Flavin Reductase Contributes to Pneumococcal Virulence by Protecting from Oxidative Stress and Mediating Adhesion and Elicits Protection Against Pneumococcal Challenge

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Pneumococcal flavin reductase (FlaR) is known to be cell-wall associated and possess age dependent antigenicity in children. This study aimed at characterizing FlaR and elucidating its involvement in pneumococcal physiology and virulence. Bioinformatic analysis of FlaR sequence identified three-conserved cysteine residues, suggesting a transition metal-binding capacity. Recombinant FlaR (rFlaR) bound Fe²⁺ and exhibited FAD-dependent NADP-reductase activity, which increased in the presence of cysteine or excess Fe²⁺ and inhibited by divalent-chelating agents. flaR mutant was highly susceptible to H₂O₂ compared to its wild type (WT) and complemented strains, suggesting a role for FlaR in pneumococcal oxidative stress resistance. Additionally, flaR mutant demonstrated significantly decreased mice mortality following intraperitoneal infection. Interestingly, lack of FlaR did not affect the extent of phagocytosis by primary mouse peritoneal macrophages but reduced adhesion to A549 cells compared to the WT and complemented strains. Noteworthy are the findings that immunization with rFlaR elicited protection in mice against intraperitoneal lethal challenge and anti-FlaR antisera neutralized bacterial virulence. Taken together, FlaR’s roles in pneumococcal physiology and virulence, combined with its lack of significant homology to human proteins, point towards rFlaR as a vaccine candidate.

Streptococcus pneumoniae is an anaerobic aero-tolerant organism whose metabolism is sensitive to the presence of oxygen¹. Both oxygen and its ultimate metabolic by-product, hydrogen peroxide², react slowly with cellular macromolecules and do not exert prominent destructive effects³. However, interaction of H₂O₂ with ferrous ions...
(Fe^{2+}) produces hydroxyl radicals by the Fenton reaction and these highly reactive oxygen species (ROS) are capable of destroying DNA, membrane lipids and proteins. In contrast to Fe^{2+}, the ferric ion (Fe^{3+}), which is the stable form of iron since it is chemically inert at neutral pH, is considered non-toxic. However, the reducing conditions in the cytoplasm can reduce Fe^{3+} to the toxic Fe^{2+} state. Among the reducing agents capable of reducing ferric iron are intracellular cysteine, reduced FAD and to a lesser extent, glutathione, thioredoxin, NADH and NADPH. In culture, pneumococci produce millimolar concentrations of H_{2}O_{2}, as part of their normal metabolism. Accordingly, S. pneumoniae is relatively resistant to the destructive effects of H_{2}O_{2} and ROS in comparison to other bacteria, even though it lacks the major H_{2}O_{2}-degrading enzyme, catalase. Two membrane associated, extracellular- and intracellular- thioredoxin pathways have been described, which prevent damage caused by ROS molecules.

Since Fe^{2+} may exert toxic effects through the Fenton reaction, the level of Fe^{2+} in all organisms is tightly controlled by its binding to specific ligands. Iron concentration in aqueous solution is extremely limited (10^{-8}–10^{-9} M) and well below the range that supports microbial growth (10^{-6} M). Iron concentration within the host is further restricted (up to 10^{-24}M) in order to suppress the generation of toxic ROS ensuring bacteriostasis at potential sites of infection. As Fe^{2+} is essential for many biological processes, scarcity of iron within the host has initiated evolutionary war between host and pathogen, where successful pathogens express various mechanisms for iron acquisition. The witholding of metals, such as iron, to effectively starve pathogens of essential elements is referred to as "nutritional immunity" and is an important facet of the innate immune system.

Intracellular Fe^{2+} in the bound state is unable to interact with H_{2}O_{2} and consequently ROS production is prevented. In eukaryotic organisms, Fe^{2+} is bound to intracellular heme-containing proteins, including ferritin, iron–sulfur proteins and extracellular iron-binding proteins, such as transferrin and lactoferrin. In prokaryotes, bacterioferritin, ferritin and Dps have similar functions. S. pneumoniae have been found to acquire free inorganic iron by the ABC transporter lipoproteins PiaA and PiaA_{30}. Pit and SPD_{1609132}. In addition S. pneumoniae binds host Fe^{2+} transporter molecules such as transferrin, lactoferrinin and haemoglobin to remove and utilize the Fe^{2+} associated with these proteins. However, the full extent of pneumococcal iron acquisition systems and their role in pneumococcal survival and virulence is not known in detail.

In our ongoing studies, we have demonstrated an age-dependent enhancement of antibody-response to a group of S. pneumoniae surface protein antigens. One of these proteins, which we have previously named pneumococcal surface immunogenic protein B (PsipB), was recently annotated in TIGR4 as flavin reductase (FlaR). In this study, we attributed functions to FlaR in Fe^{2+} binding and NADP reductase activity and demonstrated its importance to pneumococcal resistance to H_{2}O_{2}. Moreover, we found that FlaR contributes to pneumococcal virulence not only in oxygen rich environments, but also in oxygen limited environments. In line with these results, we ascribe an additional function for surface-expressed FlaR as an adhesin. Finally, FlaR elicited protective immune response in mice against S. pneumoniae, implying it can serve as a candidate vaccine.

### Results

**FlaR bioinformatic analyses.** To examine whether flavin reductase is ubiquitous among S. pneumoniae strains, the flavin reductase DNA sequence of the TIGR4 strain was compared to 29 completely sequenced genomes of S. pneumoniae. All 29 genomes were found to contain a highly similar locus to SP_RS 02775 [BLAST e-value < 10^{-76}, >97% identity, >70% query coverage (>98% coverage in 27 of the genomes)].

A BLAST search carried out with WP_000580663.1 against all S. pneumoniae RefSeq proteins annotated as "flavin reductase" retrieved 63 proteins, which were subjected to multiple sequence alignment and a phylogenetic tree was constructed. The tree was used to select a representative set of 9 divergent proteins, one from each clade of the tree (Supplementary Fig. S1). The position of a conserved flavin reductase domain is indicated, as well as tree was constructed. The tree was used to select a representative set of 9 divergent proteins, one from each clade.

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**Recombinant FlaR protein production.** Recombinant FlaR protein was overexpressed as HAT-tagged in Escherichia coli and purified under denaturing conditions. The theoretical molecular weight of the untagged protein was found to be 17.2 kDa. Separation of the HAT-tagged protein on reducing SDS-PAGE revealed a major band of 22 kDa and a minor band of 45 kDa (Fig. 1a). The upper bands were subjected to MALDI-TOF analysis and demonstrated their identity to 63 proteins, which were subjected to multiple sequence alignment and a phylogenetic tree was constructed. The tree was used to select a representative set of 9 divergent proteins, one from each clade.

A BLASTP search of WP_000580663.1 vs. RefSeq proteins excluding S. pneumoniae showed highly similar hits from other Streptococci species (e value < 10^{-52}, 36–87% identity, >70% query coverage (>98% coverage in 27 of the genomes)). A BLASTP search of WP_000580663.1 vs. RefSeq proteins excluding S. pneumoniae showed that the proteins were subjected to multiple sequence alignment and a phylogenetic tree was constructed. The tree was used to select a representative set of 9 divergent proteins, one from each clade of the tree (Supplementary Fig. S1). The position of a conserved flavin reductase domain is indicated, as well as three positions of conserved cysteines (equivalent to positions 66, 72 and 84 of WP_000580663.1). It is worth noting that cysteine was present in the three conserved positions in all 63 proteins, except 3 cases of substitution to tyrosine in the first or second position (data not shown).

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**FlaR cellular localization.** To validate the previous finding regarding the presence of FlaR on the bacterial surface, we performed flow cytometry analysis on live R6 strain probed with a polyclonal mouse anti-flaR antibody. Indeed FlaR was found to be surface expressed (Fig. 1c). In addition, polyclonal rabbit antisem(r) recognized FlaR in the cytoplasm of WU2 WT strain in two forms: at ~17 kDa (untagged flaR monomer) and 34 kDa (untagged dimer; Fig. 1d). The 34 kDa band could also be observed in the bacterial cell-wall (CW) fraction...
(Fig. 1d) in accordance with our previous study35. A third, nonspecific band (~ 50kDa) was detected in the immunoblots of both the WU2 WT and WU2ΔflaRΔkan null mutant strains (Supplementary Fig. S2). The cytoplasmic protein, malonyl-CoA:ACP transacylase (FabD), which is involved in lipid metabolism36, was used as control for successful separation between CW and cytoplasmic fractions. FabD could be visualized (33 kDa) in both the total bacterial protein extract and the cytoplasm, but not in the CW fraction. This was lately described in Mizrachi Nebenzahl et al.201637.

Refolding and Solubilization of rFlaR. The rFlaR was insoluble under physiological conditions and its purification could be obtained under denaturing conditions in buffer supplemented with 8 M urea. Different refolding conditions were tested for their ability to renature the protein. Dialysis of the denatured protein against buffers at different pH levels revealed that rFlaR could be solubilized at pH ≥ 8.0 (Fig. 2a). The presence of cysteine-rich sequence suggests that the protein has a transitional metal-binding ability38. Thus, rFlaR was dialed against PBS (pH 7.3) supplemented with 2 mM salts of either transition metal or calcium and magnesium (2 mM of each). rFlaR became fully soluble in the presence of 2 mM FeSO₄. Partial solubility could be obtained in the presence of 2 mM CoCl₂ or CuSO₄ (Fig. 2b).

FlaR is a Fe²⁺-binding protein. To further verify that rFlaR can directly bind Fe²⁺, solubilized rFlaR was incubated with ferrozine in the presence of free Fe²⁺ ions. Ferrozine is well known as an effective chelator of Fe²⁺ forming a complex that peaks at 562 nm39. During the experiment the absorbance was significantly lower

Figure 1. Expression of recombinant Flavin reductase (rFlaR). (a) rFlaR was purified using Ni-NTA beads under denaturing conditions and resolved by SDS-PAGE using sample buffer with or without β-mercaptoethanol (β-ME). (b) rFlaR was separated using SDS-PAGE. The protein was transferred to a nitrocellulose membrane and probed with rabbit antiserum raised against rFlaR (lane 1) or preimmune rabbit serum (lane 2). (c) Expression of FlaR on the surface of R6 was detected by flow cytometry using mouse anti-rFlaR antibodies. (d) Cytoplasmic (cyto) and cell wall (CW) fractions prepared from WU2 were subjected to electrophoresis and immunoblotted with rabbit anti-rFlaR antisera. Of note, this gel was cropped from Supplementary Fig. S2.
in solutions containing rFlaR, solubilized either at pH 8 or with 2 mM Fe^{2+}, in comparison to three different controls supplemented with 2 mM Fe^{2+}: 1) PBS (Fig. 3; Control; p = 0.0001); 2) PBS supplemented with bovine serum albumin (BSA) and 3) PBS supplemented with pneumococcal protein aspartate carbamoyltransferase (WP_001293838.1; ATCase)\textsuperscript{37}, not known to bind Fe^{2+}. As a positive control for Fe^{2+} binding, we have used ethylenediaminetetraacetic acid (EDTA; 10 nM), which significantly inhibited the ferrozine-iron interaction (Fig. 3, p = 0.0001). The ability of rFlaR to compete with ferrozine for Fe^{2+} binding points out that FlaR is a Fe^{2+} binding protein.

**rFlaR has NADP reductase activity.** Flavin-binding domains are found in proteins with oxidoreductase activity. Nicotinamide adenine dinucleotide (phosphate) (NAD(P)H) is used as an electron donor in the enzymatic reactions of flavin reductases\textsuperscript{38-40}. To test whether rFlaR is involved in flavin-dependent redox reactions, we performed a flavin reductase activity assay as previously described\textsuperscript{40,41}. Oxidation of NADPH, measured at A\textsubscript{340}
room temperature was found to be significantly oxidized (Fig. 6b, p = 0.0001). The reaction mixtures without individual components (w/o rFlaR, w/o FAD, w/o NADP) served as controls and no redox activity could be detected (whole graph comparisons using one way ANOVA with Dunnett’s post hoc test, ****p = 0.0001). (b) NADP-reductase activities of rFlaR refolded at the presence (rFlaR (Fe2+)) and rFlaR refolded at pH 8.0 (rFlaR (pH 8)) at the absence of Fe3+ were compared. In the control reaction mixture rFlaR was replaced with BSA. The difference between the absorbance at 340 nm after 90 min incubation and the absorbance at 340 nm at time 0 (∆A340 nm) was calculated. The results presented are the mean of three different experiments (one way ANOVA with Dunnett’s Post Hoc multiple comparisons test, n = 3 in each group, *p = 0.02; ****p = 0.0001).

**rFlaR NADP reductase activity is Fe3+ dependent.** To evaluate the importance of Fe2+ for rFlaR enzymatic activity, different divalent ion chelators were introduced into the reaction mixture. EDTA (10 nM), Nitrilotriacetic acid (NTA; 10 nM) and citrate (10 nM) all significantly inhibited rFlaR NADP reductase activity (Fig. 5a; p = 0.0001). To rule out the possibility that Fe3+ is oxidized by H2O2, NADP reductase activity was measured in the presence of the H2O2 scavenger, salicylate. Indeed, salicylate did not inhibit NADPH production by rFlaR (Fig. 5b), indicating that H2O2 does not affect FlaR activity directly.

**Fe2+ is the electron donor in the rFlaR-NADP reductase activity.** The next step was to determine the electron donor in rFlaR reductase activity. When cysteine was used as a reducing agent in the reaction mixture, reduced cysteine was applied to solutions containing either ammonium iron sulfate, or FAD or NADP. Ammonium iron sulfate incubated for 2 hours at room temperature was found to be significantly oxidized (Fig. 6b, p = 0.0001). Addition of cysteine reduced the oxidized Fe3+ to Fe2+, as detected by the ferrozine stain (Fig. 6b; p = 0.0001). Both NADP and FAD were tested in their oxidized forms and no reduction in NADP and FAD could be observed in the presence of cysteine. These findings suggest that the stimulation of rFlaR activity occurs through the recycling of Fe3+ to Fe2+ by cysteine. To verify this point, excess Fe2+ ions were supplemented into the reaction mixture containing rFlaR, FAD and NADP. Indeed, rFlaR NADP reductase activity was stimulated in the presence of Fe2+ in a concentration-dependent manner (Fig. 6c, Pearson correlation, r = 0.9947, p < 0.0001).

**FlaR contributes S. pneumoniae resistance under oxidative stress conditions.** To determine the importance of FlaR to S. pneumoniae resistance to oxidative stress, flaR null mutant (WU2ΔflaRermer) and complemented strains (WU2ΔflaRflaRermer/Kan plasmid and WU2ΔflaRermer/Kan chromosome) were constructed as
described in the Methods section. No differences in the growth rates could be observed when WU2 WT and all above strains were grown in THY under anaerobic conditions (Supplementary Fig. S3). Following a challenge with 20 mM H₂O₂, significantly reduced survival of WU2 ΔflaR Erm was measured, 44%, compared to 93%, 92% survival of WU2 WT, WU2 ΔflaRflaR/Erm/Kan plasmid and WU2 ΔflaRflaR/Erm/Kan chromosome respectively, (Fig. 7, p = 0.0001). These findings highlights the involvement of FlaR's in oxidative stress resistance. However, chromosomal complementation did not fully restore bacterial survival under oxidative stress conditions in comparison to the WT (Fig. 7, p = 0.0011).

FlaR contributes to *S. pneumoniae* virulence. To test whether FlaR contributes to *S. pneumoniae* virulence, BALB/c mice were inoculated intraperitoneally (IP) with WU2 WT or WU2 ΔflaR Kan strains. Increased survival was observed in mice inoculated with WU2 ΔflaR Kan compared to inoculation with the WT WU2, though it did not reach statistical significance (Fig. 8a). In an attempt to increase signal-to-noise responses, IP inoculation was performed using CBA/CaHN-Btkxid/J mice (CBA/N xid). This mouse strain carries a mutation in the Bruton kinase (*btk*) gene that renders it highly susceptible to *S. pneumoniae* infection. Significantly increased survival rates were observed in CBA/N xid mice inoculated IP with WU2 ΔflaR Kan compared to mice inoculated with WU2 WT (Fig. 8b, p = 0.0236).

Mechanisms underlying FlaR contribution to *S. pneumoniae* virulence. The reduced virulence in the peritoneum in the absence of FlaR may result from two different mechanisms: increased sensitivity to oxidative stress elicited by immune cells or reduced adhesion and spread in the host. To test the first possibility, the extent of phagocytosis of WU2 WT, WU2 ΔflaR Erm and WU2 ΔflaRflaR/Erm/Kan complemented strains was evaluated. Interestingly, no significant differences in the phagocytosis by primary mice macrophages were observed for all of these strains (Fig. 9a, p = 0.1190). These data suggest that the reduced pneumococcal virulence in the peritoneum in the absence of FlaR is not due to increased sensitivity to oxidative stress produced by the recruited immune cells. This is in accordance with data published recently, showing that oxidation is not the major killing mechanism of *S. pneumoniae* by phagocytic cells.

The surface expression of FlaR, presented in the previous studies (Fig. 1), encouraged testing the second mechanism of FlaR involvement in virulence i.e adhesion. As demonstrated in Fig. 9b, WU2 ΔflaR Erm mutant adhesion to lung-derived epithelial (A549) cells was significantly reduced compared to the WU2 WT (~76%; Fig. 9, p = 0.0001), WU2 ΔflaRflaR/Erm/Kan plasmid (~73%, p = 0.0001) and WU2 ΔflaRflaR/Erm/Kan chromosome (~64%, p = 0.0001). Complementation in WU2 ΔflaRflaR/Erm/Kan plasmid restored bacterial adhesion to A549 cells to about 88%, not significantly differing from the WU2 WT. The adhesion of WU2 ΔflaRflaR/Erm/Kan chromosome strain was restored only up to 65%, remaining significantly lower compared to the WU2 WT (Fig. 9b, p = 0.0011). These data suggest that FlaR is involved in bacterial adhesion to the host cells in addition to its enzymatic function. Similarly, reduced adhesion to the glioblastoma cell line U251 and the motor neuron cell...
line NSC 34 was observed with WU2ΔflaR in comparison to the WT and to the two complemented strains (data not shown).

**Vaccine potential of rFlaR.** The finding that FlaR contributes to *S. pneumoniae* virulence, combined with the lack of significant similarity to any human protein, directed us to test the vaccine potential of rFlaR. Immunization of BALB/c mice with rFlaR elicited the production anti-rFlaR antibodies. An increase in antibody levels could be observed following the secondary immunization however this increase reached significance only after the tertiary immunization. The antisera titer following the third immunization reached ~1:10000 (Supplementary Fig S4, *p* = 0.014).

In addition, significant protection against IP lethal *S. pneumoniae* challenge was observed following immunization with rFlaR in comparison to adjuvant only immunization (Fig. 10a 15 μg rFlaR + alum, *p* = 0.0388; Fig. 10b 25 μg rFlaR + alum, *p* = 0.0001; Fig. 10c 35 μg rFlaR + CFA/IFA, *p* = 0.0053). To further demonstrate rFlaR ability to elicit protective immune response we have performed a modified method of passive immunization. Bacteria were incubated *ex-vivo* with either pre-immune or anti-rFlaR antiserum. Mice were inoculated IP with bacteria pre-treated with the anti-rFlaR antiserum survived significantly longer than mice inoculated with bacteria pre-treated with pre-immune serum (Fig. 10d, *p* < 0.0001).

**Figure 6.** Participation of Fe$^{2+}$ in FlaR enzymatic activity. (a) Cysteine significantly stimulates rFlaR NADP-reductase activity (Spearman correlation, r = 1, **p** = 0.0083). (b) Solutions containing either 10 μM Fe$^{2+}$ or 200 μM NADP or 50 μM FAD were incubated with cysteine. The oxidative state of Fe$^{2+}$ was assessed with ferrozine staining and monitored at A$^{562}$ nm. The oxidative state of FAD was monitored at A$^{450}$ nm and that of NADP monitored at A$^{340}$ nm. Fe$^{2+}$, FAD and NADPH in aqueous solutions were allowed to oxidize for 2 hrs, and then cysteine was added for additional 2 hrs. Oxidation was observed only in Fe$^{2+}$ solution which was significantly reduced upon cysteine addition. The results presented are the mean of three different experiments (one way ANOVA with Dunnett’s post hoc multiple comparisons test, n = 3 in each group, **p** = 0.0001). (c) Increasing concentrations of Fe$^{2+}$ significantly enhanced rFlaR NADP-reductase activity (Pearson correlation, r = 0.9947, ****p < 0.0001).
Discussion

A putative flavin reductase (FlaR) was recently annotated in *S. pneumoniae* TIGR4 strain in the NCBI database. We have previously shown that there is an age-dependent enhancement of the antibody response to this protein. Therefore, we initially named it pneumococcal surface immunogenic protein B (PsipB), but in order to conform with the NCBI database we currently refer to this protein as FlaR. FlaR was found to be surface expressed, although it is devoid of a signal peptide or any other known export sequences. Several additional *S. pneumoniae* proteins such as α-enolase, glyceraldehyde 3-phosphate dehydrogenase, fructose-bisphosphate aldolase, glutamyl tRNA synthetase, NOX and PtsA were also shown to be cell surface localized albeit lack of known export signals. The current study further confirmed the cell surface localization of FlaR, using immunoblotting and flow cytometry.

FlaR was found to be highly conserved in different *S. pneumoniae* strains. In addition, highly conserved cysteines at positions 66, 72 and 84 of WP_000580663.1 were found in all *S. pneumoniae* sequenced flavin reductases. Multiple cysteines in a protein may enable the formation of multimers. Indeed, resolving rFlaR protein on SDS-PAGE demonstrated the existence of both dimeric and monomeric forms of FlaR. Equilibrium between the dimeric and monomeric forms may be important for FlaR enzymatic activity, as hydrogenases are known to often function as homo-dimers or multi-enzyme complexes. Moreover, multiple cysteines in a protein sequence may indicate a transitional metal ion binding capacity. Metal ions can interact with metal-binding proteins during their folding and create a local structure that initiates and directs the protein-folding process and is considered critical for its biological activity. In accordance, rFlaR could be refolded/solubilized at physiological pH only under anoxic conditions.
in the presence of Fe^{2+} and to lesser extent with Co^{2+} and Cu^{2+}. Noteworthy is the finding that rFlaR, refolded at pH 8 without Fe^{2+}, bound Fe^{2+} as detected by the ferrozine stain.

We validated the putative flavin reductase activity of rFlaR and found that NADP and FAD are necessary components for its enzymatic activity, as described for NADPH reductases. In contrast, the flavin reductase activity of ferredoxin-NADP^+ reductase from *Pseudomonas putida* was FAD-independent. The activity of NADPH-cytochrome P450 reductase is generally accompanied by measurable alterations in FAD and FMA oxidative state. Although rFlaR catalyzed NADP reduction in a FAD-dependent manner, we failed to detect the change in FAD oxidative state.

Refolding of rFlaR in the presence of Fe^{2+} enhanced its enzymatic activity in comparison to the rFlaR refolded at pH 8, possibly as a result of increased initial binding of Fe^{2+}. The ability of divalent ion chelators to inhibit rFlaR activity reinforce Fe^{2+} necessity for NADP reduction by FlaR.

The midpoint potential of Fe^{2+} in solution, as a complex with EDTA, is +0.1 V while the midpoint potential of NADP is much lower (~0.32 V), rendering reduction of NADP by Fe^{2+} improbable. However, the midpoint potential of Fe^{2+} bound to 4Fe-4S cluster of ferredoxin from *Butyrirbacterium methylotrophicum* was found to be ~0.410 V and ~0.472 V in 2Fe-2S cluster of oxidoreductase from *Paracoccus denitrificans*. Additionally, it was shown that substitution of amino acid residues around an Fe^{2+}-sulfur cluster, by site-directed mutagenesis, affects the midpoint potential of the Fe^{2+} in the cluster. The finding that under physiological conditions rFlaR remains soluble only in the presence of Fe^{2+}, impedes determination of the midpoint potential of Fe^{2+} bound to FlaR. Yet, the demonstration of Fe^{2+} dependent rFlaR NADP reductase activity indicates that the FlaR-bound Fe^{2+} midpoint potential is probably lower than that of NADP.

Mucosal surfaces lack free Fe^{2+} due to chelation by host Fe^{2+}-binding proteins. *S. pneumoniae* has several mechanisms enabling it to cope with iron deficiency, among them hijacking iron from hemin, hemoglobin or lactoferrin, or direct iron binding by ABC iron transporters Pit1B and Pit2A. A double mutant of pit1B and pit2A demonstrated growth deficiency in a cation-depleted medium, which was restored by the addition of Fe^{2+} and Cu^{2+}. In addition, the 12 mer Dpr/Dps that bind, oxidize and store ~500 Fe^{2+} atoms in its hollow cavity has been identified in streptococci. The current study revealed that FlaR belongs to the Fe^{2+} binding proteins.

rFlaR activity was found to be enhanced in the presence of increasing concentration of the reducing agent, cysteine. Our demonstration that cysteine reduced Fe^{2+} but not FAD or NADP, implies that the increase in NADPH in the presence of cysteine stemmed from Fe^{3+} to Fe^{2+} recycling, which stimulated rFlaR activity. In line is the finding that rFlaR NADP reductase activity was enhanced in a Fe^{2+} concentration-dependent manner.

Tight regulation of the free Fe^{2+} pool can be especially important for pneumococci, which produce high levels of H_{2}O_{2} but lack catalase, and hence are at high risk of damage by ROS formed via the Fenton reaction. We hypothesized that FlaR is involved in preventing the accumulation of toxic levels of free Fe^{2+} that may react with...
H₂O₂ and produce ROS. Indeed, WU2ΔflaR demonstrated marked susceptibility to H₂O₂ and complementation restored bacterial resistance to H₂O₂. Of note, the strain complemented with the plasmid demonstrated increased resistance to H₂O₂ in comparison to the strain complemented in chromosome. The explanation may be that following flaR trans-complementation, multi-copies of flaR are expressed, whereas following cis-complementation (into the same chromosome) a single copy of flaR is expressed, under the control of its native promoter. This may stem from two concomitantly occurring mechanisms: i) FlaR’s function in binding and oxidizing Fe²⁺ to protect the bacterium from the toxic effects of the Fenton reaction; ii) FlaR’s NADP reductase activity provides an additional source of NAD(P)H beside the Embden-Meyerhof-Parnas and the pentose phosphate pathways. Increased NAD(P)H levels are essential under high H₂O₂ levels for NADH oxidase activity. The action of NADH oxidase makes pyruvate available for SpxB oxidation, resulting in the production of two additional ATP molecules per one glucose molecule, beyond that available from conventional glycolysis. Moreover, the electron transport to NADPH by FlaR enzymatic activity may increase the reduction of cysteine containing enzymes to maintain redox balance thus enhancing bacterial growth and virulence.

In contrast to our finding that the presence of FlaR in the cells confer resistance to oxidative stress, Pericone et al. claimed that iron chelators had no effect on the survival of WT or a mutant lacking pyruvate oxidase (ΔspxB) following a challenge with 20 mM H₂O₂. This led the authors to suggest that the Fenton reaction does not play a major role in the killing of S. pneumoniae by H₂O₂. The mechanism by which FlaR contributes to bacterial survival under oxidative stress conditions should be further elucidated. Of note is the finding that Fe²⁺ dependent-rFlaR activity was not affected by the H₂O₂ scavenger, salicylate, suggesting that Fe²⁺ oxidation is not directly executed by H₂O₂.

The contribution of FlaR to pneumococcal virulence was tested in a mouse model of sepsis. The reduced virulence of S. pneumoniae in the peritoneum, in the absence of FlaR, could be explained by increased sensitivity to phagocytosis and oxidative stress elicited by immune cells. The phagocytic role of macrophages in S. pneumoniae engulfment and killing has been previously demonstrated for bronchoalveolar lavage derived macrophages, peritoneal macrophages and neutrophils. Interestingly, we found that FlaR absence did not significantly affect the extent of phagocytosis by primary mice peritoneal macrophages. In line, in vitro studies demonstrated that the predominant killing mechanism of pneumococci in the phagosome of neutrophils is executed by the proteolytic activity of two serine proteases rather than oxidation. The involvement of the serine proteases in pneumococcal killing was further confirmed in vivo demonstrating higher bacterial load in the spleen of mice lacking the serine proteases in a systemic infection model. These authors also emphasized that in the case of
systemic infection with *S. pneumoniae*, most of the bacteria can be found in the spleen and to a lesser extent in other organs. The peritoneum is characterized by low oxygen pressure. Thus, the role of FlaR in protection against oxidative stress could not explain the reduced virulence of *WU2ΔflaR* in this niche suggesting an additional role for FlaR in virulence. FlaR was found to be surface expressed using immunoblotting of cell-wall proteins and flow cytometry. Pre-incubation of *S. pneumoniae* with anti-FlaR antisera, which interacts only with the surface exposed FlaR, reduced bacterial virulence, reinforcing the surface localization of FlaR. Many surface proteins were found to mediate bacterial adhesion to the host. Hence, we hypothesized that FlaR may be involved in bacterial adhesion to host cells. Indeed, *WU2ΔflaR* strain was found to adhere to A549 cells significantly less than its parental WT and the two complemented strains. We have also used glioblastoma cells U251 and motor neuron cells NSC34 to establish FlaR contribution to *S. pneumoniae* adhesion and similar results were obtained. The ability of anti-rFlaR antibodies to reduce bacterial virulence also implies a direct involvement of the surface expressed FlaR in virulence and diminishes the possibility that downstream consequences of the mutation, such as upregulation of other adhesins, are responsible for the bacterial virulence.

The following findings highlight FlaR as a potential candidate vaccine: 1) possesses age dependent antigenicity; 2) is conserved among *S. pneumoniae* strains; 3) lacks significant homology to human proteins; 4) protects the bacterium against oxidative stress and 5) possibly mediates *S. pneumoniae* adhesion to the host. Indeed, FlaR elicited significant protection against IP challenge. In addition, sera obtained from immunized rabbits were able to neutralize *S. pneumoniae* virulence in the peritoneum, probably by enhancing opsonophagocytosis and/or interfering with bacterial adhesion and spread. Hence, rFlaR can be considered as a future vaccine candidate. It should be noted that proteins, such as PspA, CbpA, PavA, PavB and PhdT, which are involved in different aspects of bacterial virulence including adhesion, are known to be immunogenic and to elicit a protective immune response in experimental infection models. Furthermore, PhdT has recently been shown to elicit an immune response in phase I/II clinical trials.

In summary, this study attributes FlaR multiple functions in *S. pneumoniae* physiology and virulence: 1) Fe\(^{2+}\) binding and oxidizing to prevent the toxic effect of the Fenton reaction; ii) reducing NADP to provides an additional source of NAD(P)H, resulting in maintenance of redox balance and ATP production; and iii) mediating bacterial adhesion to the host. Importantly, we found that rFlaR can be considered for future vaccine development. Further studies should concentrate on detailed understanding of FlaRs role in pneumococcal biology and the nature of the immune protection elicited by FlaR.

**Methods**

**Inoculation and immunization of mice and ethics statements.** This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Institutional Animal Care and Use Committee of the Ben-Gurion University of the Negev, Beer Sheva, Israel (Permit number: 53:08.08).

Seven-week-old BALB/cOlaHsd (BALB/c) female mice (Harlan Laboratories, Israel) or seven-week-old CBA/ CaFIN-Btk<sup>−/−</sup> (CBA/N<sup>−/−</sup>; Jackson Laboratories, Bar Harbor, ME, USA) mice were housed in sterile conditions under 12-h light/dark cycles and fed Purina Chow and tap water ad libitum. The groups were matched for age.

BALB/c mice were inoculated IP (without anesthesia) with the *WU2 WT* (n = 8; 64 CFU) or *WU2ΔflaR<sup>kan</sup>* (n = 7; 64 CFU) strains. Survival was monitored daily. Additionally, CBA/N<sup>−/−</sup> mice were inoculated IP (without anesthesia) with *WU2 WT* (n = 10; 80 CFU) or *WU2ΔflaR<sup>kan</sup>* (n = 8; 80 CFU), and survival was monitored daily.

BALB/c mice were immunized subcutaneously with 15 µg rFlaR (rFlaR + alun n = 17; alun n = 7) or 25 µg rFlaR (rFlaR + alun n = 17; alun n = 7) mixed with 7.5 µl of the Imject Alum adjuvant (Pierce Biotechnology, Inc., Rockford, IL) or 35 µg rFlaR emulsified with complete Freund’s adjuvant (CFA) on primary immunization and boosted (days 14 and 28) with rFlaR emulsified with incomplete Freund’s adjuvant (IFA) (rFlaR + CFA/IFA n = 19; CFA/IFA only, n = 14). Mice were challenged IP on day 42, under deep anesthesia using isoﬂurane (Piramal Critical Care Inc., PA, USA) with a lethal dose of *WU2 WT* strain (10<sup>7</sup> CFU).

Ex-vivo neutralization was performed as follows: a lethal dose (~10<sup>6</sup> CFU) of *WU2 WT* strain was incubated at 37°C for 1 h with 1:10 diluted rabbit pre-immune serum or anti-rFlaR serum and subsequently inoculated IP to BALB/c mice (n = 10 in each group). Survival was monitored daily.

For all animal experiments, mice were humanely euthanized by CO<sub>2</sub> asphyxiation, as recommended by the American Veterinary Medical Association (AVMA) guidelines for euthanasia 2013, if they became moribund or been carried out.

**Chemicals and Biological Reagents.** Unless otherwise stated, all chemicals and biochemicals of highest purity available were purchased from Sigma-Aldrich Corp. (St Louis, MO). FAD and NADP were purchased from Applicheme GmbH (Darmstadt, Germany). For research involving biohazards, correct standard procedures have been carried out.

**Bacterial Strains and Growth Media.** *S. pneumoniae* serotype 3 strain *WU2*<sup>46</sup> and its derivatives, described below, were used. Pneumococci were grown on blood agar plates or in Todd-Hewitt broth supplemented with 0.5% (w/v) yeast extract (THY). Two *Escherichia coli* strains, DH5α, UltraMAX (DH5α; Invitrogen Corp, Carlsbad, CA, USA) and *E. coli* BL21(DE3)pLysS (BL21; Promega Corp, Madison, WI), were grown in Luria broth (LB), Lennox.
Cloning, Expression and Purification of rFlaR, rFabD, rATCase and antibodies production. The flaR gene was amplified from S. pneumoniae strain WU2 genomic DNA by PCR using the primers FlaR-F and FlaR-R (Supplementary Table S1, flaR pHAT expression) designed according to the sequence in the R6 strain (accession number AP04889). The amplified and BamHI-SacI (Takara Bio Inc, Shiga, Japan)-digested DNA-fragments were cloned into the pHAT expression vector (BD Biosciences Clontech, Palo Alto, CA) and transformed into DH5α E. coli cells. The vector was purified using the Qiagen High Speed Plasmid Maxi Kit (Qiagen GMBH, Hilden, Germany) and transformed into E. coli BL21. Sequencing was performed to rule out any mutation. rFlaR expression was induced by incubation for 5 h with 1 mM of IPTG at 37°C. The protein was purified under denaturing conditions using Ni-NTA agarose beads (Qiagen GMBH). This preparation was dialyzed against PBS pH 7.3, containing 2 mM FeSO₄ for 48 h with three changes of buffer (named rFlaR Fe⁺⁺). Isolation of the protein was confirmed by immunoblotting with anti-HAT (BD Biosciences Clontech) and thereafter with anti-rFlaR antibodies (described below) and trypsin in-gel digestion of the protein to derive peptides for MALDI-TOF mass spectrometry sequencing (Bruker Reflex-IV mass spectrometer Bruker-Daltonik, Bremen, Germany). Alignment to existing databases was performed using both the Mascot software package (Matrix Science Ltd, UK, http://www.matrix-science.com) and the “Profound” Program provided by the Rockefeller University.

Refolding of rFlaR was performed using 20 mM of the following buffers: Acetate pH 5.0, 2-(Nmorpholino)ethanesulfonic acid (MES), pH 6.0, phosphate pH 7.0 and pH 8.0, tris(hydroxymethyl)aminomethane pH 9.

Untagged protein was produced by amplifying the gene using the S. pneumoniae TIGR4 strain (ATCC, Rockville, MD, USA) DNA as template, using specific primers (Supplementary Table S1, flaR pET30a (+)). Digestion of pET30a (+) (Novagen, Madison, WI, USA) with NdeI and Bpi1102I restriction enzymes, prior to ligation removed all tags from the vector. Then, flaR was cloned into the vector. Protein expression was induced with IPTG in fermented E. coli BL21 followed by inclusion bodies isolation. The inclusion bodies were centrifuged and resuspended in double distilled water and frozen in aliquots. The inclusion bodies were refolded by dialysis for 1 h against a buffer adjusted to pH 11.3, containing 4.5 M urea, 40 mM Tris base and 1 mM cysteine, folowed by the addition of 0.67 mM arginine and continued dialysis overnight. The resulting solution was concentrated to 4.4 fold. This preparation was used to a preparative Superdex 200 column and elution was performed with buffer adjusted to pH 10, containing 4.5 M urea, 40 mM Tris base and 1 mM cysteine. The tubes containing the protein were pooled and dialyzed against NaHCO₃ at pH 11 followed by lyophilization. The lyophilized protein was suspended in PBS pH 8 (named FlaR pH 8).

For antibodies production, three-month-old New Zealand white rabbits and 7-week old BALB/c mice were immunized with Ni-NTA-purified rFlaR formulated with CFA in the first immunization and IFA in the two booster immunizations with two-week intervals between immunizations. The rabbits were bled following the third immunization.

FabD was cloned and expressed as a His-tagged soluble protein and antibodies were produced as previously described. ATCase is known to be involved in pyrimidine synthesis. ATCase is a cytoplasmic protein that was found to be surface exposed. ATCase (SP_RS06260) was cloned and expressed as a His-tag soluble protein. Briefly, ATCase was amplified from the R6 strains using specific primers (Supplementary Table 1, ATCase gene for pET32a+) and ligated into pET32a+ (Novagen, Madison, WI, USA) following digestion with EcoR1 and Xho1 restriction enzymes. The plasmid was transformed into E. coli BL21, and the protein was purified with Ni-NTA-purified rFlaR formulated with CFA in the first immunization and IFA in the two booster immunizations with two-week intervals between immunizations. The rabbits were bled following the third immunization.

Flow cytometry of S. pneumonia. Flow cytometry was performed as previously described. Briefly, R6 bacteria were incubated with mouse anti-rFlaR antibodies or control mouse serum, washed, and stained with Alexa Fluor 647-conjugated goat-anti-mouse-IgG (Jackson ImmunoResearch, West Grove, PA). Flow cytometry was performed using a FACSCalibur flow cytometer (Becton Dickinson, Mountain View, CA), and data were acquired and analyzed using BD CellQuest™ 3.3 software.

Isolation of S. pneumoniae Cell-Wall (CW) Proteins. CW proteins were isolated by the method of Siegel et al. Briefly, bacteria were harvested by centrifugation at 4700 g for 15 min, washed with PBS and incubated with 3 ml protoplast buffer (20% sucrose, 25 mM MgCl₂, 5 mM Tris-Cl, pH 7.4), 1000 U mutanolysin and 0.2 ml protease inhibitor cocktail for 1 h at 37°C. The soluble proteins released from the cell-wall were collected after centrifugation at 25,000 g and stored at −70°C.

Immunoblotting Analysis. Protein mixtures were separated by SDS-PAGE and transferred to nitrocellulose membranes (Bio-Rad Laboratories, Inc., Carlsbad, CA), according to manufacturer’s instructions. Immunoblotting analysis of FlaR expression was performed with rabbit anti-sera obtained against purified rFlaR and detected with Cy™3 AffiniPure Donkey Anti-Mouse IgG (H + L) (Jackson ImmunoResearch, West Grove, PA).

rFlaR NADP Reductase Assay. The reaction mixture contained 50 µM FAD, 200 µM NADP, 10 µM ammonium iron sulfate and 6 µM rFlaR in PBS buffer. NADP reduction was detected by increased absorbance at 440 nm, which does not overlap ferrous absorbance at 520 nm, as measured with a BioRad SmartSpec 3000 Spectrophotometer (Bio Rad; Oceanside, California, USA).

Detection of Iron Binding. The assay was performed as described by Da Lozzo et al., with slight modifications. Briefly, rFlaR (6 µM) was mixed with PBS containing 20 µM ammonium iron sulfate. Ferrozine (50 µM) was added to the reaction mixture and absorbance at 562 nm was measured after 10 min. BSA (6 µM), rACTase and EDTA (10 nM) were used as negative and positive controls, respectively.
Ability of Cysteine to Reduce Iron. Freshly prepared solution of 20µM ammonium iron sulfate in PBS was incubated for 2h at 25°C to allow Fe²⁺ oxidation prior to the addition of 5 mM cysteine. The extent of Fe²⁺ oxidation in PBS solution was tested with ferrozine stain, and absorbance monitored at A562 nm. The same manipulations were performed for 200µM NADP and 50µM FAD solutions in PBS, and absorbance was measured at A340 nm and at A450 nm, respectively.

Bioinformatics Analyses. All completely sequenced genomes of S. pneumoniae were downloaded from NCBI's Genome database (29 genomes as of October 2nd, 2016). The flavin reductase amino acid sequence of the TIGR4 strain (WP_000580663.1) was used as a query in tblastn and blastp searches against each of these genomes and their predicted proteins. WP_000580663.1 sequence was then used as a query in a blastp search against all RefSeq proteins annotated as “S. pneumoniae” and “flavin reductase”. The 63 retrieved protein hits were subjected to multiple sequence alignment using Tcoffee with default parameters. A representative set of 9 divergent proteins was selected from the multiple sequence alignment by inspecting a Neighbor Joining tree and choosing a representative protein from each clade. Multiple sequence alignment of these proteins was performed using Tcoffee and visualized using the Jalview software. Finally, the entire RefSeq protein database except for S. pneumoniae proteins was searched by blastp with WP_000580663.1 as a query. Only GenBank records containing the word ‘Flavin’ and having blastp e value < 0.001 were retrieved.

Preparation of WU2ΔflaR Mutant Bacteria (kanamycin or erythromycin resistance). To create null mutants of the flaR gene, the kanamycin (Kan) or erythromycin (Erm) resistance cassette was inserted into the coding sequence of flaR via homologous recombination. The transformation procedure was performed as previously described85 with minor modifications. The upstream and downstream flanking regions of flaR were amplified by PCR from DNA of WU2 strain using the primer combinations upwing-F/upwing-R (Supplementary Table S1, Upwing WU2ΔflaR), for the upstream region, and downwing-F/downwing-R, for the downstream region (Supplementary Table S1, Downwing WU2ΔflaR). The upwing 1149 bp harbors 125 bp of flaR. The downwing 582 bp harbors 68 bp of the flaR. Hence, the expected length of the flaR fragment to be deleted was 277 bp. The Kan cassette was amplified by PCR from the genome of CP125086 using the primer combinations Kan AB-F/Kan AB-R (Supplementary Table S1, Kan AB cassette). The Erm cassette was amplified by PCR from the genome of WU2ΔΔnoxAΔerm2 using the primer combinations Erm AM-F/Erm AM-R (Supplementary Table S1, Erm AM cassette). The PCR products were digested with the corresponding restriction nucleases, as specified in Supplementary Table S1, purified, ligated and transformed into WU2 in the presence of competence stimulating factor, CSP1 and CaCl₂. Transformants were selected on THY solidified with 1.5% agar, containing kanamycin (80µg/ml) or erythromycin (125µg/ml). Verification of flaR deletion was done by PCR using specific primers (Supplementary Table S1, Kan AB cassette for Kan resistance and Erm AM cassette for Erm resistance). It is worth mentioning that phenotypically, flaR mutation did not affect the size and mucosity of the colonies.

Of note, the kanamycin resistant mutant strain, WU2ΔflaRKan, was used in the mice inoculation studies. The erythromycin resistant mutant strain, WU2ΔflaRErm, was used in the susceptibility to oxidative stress experiment and the adherence to A549 cells assay.

Construction of two WU2ΔflaR(ΔflaR/Erm/Kan) complemented strains. Trans-complementation of WU2ΔflaR(ΔflaR/Erm). Complementation was performed using Gram positive compatible plasmid, pBAV1K-T5-gfp (Addgene; Cambridge, MA), bearing the kanamycin resistance cassette. The flaR gene was amplified from WU2 genome using flaR-F and flaR-R primers (Supplementary Table S1, flaR pBAV-K1 plasmid and pCPE complementation), which introduce NcoI and PstI sites. Subsequently the plasmid was digested, resulting in gfp gene deletion and retaining the constitutive T5 viral promoter. Following the ligation of flaR to pBAV1K-T5-gfp, the plasmid was transformed into E. coli DH5α, overexpressed, purified and transformed into WU2ΔflaR(ΔflaR/Erm) to produce WU2ΔflaR(ΔflaR/Erm/Kan) complemented strain. Successful insertion of pBAV1K-T5 into the chromosome was verified by PCR.

Cis-complementation of WU2ΔflaR(ΔflaR/Erm). To confirm that mutation of flaR in WU2ΔflaR(ΔflaR/Erm) introduced no polar effects, WU2ΔflaR(ΔflaR/Erm) was complemented with an intact copy of the gene using pCPE, which is a nonreplicative plasmid that allows controlled gene expression under its native promoter, following ectopic integration into the chromosome85,86. Briefly, flaR was amplified from WU2 with flaR-F and flaR-R primers (Supplementary Table S1, flaR pBAV-K1 plasmid and pCPE complementation), which introduce NcoI and PstI sites. Then the amplicons were ligated into NcoI and PstI digested vector. An aliquot of ligation mixture was transformed into Stellar™ competent cells (Clontech, Saint-Germain-en-Laye, France) as described by the manufacturer. The transformants were selected for kanamycin resistance (500µg/ml). Successful ligation was determined by colony PCR using Mal-F and pCPE-R primers (Supplementary Table S1, flaR pCPE verification), whose recognition sites are localized immediately up and downstream of the cloning site, respectively. These primers amplified 263 bp product in empty vector, while they produce a product of 913 bp in the recombinant clones (additional 650 bp represents the cloned fragment containing flaR). The recombinant plasmid was purified using a commercial kit (Qiagen), and an aliquot was transformed into WU2ΔflaR(ΔflaR/Erm) as described previously89 to produce the complemented strain WU2ΔflaR(ΔflaR/Erm/Kan) chromosome. The transformants were selected on blood agar plates supplemented with erythromycin (125µg/ml) and kanamycin (500µg/ml). Integration of flaR into the genome was confirmed by PCR (Supplementary Table S1, Kan resistance in pCPE verification).

Oxidative stress assay. Oxidative stress assays were performed as previously described85 with minor modifications. Briefly, WU2 WT, WU2ΔflaR(ΔflaR/Erm) and WU2ΔflaR(ΔflaR/Erm/Kan) plasmid and chromosome strains were grown under anaerobic conditions in 20 ml of THY medium to an OD₆₀₀ = 0.3. Five ml of culture were harvested
Primary macrophage phagocytosis assay. Mice were injected intraperitoneally with 3% thioglycolate broth. Three days later mice were euthanized, using CO₂, and their peritoneal cells were harvested by flushing the peritoneum with 10 ml of PBS. Cells were centrifuged and resuspended in complete (c) RPMI 1640 medium supplemented with 10% FCS and 2% L-glutamine. 2 × 10⁶ cells/well were plated onto 96 well culture plates. The macrophages were allowed to adhere for 9 hours and the supernatant was removed. 2.5 × 10⁶ of midlog harvested bacteria in RPMI 1640 were added to the well for 40 minutes incubation at 37 °C. Following the incubation, aliquots were taken from the supernatant and plated onto blood agar plates for enumeration. The number of residual live bacteria CFU was subtracted from the initial bacterial CFU resulting in the number of phagocytosed bacteria. The data presented is the summary of 2 biological independent experiments each performed in triplicates.

S. pneumoniae adhesion to A549 cells assay. A549 cells (lung adenocarcinoma cells; ATCC, Rockville, MD, USA) retain morphological, biochemical and immunological characteristics resembling type II lung epithelial cells⁹⁰–⁹⁲ and have been widely used as a model to study S. pneumoniae interaction with human cells⁹³,⁹⁴. A549 cells were cultured on 96-well plates (2.5 × 10⁵ cells/well) in DMEM without antibiotics. Following overnight incubation each well contained ~10⁶ A549 cells. WU2 WT, WU2ΔflaR and WU2ΔflaRΔermKan strains (~10⁶ CFU; multiplicity of infection (MOI) 10:1) were added to the cells for 1 h incubation at 37 °C. Bacteria were enumerated after the 1 h incubation and no bacterial death under the experimental conditions could be observed in WT, mutant and complemented strains (data not shown). Excess bacteria were removed, and cells detached with 0.25% trypsin-EDTA and plated onto blood agar plates for enumeration. This experiments were performed in triplicate and repeated three times.

Statistical Analysis. The Shapiro-Wilk test was used and post-hoc statistical power was calculated, respectively, to verify that small sample sized data sets assumed normal distribution and are of the sufficient size to justify the use of one-way ANOVA for parametric data, followed by the Dunnett’s test for multiple comparisons. One-tailed Student’s t-test with Welch’s correction was used for bacterial load comparisons of two groups. Data was reported as the mean ± SEM, unless stated otherwise. Pearson and Spearman correlations were used to assess the significance of change. Survival of S. pneumoniae-inoculated mice was determined using Log-rank (Mantel-Cox) test. Differences were considered significant at p < 0.05. All statistical analyses were performed with the software package in GraphPad Prism version 7 (La Jolla, CA, USA).

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Author Contributions
G.I.M., N.P., R.B., D.A.M., R.D. and Y.M.N. designed the study. G.I.M., T.K., H.N., A.A., A.O., O.L., S.A., I.M., S.D., M.P., A.A.P., D.K., B.H., T.F., M.T., R.E., A.M.M., T.J.M. and H.Y. performed the experiments. G.I.M., A.A., V.C.C. and Y.M.N. analyzed the data. G.I.M., R.B., H.Y. and Y.M.N. wrote the manuscript. All authors discussed and approved of the manuscript.

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