; in particular, inhibiting both FcγRII and FcγRIV simultaneously is required to prevent all such RBC alloantibody production (Figure 3A).

Polyclonal antibodies, such as human anti-D, contain a heterogeneous mixture of IgG subclasses, consisting of multiple epitope specificities that may work in an additive, synergistic, or even an inhibitory, manner. Because the therapeutic anti-D used for immunoprophylaxis against Rh disease is derived by pooling material from multiple human volunteers, the efficacy of anti-D could depend on numerous variables, including donor-specific and antibody-specific characteristics. For example, a recent analysis of 23 such human monoclonal antibodies revealed that glycosylation profiles influenced biological activity (31), and IgG glycosylation can vary from human to human. Moreover, studies in mice highlight additional complexities that contribute to antibody-mediated immune responses, including affinity, the RBC antigen target and/or epitope, and RBC antigen copy number (13–15, 32–34). In addition to modulating immune enhancement or suppression, subclass can also dictate which effector pathways are employed (e.g., complement, FcRs, etc.) (35, 36). Thus, additional mechanistic investigations to elucidate how polyclonal anti-D prevents RhD alloimmunization will instruct the future development of human monoclonal anti-D antibodies, such that subclass, affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s). Of note, human FcγRIIIA is the human ortholog to mouse FcγRIII; however, this affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s). Of note, human FcγRIIIA is the human ortholog to mouse FcγRIII; however, this affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s). Of note, human FcγRIIIA is the human ortholog to mouse FcγRIII; however, this affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s). Of note, human FcγRIIIA is the human ortholog to mouse FcγRIII; however, this affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s). Of note, human FcγRIIIA is the human ortholog to mouse FcγRIII; however, this affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s). Of note, human FcγRIIIA is the human ortholog to mouse FcγRIII; however, this affinity, glycosylation, and epitope recognition can be fine-tuned to exert the desired function(s).
specificity, lot-to-lot consistency), the substantial advances in antibody technology and increased understanding of immune function, now allow antibodies to be engineered to have increased affinity for the neonatal Fc receptor (FcRn), thereby facilitating an extended half-life to make passive immunization more durable. Moreover, antibodies can be engineered to inhibit antibody:FcRn interactions at the maternal-fetal interface, which is especially important for preventing hemolytic disease of the fetus and newborn mediated by alloantibodies to non-D RBC antigens (39, 40).

There are several limitations to the current studies. Although there is a clear reduction in alloantibody production upon blocking FcγRIV with 9e9, these studies would be strengthened by validating these findings in FcγRIV deficient animals or with another blocking antibody with a different binding site for FcγRIV. Additionally, passive immunization with anti-HOD IgG2c mAb followed by HOD transfusion leads to endogenous production of both IgM and IgG antibodies (13). Thus, while Fc-dependent FcγRIV ligation and signaling is blocked and signaling by IgG2c subclass antibodies is blocked, these antibodies may signal through other FcRs (e.g., FcmR) and modulate alloimmune responses (41). Finally, although the current analyses were limited to splenic antigen presenting cells, it is also likely that immune complex clearance also occurs in the liver.

The data presented herein demonstrate that FcγRIV, but neither FcγRII nor FcγRI, is required for IgG2c-mediated enhanced alloimmunization to RBCs. These results, combined with our prior findings (13), support a model whereby passive immunization with an RBC-specific IgG2c antibody increases RBC alloantibody production to the corresponding antigen through FcγRIV ligation on splenic cDCs. To apply these mechanistic insights human anti-D, the future design of monoclonal antibodies should be tailored and optimized to leverage distinct subclass effector functions (12), Fc receptor expression patterns (41), and signaling (42).

Author contributions

AQ and KH designed the studies and experiments. AQ, AM, FD, MS, and MT set up experiments, collected and processed samples, and performed data analysis. All authors participated in data interpretation, revised the manuscript, and approved of the submitted version.

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Conflict of interest

Although unrelated to the contents of this manuscript, KH has a sponsored research agreement with Alpine Immune Sciences. JZ is a consultant for Rubius Therapeutics and is the founder and CSO of Svalinn Therapeutics.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Ethics statement

The animal study was reviewed and approved by Columbia University Irving Medical Campus Institutional Animal Care and Use Committee (IACUC).

Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2022.972723/full#supplementary-material
