Isotype Switching Increases Efficacy of Antibody Protection against Staphylococcal Enterotoxin B-Induced Lethal Shock and Staphylococcus aureus Sepsis in Mice

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ABSTRACT  Staphylococcal enterotoxin B (SEB) is a potent toxin that is produced by Staphylococcus aureus strains and is classified as a category B select agent. We have previously shown that monoclonal antibody (MAb) 20B1, a murine anti-SEB IgG1, successfully treats SEB-induced lethal shock (SEBILS) and bacteremia that is caused by SEB-producing S. aureus. In this study, we have generated two isotype switch variants of the original IgG1 MAb 20B1, an IgG2a and IgG2b, both bearing the same variable region sequence, and compared their neutralizing and protective activity in in vitro and in vivo assays, respectively. All 3 isotypes demonstrated comparable affinity to SEB and comparable 50% inhibitory concentrations (IC50s) in T cell proliferation assays. In vivo, however, the IgG2a isotype variant of 20B1 exhibited significantly greater protection than IgG1 or IgG2b in murine SEB intoxication and S. aureus sepsis models. Protection was associated with downmodulation of inflammatory host response. Our data demonstrate that changing the isotype of already protective MAbs, without affecting their antigen specificity or sensitivity, can result in an enhancement of their protective ability. Isotype selection, therefore, should be carefully considered in the development of toxin-neutralizing MAbs and the design of antibody therapeutics.

IMPORTANCE  The purpose of this study was to enhance the protective efficacy of an existing, protective monoclonal antibody against staphylococcal enterotoxin B. Using two in vivo mouse models, our study demonstrates that the protective efficacy of a monoclonal antibody may be improved by inducing an isotype switch at the Fc region of an antibody, without altering the antigen specificity or sensitivity of the antibody. The development of therapeutic MAbs with higher efficacy may allow for the achievement of equal therapeutic benefit with a lower dosage. In turn, the use of lower doses may reduce the cost of these therapies, while reducing the potential for adverse side effects.

S taphylococcal enterotoxin B (SEB) is a clinically relevant staphylococcal superantigen, as well as a class B biological warfare agent. Passive immunotherapy with anti-SEB monoclonal antibodies (MAbs) has successfully neutralized the SEB toxin in vitro, as well as protected mice from SEB-induced lethal shock (SEBILS) in in vivo studies (1–6). In addition, recent work in murine infection models has demonstrated that SEB-specific MAb 20B1 successfully treats sepsis and deep-seated tissue infection that is caused by SEB-producing Staphylococcus aureus strains (7).

In recent decades, the use of MAbs has increased dramatically and represents one of the most lucrative and fast-growing classes of drugs (8). The majority of MAbs that are FDA licensed to date are used to treat either oncological or autoimmune diseases. Anti-infective antibodies, however, remain scarce, as only two such MAbs are licensed to treat infectious disease (9, 10). Although great progress has been made in antibody (Ab) technology, especially with respect to generating human and humanized antibodies, it remains unclear to what extent the constant region affects Ab efficacy. To date, 36 MAbs are FDA licensed, of which 16 are humanized, 11 are human, 6 are chimeric, and only 3 are murine.

The ability of Abs to affect host-pathogen interactions is not reliant solely on the function of their variable region, which binds to the target antigen. Their constant domains also mediate biological properties through Fc receptor (FcR) binding, complement activation, and effects on avidity and serum half-life (11). These biological properties can differ significantly between the different isotypes. For instance, the Fc portion of mouse immunoglobulin G2a (IgG2a) Abs interacts with complement components (12) and high-affinity activatory FcRs (13), whereas the Fc portion of mouse IgG1 antibodies mediates a relatively lower affinity interaction with activatory FcRs and does not stimulate FcR-mediated immune responses as effectively (14, 15). These properties have not been fully exploited for therapeutic antibody purposes, as the majority (73%) of currently licensed MAbs are of the human IgG1 isotype, which resembles mouse IgG2a in its similar effector functions. For the efficacy of anti-infective Abs, isotype selection may be of particular importance, as pathogens not only harm the host directly but also indirectly provoke immune responses by eliciting...
uncontrolled damage (16). In this regard, certain isotypes of anti-
infecive Abs could either enhance or decrease the host response
through their interaction with specific Fc receptors (17).

To investigate the effect of isotype on protective efficacy, we
generated isotype switch variants of a murine IgG1, MAB 20B1
(18). These isotype switch variants encode a variable region iden-
tical to their parent IgG1, and they differ only in their Fc binding
region. In previous studies, we have shown that passive immuno-
therapy with MAB 20B1 successfully treats SEBILS (2) and S. au-
reus-mediated sepsis (7). Our data demonstrate that isotype
switch variants of the IgG2a isotype are superior in efficacy com-
pared to the IgG1 and IgG2b isotypes. Specifically, lower doses of
the IgG2a isotype switch variant are required to enhance the sur-
vival of mice in both the SEBILS and S. aureus sepsis models.
Cytokine analysis documented that enhanced efficacy was associ-
ated with early inhibition of proinflammatory cytokines. In sum-
mary, these results provide valuable data that encourage system-
atric evaluation of specific isotypes for the development of anti-
infective MAbs.

RESULTS

Generation of IgG2a and IgG2b switch variants. Isotype switch
variants of SEB-specific IgG1 MAB 20B1 were generated. We used
the enzyme-linked immunosorbent spot assay (ELISpot) to de-
tect spontaneously arising variant cells producing new down-
stream isotypes of IgG1. IgG2a- and IgG2b-producing cells
spawned at rates of $6 \times 10^{-5}$ and $6 \times 10^{-4}$, respectively. Next, we
attempted to enrich each of the resulting two isotype variant pop-
ulations by Sib selection, which allowed us to successfully obtain
IgG2b-producing variants. However, despite successive rounds of
enrichment, the IgG2a variants remained too rare to recover
through this approach. Consequently, we used fluorescence-
activated cell sorting (FACS) with immunofluorescence staining
for surface-associated Ab to identify a cell fraction significantly
enriched for the IgG2a isotype. Small fractions of viable cells
stained for surface-associated IgG2a were isolated by cell sorting
on the basis of surface fluorescence intensity. This step enriched
the fraction of hybridomas producing IgG2a variant cells and suffi-
ciently reduced the dominant fraction of IgG1 producers. Two
rounds of soft agar cloning were performed to stabilize each vari-
ant cell type. Analysis of supernatants from the IgG2a- and IgG2b-
secreting clones revealed no trace of IgG1 by enzyme-linked im-
munosorbent assay (ELISA). Comparative alignment of
nucleotide sequences derived from each isotype variant confirmed
identical heavy- and light-chain variable sequences to the parent
IgG1 from which they were derived (data not shown).

Binding affinity of isotype switch variants to SEB. Initial assess-
ment of binding affinities by ELISA demonstrated comparable
binding to SEB by both the isotype variants and their 20B1 parent
(Pig. 1A). Standard measurement of dissociation constants ($K_d$
) using Biacore could not be performed with wild-type SEB due to
federal regulations that restrict the use of select agents. We instead
performed BLItz analysis to obtain $K_d$ values and thus investigate
whether changes to the constant regions affected the fine specific-
ity of these MABs toward SEB. This method determined the $K_d$
of 20B1 IgG1, IgG2a, and IgG2b toward SEB to be 16.25 + 1.8 nM,
66.27 + 2.7 nM, and 47.15 + 13.9 nM, respectively (Pig. 1B to D).
These $K_d$ values are considered comparable, as they are within the
experimental error range of the assay.

Inhibition of SEB-induced T-cell proliferation by MAB 20B1
is independent of IgG subclass. In vitro, we compared the ability
of all three 20B1 isotypes to inhibit SEB-induced proliferation of
murine splenocytes. All three isotypes of 20B1 demonstrated com-
parable levels of inhibition after 96 h of SEB stimulus and consis-
tently showed 90 to 100% inhibition with subnanomolar 50% inhibitory
concentrations ($IC_{50}$) (IgG1, 0.27 nM; IgG2a, 0.68 nM; and IgG2b, 1.04 nM) (Pig. 2A to C).

Isotype variants provide enhanced protection from SEB-
induced lethal shock. An in vivo murine SEBILS model was used to
test and compare the protective efficacy of the MAB 20B1 IgG1
parent and its isotype variants at equivalent doses. We have
reported earlier that MAB 20B1 IgG1 was an excellent lead candidate
to treat SEBILS (2) and SEB-secreting S. aureus sepsis (7). In this
study, various doses of MABs ranging between 50 nM and 500 nM
were compared for their efficacy. Results from these experiments
showed that switching the isotype of the parent IgG1 enhanced
protection, with relative efficacies of IgG2a > IgG2b = IgG1 (Pig.
3A to C), in a dose-dependent manner.

We further explored the protective efficacy of these isotype
variants in combination with 14G8 IgG1, an anti-SEB MAB that
we previously reported does not provide protection and does not
compete or share epitopes with MAB 20B1 (2). Improved protec-
tive efficacy was observed when MAB 20B1 IgG1 or the isotype
variants at doses as low as 50 nM were administered in combina-
tion with MAB 14G8 IgG1 in comparison to 20B1 alone (Pig. 3D).
Survival efficacy increased from 0% to 60% when 20B1 IgG1 was
combined with 14G8 IgG1, and from 10% to 40% after combined
treatment with 20B1 IgG2b and 14G8 IgG1. Increased survival was
also observed when a low dose of 25 nM of 20B1 IgG2a was admin-
istered in combination with MAB 14G8, a dose that was partially
protective (50%) when 20B1 IgG2a was used alone (data not
shown).

Enhanced protection correlated with more potent inhibition
of proinflammatory cytokines. Multiplex cytokine analysis of sera
at 2 h postchallenge showed significantly lower levels of all seven
proinflammatory cytokines examined (interleukin 6 (IL-6),
gamma interferon [IFN-$\gamma$], tumor necrosis factor alpha [TNF-$\alpha$],
IL-10, IL-1$\beta$, keratinocyte-derived chemokine/growth-related
oncogene [KC-GRO], and IL-12 p70) in mice treated with 20B1
IgG2a than in those treated with phosphate-buffered saline (PBS)
or the IgG2b isotype of MAB 20B1 (Pig. 4A to F). Levels of cyto-
kines IL-6, IFN-$\gamma$, and KC-GRO were also significantly lower after
treatment with 20B1 IgG2a than after treatment with the parental
IgG1 isotype.

Isotype switch variants have comparable effects on SEB
clearance in vivo. We further explored whether isotypes differen-
tially affect the clearance of SEB from serum. Accordingly, SEB
levels were quantified at 2 h (Pig. 5A) and 6 h (Pig. 5B) postchal-
lenge in serum and organs (liver, spleen, and kidney) of mice
treated with individual subclasses of MAB 20B1 IgG followed by
challenge with SEB. Treatment with any of the three isotypes re-
sulted in comparable clearance of SEB. As previously published,
immunocomplex formation with Abs slows the clearance of the
toxin through the kidney (19), and therefore significantly lower
clearance is observed than SEB clearance in mice treated with PBS
(20).

IgG2a isotype enhances protection from S. aureus sepsis. We
further explored the efficacy of each isotype switch variant in a
murine model of S. aureus sepsis. Previous work using this model
has shown that treatment with the 20B1 IgG1 isotype at 500 μg resulted in 80 to 100% survival, whereas all untreated mice died (7). Here, we tested a lower dose of 300 μg of each IgG subclass and compared treatment efficacies. At this lower dose, 50% of mice survived after treatment with the IgG2b isotype, 40% survived after treatment with the IgG1 isotype, and 100% survived after treatment with the IgG2a isotype (Fig. 6).

**DISCUSSION**

We conclude from our results that the constant region of anti-SEB MAbs can significantly affect toxin neutralization efficacy through modulation of the host immune response. Most importantly, the IgG2a isotype exhibits enhanced efficacy in comparison to that of other isotypes in both an SEB intoxication model and an SEB-secreting methicillin-resistant *S. aureus* (MRSA) sepsis model. Our findings underscore the importance of efforts to investigate Fc domain engineering as a strategy for optimizing therapeutic Abs. Enhancement of Ab efficacy could ultimately permit lower dosing of Abs, leading to a more reasonable pricing of this treatment modality and, therefore, substantially increase the development and use of anti-infective Abs.

Many human pathogens, including *S. aureus*, exhibit a diverse set of virulence factors, including toxins that facilitate evasion of the host’s sophisticated immune response (21–23). Many of these toxins directly damage host cells (24); however, staphylococcal enterotoxins like SEB have more of an indirect toxic effect. Specifically, they overstimulate immune cells, resulting in a cytokine storm that can ultimately kill the host (25). Historically, the view has been that Abs mediate SEB neutralization by interfering with binding of SEB to its targets, the major histocompatibility complex class II (MHC-II) molecule and the V-β chain of the T-cell receptor (TCR) (26). Interruption of this trimer formation is dependent on Ab specificity. Only high-affinity Abs or soluble V-β mimics have been successful in neutralizing SEB, largely due to the rapid and high-affinity binding between SEB and its in vivo targets, MHC-II and TCR V-β (27–29). To investigate the role of the Ab constant region in toxin neutralization, we generated IgG2a and IgG2b variants of SEB-specific murine MAb 20B1, which is of the IgG1 isotype. Isotype switching in mice progresses through the IgM ¡ IgG3 ¡ IgG1 ¡ IgG2b ¡ IgG2a spontaneous recombination process that generates Abs of different isotypes but that continue to share the identical variable region to the original Ab. Earlier studies on the generation of isotype switch variants from murine B cell lymphomas (30) have reported challenges in gener-
ating IgG2a isotypes. In this study, cytokine treatment in addition to FACS and immunofluorescence staining was required to identify and enrich populations of IgG2a-producing hybridoma cells.

The variable regions of all isotype variants were sequenced to ensure that no somatic mutation had occurred. All three isotype switch variants expressed identical variable regions, and BLItz analysis revealed comparable $K_{d}$ values in the nanomolar range for all isotypes. Of note is that in vitro T-cell assays did not demonstrate differences in neutralization efficacy, as the effective IC50 were comparable between all isotypes. This finding highlights the limitations of commonly used screening assays and is in accordance with previously published observations that SEB-specific MAbs, which neutralize the toxin in vitro, may not have the same effect in vivo (2). In our SEBILS model, significant differences in efficacy of Ab-mediated SEB neutralization were observed between the isotype switch variants. Efficacy was most potent for IgG2a, followed by IgG2b and IgG1. The IgG2a MAb also prevents SEBILS at lower doses than the IgG1 and IgG2b MAbs. As previously shown with modified capture ELISAs, MAb 20B1 binds to a single epitope on the toxin (2). Hence, only one MAb at a time can bind an SEB molecule. Furthermore, it has been shown that binding of SEB-specific MAbs to the toxin, in vivo, results in slower clearance of SEB toxin from the blood (2) because the immunocomplex is cleared not as efficiently through the glomerular filter of the kidney. Theoretically, treatment with different isotype variants, which commonly exhibit significant differences in their pharmacokinetics (31), could result in different degrees of SEB clearance. However, at least in the SEBILS model, toxin clearance is rapid (20), and therefore differences in half-life time of the various isotypes would likely not be relevant. Consequently, we found no difference in SEB levels in the sera of mice treated with different Abs. SEBILS was even more efficiently prevented when MAb 20B1 was combined with MAb 14G8, which is a SEB-specific MAb that is not protective on its own but possibly enhances the efficacy of other SEB-specific MAbs by facilitating uptake and clearance through cross-linking of Fcγ receptors (FcγR) (2). Most importantly, our data demonstrate that low doses of the IgG2a isotype prevent death from sepsis after infection with SEB-secreting MRSA. In summary, our findings support the concept that the IgG2a isotype of MAb 20B1 further enhances the efficacy of this protective Ab. Earlier studies have found that MAbs of the IgG2a isotype are more potent than other IgG isotypes in their ability to suppress tumor growth in a number of animal models (32–34). It has also been described that the IgG2a isotype has superior iodine radiolabeling characteristics compared to those of the parent IgG1 Ab (35). Furthermore, in viral infections, MAbs of the IgG2a subclass are more efficacious in providing Ab-mediated protection (36, 37). Our results are also consistent with similar studies involving isotype switch variants to the cryptococcal polysaccharide capsule (38). Those studies demonstrated that a non-protective IgG3 MAb could be converted to a protective MAb by switching the constant region. Furthermore, the protective efficacy of an IgG1 MAb was increased by isotype switching to IgG2a or IgG2b (38). Differences in efficacy have also been observed in Abs that treat HIV infections, where switch variants of human anti-CD4-binding site MAb F105 differed in their ability to neutralize infection (39, 40).

In a study that generated IgG2a and IgG2b variants of the Bacillus anthracis protective antigen-binding IgG1 MAb 19D9 (41), variants expressed identical variable regions and affinities to the

![Graph](https://example.com/graph.png)
target. Their readout of efficacy similarly determined that neutralization activity was highest by IgG2a, followed by IgG2b and IgG1. In that study, it was found that Ab protection required engagement of FcγR. Engagement of FcγR is not essential for neutralization of SEB, as Fab fragments of MAb 20B1 also prevent death of mice from SEBILS (unpublished data), and a humanized MAb variant of MAb 20B1 containing a mutated constant region that prevents FcγR interaction is also effective (42). Enhanced protection in our SEBILS model correlated with inhibition of proinflammatory cytokines. We hypothesize that these effects are indeed mediated through engagement of the different FcγRs or even different degrees of complement activation (12, 13). We, however, did not pursue experiments in FcγR-deficient mice, as they are not sensitive to SEBILS. It is noteworthy that isotype switching can also result in loss of protective activity due to a change in binding of MAb to its target, as recently described with anticapsular Abs to anthrax (43). These and other findings emphasize that changes distant from the Ag binding site can have an effect on Ab efficacy due to cross-domain interrelationships, whereby constant region changes may result in significant overall structural differences (43, 44).

Most FDA-licensed MAbs are human, humanized, or chimeric IgG1. The first MAb to be licensed by the FDA is a murine anti-CD3 IgG2a that is still being used. Currently, only three murine MAbs (two IgG2a and one IgG1) are licensed. One reason why isotype subclasses have not been consistently explored in preclinical trials may be that most novel therapeutic Abs are either humanized or chimeric. Given the differences between the FcγR of mice and humans, it has been difficult to study the interaction of human Ab Fc regions with FcγRs of mice. Newer studies with mice that express human FcγRs promise to elucidate the contribution of FcγR-mediated pathways to the neutralizing activity of MAbs (45). Our findings highlight unique properties of specific isotypes and underscore the importance of isotype selection in the design of antibodies for therapeutic applications.

MATERIALS AND METHODS

*S. aureus* toxins. SEB toxin was purchased from Toxin Technology (Sarasota, FL). SEB toxin was handled in the laboratory in accordance with the Centers for Disease Control and Prevention biosafety regulations. A single colony of clinical MRSA strain 38 (46) streaked on a brain heart infusion (BHI) agar plate was transferred to 50 ml liquid BHI medium grown overnight at 37°C in a shaking incubator. Bacteria were centrifuged at 2.9 × g at 4°C and washed in sterile PBS. Inocula of MRSA were prepared in sterile PBS, and dilutions were plated to verify CFU.

Isolation of isotype-switched variant hybridomas. The development of murine MAb 20B1 IgG1 and MAb 14G8 IgG1 that bind to SEB has been described earlier (2). MAbs 20B1 IgG2a and IgG2b isotype switch variants were identified using the ELISPOT and isolated by the Sib selection tech-
Briefly, 5 × 10^6 cells were treated with lipopolysaccharide (LPS) and/or interleukin 4 (IL-4) and further examined for IgG2a or IgG2b switch variants using ELISPOT after 3 days. A dissecting microscope was used to count spots, and then the median frequencies of switching were calculated. The corresponding well with the highest number of spots was subsequently plated out in a new 96-well plate at progressively lower cell densities. Those plates were then screened again for the frequency of switch variants. Later, cell sorting for specific isotypes was performed by FACS analysis using a FACSAria (BD) and coherent sapphire with a 100-mm quartz nozzle as described elsewhere (41). Soft agar subcloning was performed and random clones were picked and characterized further for specific isotypes (49).

High-scale MAb production and purification. Large-scale production of 20B1 MAbs (IgG1, IgG2b, and IgG2a) was performed in the CEL-Line bioreactor CL 1000 (Argos Technologies) according to the manufacturer’s instructions. Briefly, 2.5 × 10^7 viable hybridoma cells were inoculated into the lower cell compartment followed by ~800 to 900 ml of 5% Dulbecco’s modified Eagle’s medium (DMEM) into the upper chamber. Hybridoma cells were monitored and their supernatants were harvested every week. The concentration of MAb was quantified by ELISA, purified by protein G chromatography (Thermo, Fisher Scientific), and sterilized by filtering through a 0.2-mm-pore-size syringe filter (Thermo Scientific).

Relative affinity and kinetic measurements. SEB binding analysis to MAbs was established by decreasing titers of MAb in an ELISA. Briefly, ELISAs were performed by coating a 96-well plate with SEB (0.5 μg/ml), followed by the serial dilution of MAb 20B1 (IgG1 or IgG2a or IgG2b) and further detected by alkaline phosphatase-conjugated anti-mouse isotype MAb. These experiments were done in triplicates. Kinetic rate constants to compare binding of different isotypes of MAbs 20B1 to SEB were measured using one-way ANOVA followed by Tukey’s post hoc multiple-comparison test to compare the mean of each data set with the mean of other datasets using GraphPad Prism 6 software, and data were considered significant as follows: *, P < 0.05; **, P < 0.01; ***, P < 0.001. NS, not significant.
sured using a Forte Bio BLItz instrument (Forte Bio, Menlo Park, CA). Biotinylated SEB was immobilized on a streptavidin sensor, and MAbs were allowed to bind SEB at different dilutions (0 nM to 100 nM). The final MAb-SEB binding was analyzed using Forte Bio’s analysis software to determine $R_{equilibrium}$ as described by the manufacturer.

**T-cell proliferation assay.** Splenocytes (1 x 10^5 per well) from 6- to 8-week-old female BALB/c mice were seeded in a 96-well plate, with a final volume of 100 μl. Serial dilutions of isotype variants of MAb 20B1 were added starting from 100 μg/ml followed by SEB (25 ng/ml) and incubated for 96 h. Experiments were performed in triplicates. T-cell proliferation was measured using the Via-Light HS cell proliferation kit (Lonza, USA) according to the manufacturer’s instructions. IC_{50} values were determined from concentration-response curve analysis using GraphPad Prism version 6.

**Animal experiments.** All animal experiments were carried out with the approval of the Animal Institute Committee (AIC), in accordance with the rules and regulations set forth by the Albert Einstein College of Medicine. Six- to eight-week-old female BALB/c mice were purchased from National Cancer Institute (Bethesda, MD). Protective efficacy of isotype switch variants of MAb 20B1 was tested in vivo in the SEB-induced lethal shock (SEBILS) model as previously described (2). Mice (n = 5) were treated intraperitoneally (i.p.) at indicated doses of isotype variants of MAb 20B1 (50 μg, 100 μg, 250 μg, or 500 μg) 10 min prior to challenge with SEB (20 μg) and D-galactosamine (25 mg). Survival of mice was monitored for 5 days. In another subset of experiments, blood was drawn after 2 h from mice (n = 5) treated with a low dose of MAb 20B1 isotypes (50 μg) followed by challenge with SEB. Sera were analyzed for mouse cytokines using a proinflammatory 7 ultrasensitive kit (Meso Scale Discovery, Gaithersburg, MD) as per the manufacturer’s instructions. Data were further analyzed using one-way analysis of variance (ANOVA) followed by Tukey’s post hoc multiple comparison tests using GraphPad Prism 6. All samples were measured in duplicates. SEB clearance from blood, kidney, spleen, and liver was measured at 2 h and 6 h after mice were challenged with SEB and treated with 500 μg of MAb 20B1 isotypes. Protective efficacy of MAb 20B1 isotypes was further investigated in the *S. aureus* sepsis model induced by intravenous injection (i.v.) of 5 x 10^7 CFUs of MRSA strain 38 as previously described (7). Mice were given prophylactic treatment of MAb 20B1 isotypes (300 μg) 2 h prior to the SEB-secreting MRSA infection. Control mice were treated with PBS only.

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Varshney et al.

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