A targeted boost-and-sort immunization strategy using *Escherichia coli* BamA identifies rare growth inhibitory antibodies

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Outer membrane proteins (OMPs) in Gram-negative bacteria are essential for a number of cellular functions including nutrient transport and drug efflux. *Escherichia coli* BamA is an essential component of the OMP β-barrel assembly machinery and a potential novel antibacterial target that has been proposed to undergo large (~15 Å) conformational changes. Here, we explored methods to isolate anti-BamA monoclonal antibodies (mAbs) that might alter the function of this OMP and ultimately lead to bacterial growth inhibition. We first optimized traditional immunization approaches but failed to identify mAbs that altered cell growth after screening >3000 hybridomas. We then developed a “targeted boost-and-sort” strategy that combines bacterial cell immunizations, purified BamA protein boosts, and single hybridoma cell sorting using amphipol-reconstituted BamA antigen. This unique workflow improves the discovery efficiency of FACS + mAbs by >600-fold and enabled the identification of rare anti-BamA mAbs with bacterial growth inhibitory activity in the presence of a truncated lipopolysaccharide layer. These mAbs represent novel tools for dissecting the BamA-mediated mechanism of β-barrel folding and our workflow establishes a new template for the efficient discovery of novel mAbs against other highly dynamic membrane proteins.

The emergence of multi-drug resistant bacteria is a global health crisis that demands the discovery of new antibiotics, which requires a better understanding of microbial physiology and vulnerabilities. Integral outer membrane proteins (OMPs) reside within the distinctive outer membrane (OM) of Gram-negative bacteria where they perform diverse functions critical for bacterial survival. OMP functions include roles in active transport and diffusion of nutrients, efflux of toxic molecules, and construction of the OM permeability barrier. OMPs assume a common fold known as the β-barrel and are inserted into the OM by the essential β-barrel assembly machinery (BAM) complex, of which BamA is itself an OMP and a central component of the BAM complex. For the BAM complex to accomplish its essential OMP-foldase function, it has been postulated to undergo rapid and extensive conformational changes. Recent x-ray crystal and electron microscopy structural studies have suggested that the BamA β-barrel can assume inward- and outward-facing states that differ by a 65° rotation and 15 Å shift within the extracellular exit pore region. In fact, the extent of conformational change proposed for BamA is among the largest described for any previously reported membrane protein structure to date. Obtaining a detailed...
mechanistic understanding of essential OMPs like BamA may ultimately enable the design of future therapeutics targeting Gram-negative bacteria.

To date, genetic tools have greatly facilitated mechanistic studies of BamA, but gene knockouts or mutations do not necessarily mimic the effects produced by potential small molecule or protein-based inhibitors. The discovery of a peptidomimetic that interacts with many OMPs, including BamA, and possesses antimicrobial activity further highlights the potential of OMP-targeted therapeutics. However, potent and highly selective pharmacological tools to modulate the function of BamA have not yet been reported. Here, we hypothesized that antibody-mediated interference through an extracellular epitope of BamA could inhibit the essential OMP-folding function, resulting in bacterial growth inhibition and cell death. To our knowledge, this proposed mechanism of direct bacterial growth inhibition by a naked anti-OMP mAb has not been previously demonstrated on any pathogenic or laboratory strain.

The use of monoclonal antibodies (mAbs) have revolutionized the study of mammalian membrane protein function through Western- and FACS-based expression profiling, as chaperones for high-resolution structural studies, and, in rare cases, as proof-of-concept agonists or antagonists. For bacterial OMPs, while mAbs useful for Western or FACS analysis can often readily be obtained, remarkably few agonists or antagonists have been described, potentially due to masking of functional epitopes by lipopolysaccharide (LPS). Limitations in commonly used mAb discovery methods can also partially account for this gap. First, antibody phage display selections against bacterial OMPs can be challenging due to the high homology between the target OMP and host proteins in the strain used to amplify the libraries. Second, immunization-based methods are resource intensive and inefficient, and replete with uncertainties related to antigen format (summarized in Supplementary Table 1). While highly desirable, the production of sufficient purified full-length membrane protein (MP) antigens stabilized in various matrices (e.g., detergent, liposomes, nanodiscs, amphipols or styrene-maleic acid polymers) remains challenging, resource-intensive, and may not guarantee targeting of the native form of the MP particularly if mutagenesis is required for recombinant expression or stable purification. Finally, native antigen formats, such as membrane fractions, membrane vesicles, or whole cells, often give rise to significant titers against non-target proteins, thus leading to high backgrounds and especially low mAb discovery efficiencies.

Adding to the aforementioned complexities, the distinct structural and functional characteristics of OMPs like BamA may further confound mAb discovery efforts. Specifically, the extracellular epitopes on many OMPs are inherently dynamic, may change conformation depending on the lipid or host environment, and may be masked by LPS. Additionally, mAbs capable of functionally modulating an OMP into an activated or inhibited state have not been reported to date, and are therefore likely to be rare. Faced with these challenges, we endeavored to streamline anti-BamA mAb discovery by optimizing antigen formats (reconstitution matrices and native whole cells), adjuvants, immunization strategy, BamA-specific cell sorting, and high-throughput hybridoma purification. Notably, while our extensive traditional mAb discovery efforts failed to identify any functional anti-BamA mAbs, our optimized workflow improved the efficiency of FACS-anti-BamA mAb generation by >600-fold and enabled the discovery of rare anti-BamA mAbs with direct growth inhibitory activity in the presence of a truncated LPS layer. These growth inhibitory anti-BamA mAbs will enable deeper mechanistic studies of BamA function and further interrogation of BamA as a novel therapeutic target.

Results Evaluation of adjuvants and matrices with purified MP antigen. After we established methods to produce high-quality recombinant E. coli BamA reconstituted into a non-detergent amphipol matrix (see Methods), we set out to evaluate the impact of two commonly used adjuvants on the success of mAb discovery against this dynamic and essential OMP. Balb/c mice were immunized with BamA-amphipol using either CFA (Complete Freund’s Adjuvant) or Ribi adjuvants. All mice showed a strong immune response against BamA as measured by ELISA with the CFA adjuvant yielding antibody titers ~10× lower compared to the Ribi adjuvant (Fig. 1a). To monitor binding to extracellular epitopes of BamA, we used a FACS-based E. coli whole cell binding assay. However, because Gram-negative bacteria like E. coli possess LPS in their outer membrane which hinders access of antibodies to bacterial OMPs, we screened the polyclonal antibodies (pAb) by FACS on K-12 E. coli and a ΔwaaD E. coli mutant strain, which possesses the most truncated form of LPS that still supports bacterial growth (Supplementary Fig. 1). As predicted, we observed increased FACS signal with the ΔwaaD strain compared to K-12 E. coli, confirming that elaborated LPS can prevent binding of most antibodies to OMPs like BamA (data from Ribi immunized mice are shown in Fig. 1b). Nevertheless, all pAbs showed some FACS binding to the K-12 strain, suggesting that certain mAbs are able to bind BamA in the presence of the K-12 core-LPS (Fig. 1b). Notably, immunizations of the BamA-amphipol complex with Ribi adjuvant yielded higher FACS signals compared to those with the CFA adjuvant (Fig. 1c), consistent with the observed ELISA results (Fig. 1a). Therefore, Ribi was the adjuvant selected for all subsequent immunizations.

In favorable cases, recombinant purified MPs can be prepared in a range of solubilizing matrices. We next tested the effect of select protein matrices on the anti-BamA antibody response, in part, motivated by the hypothesis that different matrices may present more membrane proximal epitopes. Using Ribi adjuvant, we immunized mice with amphipol-reconstituted BamA or a short-chain detergent-solubilized BamA antigen and generated hybridomas from both groups. Because we observed that immunizations with amphipol-reconstituted BamA generated some non-specific antibodies against the amphipol matrix, we performed all ELISAs with detergent-solubilized BamA in order to decrease non-specific binding. In the short-chain octyl-glucoside detergent-solubilized format, intended to promote BamA conformational dynamics and maximum exposure of extracellular epitopes, only 54 out of 1299 IgG+ supernatants (4.2%) bound to BamA by ELISA. In the amphipol format, chosen because of its higher recovery yields compared to liposome incorporation, 375 out of 1700 IgG+ supernatants (22.1%) bound to BamA by ELISA (Fig. 1d). Intuitively, the 5-fold increase in antigen+ (Ag+) hybridomas could be due to greater in vivo stability of the amphipol format over the detergent-solubilized material.
Subsequent FACS screening of the panel from the BamA-amphiphol immunized group identified that ~2.4% of the IgG+ supernatants were FACS+ on the ΔwaaD strain, including several clones that also bound to the K-12 strain (Fig. 1d). As we ultimately sought to identify potentially growth inhibitory mAbs, we tested the ability of purified FACS+ mAbs to inhibit growth of the ΔwaaD mutant without any additional immune system components such as complement. However, despite robust immune responses that led to the discovery of many FACS+ anti-BamA mAbs, none of these mAbs were biologically active.

Combining native and recombinant antigen formats. We hypothesized that our failure to obtain inhibitory anti-BamA mAbs might be due to a limited sampling of the BamA extracellular epitopes or poor targeting of the OM-relevant conformations of BamA. Indeed, recent high-resolution structural studies have posited that BamA is a highly dynamic MP that likely exists in multiple conformations relevant for its function3,5,6. To explore a larger range of potentially relevant BamA conformations during antibody generation, and also provide a more relevant OM context for presentation, we revised our immunization strategy to employ whole bacterial cells (Fig. 2a). In this respect, it is particularly important to note that BamA is estimated to be present at only ~1500–4000 copies per cell, and therefore represents a minor fraction of the total cellular protein in E. coli26,27. Accordingly, we also switched from mice to rat immunizations because rats can tolerate larger doses of whole bacteria and a higher number of B cells can be recovered post-immunization. We reasoned that higher doses of bacteria and more recovered B cells should enhance our initial sample size to aid in the recovery of potentially rare functional anti-BamA mAbs.

Following the above rationale, after five sub-lethal injections of live K-12 E. coli, all immunized animals failed to show significant pAb binding to BamA by ELISA (solid lines in Fig. 2a). However, an anti-BamA response against purified BamA could be detected by Western blot, suggesting the presence of some anti-BamA titers and antibodies (Supplementary Fig. 2). To boost the recovery of BamA-specific antibody responses, we next challenged these same rats with amphipol-reconstituted purified BamA, but without the use of adjuvant in order to avoid initiating a new immune response. After the two injections of only 10 μg of purified protein antigen per rat, we observed a dramatic increase in the anti-BamA titers (dashed lines in Fig. 2a). We assayed the pAbs by FACS, but as expected, were unable to verify a BamA-specific FACS response due to the high background of anti-β-lactam antibodies against other components of the bacteria (Supplementary Fig. 3). Nevertheless, we wanted to determine whether this new hybrid immunization strategy that produced high anti-BamA-specific titers led to mAbs with distinct biological activity. We therefore tested the ability of pAbs from before and after the BamA protein boost to inhibit bacterial cell growth. Remarkably, pAbs from several rats slightly reduced bacterial growth only after the BamA-amphipol protein boost (black vs. white bars in Fig. 2c, p < 0.01). Because this weak growth effect was only observed after boosting with amphipol-reconstituted BamA, we predicted that the inhibitory effect was likely due to a specific anti-BamA response and that only a small fraction of the anti-BamA mAb pool was functional.
MP-specific sorting and high-throughput production of hybridomas. Since the percentage of IgG+ and Ag+ hybridomas are often extremely low, especially for complex and dynamic MP antigens like BamA, it is labor and resource intensive to generate a large panel of Ag+ clones for functional screening. Furthermore, traditional hybridomas do not display IgG on the cell surface, and hence, cannot be directly sorted for antigen-specific binding. To overcome these traditional bottlenecks to mAb discovery, we hypothesized that we could employ amphipol-reconstituted BamA to directly sort BamA+ hybridomas generated with a specialized myeloma fusion partner (SP2ab) that enables the production of both secreted and surface displayed IgG28. Hybridomas generate transcripts for both the secreted and membrane bound forms of IgG via alternative splicing. However, in the absence of co-expression of the Igα and Igβ (both components of the B-cell receptor complex) the membrane bound form of IgG is not stably displayed on the cell surface. The SP2ab fusion partner expresses both Igα and Igβ to enable surface display of IgG in addition to IgG secretion. To first verify that fluorescently labeled amphipol-reconstituted BamA could detect Ag+ cells, we generated a CHO cell line that transiently expressed a murine anti-BamA antibody on the surface via a glycosylphosphatidylinositol anchor29. Phycoerythrin (PE)-labeled BamA successfully stained the CHO cells with surface IgG, but not control CHO cells (Fig. 3a). Having validated that nanogram quantities of fluorescently labeled BamA should enable staining of BamA-specific cells, we generated hybridomas from the above immunized rats using the SP2ab fusion partner and sorted double positive (IgG+ and BamA+) hybridomas into individual wells of 96-well plates using a total of only 5 μg of PE-labeled amphipol-reconstituted BamA material (Fig. 3b).

Most robust functional assays utilized in successful mAb discovery efforts, like our bacterial growth inhibition assay, require concentrated and purified mAbs for direct testing; and this represents a significant technical and practical bottleneck when there is a need to evaluate large mAb panels. To overcome this traditional screening impediment, we developed an efficient, high-throughput hybridoma culture and purification process. In brief, single BamA-sorted hybridoma cells were grown for one week in plates followed by another ten days in deep well blocks in a shaking incubator. Supernatants were then purified using an automated 1 mL Protein A purification process to yield final purified IgGs in a smaller volume (400 μL) for functional screening. The final purified mAb concentrations exceeded 100 μg/mL in ~95% of the samples with total yields of ~100 μg (Fig. 4a). We subsequently screened roughly one quarter of 1632 purified hybridomas by ELISA and FACS to quantify our antigen-specific sorting efficiency. Strikingly, 88.5% of all tested hybridomas were BamA ELISA+ and 53% were ΔwaaD FACS+ (Fig. 4b). These 5 and 20-fold efficiency increases in identifying ELISA+ and FACS+ mAbs over our more traditional discovery methods described above (Fig. 1d) highlights the efficiency and robustness of our
Identification and characterization of growth inhibitory anti-BamA mAbs. To first characterize mAbs isolated from our workflow described above, epitope binning and affinity measurements were performed on a select panel of anti-BamA rat mAbs to characterize their binding properties (see below). For comparison, we also included several non-inhibitory mouse anti-BamA mAbs isolated from our traditional protein-only based hybridoma campaign (described above). Overall, the affinities of the mAbs ranged from 50 pM to 100 nM, with no clear correlation between affinity and discovery method (Table 1). In contrast to the high milligram amounts of purified BamA protein consumed by our traditional discovery efforts (Fig. 1d), this key result establishes that combining native and readily available BamA in the form of whole live bacteria with only a minimal amount of purified protein antigen for boosting and Ag-specific sorting (10^5 μg in total) can deliver a large panel of high-affinity and highly-selective mAbs (Fig. 4b).

We next wished to determine if any of the mAbs we isolated inhibited the growth of E. coli considering that no anti-OMP antibodies with growth-inhibitory properties have been described to date. As such, all 1632 purified mAbs isolated from the targeted boost-and-sort approach described above were screened in a bacterial growth assay using both K-12 and ΔwaaD E. coli. Consistent with the results from all of our previous anti-BamA discovery efforts, 95% of the screened rat mAbs had no (or minimal) impact on bacterial growth of the ΔwaaD or K-12 E. coli strain. In contrast, and most remarkably, 5% of the rat mAbs partially or fully inhibited growth of the ΔwaaD E. coli strain. To further characterize these mAbs, we measured the change in the number of viable bacteria after growth in the presence of a subset of these antibodies. We observed a ~50-fold increase in colony forming units (CFUs) when the ΔwaaD E. coli strain was grown in the presence of buffer or non-inhibitory mAbs (black and blue bars in Fig. 5b). Bacterial growth in the presence of several partially inhibitory mAbs led to...
no change in CFU, suggesting these mAbs are bacteriostatic (orange bars in Fig. 5b, \( p < 0.001 \)). Finally, and most remarkably, growth in the presence of the full inhibitors resulted in ~50–100-fold decrease in CFUs, suggesting that these mAbs are bactericidal (pink bars in Fig. 5b, \( p < 0.001 \)). These data collectively suggest that we have identified naked anti-OMP mAbs capable of inhibiting growth of an \( E. \ coli \Delta waaD \) strain in the absence of other immune effectors.

We found that the majority of the inhibitory anti-BamA mAbs occupied a single epitope bin on BamA (Fig. 5c). Incredibly, we note that several of the non-functional mouse mAbs also binned within this same epitope, suggesting that our targeted boost-and-sort discovery method is capable of more in-depth sampling of a given epitope than traditional mAb discovery efforts. To confirm that the unique inhibitory mAbs were specific to \( E. \ coli \) BamA, we tested and observed that there was no cross-reactivity by ELISA with the closely related BamA homologs from \( Klebsiella \ pneumonia \) and \( Enterobacter \ cloacae \) (Table 1). Unfortunately, none of the inhibitory anti-BamA mAbs bound to the K-12 strain by FACS, reconfirming that LPS likely masks the relevant or susceptible BamA epitope in the context of the K-12 strain. Despite this shortcoming, these inhibitory mAbs do validate the hypothesis that an anti-BamA antibody can result in direct growth inhibition of bacteria. This critical observation validates our key screening approach to use the \( \Delta waaD \) strain of \( E. \ coli \) in order to maximize the potential to interrogate extracellular epitopes on BamA by mAbs. Moreover, these context-dependent mAbs represent valuable research tools to study BamA and BAM complex function by dissecting the mechanism of action of these rare growth inhibitory anti-BamA mAbs\(^{30}\).

**Discussion**

Previous discovery efforts, including our own described above, have highlighted that solely relying on recombinant MPs as antigens can limit the success of an antibody discovery campaign. For example, different detergents or lipids can alter epitope conformations or global structures to generate proteins in non-native states\(^{31,32}\). Not

| % Growth inhibition | # of mAbs | KD range (nM) |
|---------------------|-----------|--------------|
| <50%                | 1554      | n.d.         |
| 50–75%              | 59        | ~0.05–100    |
| 76–90%              | 12        | ~0.1–20      |
| >90%                | 7         | ~0.1–40      |

Table 1. Summary of functional properties of anti-BamA mAbs. Affinity values for a subset of the anti-BamA mAbs as determined by Wasatch. All inhibitory mAbs bind by ELISA to only \( E. \ coli \) BamA and not \( Entero \) or \( Kleb \) BamA.
surprisingly, relying solely on recombinant MP antigens to generate mAbs can lead to the discovery of detergent- and/or lipid-sensitive binders that fail to react with the native protein. We observed that using whole bacteria to prime the animals followed by boosting with µg amounts of recombinant OMP protein enables robust generation of FACS+ pAbs and mAbs (Fig. 2). In total, we used only ~105 µg of recombinant MP antigen, 100 µg for boosting and 5 µg for sorting, to discover the first known anti-BamA mAbs with growth inhibitory activity. Our targeted boost-and-sort strategy is therefore particularly attractive for other dynamic OMP targets such as TonB-dependent receptors, autotransporters, or other virulence factors. Our method also alleviates the significant challenges and resources associated with generating high milligram amounts of recombinant material that are often required to support successful traditional mAb discovery workflows. It is therefore noteworthy that mAbs discovered from the targeted boost-and-sort method had comparably high-affinity (pM–nM range) and selectivity profiles to those mAbs that were discovered through more traditional, resource-intensive protein-based immunizations.

Whole cell-based immunizations are known to generate high levels of mAbs against non-target proteins, and thus, downstream mAb screening becomes extremely resource intensive. For the first time, to our knowledge, we established that sorting for Ag-specific hybridomas using amphipol-reconstituted MPs dramatically improves antibody discovery efforts by enriching the initial complex pool for antigen binding. Notably, sorting hybridomas with amphipol-reconstituted BamA greatly enriched the mAb pool for FACS binders (~53% of enriched IgG+ clones compared to 2.4% of non-enriched IgG+ clones), which also extended the epitope coverage across the target and likely enabled the identification of the rare bacteriostatic and bactericidal mAbs. We highlight the critical importance of having the target MP reconstituted in an appropriate matrix, since Ag-based sorting with detergent-solubilized BamA led to a loss of cell viability. Additionally, the amphipol matrix successfully masks the hydrophobic regions of BamA to help avoid the sorting of non-specific mAbs. A variety of reconstitution methods are emerging to generate stabilized recombinant MPs which should greatly extend our methods to other MP targets; and in fact, we have already validated that several of these reconstitution formats also enable sorting of Ag-specific cells.

We had hypothesized that an antibody might affect the function of BamA and result in bacterial growth inhibition by binding to an extracellular epitope and stabilize either an open, intermediate or closed conformation of BamA. Notably our own traditional protein antigen-only based mAb discovery approach as well as previous reports on BamA vaccine efforts had failed to identify such an antibody. By contrast, using our new high-throughput targeted boost-and-sort platform, we rapidly identified BamA antagonist mAbs at a rate of 0.4% (7 inhibitors out of 1632 mAbs screened) that could completely inhibit growth of ΔwaaD E. coli. We found that many non-inhibitory and partially inhibitory anti-BamA mAbs bin in the same epitope as the bactericidal mAb clones, suggesting small molecular differences in BamA binding can substantially impact antibody function. This key result suggests that broad coverage of a single epitope with a diverse panel of mAbs may be required to obtain functional mAbs against highly dynamic or especially challenging MP targets. While the present bactericidal mAbs were unable to bind their epitope in the presence of an intact LPS layer, future structural characterization of select mAbs may provide a starting point to engineer smaller scaffolds or peptides with the ability to penetrate the LPS layer and antagonize BamA.

The conformational stabilization of mammalian MPs via mutagenesis or other methods can enable direct discovery of agonist or antagonist mAbs. However, such stabilization methods typically require an existing tool compound to validate a given locked conformation, whereas for many MP targets (including BamA), such tools do not exist. Here, without such pre-existing tools, our efficient targeted boost-and-sort workflows robustly identified rare anti-BamA mAbs with bacterial growth inhibition activity, where our traditional methods failed to do so. One can therefore envision that our method may directly translate to more traditional targets like GPCRs; especially since numerous expression and stabilization methods to facilitate the production of ~100 µg levels of these high-value recombinant proteins have emerged in recent years. Moreover, GPCRs and other mammalian targets are not masked by a surrounding LPS layer like BamA or other OMPs, so one might even anticipate a high discovery rate for binding antibodies (~50% FACS+) and rare modulators. Overall, we anticipate that the methods described herein will provide a resource-efficient approach for identifying novel mAb tools against additional MP targets to reveal new biological insights and enable the discovery of new therapeutics.

Methods

Protein expression and purification. The BamA gene lacking the N-terminal signal sequence and POTRA domains but encoding the beta-barrel outer membrane component from Escherichia coli (V418-W810) was cloned into a modified pET-52b expression vector containing a N-terminal His6 and Avi-tag. Protein was overexpressed using E. coli host BL21(DE3) grown by fermentation at 17°C for 64 hours in autoinduction media. Cells expressing E. coli BamA were harvested and resuspended in buffer A (50 mM Tris pH 8.0, 500 mM NaCl, 0.1 mM TCEP), supplemented with Complete Protease Inhibitors (Roche), 1 mM PMSF and DNase. Following cell lysis by microfluidization, the soluble BamA protein in inclusion bodies were harvested by low speed centrifugation (3,000 g) at 4°C for 20 minutes and resuspended by brief vortexing in denaturing buffer B (50 mM Tris pH 8.0, 6 M GuHCl, 1 mM TCEP) followed by gentle agitation on a platform shaker for 1 hour at 22°C. Insoluble debris and residual membrane fractions were pelleted by ultracentrifugation at >100,000 g at 4°C for 1 hour using a Ti45 rotor. Supernatant was collected, protein concentration was determined (absorbance 280 nm), and the total protein concentration was adjusted to 1 mg/mL using denaturing buffer B. 40 mg of BamA was refolded by rapid dilution into a solution containing buffer C (1% (wt/v) zwittergent 3–12, 50 mM Tris pH 8.0, 0.1 mM TCEP) which was being mixed on a stir plate at 22°C. The refolding reaction mixture was subsequently set at 22°C with-out stirring for 2 days, then BamA was captured by batch binding using Ni-NTA resin. Resin was captured by gravity flow and washed with 5 column volumes of buffer D (1% (wt/v) zwittergent 3–12, 50 mM Tris pH 8.0) and eluted with 15 column volumes of the same buffer containing 300 mM imidazole pH 8.0. The eluted protein was
After washing twice with PBS, cells were run on a BD LSRFortessa™ X-20 cell analyzer and data analyses was applied over a Superdex 200 (26/60) column in buffer E (50 mM Tris pH 8, 100 mM NaCl). Protein reconstituted ImmunoResearch) reconstituted in buffer F; the PE-streptavidin-biotin-BamA-amphipol complex was incubated buffer E was incubated with a stock solution of 2 mg/mL A8-35 amphipol (Anatrace) at 22 °C for 1 hour and then Superdex 200 (26/60) in buffer E as described above in order to remove free biotin and other reaction components. When protein was required in amphipol (either non-biotinylated or biotinylated), BamA at 1 mg/mL in the Superdex 200 (26/60) for ELISA or antigen-based sorting, an in vitro biotinylation reaction using BirA enzyme targeting the N-terminal Avi tag was first carried out according to the manufacturers suggestions (Avidity); BamA was then run over the Superdex 200 (26/60) in buffer E as described above in order to remove free biotin and other reaction components. When protein was required in amphipol (either non-biotinylated or biotinylated), BamA at 1 mg/mL in buffer E was incubated with a stock solution of 2 mg/mL A8-35 amphipol (Anatrace) at 22 °C for 1 hour and then applied over a Superdex 200 (26/60) column in buffer F (50 mM Tris pH 8, 100 mM NaCl). Protein reconstituted in A8-35 amphipol for immunization was concentrated to 1 mg/mL using a centrifugal device (10 K MWKO; Millipore). When PE-labeled protein was required for antigen-specific sorting, BamA at 1 mg/mL (biotinylated and reconstituted in A8-35 amphipol in buffer F) was mixed 1:1 (vol/vol) with PE-streptavidin (Jackson ImmunoResearch) reconstituted in buffer F; the PE-streptavidin-biotin-BamA-amphipol complex was incubated at 22 °C for at least 15 minutes prior to use.

Generation of murine hybridomas. All animal study designs for the mouse and rat immunizations were reviewed and approved by the Genentech Institutional Animal Care and Use Committee prior to the start of this work. All animal work was performed in accordance with relevant guidelines and regulations. Balb/c mice (Charles River Laboratories, Hollister, CA) were immunized with detergent-solubilized or amphipol-reconstituted E. coli BamA protein and either CFA or Ribi adjuvant (Sigma). pAbs were purified by Protein A and assayed by ELISA, FACS, and inhibition of bacterial growth as described below. Hybridoma fusions were performed as previously described and supernatants were screened for protein binding by ELISA38. All ELISA positive clones were purified and screened for inhibition of bacterial growth.

Generation of rat hybridomas. Sprague Dawley rats (Charles River Laboratories, Hollister, CA) were immunized with E. Coli K12 cells in PBS (10⁵–10⁶ cfu via intravenous injection). The rats were then boosted four times with E. Coli K12 cells and a first set of sera was isolated (bleed #1). Protein A purified pAbs were then evaluated for binding by ELISA and inhibition of bacterial growth. Rats were further boosted twice with 10 µg E. coli BamA protein in amphipol and a second set of sera was isolated for analysis (bleed #2). To generate monoclonal antibodies, hybridoma fusions were performed as previously described except with a myeloma partner SP2ab that enables surface display of IgG cell28,38. After HAT selection in ClonaCell-HY Medium C (StemCell Technologies) for 4 days, hybridomas were stained with a cocktail of FITC-conjugated anti-rat IgG1/IgG2a/IgG2b mAbs (1:100 dilution; Bethyl Laboratories) and PE-conjugated BamA protein antigen (5 µg/mL). Samples were sorted on a FACSaria II cell sorter (BD Biosciences). Gating strategy to identify hybridoma cells was first based on size (FSC/SSC). After dead-cell exclusion with Propidium iodide (Sigma-Aldrich P-4864), IgG+ “BamA”+ single cell hybridomas were sorted into 96-well tissue-culture plates containing 200 µl of ClonaCell-HY Medium E (StemCell Technologies). Profiles were analyzed by FlowJo v.9.7.7 software. Sorted cells were cultured for seven days.

Hybridoma culture and purification. A semi-automated high throughput process for hybridoma culturing and antibody purification will be described elsewhere (Nakamura G., et al. manuscript in preparation). Briefly, individual hybridoma clones were transferred to 96-well tissue culture plates and cultured for 6–8 days. Cells were then transferred to 96-deep round well boxes and cultured for an additional 10 days in continuous orbital shaking incubators at 37 °C degrees. Supernatants were then harvested via centrifugation and IgG was purified from supernatants using MabSelect SuRe (GE Healthcare, Piscataway, NJ, USA). The eluted IgGs in 400 µL final volume were neutralized and normalized to 50–150 µg/mL prior to subsequent functional assay testing. IgG concentrations were determined by OD280 measurements.

ELISA assays. Antibodies were screened by capture ELISA. Briefly, biotinylated detergent-solubilized BamA protein, diluted in assay buffer (PBS + 1.5% bOG + 0.5% BSA) was added to 384 well streptavidin-coated plates (Thermo Scientific USA) and incubated at 25 °C for 1 h. The plates were washed with wash buffer (PBS + 1.5% bOG). Hybridoma supernatants or purified antibodies were added to the wells and incubated at 25 °C for 1 h. The plates were washed with wash buffer. The captured antibody was detected with goat anti-rat HRP secondary antibody (Bethyl Laboratories USA). The plates were incubated at 25 °C for 1 h, washed with wash buffer and developed with TMB solution (Surmodics, USA). Plates were read at 630 nm.

Mammalian flow cytometry assays. The variable light chain and variable heavy chain sequences of a murine anti-BamA hybridoma were determined using 5’ RACE followed by sequencing of the PCR amplified products. cDNAs encoded for both variable domains were gene synthesized (Genewiz) and cloned into a mammalian surface display vector. This vector contains the light chain connect to heavy chain via an IRES sequence and has a glycosylphosphatidylinositol linkage signal (GPI anchor) derived from human decay-accelerating factor fused to the C-terminus of the heavy chain to enable surface display of the IgG39. CHO cells that were transiently transfected with the cDNA encoding for the membrane-bound anti-BamA antibody were stained with streptavidin PE-conjugated biotinylated BamA protein antigen in 200 ul FACS staining buffer (PBS with 0.5% BSA and 2 mM EDTA) at a final concentration of 5 µg BamA/µl staining volume. Cells were stained for 15 min at 4 °C. After washing twice with PBS, cells were run on a BD LSRFortessa™ X-20 cell analyzer and data analyses were performed using FlowJo v.9.7.7 software.
**Bacterial flow cytometry assays.** An *E. coli* K12 glycerol stock was thawed in LB broth and incubated at 37 °C until reaching exponential phase of growth. Bacterial cells were then pelleted, re-suspended in ice-cold PBS with 1% BSA, and normalized to an OD600 of 0.5. Bacterial suspensions were diluted 1:2.5 in ice-cold PBS, 1% BSA and 2 × 10^8 CFU were incubated with 10 μg/mL mAb or 40 μg/mL pAb at 4 °C for 1 h under gentle agitation. Bacterial cells were then washed three times in ice-cold PBS, 1% BSA and labeled using Alexa 488 anti-mouse IgG (H + L) (1:1000; Invitrogen) or Alexa 488 anti-rat IgG (H + L) (1:1000; Invitrogen) for 30 min at 4 °C. Bacterial cells were then washed three times with ice-cold PBS, and fixed by adding 1 vol. of 2% w/v paraformaldehyde in PBS. Samples were run on a BD FACSCalibur and data were analyzed using FlowJo software.

**Bacterial growth assay.** A kanamycin deletion-insertion mutation of *waaD* was obtained from the Keio collection44. The genotype of the parental BW25113 strain is: rmmB3 A LacZ4787 hsdR514 (ΔrhaBAD)567 Δ(rhaBAD)568 rph-1. The screening strain, BW25113 ΔwaaD or parental BW25113, was grown to log phase in MHB II supplemented with 0.002% Tween-80 and 0.05% BSA, diluted to a final OD600 of 0.01 in sterile round-bottom 96-well plates. Antibodies were initially screened for activity at 10 μg/mL and growth inhibition was measured after 4 h of static incubation at 37 °C. A plate reader measured bacterial growth via optical density at 600 nm after shaking the plate for 25 s. For analysis of viable bacteria, colony-forming units (CFUs) were determined by plating cells on agar plates both prior to antibody addition and after 4 h of static incubation at 37 °C with 10 μg/mL antibody.

**Epitope binning and affinity characterization.** Epitope bins were determined by 96 × 96 array-based SPR imaging system (Carterra, USA) using classical sandwich method. Purified antibodies were diluted at 20 μg/mL in 10 mM sodium acetate buffer pH 4.5. Using amine coupling, anti-CP16 was directly immobilized onto a SPR sensorprism CM200 M chip (XanTec Bioanalytics, Germany) using a Continuous Flow Microspotter (Carterra, USA) to create an array of 96 antibodies. For antibody binning, printed chip was loaded on IBIS MX96 SPRi (Carterra USA) and *E. coli* BamA protein, diluted to 50 nM in 1.5% bOG HBS-P buffer, was injected over the chip at 25 °C followed by a second injection of purified antibody, diluted at 20 μg/mL in 1.5% bOG HBS-P buffer, to make a sandwich. The epitope binning data was processed using Wasatch binning software tool.

**Statistics.** All experiments involving bacterial growth were analyzed via the unpaired Student’s t test using Prism 7.0 (Graphpad Software) and compared mAb treated samples to the buffer controls.

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Author Contributions
R.V., Z.L., N.C., J.V., K.M., S.P., J.C., P.L., S.L., K.S., J.B., I.Z., M.A., P.S., R.J., and J.K. performed experiments and analyzed data. Y.G.M., L.C.A., C.K., M.X., A.G., P.S., G.N., and D.S. designed experiments and analyzed data. S.R., J.P., and J.K. performed experiments and analyzed data. Y.G.M., L.C.A., C.K., M.X., A.G., P.S., G.N., and D.S. designed experiments and analyzed data. S.R., J.P., and J.K. wrote the manuscript.

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