

Engineered human antibodies for the opsonization and killing of Staphylococcus aureus

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Gram-positive organisms with their thick envelope cannot be lysed by complement alone. Nonetheless, antibody-binding on the surface can recruit complement and mark these invaders for uptake and killing by phagocytes, a process known as opsonophagocytosis. The crystallizable fragment of immunoglobulins (Fcy) is key for complement recruitment. The cell surface of S. aureus is coated with Staphylococcal protein A (SpA). SpA captures the Fcy domain of IgG and interferes with opsonization by anti-S. aureus antibodies. In principle, the Fcy domain of therapeutic antibodies could be engineered to avoid the inhibitory activity of SpA. However, the SpA-binding site on Fcy overlaps with that of the neonatal Fc receptor (FcRn), an interaction that is critical for prolonging the half-life of serum IgG. This evolutionary adaptation poses a challenge for the exploration of Fcy mutants that can both weaken SpA-IgG interactions and retain stability. Here, we use both wildtype and transgenic human FcRn mice to identify antibodies with enhanced half-life and increased opsonophagocytic killing in models of S. aureus infection and demonstrate that antibody-based immunotherapy can be improved by modifying Fcy. Our experiments also show that by competing for FcRn-binding, staphylococci effectively reduce the half-life of antibodies during infection. These observations may have profound impact in treating cancer, autoimmune, and asthma patients colonized or infected with S. aureus and undergoing monoclonal antibody treatment.

engineered antibody | C1q | Staphylococcus aureus | FcRn | immune evasion

mmunoglobulin G (IgG) accounts for about 75% of serum antibodies in humans and consists of four subclasses, IgG1 (66%), IgG2 (23%), IgG3 (7%), and IgG4 (4%) (1). The effector functions of IgG are contributed by the crystallizable fragment (Fc γ). Antibody engagement with Fc γ receptors (Fc γ Rs) on the surface of professional leukocytes enhances phagocytic uptake of marked pathogens, while Fc recruitment of complement component C1q initiates the activation of the classical complement pathway (2). IgG1 and IgG3 display the highest affinity toward FcyRs; IgG3 displays the strongest interaction with C1q followed by IgG1 (1). Human IgG1 (hIgG1) is often preferred for the development of therapeutic antibodies over hIgG3 owing to its stability (1). Following pinocytosis, IgG1 binding to the neonatal Fc receptor (FcRn) at the slightly acidic pH of endosomes allows for its recycling to the cell surface and its release from FcRn at the neutral extracellular pH (1). IgG3 fails to escape lysosomal degradation because of altered binding to FcRn and competition with IgG1 (3).

C1q is the recognition module of C1 complement complex that includes the serine proteases C1r and C1s. The weak affinity of C1q for monomeric IgG increases when its six globular domains interact with IgG hexamers (4-6), a clustering that is facilitated by IgG binding to cell surface antigens. The ensuing activation of C1r and C1s results in the generation of C3 convertases that catalyze the covalent modification of cell-surface determinants with opsonins such as C3b (2, 7). Most cells marked with C3-derived opsonins can be lysed by the membrane attack complex (MAC) (i.e., lytic pores made of five different complement

proteins) (7). Because of their thick peptidoglycan, gram-positive bacteria cannot be lysed by MAC (8) but can be taken up in a complement receptor (CR)-dependent manner by phagocytes (7). Bacteria have evolved immune evasion strategies to escape opsonization by antibodies and complement and uptake by phagocytes (9). The conserved Staphylococcal protein A (SpA) and Staphylococcal binder of immunoglobulin (Sbi) contain five and two immunoglobulin-binding domains (IgBDs), respectively (10). The IgBDs fold into triple-helical bundles that associate with the Fcy domain of human and vertebrate immunoglobulins (11, 12). The extensive tethering of SpA to peptidoglycan results in the capture and inhibition of IgG at the cell surface (13). SpA interaction with Fcy also interferes with FcRn-binding (3, 14). However, the physiological implications of this interaction are not known. Lastly, SpA binds with the variant heavy chain of V_H3-IgM, -IgG, -IgD, and -IgE (10). When released from the bacterial envelope (15), SpA binding to V_H3-IgM B cell receptors promotes B-cell expansion and the transient flooding of nonspecific V_H 3-IgM and V_H 3-IgG (16–18).

Humans and vertebrates develop antibodies against molecular determinants of Staphylococcus aureus yet are unable to generate antibodies that bind to and neutralize the IgBDs of SpA and Sbi (17, 19, 20). By immunizing animals with a SpA variant that

Significance

Staphylococcus aureus invariably acquires resistance mechanisms against new antibiotics. The persistent colonization with S. aureus is the key risk factor for invasive disease and a driver for the evolution of antibiotic resistant isolates. Anti-S. aureus antibodies that could promote decolonization, prevent infection, or treat disease would alleviate the selection for drug resistance. The successful development of such antibodies is complicated by Staphylococcal protein A (SpA) in the envelope of S. aureus. SpA captures immunoglobulins via their constant region, preventing antibodies from initiating anti-staphylococcal activities. Here, we demonstrate that therapeutic anti-S. aureus antibodies can be engineered to avoid sequestration by SpA. Such antibodies display extended half-lives and improve bacterial uptake and killing by immune cells.

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Competing interest statement: The authors are inventors of patents describing therapeutic antibodies against Staphylococcus aureus. D.M. declares a competing financial interest as a founder of ImmunArtes LLC, a University of Chicago startup company that aims to develop vaccines and therapies against S. aureus infections.

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cannot bind IgG (19), we isolated earlier the mouse monoclonal antibody 3F6 with SpA-specific complementarity determining region (CDR) and introduced these sequences into hIgG1 to yield recombinant 3F6-hIgG1 (21). Here, we introduce amino acid substitutions in the Fc γ domain of 3F6-hIgG1 to block SpA- and Sbi-binding and generalize this approach using Tefibazumab (Tefi), a humanized antibody with specificity to the surface virulence factor, ClfA (22). The Fc γ -engineered antibodies were evaluated for their ability to bind C1q and FcRn in vitro and tested for blood half-life and opsonophagocytic activity of *S. aureus* in several models of disease.

Results

Engineering the Fcy Domain of 3F6-hlgG1 to Escape SpA- and Sbi-Binding. Although 3F6-hIgG1 can neutralize SpA via its 3F6 paratope (23), its Fcy is still a ligand for SpA and Sbi. Each IgBD of SpA contains one 3F6 epitope and one Fcy-binding site. Thus, full-length SpA contains 10 possible binding sites for 3F6 with two distinct binding domains (23). Half-maximal effective concentration (EC₅₀) values measured by enzyme-linked immunosorbent assay (ELISA) represent the sum of all these interactions. The EC50 value of 3F6-hIgG1 for SpA was 4.11 times lower than for SpAKKAA (Fig. 1A and SI Appendix, Table S1). In this study, SpA and SpA_{KKAA} refer to antigens composed of the five IgBDs of SpA. SpA_{KKAA} has been modified with 20 amino acid substitutions to disrupt Fc γ and V_H3 antigen-binding fragment (Fab) binding; in the conventional IgBD nomenclature, these residues correspond to Q9,Q10 changed to K9,K10 and D36,D37 changed to A36,A37 (19). When offered to wild-type S. aureus strain Newman (Newman) or the isogenic mutant strain with the spa_{KKAA} sequence replacing the wild-type spa gene on the chromosome (spa_{KKAA}) , lower concentrations of 3F6-hIgG1 were required to saturate binding sites on spa_{K-1} KAA bacteria as compared to Newman (Fig. 1B and SI Appendix, Table S1). Of note, Newman and the spa_{KKAA} variant strain produce the same amount of protein A (13). No binding was observed with the Δspa control strain deleted for the spa gene (Fig. 1B). When 3F6-hIgG1 was coated to ELISA plates, SpA and SpAKKAA displayed distinct binding saturation (relative Bmax for SpA 1.911 versus SpA_{KKAA} 0.771) (Fig. 1C and SI Appendix, Table S1); this is in agreement with the notion that SpA contains twice as many binding sites for 3F6 as compared to SpA_{KKAA}. The Fc interaction with SpA can be gleaned from the atomic structure of SpA fragments bound to human Fcy (11, 12). Four hydrogen bonds promote interactions between residues Q9, Q10, N11, and Y14 of SpA IgBD-B [also conserved in Sbi (23)] and residues S254, Q311, N434, and L432 of hIgG1, respectively. Single and combined amino acid substitutions (S²⁵⁴A, Q³¹¹E, L^{432} S, or N⁴³⁴P) were generated yielding antibody variants 3F6hIgG1^A, 3F6-hIgG1^E, 3F6-hIgG1^S, 3F6-hIgG1^P, and 3F6-hIg-G1^{AESP}. A sixth variant 3F6-hIgG1^R with amino acid substitution $\mathrm{H}^{435}\mathrm{R}$ was generated, because the common IgG3 allotype with arginine at position 435 avoids SpA-binding (3, 24, 25). Of note, the same allotype also accounts for altered binding to FcRn (3). The substitutions ($S^{254}A$, $Q^{311}E$, $L^{432}S$, $N^{434}P$, and $H^{435}R$) did not affect CDR-binding to SpAKKAA (SI Appendix, Fig. S1A). Next, immobilized SpAKKAA was incubated with test antibodies to form immune complexes (IC), and histidine-tagged SpAAA (His-SpAAA, a variant that can only bind hIgG1-Fc) (19) was added to preformed ICs. Bound His-SpAAA was detected with a nickelactivated derivative of horseradish peroxidase (HRP). SpA_{AA} could no longer bind to ICs formed between 3F6-hIgG1^{AESP} or 3F6-hIgG1^R (Fig. 1D). Unlike 3F6-hIgG1, 3F6-hIgG1^{AESP} and $3F6-hIgG1^R$ interacted similarly with SpA and SpA_{KKAA} (Fig. 1*E* and SI Appendix, Table S1), with Sbi and Sbi_{KKAA} (SI Appendix, Fig. S1B), and with Newman and spa_{KKAA} bacteria (Fig. 1F and SI Appendix, Table S1). As a control, test antibodies did not interact with Δspa bacteria (SI Appendix, Fig. S1C). When 3F6-hIgG1^{AESP} and 3F6-hIgG1^R were coated to ELISA plates, the binding curves for both SpA and SpAKKAA exhibited similar saturation patterns (relative Bmax 0.969 versus 0.870) and similar EC₅₀ values (Fig. 1 G and H and SI Appendix, Table S1) in contrast to 3F6-hIgG1 (Fig. 1C). K_D measurements were also obtained using surface plasmon resonance (SPR) to compare antigen (SpA_{KKAA})antibody interactions (SI Appendix, Fig. S1D). None of the substitutions in the Fc domain interfered with epitope binding; $K_{\rm D}$ values of 45, 38, and 29 pM were obtained for 3 F6-hIgG1, 3 EC₅₀ values obtained by ELISA (SI Appendix, Table S1). Together, the experiments show that SpA associates with both the Fc and CDR of 3F6-hIgG1 and that the antibody can be engineered to redirect SpA binding to CDR exclusively. To corroborate these results, the AESP and R substitutions were introduced in the sequence of Tefi, a hIgG1 monoclonal antibody that binds the A domain of Clumping factor A (ClfA-A) of S. aureus. The Fc substitutions did not significantly alter the binding to ClfA-A (*SI Appendix*, Fig. S1*E*), and Spa interacted only marginally with the modified Tefi^{AESP} and Tefi^R antibodies possibly through V_{H3} residues (Fig. 1*I* and *SI Appendix*, Table S1). Similarly, Sbi interacted marginally with Tefi^R (Fig. 1*J* and *SI Appendix*, Table S1).

Engineering the Fcy Domain of 3F6-hlgG1 to Restore Complement Activation. SpA binding to Fcy prevents Fc-Fc association and, consequently, the formation of antibody hexamers needed for the recruitment of C1q to the bacterial surface (26). This was reflected by a 23-time-higher EC₅₀ value for 3F6-hIgG1 interaction with hC1q in the presence of SpA as compared to SpA_{K-1} KAA (Fig. 2A and SI Appendix, Table S2). Similarly, SpA but not SpAKKAA reduced interactions between the mouse IgG2a antibody, 3F6-mIgG2a, and mouse C1q (SI Appendix, Fig. S2A and Table S2). As predicted, SpA no longer blocked the association of hC1q with Fcy-engineered 3F6-hIgG1^{AESP} and 3F6-hIgG1^R; in fact, hC1q binding to the modified AESP/R antibodies increased in the presence of SpA or SpA_{KKAA}, as compared to hC1q binding to the unmodified 3F6-hIgG1 in complex with SpA_{KKAA} (Fig. 2A and SI Appendix, Table S2). When tested with staphylococci, recruitment of purified hC1q was dramatically increased following opsonization of Newman cells with 3F6-hIg-G1^{AESP} and 3F6-hIgG1^R; only a modest hC1q increase was observed in the presence of 3F6-hIgG1 as compared to hIgG1 (Fig. 2B). All three antibodies (3F6-hIgG1, 3F6-hIgG1^{AESP}, and 3F6-hIgG1^R) recruited C1q equally well when incubated with spa_{KKAA} cells. As a control, hC1q binding was reduced to background levels for Δspa bacteria (Fig. 2B). Similar results were obtained when normal human serum (NHS) was used as the source of C1q (SI Appendix, Fig. S2B). hC1q binding to Tefi was also reduced in the presence of SpA, but not of SpAKKAA (Fig. 2*C* and *SI Appendix*, Table S2). SpA did not block hC1q binding to Tefi^{AESP} and Tefi^R; of note, Tefi^{AESP} establishes weaker interactions with hC1q as compared to Tefi/Tefi^R (Fig. 2*C*). To evaluate whether C1q recruitment correlated with complement activation, rates of C3a and C4d production were recorded (Fig. 2 D and E and SI Appendix, Fig. S2 C and D). Upon incubation with bacteria, the rate of C3a production increased in the presence of 3F6-hIgG1 versus control hIgG1 and was further accelerated in the presence of 3F6-hIgG1^{RESP} and 3F6-hIgG1^R (Fig. 2D). C4d levels increased similarly within the first 10 min of interaction with staphylococci (SI Appendix, Fig. S2C). C3a production occurred at the same rates when using C1q-depleted serum (*SI Appendix*, Fig. S2D). C3a production was also increased in the presence of Tefi^R; in this assay, unmodified Tefi behaved like the isotype control, hIgG1 (Fig. 2*E*).

Next, we examined antibody interactions with $Fc\gamma Rs$. IgGbinding sites for SpA and $Fc\gamma Rs$ (at least $Fc\gamma I$ and $Fc\gamma RIIa$) in Fc do not overlap (14, 27). Thus, unlike with C1q, SpA is not



Fig. 1. hIgG1-Fc variants that escape SpA- and Sbi-binding. Antigen–antibody interactions were measured by ELISA. ELISA plates were coated as indicated on the figure panels: recombinant histidine-tagged SpA/SpA_{KKAA}/Sbi/Sbi_{KKAA} proteins (*A*, *E*, *I*, and *J*); Newman/*spa_{KKAA}* bacteria (*B* and *F*); 3F6 or its variants (*C*, *G*, and *H*); or ICs that were preformed between antigen (SpA_{KKAA}) and antibody variants (with single or quadruple substitutions in the Fc region) as shown in the drawing (*D*). Bound antibodies and proteins were detected with HRP-conjugated secondary antibody and His-HRP, respectively, and absorbances were recorded at 450 nm (A₄₅₀). Data are presented as mean \pm SEM (n = 3 assays) and are representative of at least two independent experiments.

thought to affect interactions between human Fc γ Rs (hFc γ Rs) and IgG. This was confirmed for six hFc γ Rs (*SI Appendix*, Fig. S2*E*). In fact, hFc γ Rs incubated with antibodies, including the non-V_H3 control IgG1, interacted with bacterial cells in a SpAdependent manner (*SI Appendix*, Fig. S2 *E* and *F*). All engineered antibodies interacted similarly with Fc γ Rs in this assay (*SI Appendix*, Fig. S2*E*). In conclusion, SpA blocks hC1q but not hFc γ R interactions with ICs. hC1q recruitment can be restored by introducing defined amino acid substitutions in the Fc region that do not affect interactions with antigen and hFc γ Rs. Engineering the $Fc\gamma$ Domain of 3F6-hlgG1 Improves Its Antistaphylococcal Activity. The therapeutic potential of antibodies can be evaluated in a mouse model of bloodstream infection (28). Consistent with previous results (29), animals that received 3F6-hlgG1 harbored reduced bacterial burdens (Fig. 3*A*) and fewer abscess lesions (Fig. 3*B*) in kidneys following challenge with methicillin-resistant *Staphylococcus aureus* (MRSA) strain MW2, as compared to animals that received hlgG1. Total bacterial loads were further reduced in animals treated with 3F6-hlgG1^R and 3F6-hlgG1^{AESP} (Fig. 3*A* and *B*). At day 15



Fig. 2. hlgG1-Fc variants that restore C1q-binding in the presence of SpA. (A–C) Antibody interactions with purified (human) hC1q were assessed in the presence of mock (PBS), SpA, or SpA_{KKAA} (A and C) or in the presence of bacteria: wild-type Newman or its variants, Δspa or spa_{KKAA} (B). ELISA plates were coated with serially diluted test antibodies (A and C) or bacteria (B). Bound hC1q was detected with HRP-conjugated secondary antibody, and absorbances were recorded at 450 nm (A_{450}). (D and E) Quantification of C3a production over time following incubation with bacteria (Newman 1 × 10⁸ CFU/mL) and 3F6 (D) or Tefi (E) test antibodies. NHS was used as the source of opsonins. In all panels, data are presented as mean ± SEM (n = 3 assays) and differences identified in B, D, and E by two-way ANOVA with Bonferroni posttests (**P < 0.01; *P < 0.05). One representative of at least two independent experiments is shown.

postinfection, serum concentrations of 3F6-hIgG1^R were significantly higher than that of 3F6-hIgG1 (SI Appendix, Fig. S3A). Administration of Tefi in this animal model afforded no protection; nonetheless, the single H⁴³⁵R substitution in Tefi-Fc boosted the activity of this antibody as reflected by a small but significant reduction of bacterial loads in kidney tissues (Fig. 3 C and D). Serum concentrations of Tefi^R were significantly higher than that of Tefi 15 d post infection (SI Appendix, Fig. S3B). To assess the opsonophagocytic killing (OPK) potential of test antibodies in humans, we developed earlier a whole-blood killing assay using freshly drawn anticoagulated blood from healthy volunteers. The canonical HL60-based OPK assay lacks the hemostasis factors (prothrombin and fibrinogen) that are exploited by S. aureus to agglutinate and avoid phagocytic uptake (30). Addition of 3F6-hIg $G1^{AESP}$ and 3F6-hIg $G1^{R}$ to anticoagulated human blood resulted in increased killing of MRSA isolates MW2 and USA300 as compared to addition of 3F6-hIgG1 (Fig. 3E and SI Appendix, Fig. S3C). As observed with animals (Fig. 3C), Tefi displayed no antistaphylococcal activity in human blood, but significant OPK activity was observed with Tefi^R (Fig. 3F). Clearly, the lack of OPK activity correlated with the presence of protein A in the envelope of bacteria since Tefi promoted the killing of spa_{KK4A} but not of Newman (spa⁺) in blood (SI Appendix, Fig. S3D). As a control, Newman and its spa_{KK44} variant displayed similar amounts of ClfA in their envelope (SI Appendix, Fig. S3E).

The whole-blood killing assay was repeated in the presence of anti-CR3 or anti-CR4 blocking antibodies. Myeloid cells produce various amounts of CR3 and CR4, and both share the ligand iC3b, the activation product of C3 on opsonized targets (7). Blocking CR4 had little effect on bacterial replication in general (Fig. 3*G*). Blocking CR3 resulted in increased bacterial replication in mock (hIgG1) samples, in agreement with the notion that CR3 is critical for the clearance of complement-opsonized *S. aureus* by effector cells (31–33). The OPK activity of all 3F6 antibodies was lost in the presence of anti-CR3, suggesting that these antibodies operate by enhancing the natural defenses of the host (Fig. 3*G*).

SpA Affects the Half-life of Antibodies In Vivo. Atomic structures of protein complexes revealed overlapping binding sites for SpA and FcRn with IgG-Fc, and in vitro binding studies demonstrated that SpA and FcRn are competitive ligands for IgG-Fc (3, 11, 14, 34). Nonetheless, the physiological significance of this competitive interaction is not known but implies that SpA could affect the half-life of antibodies during infection (SI Appendix, Fig. S3 A and B). SpA but not SpA_{KKAA} blocked interactions between hIgG1, hIgG2, and hIgG4 and hFcRn at pH 6.0 (Fig. 4A and SI Appendix, Table S2); hIgG3 that does not bind SpA was omitted from the analysis (SI Appendix, Fig. S4A). SpA also blocked interactions between mouse IgG (mIgG) and mouse FcRn (mFcRn) at pH 6.0 (Fig. 4B and SI Appendix, Table S2). The inhibitory activity correlated with SpA affinity for mIgG3 > mIgG2a > mIgG2b (SpA does not bind mIgG1) (SI Appendix, Fig. S4B and Table S1). Next, mice (n = 4) were injected with purified proteins (SpA/SpAKKAA) or bacteria (Newman/spa_{KK4A}), the latter of which mimicked the bloodstream infection displayed in Fig. 3A. A single injection of SpA in the bloodstream but not of SpAKKAA resulted in the transient



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Fig. 3. Antistaphylococcal activity of 3F6 and Tefi variants. (*A* and *B*) Enumeration of bacterial loads in kidney tissues (CFU) (*A*) and of tissue surface abscesses following 15 d infection with *S. aureus* MW2 of BALB/c mice (n = 20 animals, from two independent experiments) pretreated with antibody hlgG1, 3F6-hlgG1, 3F6-hlgG1^R, or 3F6-hlgG1^{AESP} at 5 mg.kg (body weight). (*C* and *D*) Enumeration of bacterial loads (C) and kidney surface abscesses (*D*) as described in *A* and *B* in animals pre-treated with antibody hlgG1, Tefi, or Tefi^R (n = 16 to 20 mice from two independent experiments). Data in *A–D* are presented as mean \pm SEM (*E–G*) The OPK activity of antibodies toward *S. aureus* MW2 was analyzed using anticoagulated freshly drawn human blood. Test antibodies in *E* included hlgG1, 3F6-hlgG1^R, and 3F6-hlgG1^R, and 3F6-hlgG1^{AESP} (n = 12 donors) and, in *F*, hlgG1, Tefi, and Tefi^R (n = 6 donors). In *G*, OPK (α -CR3) or anti-CR4 (α -CR4) (n = 6 donors). Data were plotted as the average \pm SEM of CFUs after 60-min incubation in blood compared to CFUs of inoculum (set as 100%). (*H*) Color code for test antibodies used in this figure. Significant differences were identified in *A–F* by one-way ANOVA with Tukey's multiple-comparison test and in *G* by two-way ANOVA with Bonferroni posttests (**P* < 0.05; ***P* < 0.01).

but noticeable reduction of mIgG2a and mIgG3 (Fig. 4C). Similarly, levels of mIgG2a and mIgG3 dropped in animals infected with Newman but not with spa_{KKAA} (Fig. 4D). Levels of mIgG1 and mIgG2b that are poor ligands for SpA were not affected (SI Appendix, Fig. S4 C and D). Rising levels of mIgG after day 5 of infection with Newman (Fig. 4D and SI Appendix, Fig. S4D) were attributed to the B-cell superantigen activity of SpA (i.e., the expansion of V_H3 antibodies as measured using antigen SpA_{KK}) (SI Appendix, Fig. S4E) (18). To study the effect of SpA on human antibodies, µ chain gene mutation (µMT) mice that lack mature B cells and the ability to produce antibodies (35) were injected with 100 µg hIgG1 or hIgG3, 12 h prior to intravenous challenge with phosphate buffered saline (PBS) (mock), Newman, or spa_{KK4A} (Fig. 4E). As compared to mock treatment, serum concentrations of hIgG1 and hIgG3 decreased slightly upon infection, presumably because S. aureus secretes the V8 protease that cleaves all hIgG (36, 37). Nonetheless, hIgG1 levels decreased much more rapidly in animals infected with Newman as compared to spa_{KKAA} (Fig. 4E). Such a difference was not observed for IgG3 since hIgG3-Fc is not a ligand for SpA (Fig. 4E). Next, we asked whether SpA alters the half-life of therapeutic antibodies through Fc interactions. 3F6-hIgG1 and the

engineered variants AESP and R were mixed with PBS, SpA, or SpA_{KKAA} and injected in wild-type mice (Fig. 4F). Serum concentrations of all three antibodies remained stable when injected in PBS (Fig. 4F). In contrast, serum concentrations of 3F6-hIgG1 decreased precipitously when injected with SpA (Fig. 4F). When SpA-Fc interactions were circumvented, such as when 3F6-hIgG1 was coinjected with SpAKKAA or engineered variants (AESP/R) were coinjected with SpA/SpA_{KKAA}, antibody clearance was less pronounced (Fig. 4F). We attribute this slower turnover to the uptake of productive antigen-antibody IC by phagocytes. This notion is supported by the increased antibody responses to 3F6-hIgG1 and its variants (SI Appendix, Fig. S4F). Injection of SpA with 3F6-hIgG1 that results in both productive (paratope-bound) and illegitimate (Fc-bound) complexes dampened antibody responses to 3F6-hIgG1 (SI Appendix, Fig. S4F). To further validate these observations, we used the therapeutic antibody Tefi that interacts with SpA though hIgG1-Fc exclusively. Coinjection of Tefi with SpA in animals resulted in enhanced clearance of the antibody; coinjection with SpAKKAA had no effect on the half-life of Tefi (Fig. 4G). Animals developed antibodies against Tefi injected with SpA but not with SpA_{KKAA} (SI Appendix, Fig. S4G). When tested in vitro, Tefi



Fig. 4. SpA influences the stability of antibodies in vivo. (*A* and *B*) Human and mouse hFcRn/mFcRn binding to human (*A*) and mouse (*B*) IgG at pH 6.0 was measured in the presence of PBS, SpA, or SpA_{KKAA} by ELISA (n = 3 assays). ELISA plates were coated with hIgG, mIgG, or mock (PBS). Bound biotinylated FcRn was detected with HRP-conjugated streptavidin, and absorbances were recorded at 450 nm (A₄₅₀). (*C*–*D*) Fold change over time of the serum concentration of mIgG2a and mIgG3 following injection of PBS, SpA, or SpA_{KKAA} (*C*) and infection with mock (PBS), Newman, or its *spa_{KKAA}* variant in BALB/c mice (*D*) (n = 4 animals per group). (*E*) hIgG1 and hIgG3 concentrations over time in µMT mice infected with mock (PBS), Newman, or *spa_{KKAA}* 12 h after administration of hIgG1/3 (n = 4 mice per group). (*F* and *G*) Serum concentrations over time of test antibodies following injection of PBS, SpA, or SpA_{KKAA} in BALB/c mice (n = 4 mice per group). (*H*) hFcRn binding to Tefi at pH 6.0 was measured in the presence of PBS, SpA, or SpA_{KKAA} by ELISA (n = 3assays). Bound FcRn was detected as in *A*. (*I*) Serum concentrations over time of test antibodies in BALB/c mice (n = 5 mice per group). (*A*) hFcRn binding to Tefi at pH 6.0 was measured in the presence of PBS, SpA, or SpA_{KKAA} by ELISA (n = 3assays). Bound FcRn was detected as in *A*. (*I*) Serum concentrations over time of test antibodies in BALB/c mice (n = 5 mice per group). Animals were infected with the Newman variants $\Delta clfa-spa_{KKAA}$ or PBS (mock) treated 12 h after the administration of test antibodies. (*J*) ICs of test antibodies Tefi/Tefi^R and cognate antigen ClfA-A were injected in BALB/c mice (n = 5 mice per group) along with PBS, SpA, or SpA_{KKAA}, and the remaining fraction of test antibodies was measured over time. Data are presented as mean \pm SEM and representative of at least two independent experiments (*A*, *B*, and *H*). Significant differences were identified by two-way ANOVA with Bonferr

interaction with hFcRn at pH 6.0 was inhibited by SpA but not by SpA_{KKAA} (Fig. 4*H* and *SI Appendix*, Table S2). To examine the impact of infection, animals treated with Tefi were challenged 12 h later with the isogenic $\Delta clfA$ -spa⁺ and $\Delta clfA$ -spa_{KKAA} variants or PBS (mock control). The *clfA* gene was deleted to eliminate antibody turnover caused by productive Tefi-ClfA ICs. Serum levels of Tefi dropped much more rapidly in animals infected with the *spa*⁺ ($\Delta clfA$ -spa₊) strain as compared to the *spa* defective strain ($\Delta clfA$ -spa_{KKAA}) (Fig. 4*I*). ICs formed between the ClfA-A antigen and Tefi or Tefi^R were coinjected in animals with SpA, SpA_{KKAA}, or PBS. The fastest clearance was observed for the ClfA-A-Tefi complex when injected with SpA (Fig. 4*J*), further emphasizing that Fc-SpA interactions reduce antibodies' half-life even when the antibody is bound to cognate antigen (Fig. 4*J*).

Engineering 3F6 Antibodies for Human Use. Ig-Fc mutations that decrease SpA-binding may also alter interactions with FcRn and the trade-off for escaping protein A-mediated immune evasion may be reduced antibody half-life. To investigate this possibility, interactions between purified antibodies and FcRn were first measured by ELISA. 3F6-hIgG1 and 3F6-hIgG1^R bound mFcRn similarly at pH 6.0 and 7.0, while 3F6-hIgG1^{AESP} displayed lower binding at both pH (*SI Appendix*, Fig. S5*A*). As compared to hIgG1or 3F6-hIgG1, 3F6-hIgG1^R and 3F6-hIgG1^{AESP} displayed a weaker association with hFcRn at pH6.0 (Fig. 5*A* and *SI Appendix*, Table S1). *K*_D values for hFcRn at pH 6.0 were also obtained using SPR and measured as 538 nM for 3F6-hIgG1, 1,189 nM for 3F6-hIgG1^R, and 9,825 nM for 3F6-hIgG1^{AESP},

suggesting that 3F6-hIgG1^{AESP} may have a shorter serum halflife (SI Appendix, Fig. S5B). An earlier structure-guided design suggested that substitutions at residues 256, 286, 307, 311, and 378 may ameliorate antibody-FcRn interactions (38). Combinations RV ($T^{307}R/A^{378}V$), QVV ($T^{307}Q/Q^{311}V/A^{378}V$), and DDRVV ($T^{256}D/N^{286}D/T^{307}R/Q^{311}V/A^{378}V$) were introduced in four test antibodies 3F6-hIgG1^{AESP-RV}, 3F6-hIgG1^{R-RV}, 3F6-hIgG1^{R-QVV} and 3F6-hIgG1^{R-DDRVV} (*SI Appendix*, Fig. S5*C*). The substitutions did not interfere with hC1q/SpA-binding (Fig. 5B) and, except for R-DDRVV, did not alter antibody binding to Newman or spa_{KKAA} (SI Appendix, Fig. S5 D-F). When subjected to size exclusion chromatography, all 3F6 variants eluted as monomers (SI Appendix, Fig. S5G). QVV and DDRVV substitutions markedly increased 3F6-hIgG1^R binding to hFcRn at pH 6.0 but not at pH 7.0 (Fig. 5C). 3F6-hIgG1^{R-QVV} and 3F6hIgG1^{R-DDRVV} interacted strongly with mFcRn at pH 7.0 (Fig. 5D), which correlated with their rapid turnover (Fig. 5E and SI Appendix, Table S3) and increased immunogenicity in BALB/c mice (*SI Appendix*, Fig. S5H). The reduced half-life of 3F6-hIgG1^{R-QVV} in mice abrogated any gain of protection against MRSA observed with 3F6-hIg $G1^R$ (Fig. 5 F and G). Nonetheless, when OPK activity was tested in whole human blood, 3F6-hIgG1^{R-QVV} and 3F6-hIgG1^{R-DDRVV} performed better than 3F6-hIgG1 and similarly to 3F6-hIgG1^{AESP} and 3F6-hIgG1^R (Fig. 5H). Next, humanized FcRn Tg32 mice (39) were used to measure the half-life and therapeutic activity of test antibodies. 3F6-hIgG1^{AESP} that failed to bind hFcRn in vitro was rapidly turned over in Tg32 animals, while R435 variants behaved as the parent 3F6 antibody (Fig. 6A).



Fig. 5. Amino acid substitutions for improved interactions with hFcRn. (*A–D*) ELISA-binding curves (n = 3 assays) of hFcRn (at pH 6.0 and 7.0) (*A* and *C*), hC1q (*B*), or mFcRn (pH 6.0 and 7.0) (*D*) to test antibody variants including control hIgG1 and 3F6-hIgG1. ELISA plates were coated with serially diluted test antibodies (color code in *f*). (*B*) ELISA-binding curves of purified hC1q to test antibodies were assessed in the presence of SpA or SpA_{KKAA} (n = 3 assays). Binding curves were mostly superimposable. (*E*) Serum concentrations over time of 3F6-hIgG1 and variants (AESP, R, R-QVV, and R-DDRVV) following injection in BALB/c mice (n = 4 mice per group). (*F* and G) Enumeration of bacterial loads (*F*) and kidney surface abscesses (G) as described in Fig. 3 in animals pretreated with antibody hIgG1, 3F6-hIgG1^R, or 3F6-hIgG1^{R-QVV} (n = 10 to 20 mice). Data are presented as mean ± SEM and analyzed with one-way ANOVA with Tukey's multiple-comparison test (***P* < 0.01; **P* < 0.05). (*H*) OPK activity of test antibodies (color code in *f*) toward *S. aureus* MW2 in anticoagulated freshly drawn human blood (n = 12 donors). Data were plotted as described in Fig. 3 and significance calculated with one-way ANOVA and Tukey's multiple-comparison test (**P* < 0.05; ***P* < 0.01). (*f*) Color code for test antibodies used in this figure. Data are presented as mean ± SEM and representative of two independent experiments (*A–D*).

Accordingly, Tg32 mice did not develop strong antibody responses against test antibodies (*SI Appendix*, Fig. S6S2). Since mIgG interacts poorly with hFcRn, Tg32 mice were also pretreated with hIgG to better mimic human conditions (40). All engineered antibodies retained high serum concentrations with the slowest turnover observed for 3F6-hIgG1^{R-QVV} (Fig. 6*B* and *SI Appendix*, Table S3) and were further tested for efficacy (Fig. 6*C* and *D*). In Tg32 mice unlike in BALB/c animals, 3F6-hIgG1 protected partially by reducing bacterial loads (but not abscess lesions) as compared to hIgG1 (Fig. 6 *C* and *D*). Whereas in BALB/c animals, 3F6-hIgG1^{R-QVV} provided less protection than 3F6-hIgG1^R against MRSA challenge (Fig. 5 *F* and *G*); in Tg32 mice, this difference was no longer observed. In fact, both 3F6-hIgG1^R, with a small but statistical improvement over 3F6-hIgG1 (Fig. 6 *C* and *D*). Therapeutic superiority correlated with enhanced serum concentration of antibodies (Fig. 6*E*).

Discussion

S. aureus is responsible for both healthcare- and communityassociated infections (41, 42). Infections are treated with antibiotics. However, the treatment of S. aureus infections can be complicated by disease severity, high recurrence, a dwindling antibiotic arsenal against methicillin-resistant strains, and treatment length (43). Some infections such as skin and soft-tissue collections also require incision and drainage. Failure to drain abscesses delays resolution and favors the development of antibiotic resistance (44). For all these reasons, it seems reasonable to expand efforts toward the development of antibodies that could aid in the treatment of S. aureus diseases. Several envelope determinants have been considered as targets of opsonophagocytic antibodies but failed in clinical trials (43). None of these approaches accounted for the ability of SpA to bind the Fc region of IgG, the effector region of antibodies that interacts with FcyRs, C1q, and FcRn. The possibility that SpA may affect all effector functions of antibodies and the

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Fig. 6. Antibody efficacy in the humanized FcRn (Tg32) mice. (*A* and *B*) Serum concentrations over time of 3F6-hlgG1^R and variants R-QVV and R-DDRVV following injection (5 mg/kg) in Tg32 mice without (*A*) or with (*B*) pretreatment with pooled hlgG at 100 mg/kg (n = 5 mice per group). (*C* and *D*) Enumeration of bacterial loads (*C*) and kidney surface abscesses (*D*) as described in Fig. 3 in Tg32 animals pretreated with antibody hlgG1, 3F6-hlgG1^R, 3F6-hlgG1^R. ^{QVV}, or 3F6-hlgG1^{R-DRVV} challenged for 15 d with *S. aureus* MW2 (n = 13 to 17 mice). (*E*) Serum concentration of test antibodies that remained after 15 d infection with *S. aureus* MW2. Sera were from animals shown in *C.* (*F*) Color code for test antibodies used in this figure. Data are presented as mean \pm SEM and analyzed with one-way ANOVA and Tukey's multiple-comparison test in *C–E* (***P* < 0.01; **P* < 0.05).

implication for the therapeutic activity of antibodies has not been addressed.

Here, we report that SpA binding to IgG-Fc in vitro obstructs further association with C1q and FcRn but not with recombinant Fc γ Rs. The Fc region of antibodies with CDRs directed against SpA (3F6) or ClfA (Tefi) were engineered to prevent SpA and Sbi interference with C1q recruitment. Antibody variants with AESP and R substitutions that disrupted SpA/Sbi interference were found to improve the therapeutic and OPK activities of 3F6 and Tefi antibodies in a mouse model of MRSA bloodstream infection and in human whole blood, respectively. Further, blocking CR3 prevented *S. aureus* killing in whole blood, suggesting that 3F6 variants enhanced iC3b opsonization to promote phagocytic uptake. We propose that the OPK activity of 3F6 candidate antibodies correlates with enhanced activation of C1q at the bacterial cell surface. We cannot rule out the involvement of other effector or complement activities.

Our experiments confirmed that SpA and FcRn compete for Fc binding in a manner that correlated with binding affinity to antibodies. SpA interacts with hIgG1 and mIgG2 with high affinity but fails to bind the Fc fragment of most other immunoglobulins including hIgG3, IgM, IgE, IgA, and mIgG1 that do not interact with FcRn and display shorter half-lives in vivo (3, 40, 45, 46). By competing for FcRn-binding, SpA effectively reduced the half-life of hIgG1 and mIgG2. Depletion was very effective and observed following injection of SpA in animals as well as during infection with S. aureus. We reported earlier that SpA is key to the colonization success of S. aureus in a mouse model (21, 47), implying that colonization could also accelerate the turnover of hIgG1. This evolutionary adaptation represents a challenge for the engineering of therapeutic antibodies. Breaking apart Fc-SpA interactions restores C1q recruitment at the cost of reducing the half-life of antibody. Unsurprisingly, the 3F6-hIgG1^R and 3F6-hIgG1^{AESP} variants displayed a weaker association with hFcRn at pH6.0 as compared to3F6-hIgG1. To circumvent this problem, amino acid substitutions shown to enhance hFcRnbinding (38) were introduced in the sequences of 3F6 variants. As the substitutions resulted in increased binding to mFcRn at 7.0 (indicative of short half-life), transgenic Tg32 animals with an mFcRn α -chain replaced with the hFcRn α -chain (39) were used for testing. Predictably, 3F6-hIgG1^{AESP} was rapidly turned over in Tg32 mice; surprisingly, 3F6-hIgG1^R displayed a similar half-life as 3F6-hIgG1. Variant 3F6-hIgG1^{R-QVV} was even more stable than 3F6-hIgG1; the variant afforded significant protection against bacterial burdens and abscess lesions in animal-infected tissues. We conclude that R-QVV substitutions may be best suited for the development of antibodies against *S. aureus* in humans.

All Fc γ Rs in their soluble form were found to interact with bacteria via nonspecific SpA–Fc complexes. It is unclear whether such interactions also occur between bacteria and immune cells and what the outcome of such interactions may be. Possibly, *S. aureus* exploits such interactions to its advantage. SpA released from bacteria and bound to Fc or to legitimate IgG–antigen complexes may also bind Fc γ Rs for uptake; once in the endocytic compartment, SpA will obstruct further FcRn binding. In antigen-presenting cells, FcRn interactions with large IgG ICs has been shown to mediate the secretion of inflammatory cytokines and enhance intracellular routing for further antigen processing (48). Thus, SpA-mediated FcRn avoidance may contribute multiple immune evasion strategies, including reduced half-life of hIgG1 antibodies and reduced immune responses.

In conclusion, it may be possible to develop therapeutic antibodies against surface determinants of *S. aureus*. Such determinants should be abundant, conserved, preferentially key to the pathogenic program of *S. aureus*, and the antibodies should be designed to escape neutralization by SpA and other proteins, such as Sbi that have evolved to bind Ig-Fc (10). SpA represents an ideal target for such antibodies.

Materials and Methods

Ethics Statement. The University of Chicago's Institutional Review Board reviewed, approved, and supervised the protocol used for all experiments utilizing blood from human volunteers, and informed-consent forms were obtained from all participants. Animal research was performed in accordance with institutional guidelines following experimental protocol review, approval, and supervision by the Institutional Animal Care and Use Committee at The University of Chicago. Experiments with *S. aureus* were performed in biosafety level 2 (BSL2)/animal BSL2 containment upon review by The University of Chicago Institutional Biosafety Committee.

Bacterial Strains, Mammalian Cell Lines, and Growth Media. *S. aureus* strains Newman and its variants (*spa_{KK}*, *spa_{AA}*, *spa_{KKAA}*, *Δspa*, *ΔclfA*-*spa*⁺, and *ΔclfAspa_{KKAA}*), USA300 LAC and its variant *Δspa*, and MW2 were grown in tryptic soy broth or agar at 37 °C. *Escherichia coli* strain DH5 α and BL21 (DE3) were grown in Luria broth or agar with 100 µg/mL ampicillin at 37 °C. Suspension serum-free adapted FreeStyleTM 293-F (human embryonic kidney [HEK]-293F) cells were cultured in FreeStyleTM 293 Expression Medium (Life Technologies) and maintained in a 5% CO₂ humidified incubator at 37 °C.

Construction, Expression, and Purification of 3F6-hlgG1 and Tefi Variants. Recombinant 3F6-hlgG1 cloned into the expression vector pVITRO1-102.1F10-IgG1/ $\!\lambda$ (Addgene no. 50366) was as described previously (29) and served as a template for further mutagenesis. The heavy- and light-chain genes of Tefi were synthesized from published sequences (22, 49) by Integrated DNA Technologies, Inc (IDT) and swapped into the pVITRO1 plasmid encoding 3F6-hIgG1. Primers 5'-AACCGCTACACGCAGAAGAGCCTCTC-3' and 5'-GTAGCGGTTGTGCA-GAGCCTCATGCAT-3' were used to generate the Tefi^R and 3F6-hlgG1^R variants. For other variants, partial Fc genes with specific mutation(s) were synthesized by IDT and swapped into the vector encoding Tefi or 3F6-hlgG1 using the polymerase incomplete primer extension (PIPE) method (21). PIPE products were transfected into HEK-293F cells using polyethylenimine and transfectants selected with hygromycin B (400 µg/mL) before expansion using Triple-Flask Cell Culture Flasks (Thermo Fisher Scientific). Antibodies were affinity purified from supernatants of expanded cultures on protein G resin (Gen-Script) and dialyzed against PBS as described (29). Dialyzed antibodies were separated by size exclusion high performance liquid chromatography using a Tricorn Superose 6 Increase 10/300 GL column. Briefly, up to 200 µL of antibody suspension at concentrations ranging between 2 and 3 mg/mL was injected in PBS pH 7.2 over the column. PBS at pH 7.2 was used as the eluent and run at 0.4 mL/min.

Enzyme-Linked Immunosorbent Assays. Binding measurements were performed in microtiter plates (Nunc Maxisorp) coated with either 10 nM test antibodies, 30 nM recombinant proteins (SpA_{KKAA}, SpA, Sbi, or Sbi_{KKAA}), or 1 \times 10⁶ colony-forming units (CFU) of S. aureus bacteria, in 0.1 M carbonate buffer (pH 9.5) at 4 °C overnight or at 37 °C for 2 h. Wells were blocked by 2% (wt/ vol) bovine serum albumin (BSA) before incubation with serial concentrations of test antibodies or His-tagged proteins. Binding capacity was quantified following incubation with HRP-conjugated goat anti-hlgG (Promega, W403B, 1:15,000) or with the HisProbe-HRP conjugate (Thermo Fisher Scientific, 15165). For competition experiments with C1q, microtiter plates were coated with serial dilutions of purified test antibodies overnight. After blocking, recombinant SpA or SpA $_{KKAA}$ was added along with 20 $\mu g/mL$ human or mouse C1g (CompTech, A099 and M099), and C1g binding was detected with HRP-conjugated sheep anti-C1q (Bio-Rad, 2221-5004P). The molar ratio of test antibody over recombinant SpA/SpA_{KKAA} was 5:1. To measure the binding of hC1q and FcyRs to bacteria, microtiter plates were first coated with bacteria at 37 °C for 2 h and blocked with BSA before adding 10 µg/mL test antibody and either 20 µg/mL hC1q, 10% NHS (CompTech), or 2 µg/mL His-tagged hFcγRs (SinoBiological). Bound hC1q binding was detected with HRP-conjugated sheep anti-C1q (Bio-Rad, 2221–5004P) and bound FcyR with the HisProbe-HRP conjugate (Thermo Fisher Scientific, 15165). For competition experiments with hFcRn or mFcRn, microtiter plates were coated with serial dilutions of purified test antibodies overnight. After blocking, 30 nM recombinant SpA or SpAKKAA was added to each well at either pH 6.0 or pH 7.0 along with 2 μ g/mL biotinylated FcRn (Immunitrack). Bound FcRn was detected with HRP-conjugated streptavidin (New England Biolabs). All plates were developed using Opt enzyme immunoassay (EIA) reagent (BD Biosciences). Recombinant SpA/Sbi proteins and their variants were purified from E. coli as described (19). Values of dissociation constants reported in this study are listed in SI Appendix, Tables S1 and S2.

SPR Analysis. SPR experiments were performed with a Biacore 8K instrument by the Biophysics Core of the University of Illinois Chicago. Ligands were bound to sensor chip nitrilotriacetic acid. Analytes were provided in PBS containing 0.05% Tween-20 and 1% BSA at indicated pH. The sensor-chip surfaces were activated with 2 mM nickel sulfate and regenerated with 300 mM ethylenediamine tetraacetic acid, respectively. In total, 500 nM of test articles (SpA, SpAKKAA, and hFcRn) were immobilized at a flow rate of 25 μ L/min. To measure interactions with SpA or SpA_{KKAA} ligands, analytes (3F6-hlgG1 and its variants) were used at a concentration range of 10 to 0.045 nM in a threefold serial dilution. To measure interactions with hFcRn, analytes were used at a concentration range of 2.5 to 0.02 μ M with a threefold serial dilution. The association rates were measured at a constitution constants were determined from two independent experiments.

C3a Release Assay and C4d Measurement. In total, 1×10^8 CFU of live bacteria were suspended in 900 µL GVB++ buffer (CompTech, B100) with 100 µL NHS or C1q-depleted human serum (CompTech, A300) and incubated at 37 °C for 10, 30, or 60 min. Complement activation was quenched by adding 1 mL gelatin veronal sodium buffer with EDTA (CompTech, B105), and the samples were spun at 14,000 × g for 5 min. Amounts of C3a in supernatants were determined using the MicroVue C3a Plus EIA kit (Quidel, A031) according to the manufacturer's instructions. To measure C4d deposition in the envelope of staphylococci, reaction mixtures were removed after 0 and 10 min, spun at 14,000 × g for 5 min at 37 °C. C4d levels in cell lysates were determined using the MicroVue C4d EIA kit (Quidel, A009) according to the manufacturer's instructions.

Measuring Serum Concentrations of Antibodies in Mice. To determine the effect of SpA on mIgG, BALB/c mice (6 to 7 wk of age) obtained from Jackson Laboratory were injected into the peritoneal cavity with 100 μ g recombinant, endotoxin-free, vaccine-grade SpA or SpAKKAA or into the periorbital venous plexus with 1×10^7 CFU of live bacteria. After 0, 4, and 6 h and 1, 2, 3, 5, 8, 9, 11, and 15 d, periorbital venous blood was obtained, and serum samples analyzed by ELISA. Briefly, microtiter plates were coated overnight with $10,000 \times$ dilution of serum sample. After blocking, wells were incubated with goat anti-mlgG1, -mlgG2a, -mlgG2b, or mlgG3 (Novus Biologicals). Serum mlgG concentrations were calculated using a standard curve of purified mIgG1, mlgG2a, mlgG2b, and mlgG3 (Biolegend) at a range of 1 to 1,500 ng/mL Microtiter plates coated with recombinant SpA_{KK} (100 ng/well) were used to measure the V_H3-clonal mIgG in serum samples. To investigate the impact of infection on hIgG, 100 μ g of purified hIgG1 or hIgG3 (Sigma-Aldrich) were first injected into the peritoneal cavity of μ MT mice (6 to 7 wk of age). A total 16 h later, animals were infected with 1×10^7 CFU of live bacteria or mock (PBS) into the periorbital venous plexus. Serum samples were collected at 4 h and 3 and 11 d after antibody injection, and serum concentrations of hIgG were calculated using a standard curve of purified hIgG1 and hIgG3 at a range of 1 to 50 ng/mL. To examine the effect of soluble SpA/SpA $_{\mbox{\scriptsize KKAA}}$ on test the rapeutic antibodies, BALB/c mice (6 to 7 wk of age) were injected into the peritoneal cavity with 100 µg test antibody plus 4 µg recombinant, endotoxin-free, vaccine-grade SpA/SpA_{KKAA} (molar ratio 5:1). Control mice were injected with 100 μg test antibody plus PBS. After 4 h and 1, 2, 3, 7, 11, and 15 d, serum samples were collected, and serum concentrations of test antibody were detected in microtiter plates coated with matched antigens (SpA $_{KKAA}$ or ClfA-A). To examine the impact of infection on therapeutic antibodies, 100 µg of test antibodies Tefi/Tefi^R were first injected into the peritoneal cavity of BALB/c mice (6 to 7 wk of age). After 16 h, animals were challenged intravenously with mock (PBS) or 1×10^7 CFU of $\Delta clfA$ -spa⁺ or $\Delta clfA$ -spa_{KKAA}. Serum concentrations of test antibody were measured following serum sampling at 4 h and 1, 2, 3, 7, 11, and 15 d post infection using microtiter plates coated with ClfA-A (19). Tefi/Tefi^R concentrations in samples were calculated using a standard curve of purified Tefi/Tefi^R diluted in the mouse serum at a range of 0.1 to 150 ng/mL. To measure the persistence of 3F6-hlgG1 and its variants in Tg32 mice (B6.Cg-Fcgrt^{tm1Dcr} Tg(FCGRT)32Dcr/DcrJ), animals (n = 5, 6 to 7 wk of age) were either pretreated or not with hIgG (100 mg/kg body weight) before 100 µg test antibody was delivered by intraperitoneal injection. After 4 h and 1, 2, 3, 7, 11, 15, 24, and 31 d, periorbital venous blood was obtained, and test antibody in serum samples was captured by SpA_{KKAA} -coated plates. Serum antibody concentrations were calculated using a standard curve of each test antibody diluted in mouse serum at a range of 1 to 500 ng/mL Pharmacokinetic (PK) parameters were determined for each group of mice with a noncompartmental analysis model using PKSolver (50).

Therapeutic Evaluation of Test Antibodies in Animals. Animals (6 to 8 wk of age) were treated with 100 μ g of indicated test antibody 12 to 18 h before challenge with *S. aureus* USA400 (MW2). Bacterial cultures were grown to an absorbance at 600 nm of 0.42, and bacteria were washed in PBS once and

adjusted to a suspension of 6.5×10^7 CFU/mL. In total, $100 \,\mu$ L of this suspension was injected into the periorbital venous plexus of animals anesthetized with ketamine/xylazine (100 and 20 mg/kg). On day 15, mice were killed by carbon dioxide inhalation and necropsied to remove kidneys. Surface abscesses visible on intact kidneys were enumerated. The second kidney was weighed, homogenized, serially diluted, and plated on agar to count bacterial burden in tissues (CFU per gram of tissue).

Staphylococcal Survival in Human Blood. Test or control antibody (2.5 μ g) was added to 0.5 mL of freshly drawn human blood anticoagulated with 5 μ g/mL desirudin or heparin. Where indicated, 5 μ g/mL of anti-CR3 or anti-CR4 (BioLegend; Catalog no. 301302 or Catalog no. 301616) was added to the blood for 10 min prior to infection with bacteria. At time 0, a 50- μ l bacterial suspension in PBS (5 × 10⁶ CFU) was added to the blood, and tubes were incubated for 0 or 60 min at 37 °C. 0.5 mL streptokinase lysis buffer (PBS containing 0.5% saponin, 100 U streptokinase, 50 μ g trypsin, 1 μ g DNase, and 5 μ g RNase)

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was added to each sample for 10 min at $37 \,^{\circ}$ C prior to plating on agar for CFU enumeration. Assays were performed in duplicate and repeated for reproducibility.

Statistical Analyses. All statistical analyses were performed using GraphPad Prism version 7.04 (GraphPad Software, Inc., La Jolla). Statistically significant differences were calculated by using statistical methods, as indicated. *P* values of <0.05 were considered significant.

Data Availability. All study data are included in the article and/or SI Appendix.

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