IgG hexamers initiate complement-dependent acute lung injury

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**Graphical abstract**

Pathogenesis of alloantibody-mediated acute lung injury

Alloantibodies bind multiple alloantigens to deposit onto endothelial surfaces at densities enabling Fc:Fc interactions

IgG hexamers assemble and activate C1 complexes to trigger excessive classical complement activation, initiating injury

IgG antibody, MHC antigen, Fc:Fc interaction, IgG hexamer assembly, C1 complex activation

Excessive complement activation on endothelium

Inhibition of Fc:Fc interactions with SpA-B

Therapeutic strategies showing efficacy in model

Inhibition of classical complement activation with CSL777 ‘decoy’ Fc hexamers

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IgG hexamers initiate complement-dependent acute lung injury

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Abstract

Antibodies can initiate lung injury in a variety of disease states such as autoimmunity, transfusion reactions, or after organ transplantation, but the key factors determining in vivo pathogenicity of injury-inducing antibodies are unclear. Harmful antibodies often activate the complement cascade. A model for how IgG antibodies trigger complement activation involves interactions between IgG Fc domains driving assembly of IgG hexamer structures that activate C1 complexes. The importance of IgG hexamers in initiating injury responses was unclear, so we tested their relevance in a mouse model of alloantibody and complement-mediated acute lung injury. We used three approaches to block alloantibody hexamerization (antibody carbamylation, the K439E Fc mutation, or treatment with domain B from Staphylococcal protein A), all of which reduced acute lung injury. Conversely, Fc mutations promoting spontaneous hexamerization made a harmful alloantibody into a more potent inducer of acute lung injury and rendered an innocuous alloantibody pathogenic. Treatment with a recombinant Fc hexamer ‘decoy’ therapeutic protected mice from lung injury, including in a model with transgenic human FCGR2A expression that exacerbated pathology. These results indicate an in vivo role of IgG hexamerization in initiating acute lung injury and the potential for therapeutics that inhibit or mimic hexamerization to treat antibody-mediated diseases.

Brief summary

IgG antibodies can form hexamers. This study demonstrates the importance of IgG hexamer assembly in determining the ability of alloreactive IgG to trigger acute lung injury.
Introduction

Antibodies and the complement cascade mediate protective immunity but can both become misdirected to cause harm in autoimmune and alloimmune diseases. Some antibodies direct activation of the complement cascade at their targets, an event associated with severe pathology in disease states including several forms of transfusion reactions (1, 2), immune rejection after solid organ transplantation (3), and complications of pregnancy (4). Complement-activating alloantibodies are known mediators of transfusion-related acute lung injury (TRALI) (5, 6), a leading cause of transfusion-related deaths (7), and are linked to particularly poor outcomes following solid organ transplantation (3, 8). Complement activation by autoreactive antibodies also contributes to pathogenesis of forms of autoimmune hemolytic anemia (9), small vessel vasculitis (10), and neurological autoimmune disease (11).

Immunoglobulin G (IgG) antibodies are the most prevalent type of antibody in circulation and complement-activating alloantibodies are frequently IgG class. IgG antibodies achieve complement activation through recruiting and activating C1 complexes, each of which contain six Fc-binding domains (12, 13). A theory for how IgG achieves C1 complex activation involves groups of six IgG antibodies interacting through their Fc domains to form IgG hexamers (14). This theory recently gained experimental support from direct imaging of IgG1 and IgG3 hexamer assembly on antigenic liposomes (15, 16), with in vitro studies connecting IgG hexamerization to increased complement deposition on target surfaces (15, 17, 18). However, it is currently unclear whether IgG hexamer assembly is important in vivo in the pathogenesis of complement-dependent forms of alloantibody-mediated disease.

Here, we report testing of interventions that exploit IgG hexamerization in a mouse model of acute lung injury driven by alloantibody deposition in the pulmonary microvasculature, a process that drives pathology in forms of both TRALI and antibody-mediated rejection (AbMR) of lung transplants (6). Our results identify key molecular events driving alloantibody-mediated pathophysiology in vivo. We also demonstrate preclinical efficacy of new therapeutic approaches that prevent pathology of complement-dependent organ damage caused by alloantibodies, serving as a rationale to pursue translational studies in human alloantibody driven disease.
Results

Alloantibodies are prevalent but not always harmful, so determining whether alloantibodies are ‘clinically significant’ is a frequent conundrum in transfusion and transplantation medicine. Reflecting this clinical challenge, of the many mouse monoclonal alloantibodies targeting major histocompatibility complex (MHC) class I antigens, only clone 34-1-2S triggers acute lung injury when microgram quantities are intravenously injected into mice (19, 20). In addition, only mice expressing the H-2^d^ MHC class I haplotype are known to be susceptible to injury caused by the 34-1-2S antibody (5, 6). Curiously, the 34-1-2S antibody does not readily cause injury in H-2^b^ mice, including the widely used C57BL/6 (B6) strain, despite the fact that it binds to MHC class I antigens expressed by H-2^b^ mice (5, 6). We aimed to improve our understanding of the factors determining the ability of antibodies to cause injury in both this widely used model and more generally in antibody-mediated disease states.

We measured the binding affinity of 34-1-2S antibody to each of the classical MHC class I antigens present on injury-resistant H-2^b^ B6 mice and injury-susceptible H-2^d^ mice (Figure 1A). Of the three MHC class I antigens in the H-2 locus (K, D, and L), B6 mice only express K^b^ and D^b^, and we detected binding of 34-1-2S to K^b^ but not D^b^. In contrast, we detected binding of 34-1-2S to all three MHC class I antigens from H-2^d^ mice, with high affinity binding to K^d^ and D^d^, and weak binding to L^d^ (Figure 1B). Other MHC class I antibodies (clones AF6-88.5.5.3, 20-8-4S, SF1.1.10, 30-5-7S and 34-5-8S), which do not readily induce injury (5)) each bound to only one MHC class I antigen from each MHC type (Supplemental Figure 1).

Together, the above findings led us to the hypothesis that the ability of 34-1-2S to induce lung injury in H-2^d^ mice is a function of increased density of bound antibody in H-2^d^ animals resulting from 34-1-2S simultaneously binding K^d^, D^d^, and possibly L^d^. This hypothesis was tested by injecting 34-1-2S antibody into B6.ConK^d^-on mice, which express K^d^ but do not express D^d^ or L^d^ (21). B6.H2^d^ mice expressing the full complement of MHC class I antigens recognized by 34-1-2S (K^d^, D^d^, and L^d^) were used as background-matched positive controls for susceptibility to injury (Figure 1C). In contrast to B6.H2^d^ mice, B6.ConK^d^-on mice did not develop lung injury (Figure 1, D and E). These data are consistent with 34-1-2S antibody causing injury in H2^d^ mice through high affinity binding to multiple MHC class I antigens.

Engagement of multiple antigens can permit high density antibody deposition, an event associated with classical complement activation. Complement activation has been implicated in pathogenesis of acute lung injury caused by 34-1-2S antibody, but previous studies have not determined whether injury in this model is directly triggered by antibody-mediated complement activation via the classical pathway (5, 6).
To test whether 34-1-2S-induced injury requires classical complement activation, we bred mice expressing the \(H2^d\) susceptibility locus with mice lacking C1qa (22), a protein that is necessary for classical complement activation as it is one of the three proteins which make up each of the six Fc-binding C1q subcomponents in each C1 complex (Figure 2A).

Relative to B6.\(H2^d\):C1qa\(^{+/+}\) littermate controls, C1qa-deficient B6.\(H2^d\):C1qa\(^{-/-}\) mice were resistant to alloantibody-mediated acute lung injury and mortality (Figure 2, B-D). Mice lacking C1qa were also protected from deposition of complement component C3 split products on the endothelium of pulmonary capillaries (Figure 2E). Staining for C1qa in lungs confirmed absence of C1qa protein in knockout mice, with intense C1 complex deposition seen around pulmonary arterioles in C1qa-expressing mice injected with 34-1-2S antibody (Figure 2F).

To identify the microanatomical site of classical complement activation, we stained lungs of mice injected with 34-1-2S for the complement C4 split products C4b and C4d, which form covalent bonds with proteins at sites of classical complement activation. We observed strong positivity for C4b/d highlighting the endothelium of medium and small-sized pulmonary arterioles in B6.\(H2^d\):C1qa\(^{+/+}\) mice injected with 34-1-2S, but not in B6.\(H2^d\):C1qa\(^{-/-}\) mice (Figure 2G, and Supplemental Movie 1). Together, these results indicate that 34-1-2S causes acute lung injury because this antibody is deposited onto the pulmonary arteriolar endothelium at densities sufficient to trigger excessive classical complement activation directed at the walls of these blood vessels.

Dense binding to membrane-expressed antigens would be expected to facilitate IgG Fc:Fc interactions and IgG hexamer assembly. IgG hexamers are potent activators of C1 complexes in vitro (15), and are further implicated in classical complement activation by models for C1 complex activation involving shifting of its six Fc-binding C1q subcomponents into a regular hexagonal configuration (Figure 2A and Figure 3A) (12, 13, 23). We therefore hypothesized that 34-1-2S assembles into hexamers on the pulmonary endothelial surface of susceptible mice to trigger complement-dependent acute lung injury.

Imaging methods cannot currently resolve IgG hexamers in vivo, but recent studies have developed methods for inhibiting IgG hexamerization. One approach to impair IgG hexamer assembly is to carbamylate antibodies, converting lysine residues to homocitrullines to alter charge densities in IgG Fc regions, inhibiting Fc:Fc interactions and IgG hexamer assembly (Figure 3B) (24). Mice treated with carbamylated 34-1-2S showed greatly reduced acute lung injury responses compared to littermate controls treated with non-carbamylated 34-1-2S (Figure 3, C and D, and Supplemental Figure 2, A-D). Carbamylated 34-1-2S retained its ability to bind antigens and become deposited in lungs but, in
contrast to unchanged 34-1-2S, did not induce complement C3b/d deposition in the pulmonary microvasculature (Figure 3E, and Supplemental Figure 2, E and F).

Lysine residues are present on regions of IgG outside of the Fc:Fc interaction interface (illustrated in Figure 3B), including at the Fc:C1q interaction site (25), so we pursued a more targeted strategy for inhibition of IgG hexamer assembly. We determined the sequence of both heavy and light chain complementary-determining regions and engineered a chimeric antibody with the Fab domain of 34-1-2S fused in frame to human IgG1 (hlgG1-34-1-2S). To test whether hlgG1-34-1-2S causes injury through hexamerization, we also expressed this antibody with an Fc point mutation that inhibits Fc:Fc interactions required for IgG hexamer assembly (K439E) (Figure 3F) (15). Like mouse IgG2a 34-1-2S, hlgG1-34-1-2S injections caused acute lung injury (Figure 3, G and H, and Supplemental Figure 3, A and B). Lung vascular permeability and pulmonary edema responses were reduced by the K439E mutation (Figure 3, G and H), as was complement C4b/d deposition in lungs (Fig 3I), lending further support to a role for Fc:Fc interactions and hexamerization in the pathogenesis of this disease model.

We also tested a strategy for pharmacologic inhibition of Fc:Fc interactions by mixing hlgG1-34-1-2S with recombinant B domains from Staphylococcus aureus protein A (SpA-B), which bind to IgG antibodies near to Fc:Fc interaction sites and inhibit hexamer assembly and complement activation by antibodies targeting bacterial antigens (Figure 3J) (17, 26). We hypothesized that these properties of SpA-B, which likely evolved as part of an immune evasion strategy, might be harnessed to prevent hlgG1-34-1-2S from causing acute lung injury. Adding SpA-B to hlgG1-34-1-2S reduced its ability to both induce acute lung injury (Figure 3, K and L) and cause complement C4b/d deposition within pulmonary arterioles (Figure 3M). These findings provide a third line of evidence that Fc:Fc interactions leading to hexamer assembly are important for the injury response caused by this alloantibody.

Turning to hexamer gain of function experiments, the introduction of three mutations into the Fc domain of hlgG1 (RGY mutations: E345R, E430G, S440Y) has yielded antibodies capable of off-target hexamer assembly as well as increased on-target hexamerization (15) (Figure 4A). We hypothesized that RGY-mutated 34-1-2S (RGY-hlgG1-34-1-2S) would have enhanced ability to cause acute lung injury due to increased IgG hexamer formation. We produced RGY-hlgG1-34-1-2S and confirmed its ability to spontaneously assemble into hexamers in solution (Figure 4B). Consistent with a role for alloantibody hexamerization in driving injury, RGY-hlgG1-34-1-2S showed increased potency in triggering acute lung injury relative to hlgG1-34-1-2S (Figure 4, C and D). Likely due to the ability of hlgG1-34-1-2S to form hexamers and cause complement-dependent injury when injected at 1 mg/kg, effects of the RGY mutations were only apparent at a lower dose (0.3 mg/kg), at which unaltered hlgG1-34-1-2S did not provoke complement C4b/d deposition in lungs (Figure 4E). Consistent with
binding to multiple antigens being a requirement for alloantibody-mediated acute lung injury, a novel chimeric hlgG1 antibody binding a single MHC class I antigen (hlgG1-Kd1, targeting K\(^d\)), did not provoke injury when injected into B6.H2\(^d\) mice (Figure 4F, G). However, introduction of the RGY mutations promoting hexamerization into this innocuous antibody resulted in a modified version (RGY-hlgG1-Kd1) that was able to provoke increases in lung vascular permeability and edema (Figure 4, F and G). These findings demonstrate that mutations promoting IgG hexamer assembly can increase pathogenicity of antibodies at doses that do not normally result in sufficiently dense binding to trigger complement-dependent injury responses.

Another approach for therapeutic exploitation of IgG hexamerization involves use of Fc hexamers as ‘decoy’ treatments. These therapeutic candidates are under investigation as recombinant alternatives to plasma-derived intravenous or subcutaneous immunoglobulin (IVIg or SCIg) treatments that are used in management of autoimmune and alloimmune diseases (27). We hypothesized that due to its ability to inhibit classical complement activation (11, 28), the Fc hexamer ‘decoy’ treatment CSL777 (previously Fc-µTP-L309C) would be effective in preventing alloantibody-mediated acute lung injury.

We randomized mice to receive either CSL777, SCIg (IgPro20, a human plasma-derived immunoglobulin product that is currently used to treat antibody-mediated diseases), or vehicle controls prior to injection with 34-1-2S (Figure 5, A and B). Treatment with CSL777 protected mice from developing 34-1-2S-induced lung vascular permeability and pulmonary edema responses at all doses tested, whereas treatment with SCIg only had a partial effect on alloantibody-induced acute lung injury responses (Figure 5, C-F). CSL777 treated mice lacked alloantibody-mediated deposition of complement C4 split products on pulmonary arterioles, whereas arteriolar endothelial C4b/d deposition was still observed in lungs of SCIg-treated mice after 34-1-2S antibody injections (Figure 5, G and H). Recombinant Fc hexamer therapeutics such as CSL777 might therefore be useful for prevention or treatment of complement-dependent forms of alloantibody-mediated organ injury.

Unlike humans, mice do not express the Fcy receptor FCGR2A (FcγRIIA, CD32A), negatively impacting the predictive value of mouse models for studying human antibody-mediated diseases (29, 30). To test whether our previous findings held up in a system involving FCGR2A-driven pathology, we crossed existing mouse lines to generate 34-1-2S-mediated injury-susceptible H-2\(^d\) mice expressing a human FCGR2A (hFCGR2A) transgene (B6.H2\(^d\):hFCGR2A\(^{Tg^0}\)). In response to hlgG1-34-1-2S injections, mice expressing hFCGR2A developed similar levels of lung injury relative to littermates without hFCGR2A expression but displayed a survival disadvantage (Figure 6, A-C). B6.H2\(^d\):hFCGR2A\(^{Tg^0}\) mice expressed hFCGR2A on platelets, became more thrombocytopenic than littermates lacking hFCGR2A after hlgG1-34-1-2S injections, and occasionally died before onset of
pulmonary edema (Supplemental Figure 3, A-G). Based on these observations, we hypothesized that hFCGR2A expression was enhancing intravascular immunothrombotic responses that occur within 5-10 minutes of 34-1-2S injections (6), causing fatal hypoxic respiratory failure by impairing lung microvascular perfusion and increasing alveolar dead space. Supporting this hypothesis, within 20 minutes of hIgG1-34-1-2S injections, hFCGR2A transgenics had increased sequestration of platelets and neutrophils in their pulmonary microvasculature (Figure 6, D and E), and rapidly became more hypoxic than littermates without hFCGR2A expression (Figure 6H). B6.H2d:hFCGR2ATg<sub>0</sub> mice developing particularly severe hypoxemia (SpO<sub>2</sub> readings <40%) did not survive until 2 hours after antibody injections (Supplemental Figure 3H). Surviving hFCGR2A transgenics went on to develop increased lung vascular permeability responses compared to littermates without hFCGR2A expression, indicating that given time to develop, responses to FCGR2A engagement also exacerbate pulmonary edema (Supplemental Figure 3, I and J), as recently described in a similar model (30). Classical complement activation was still critical for pathogenesis in the presence of hFCGR2A, as knockout of C1qa protected mice expressing hFCGR2A from lung injury and mortality (Figure 6, I-K).

Consistent with retained importance of complement for initiation of injury responses in mice expressing FCGR2A, blocking hexamerization with SpA-B, or pre-treating mice with CSL777 hexamer ‘decoys’ were protective in B6.H2d:hFCGR2ATg<sub>0</sub> mice given hlgG1-34-1-2S injections (Figure 7, A-F). These results provide evidence that strategies to inhibit or mimic IgG hexamerization can maintain efficacy in the presence of increased severity of pathophysiology due to FCGR2A engagement.
Discussion

This study advances our understanding of immunology in three areas. Our work elucidates the molecular determinants of susceptibility in a widely used inflammation model. To our knowledge, our experiments represent the first in vivo evidence that alloantibody hexamerization is important for pathophysiology. In addition, we show that two experimental therapeutic approaches that target antibody hexamerization can reduce alloantibody-mediated organ injury.

The initial aim of our study was to solve the mystery of why injections of 34-1-2S antibody (but not other monoclonal MHC class I antibodies) cause striking pathophysiology in H-2^d^ mice (but not in mice with other haplotypes) in what has become a widely-used inflammation model (6, 19, 20, 30–40). Our results provide an explanation for this pattern of susceptibility - high affinity binding to multiple MHC class I antigens on the pulmonary endothelium of mice with the H-2^d^ haplotype facilitates sufficiently dense alloantibody deposition for IgG hexamer assembly to occur. These IgG hexamers then direct classical complement activation onto the pulmonary endothelial surface, initiating the excessive leukocyte and platelet responses that have been reported in previous studies to cause acute lung injury in this model (6, 19, 20). Antibody/haplotype combinations in which single MHC class I antigen is targeted do not enable sufficiently dense antibody binding for IgG hexamer assembly to occur, unless Fc mutations are introduced into the antibody to increase hexamerization. Our findings imply that both lymphocyte crossmatch and single antigen bead assays for detecting complement-fixing antibodies may lack sensitivity for detecting antibodies that activate complement in vivo. The mobility, density and diversity of antigens on the lymphocytes or solid-phase beads used in these assays does not exactly resemble those on endothelial cell surfaces targeted by donor-specific antibodies in vivo, and our results indicate that each of these factors determines complement-fixing capability of antibodies. Conversely, antigen density on beads exceeding that found in vivo may give false positive findings of complement-fixing antibody responses.

By demonstrating that IgG hexamers are important in pathophysiology and represent therapeutic targets in vivo, our work builds on in vitro studies implicating antibody hexamerization in complement activation by antibodies targeting antigens on liposomes, lymphoma cells or bacterial membranes (15–18, 24, 41). Further clinical translation will require determination of the importance of IgG hexamers in more complex models of diseases involving complement-activating alloantibodies (e.g. AbMR, TRALI, hemolytic transfusion reactions, and hemolytic disease of the fetus and newborn) or autoantibodies (e.g. warm autoimmune hemolytic anemia, antiphospholipid syndrome, myasthenia gravis and neuromyelitis optica). As SpA-B does not inhibit IgG3-mediated complement activation but IgG3 can
assemble into hexamers (15, 16), it will also be important to develop strategies to inhibit IgG3 hexamerization to examine the therapeutic potential of targeting hexamers formed by IgG3 antibodies.

Our results also provide new insights into the modes of action of past, present, and potential future therapeutics. Full length staphylococcal protein A (SpA) has been used as a therapeutic in the form of a now-discontinued extracorporeal immunoadsorption product (ProSORBA column). Efficacy of SpA immunoadsorption has been observed in patients with symptoms unchanged by plasma exchange, an effect ascribed to leakage of SpA from columns into the bloodstream of patients resulting in B cell depletion caused by the action of SpA as a B cell receptor super agonist (42). Purified SpA infusions (PRTX-100) were subsequently studied in early-stage clinical trials before abandonment for financial reasons (43). Our results suggest that there may be settings where therapeutics based on the SpA-B subdomain of SpA have efficacy through preventing complement activation without risk of adverse effects related to immune complex formation and B cell super agonism caused by immunoglobulin polyvalency of full-length SpA. Donor-derived immunoglobulin products (e.g. IVIg and SCIg) are currently used to treat antibody-mediated disease flares. Our observation that SCIg reduces injury responses but does not prevent classical complement activation in vivo is concordant with previous studies concluding that immunoglobulin therapeutics act on downstream mediators in vitro and in vivo (35, 44). CSL777 is an attractive potential future therapeutic for treatment of alloantibody-mediated diseases as it showed efficacy in our models, and had a mode of action that would be anticipated to prevent complement activation by both IgM and IgG antibodies as well as pathophysiology resulting from Fcγ receptors (11, 27, 28). CSL777 also lacks issues with use of donor-derived products related to sourcing, purification and concentration for injections (27, 45).

In conclusion, this study provides evidence that IgG hexamers can be important triggers of complement-dependent pathophysiology in vivo. Our preclinical studies support further investigation of IgG hexamerization inhibitors and IgG hexamer ‘decoy’ therapeutics for use in preventing disease states caused by antibodies and complement activation.
Materials and methods

Sex as a biological variable. Male mice were used as female mice are not susceptible to 34-1-2S-mediated injury (5).

Animals. B6.C-H2<sup>d</sup>/bByJ (B6.H2<sup>d</sup>) mice (Cat# 000359) (46) and B6(Cg)-C1qa<sup>tm1d(EUCOMM)Wtsi/TennJ</sup> (C1qa<sup>-/-</sup>) mice (Cat# 031675) (22) originated from the Jackson Laboratory. B6-background mice were bred with B6.H2<sup>d</sup> mice and progeny were crossed to produce mice with homozygous expression of H2<sup>d</sup> MHC antigens for use in experiments (6). B6.ConK4<sup>-/-</sup> and B6.Tg(CD2-Tcra,-Tcrb)75Bucy (TCR75) mice were provided by James Zimring (University of Virginia, Charlottesville, VA) (21, 47). BALB/c mice were from Charles River Laboratories (Cat# 028). B6;SJL-Tg(FCGR2A)11Mkz/J mice (expressing human FCGR2A isoform R131, Jackson Laboratory Cat# 003542) (29) were backcrossed to B6 congenicity (48). Mice were studied at 8-16 weeks of age after maintenance in the UCSF specific pathogen-free animal facility for at least 2 weeks.

Surface plasmon resonance. Binding affinities were determined by injecting serial dilutions (0.5-200 nM) of MHC class I monomers (MBL International Cat#: TB-5001-M (K<sup>b</sup> presenting SIINFEKL); TB-5008-M (D<sup>b</sup> presenting RAHYNIVTF); TB-M552-M (K<sup>d</sup> presenting VYLKTNVFL); TB-M536-M (D<sup>d</sup> presenting IGPGRAFYA); TB-M521-M (L<sup>d</sup> presenting SPSYVYHQF)) over monoclonal antibodies bound via amine coupling to SensEye G Easy2Spot sensors (Ssens Cat# 1-09-04-006), assayed in triplicate with an IBIS MX96 SPR imager.

Antibody-mediated acute lung injury model. As previously described, mice were given i.p. injections of LPS (Sigma Aldrich Cat# L2880, 0.1 mg/kg) for ‘priming’ needed to render barrier-housed mice responsive to antibody injections (19). At 24 hours after LPS priming, mice were anesthetized (60 mg/kg ketamine + 40 mg/kg xylazine i.p.) and the indicated antibody treatments were injected into the jugular vein over the course of 1 minute (at 1 mg/kg unless otherwise stated). Antibodies were from BioXCell (mlgG2a isotype control clone C1.18.4 Cat# BE0085, anti-MHC class I clone 34-1-2S Cat# BE0180, hlgG1 isotype control Cat# BE0297) or newly produced (described below). Lung vascular permeability was measured by giving each mouse 0.01 KBq of <sup>131</sup>I-conjugated albumin (Iso-Tex Diagnostics, NDC:50914-7731) together with i.v. antibody injections, collecting lungs and blood samples 2 hours later or at cessation of breathing for radioactivity measurements used to quantify volume of extravasated plasma in lungs (lung vascular permeability). Wet-dry weight ratios of lungs and blood were used to calculate excess lung water volumes (6).

Immunofluorescence imaging. Cryosections were made at 200 µm or 400 µm thickness from lungs fixed by inflation with and immersion in 1% formaldehyde in PBS, as previously described (6). Sections
were incubated overnight with antibodies targeting C3b/d (clone 11H9, Novus Cat# NB200-540), C1qa (clone 4.8, Abcam Cat# ab182451), C4b/d (clone 16D2, Novus Cat# NB200-541), Scgb1a1 (Sigma-Aldrich Cat# ABS1673), CD41 (clone MWReg30, Biolegend Cat# 133902), or S100a8 (R&D Systems Cat# AF3059) together with a FITC-conjugated antibody raised against Acta2 (clone 1A4, Sigma-Aldrich Cat# F3777), all at 1:500 with 5% normal donkey serum, 0.1% bovine serum albumin and 0.3% triton X-100 in phosphate-buffered saline (PBS). After washing, Cy3 or Alexa Fluor 647-conjugated cross-adsorbed polyclonal secondary antibodies targeting rat, rabbit, goat and/or mouse IgG (Jackson Immunoresearch Cat# 712-165-153, Cat# 711-165-152, Cat# 705-605-147 and/or Cat# 115-605-206) were incubated with sections at 1:500 in PBS + 0.3% triton X-100 overnight. After additional washes, sections were either mounted in Vectashield (Vector Laboratories Cat# H-1700) for standard confocal imaging on a Nikon A1r microscope, or cleared after staining using the EZ clear protocol (49) for 3D imaging with a Nikon AZ100M confocal microscope.

**Antibody carbamylation.** Carbamylation of 34-1-2S was achieved by incubating 1 mg of antibody in PBS + 0.1 M KOCN for 1 hour at 37°C before buffer exchange back into PBS (24). Control 34-1-2S was subjected to the same process with omission of KOCN.

**Antibody sequencing and engineering.** The 34-1-2S hybridoma was purchased from ATCC (Cat# HB-79). To generate the Kd1 hybridoma, B6 mice were injected i.v. with $3 \times 10^6$ CD4+ T cells from TCR75 mice transgenic for a T cell receptor (TCR) specific for a peptide from K^d^ presented by IA^b^, and i.p. with $5 \times 10^6$ Con-K^d^-on splenocytes, resulting in an extreme B cell antibody response directed at an immunodominant peptide from K^d^, the only mismatched antigen between donor and recipient. Three days after a boost with an additional i.p. injection with Con-K^d^-on splenocytes, splenocytes from the sensitized recipient were fused with a myeloma cell line as previously described (50), and monoclonal antibodies specific for K^d^ were identified using Con-K^d^-on splenocytes as targets (21).

Monoclonal antibody aliquots were digested with either peptidyl-Asp metalloendopeptidase, chymotrypsin, elastase, trypsin, or pepsin enzymes. Peptides were then assayed using liquid chromatography coupled to tandem mass spectrometry for sequencing of variable fragments (Bioinformatics Solutions Inc.). Amino acid sequences determined were, for 34-1-2S:

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EVQLQSGAELFVPRGASVQLSLTCTASGFNLKDDYMFWVKQRPEQRLIYATYSLDSGVPKRFSGSRSG
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(DI)MTQQSPSSLSAELGERVSQTLRCSQDASNLNWLQEPDGTIKRLIYATYSLLDSGVPKRFSGSRSGS

DYSLTISSEDFVDDYCLQASSPYFTGGGTKEIK (light chain variable region); and for Kd1:

```
EVLLVESGGGLVKPGRGLSCLCAASGFTFRTYAMSWVRQTPEKRELVATIGDDGSYTFYPDNVKGR
```
FTISRDNAKNLQLMRHKSEDATIYFCARDGLFAYWQGTVSA (heavy chain variable region) and:
DIQMTIQSPSSLSAGGGKVITCKASQDIKKNIAWYQYKPGKPRLIHYTSLQPGLISSRFSGSGR
DYSFSISNLPEPIATYYCLQYDSLLYTFGGGTKLEIK (light chain variable region). Correct
identification of variable domains was confirmed by performing sequencing of the products of 5’ rapid
amplification of cDNA ends (5’-RACE) for heavy and light chain mRNA using RNA isolated from
hybridomas. The isolated 5’-RACE amplicons contained open reading frames that encoded the above
peptides sequenced by mass spectrometry. These sequences were codon-optimized and antibodies
were expressed as chimeric hIgG1 with or without Fc point mutations in a HEK293 cell system and
purified using protein A and buffer exchange (Absolute Antibody).

Pharmacologic treatments. Recombinant subdomain B from Staphylococcus aureus (SpA-B, amino acid sequence:
HHHHHADNKFNKIQNNAFYEILHLPNLNEEQRNGFIQLDDPSQSANNLAEMKLNDAQPK, His-tag added for purification) was produced in an E. coli system by Genscript and supplied in protein
storage buffer (50 mM Tris-HCl, 150 mM NaCl, 10% Glycerol, pH 8.0). At 1-2 hours before i.v. injection,
SpA-B (3 mg/kg) or vehicle were mixed with hIgG1-34-1-2S resulting in a 30% vehicle, 70% PBS
mixture. Trial formulations of CSL777 (in PBS vehicle), as well as clinical-grade IgPro20 (Hizentra™)
and the proprietary vehicle for IgPro20 were provided by Rolf Spirig (CSL Behring, Bern, Switzerland).
Mice were given i.p. injections of CSL777, IgPro20 or relevant vehicle 1-2 hours before i.v. injections of antibodies at stated doses.

Negative stain electron microscopy. Antibody samples were diluted to 0.01 mg/ml in 25 mM HEPES,
100 mM NaCl and added to carbon-coated grids (TedPella Cat# 01702-F, manually coated with 20 nm
carbon using a Cressington 208 Sputter Coater). Sample-coated grids were stained using 0.75% uranyl
formate and imaged on an FEI Tecnai T12 transmission electron microscope.

Pulse oximetry. Peripheral blood oxygen saturation (SpO₂) was measured using a MouseOx+ pulse
oximeter with collar sensor (Starr Life Sciences), with mice breathing room air during recordings.
Baseline measurements were taken one day after LPS priming. Mice were then given an i.v. injection of
hIgG1-34-1-2S under brief isoflurane anesthesia, followed by additional readings taken by averaging
recordings collected over 1 minute sampling periods every 5 minutes over the following 20 minutes.

Experimental design. Within-cage randomization was used for group allocations in studies testing
exogenous treatments. Littermate controls from heterozygous crosses were used to test the effect of
C1qa knockout and hFCGR2A expression. Congenic animals housed in the same room were used to
study haplotype effects. Handlers were blinded to group allocations during experiments. Group numbers (n) and analysis approaches were predetermined before initiation of experiments.

Statistics. Statistical tests used on each dataset are described in figure legends. P values and multiplicity-adjusted \( P \) values (\( P_{\text{adj}} \)) above 0.0001 are reported in figures, with asterisks used to highlight \( P \) values less than 0.05 (* = \( P<0.05 \), ** = \( P<0.01 \), *** = \( P<0.0001 \)).

Software. GraphPad Prism and InVivoStat were used for graphing and statistical analysis. UCSF ChimeraX was used for molecular graphics (51). Imaris was used to render fluorescence micrographs and ImageJ was used to process electron microscopy data.

Study approval. Procedures received ethical approval from the UCSF institutional animal care and usage committee.

Data availability. Values for all data points in graphs are reported in the Supporting Data Values file.
Author contributions

SJC, JCZ and MRL conceptualized research. SJC, YS, JJT, NK, DPB and AEHB contributed to data acquisition and analysis. SJC, MRL, AEHB, GV, ÉB, RS and JCZ developed methodology. SJC, JCZ and MRL acquired funding. SJC, JCZ and MRL wrote the original draft. All authors contributed to review and editing.

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Conflict of interest statement

RS is an employee of CSL Behring, Bern, Switzerland. The authors have no additional conflicts of interest.

Supplemental materials

Supplemental materials and methods.
Supplemental Movie 1. Complement C4 split product deposition in the pulmonary vasculature.
Supplemental Figure 1. Binding of MHC class I monoclonal antibodies to MHC class I monomers.
Supplemental Figure 2. Extended characterization of effects of 34-1-2S carbamylation.
Supplemental Figure 3. Extended characterization of effects of hIgG1-34-1-2S injections and hFCGR2A expression.
Supporting data values file.
References


Figure 1. The 34-1-2S alloantibody binds to multiple MHC class I antigens to trigger acute lung injury.

(A). Schematic showing approach for measuring affinity of the MHC class I alloantibody 34-1-2S for MHC class I monomers using surface plasmon resonance (SPR).

(B). SPR sensorgrams showing detection of binding of 34-1-2S to the K\textsuperscript{b} MHC class I antigen from H2\textsuperscript{b} mice and the K\textsuperscript{d}, D\textsuperscript{d} and L\textsuperscript{d} antigens from mice with the H2\textsuperscript{d} haplotype.

(C). Classical MHC class I antigens present in B6, B6.H2\textsuperscript{d} and B6.Con-K\textsuperscript{d}-on mice with summary of results from (B).

(D). Lung vascular permeability and (E). excess lung water measurements from LPS-primed B6.H2\textsuperscript{d} mice given intravenous (i.v.) doses of either 34-1-2S or isotype control, versus B6.Con-K\textsuperscript{d}-on mice given i.v. 34-1-2S.

Depictions of IgG and MHC class I in (A-C) are based on protein data bank (PDB) entries 1HZH and 1RK1. (B, D and E) show means ± standard errors. \( P \) values adjusted for multiple comparisons (\( P_{\text{adj.}} \)) in (D and F) were from an ordinary one-way ANOVA with Dunnett's test for differences relative to B6.H2\textsuperscript{d} + 34-1-2S i.v. group, with data log\(_{10}\)-transformed prior to analysis.
Figure 2. Classical complement activation on the pulmonary endothelium initiates 34-1-2S-induced acute lung injury.

(A) Molecular model of C1 complex based on small angle scattering database entry SASDB38 (12).

(B) Lung vascular permeability and (C) excess lung water measurements from LPS-primed B6.H2\textsuperscript{d}:C1qa\textsuperscript{-/-} mice and B6.H2\textsuperscript{d}:C1qa\textsuperscript{+/+} littermates given i.v. 34-1-2S at 1 mg/kg. Horizontal gray lines are standard deviations of values from 'no injury' controls (B6.H2\textsuperscript{d} mice given LPS i.p. + mlgG2a isotype control i.v.).

(D) Survival of LPS-primed B6.H2\textsuperscript{d}:C1qa\textsuperscript{-/-} mice and B6.H2\textsuperscript{d}:C1qa\textsuperscript{+/+} littermates given i.v. 34-1-2S at 4.5 mg/kg.

(E) Immunofluorescence staining for complement C3, (F) C1qa or (G) C4/C4b/C4d (red) as well as Acta2 (α-smooth muscle actin, cyan) and, in (G), Scgb1a1 (club cell secretory protein, magenta) in lung sections from LPS-primed B6.H2\textsuperscript{d}:C1qa\textsuperscript{-/-} mice and B6.H2\textsuperscript{d}:C1qa\textsuperscript{+/+} mice fixed 5 minutes after i.v. 34-1-2S at 1 mg/kg. Images in (G) are maximum intensity projections sampling 240 µm from a cleared thick section. White arrowheads point to arterioles positive for complement components.

(B and C) show means ± standard errors. P values are from unpaired two-tailed t-tests on log\textsubscript{10}-transformed data (B and C) or log-rank test (D), with group n=12 or for immunofluorescence staining representative of n=3 samples per group.
Figure 3. Inhibiting IgG hexamer assembly reduces 34-1-2S-induced acute lung injury responses.
(A) Molecular models of IgG hexamers based on PDB entry 1HZH, showing Fc:Fc and Fc:C1q interaction sites.
(B) Molecular model showing lysine residues in mouse IgG2a (mlgG2a), PDB entry 1IGT.
(C) Lung vascular permeability, (D) excess lung water measurements and (E) lung complement C3 and mlgG immunostains from LPS-primed BALB/c mice after i.v. injection of carbamylated or control 34-1-2S.
(F) Molecular model showing location of Fc domain lysine 439 (K439) in human IgG1 (hlgG1), PDB entry 1HZH.
(G) Lung vascular permeability, (H) excess lung water measurements and (I) lung complement C4/C4b/C4d and Acta2 immunostains from LPS-primed B6.H2d mice after i.v. injection with hlgG1-34-1-2S or hlgG1-34-1-2S with K439E mutation.
(J) Molecular model showing binding site of SpA-B to Fc domain of human IgG1 (hlgG1), PDB entries 1HZH and 5U4Y.
(K) Lung vascular permeability, (L) excess lung water measurements and (M) lung complement C4/C4b/C4d and Acta2 immunostains from LPS-primed B6.H2d mice after i.v. injection with hlgG1-34-1-2S and 75 µg SpA-B or vehicle control.
Samples for injury measurements were collected at 2 hours after antibody injections and lungs were fixed for immunostaining at 5 minutes after antibody injections. Graphs show means ± standard errors, with horizontal gray lines showing 95% confidence intervals of measurements from ‘no injury’ control mice given LPS and non-reactive isotype antibodies. $P$ values are from unpaired two-tailed t-tests on log$_{10}$-transformed data, with group $n=8$ (C, D), $n=12$ (G, H, K, L) or representative of 3 samples per group (E, I, M).
Figure 4. Fc mutations promoting IgG hexamer assembly increase in vivo pathogenicity of alloantibodies.

(A) Molecular model showing amino acids mutated in RGY-hlgG1 antibodies, based on PDB entry 1HZH.

(B) Negative stain electron micrographs showing single hlgG1-34-1-2S molecules and spontaneous solution-phase hexamers formed by RGY-hlgG1-34-1-2S (colored overlay highlights structures in expanded images).

(C) Lung vascular permeability and (D) excess lung water measurements from LPS-primed B6.H2d mice injected with control or RGY-mutated hlgG1-34-1-2S monoclonal antibodies (mAbs) at i.v. doses of either 0.3 or 1 mg/kg.

(E) Immunofluorescence staining showing pulmonary arterioles stained for complement C4/C4b/C4d (red) and Acta2 (cyan) in lung sections from LPS-primed B6.H2d mice given indicated treatments, representative of 3 samples per group fixed 5 minutes after antibody injections.

(F) Lung vascular permeability and (G) excess lung water measurements from LPS-primed B6.H2d mice injected with control or RGY-mutated hlgG1-Kd1 (a novel mAb targeting only the H-2Kd MHC class I antigen) at 1 mg/kg i.v.

Graphs show means ± standard errors with horizontal line representing 95% confidence intervals from ‘no injury’ controls (LPS-primed B6.H2d mice given hlgG1 isotype control i.v.). Log10-transformed data were analyzed using an ordinary two-way ANOVA with Šídák’s multiple comparisons test for effect of Fc mutation within dose level (C and D) or unpaired two-tailed t-test (F and G), with group n=12.
Figure 5. Treatment with recombinant Fc hexamer decoys prevents alloantibody-mediated acute lung injury.

(A) Molecular representation of CSL777, an investigational recombinant Fc hexamer ‘decoy’ treatment which inhibits classical complement activation, based on PDB entry 7X13 (52).

(B) Molecular representation of SCIg, a pooled human immunoglobulin therapeutic with anti-inflammatory properties at high doses, based on PDB entry 1HZH.

(C) Lung vascular permeability and (D) excess lung water measurements from LPS-primed BALB/c mice given i.p. vehicle or CSL777 at indicated doses 1 hour before i.v. injection with 34-1-2S or mlgG2a isotype control.

(E) Lung vascular permeability and (F) excess lung water measurements from LPS-primed BALB/c mice given i.p. vehicle or 2000 mg/kg SCIg 1 hour before i.v. injection with 34-1-2S or mlgG2a isotype control.

(G and H) Immunofluorescence showing pulmonary arterioles stained for complement C4/C4b/C4d (red) and Acta2 (cyan) in lung sections from LPS-primed BALB/c mice given indicated treatments, representative of 3 samples per group fixed 5 minutes after antibody injections. White arrowheads point to arterioles with endothelial positivity for C4/C4b/C4d.

Graphs show means ± standard errors with horizontal line representing 95% confidence intervals of data from ‘no injury’ controls (from vehicle + isotype control group). Log$_{10}$-transformed data were analyzed using either (C and D) an ordinary one-way ANOVA with $P$ values from (C and D): Dunnett’s multiple comparisons test for difference relative to vehicle + 34-1-2S group, or (E and F): two-tailed unpaired t-test, with group n=10.
Figure 6. Acute lung injury is complement-dependent in a model incorporating human FCGR2A-mediated pathology. 

(A) Lung vascular permeability, (B) excess lung water and (C) survival readouts from LPS-primed B6.H2d:hFCGR2aTg0 mice and littermate controls lacking hFCGR2A expression given i.v. hlgG1-34-1-2S at 1 mg/kg. 

(D) Immunofluorescence imaging of platelet sequestration (CD41, red, with Acta2 in cyan) and (E) neutrophils (S100a8, red, with Acta2 in cyan) in lungs from LPS-primed B6.H2d:hFCGR2aTg0 mice and littermates without hFCGR2A expression fixed at 20 minutes after hlgG1-34-1-2S injections, quantified in (F and G). 

(H) SpO2 measurements from LPS-primed B6.H2d:hFCGR2aTg0 mice and littermate controls without hFCGR2A expression before and after hlgG1-34-1-2S injections. 

(I) Lung vascular permeability, (J) excess lung water and (K) survival readouts from LPS-primed B6.H2d:C1qa+/+ and B6.H2d:C1qa-/- mice, as well as littermates of each genotype expressing hFCGR2A, given i.v. hlgG1-34-1-2S at 1 mg/kg. 

(A, B, F, G, H, I and J) show means ± standard errors with horizontal gray lines showing values from ‘no injury’ controls (baseline readings or values from B6.H2d mice given LPS i.p. + hlgG1 isotype control i.v.) and were log10-transformed prior to analysis. P values are from: (A, B, F and G) unpaired, two t-tests; (I and J) two-way ANOVA with Šídák’s multiple comparisons tests; (C and K) log-rank test; or (H) a two-way repeated measures mixed model approach with tests for main effect of genotype, and for post-baseline effects of genotype within time levels with Holm’s adjustment for multiple comparisons, with group n=4 (D-G), n=10 (H), or n=12 (other graphs).
Figure 7. Approaches to inhibit or mimic IgG hexamerization reduce antibody-mediated acute lung injury in mice expressing human FCGR2A.

(A) Lung vascular permeability, (B) excess lung water and (C) survival readouts from LPS-primed B6. H2d: hFCGR2aTg0 mice given i.v. hlgG1-34-1-2S at 1 mg/kg with either IgG hexamerization inhibitor SpA-B or vehicle.

(D) Lung vascular permeability, (E) excess lung water and (F) survival readouts from LPS-primed B6. H2d: hFCGR2aTg0 mice given either vehicle or 50 mg/kg CSL777 i.p. before i.v. hlgG1-34-1-2S at 1 mg/kg.

(A, B, D and E) show means ± standard errors with horizontal gray lines showing standard deviations of values from ‘no injury’ controls (B6. H2d mice given LPS i.p. + hlgG1 isotype control i.v.) and were log10-transformed prior to analysis. P values are from: (A, B, D and E) unpaired, two tailed t-tests 12 (other graphs); or (C and F) log-rank test, with group n=12.