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Influence of serogroup B meningococcal vaccine antigens on growth and survival of the meningococcus in vitro and in ex vivo and in vivo models of infection

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Abstract
A novel vaccine against serogroup B meningococcal disease - containing a combination of protein antigens identified by reverse vaccinology: fHBP fused to GNA2091, GNA2132 fused to GNA1030, and NadA - is currently in Phase III clinical trials. In order to determine the role of these antigens in the growth, survival and fitness of the meningococcus, we generated a mutant lacking the expression of all five protein antigens (5KO), a mutant lacking the three main antigens (fHBP, GNA2132 and NadA; 3KO), as well as strains lacking the single antigens. Our results show that abrogation of expression of these antigens in Neisseria meningitidis results in reduced growth in vitro, increased sensitivity of the bacterium to stresses it may encounter in the host, as well as reduced fitness in ex vivo models of infection and in an in vivo infant rat competitive index assay. These results support a multivalent vaccine approach, which was undertaken to strengthen the protective activity of the vaccine antigens, increase the breadth of MenB strains targeted by the vaccine, and limit the potential for selection of vaccine escape mutants.

Keywords: meningococcal vaccine antigens; reverse vaccinology; fHBP; NadA; GNA2132; GNA2091; GNA1030.

Running title: Role of meningococcal vaccine antigens in survival and fitness
1. Introduction

*Neisseria meningitidis* (the meningococcus) is an encapsulated Gram-negative diplococcus that colonizes the nasopharyngeal mucosa of approximately 5-10% of the population. However, the bacterium can also cross the epithelial layer into the bloodstream, leading to septicemia and/or meningitis [1]. Disease caused by *N. meningitidis* results in significant morbidity and mortality worldwide, with the majority of this disease being caused by five meningococcal serogroups (serogroups A, B, C, W-135 and Y). Conventional vaccinology approaches have led to the development of effective capsular-polysaccharide based vaccines against meningococcal serogroups A, C, W-135, Y [reviewed in 2]. However, there is still no broadly protective vaccine available against meningococcal serogroup B (MenB) disease, which is the main cause of meningococcal disease in many countries (approximately one-third of disease in the US [3] and more than half in Europe [4]). MenB vaccine candidates identified by conventional approaches have proven to be either (1) similar to self-antigens and as such poorly immunogenic (e.g. the capsule polysaccharide) or (2) hypervariable and/or poorly conserved between the diverse strains that cause endemic MenB disease (e.g. antigenically and phase variable outer membrane proteins and LPS) [5]. As a result of these limitations, the only currently available MenB vaccines are the “tailor-made” outer membrane vesicle (OMV) vaccines that have been developed and successfully used in New Zealand, Cuba and Norway to control epidemics that have primarily been caused by a single strain [6].

Recently, a multivalent recombinant vaccine against MenB, developed using genome-based "Reverse Vaccinology", has entered phase III clinical trials. The MenB vaccine contains 2 fusion proteins and one single polypeptide: factor H–binding protein (fHBP; formerly GNA1870) fused to GNA2091; GNA2132 fused to GNA1030; and NadA (formerly GNA1994). These antigens were selected based on their ability to induce broad protection inferred by serum bactericidal activity (SBA) assays or by passive protection in infant rat or mouse models [7]. NadA, fHBP and GNA2132 are considered the main antigens, whereas GNA1030 and GNA2091 are regarded as accessory proteins, due to a less pronounced protective activity. The recombinant antigens are formulated with alum, with or without OMV from the New Zealand MeNZB™ vaccine strain that contains PorA 1.4 as the main immunogen [8]. When tested against a panel of 85 MenB strains representative of the global population of disease-causing strains, the combination of the recombinant antigens induced bactericidal antibodies in mice against 78% and 90% of strains, when administered with the adjuvants aluminum hydroxide and MF59 (an oil-in-water emulsion), respectively [7]. Phase II clinical results in infants indicated that this vaccine was well tolerated and induced a protective immune response against three diverse MenB strains in 89–96% of subjects following three vaccinations, and 93–100% after four vaccinations [9].
fHBP is a *Neisseria*-specific surface lipoprotein that binds human factor H (fH), a key inhibitor of the complement alternative pathway (AP). Binding of human fH to fHBP on the meningococcal surface allows the pathogen to evade complement-mediated killing by the innate immune system [10, 11]. As a result, fHBP is important for survival of bacteria in human serum and blood, and is involved in protection against the antimicrobial peptide LL-37 via an unknown mechanism [10, 12, 13]. fHBP is expressed by all *N. meningitidis* strains studied to date [14] and can be classified into three variants (or two subfamilies) that are not cross protective [14, 15]. fHBP variant 1, present in the MenB vaccine, is present in approximately 50-60% of disease-producing group B isolates [16, 17].

NadA is a surface-exposed trimeric autotransporter adhesin that is involved in adhesion and invasion of *N. meningitidis* into human epithelial cells [18, 19]. Additional studies have shown that NadA is also involved in tissue and blood invasion as well as interacting with and stimulating immune cells during infection [20-22]. This antigen clusters into 3 main variants in virulent strains [16], and induces bactericidal antibodies that are not influenced by variant diversity. The *nadA* gene is present in approximately 50% of meningococcal isolates and is almost always present in three of the main hypervirulent lineages assigned by multilocus sequence type (MLST) [18, 23].

GNA2132 has recently been described as a heparin binding protein of *N. meningitidis* [24]. This antigen binds heparin at an arginine rich region and this interaction may increase bacterial serum resistance due to the potential interactions of heparin with components of the complement pathway, such as fH [24]. GNA2132 is cleaved by 2 different proteases, the *Neisserial* autotransporter lipoprotein NalP and human lactoferrin (Lf), and the functional significance of these cleavages is being studied. There are several variants of this protein [16] that are cross protective [7, 25].

To further increase the immunogenicity of the recombinant antigens, protein-protein fusions of fHBP and GNA2132 were generated with GNA2091 and GNA1030, respectively. GNA2091 was selected for its ability to induce protection in an mouse protection model, and GNA1030 for its ability to induce bactericidal antibodies against several strains. These proteins are well conserved in *N. meningitidis* strains, but are less well functionally characterized. The fusions with these proteins induce immune responses in SBA assays that are generally more potent than those induced by the individual antigens [7].

The inclusion of multiple antigens in the MenB vaccine should allow the breadth of MenB strains targeted by the vaccine to be increased, strengthen the protective activity of the vaccine, and reduce the risk of selecting for vaccine escape mutants (i.e. bacteria that have a mutation in a gene encoding an antigen that could allow them to avoid vaccine-induced bactericidal antibodies). To determine the contribution of each of the vaccine antigens in the growth and survival of *N.
meningitidis, and to evaluate the outcome of possible escape mutants lacking these vaccine antigens, MC58 derivative strains were generated that lacks the expression of all five antigens contained in the vaccine, a combination of the three main antigens, or each of the single antigens. These strains were analysed with respect to growth and survival in various media and stress conditions (including antimicrobial peptide, oxidative and envelope stress assays), in \textit{ex vivo} human serum and whole blood models of meningococcal bacteremia and \textit{in vivo} in the infant rat model of infection.

2. Materials and Methods

2.1. Bacterial strains and culture conditions

\textit{N. meningitidis} strains used in this study are described in Table 1. \textit{N. meningitidis} strains were routinely grown on GC (Difco) agar supplemented with Kellogg's supplement I, or on Mueller Hinton (MH) agar (Difco) at 37°C/5% CO$_2$ overnight. For liquid cultures, overnight growth was used to inoculate GC broth supplemented with Kellogg's supplement I, 12.5 µM Fe(NO$_3$)$_3$, Catlin6 broth [26], MH broth or MH broth supplemented with 0.25% glucose. When required, tetracycline, erythromycin, chloramphenicol and kanamycin antibiotics were used at final concentrations of 2 µg/ml, 5 µg/ml, 5 µg/ml and 100 µg/ml, respectively. \textit{Escherichia coli} strains used for cloning were cultured in Luria-Bertani (LB) broth or on LB agar (Difco). When required, erythromycin and tetracycline were used at a final concentration of 5 µg/ml and 10 µg/ml, respectively and chloramphenicol and kanamycin at a final concentration of 20 µg/ml.

Growth experiments were performed using a suspension of several colonies of bacteria taken from overnight plates, which were adjusted to an initial optical density at 600 nm (OD$_{600}$) of 0.05 in 7 ml of broth in 14 ml tubes. The OD$_{600}$ was then followed over time directly from the culture tubes. When shown, colony forming units (CFU) were also determined. Samples were taken at various times, from cultures grown in parallel, diluted and plated. Experiments were performed in triplicate and on several occasions.

2.2. Recombinant DNA techniques

Recombinant DNA techniques were routinely performed as described by Sambrook \textit{et al.} [27]. Plasmid DNA preparations and purification of DNA fragments from PCR samples and agarose were performed using QIAGEN kits according to the manufacturer’s instructions. PCR amplification was carried out in a 50 µl reaction consisting of 1x reaction buffer, 1.5 mM MgCl$_2$, 0.2 mM dNTPs, 1 unit of Platinum Taq polymerase (Invitrogen), and 10 pmol of each forward and reverse primer, an appropriate amount of template DNA (chromosomal or plasmid DNA, or DNA from a single colony resuspended and boiled in 100 µl water) and sterilized deionised water. All
2.3. Construction of knockout and complemented N. meningitidis strains

In order to generate N. meningitidis mutant strains lacking the antigens contained in the MenB vaccine, five knockout plasmids were constructed to enable the deletion of all or part of the respective antigen-coding genes and replacement by allelic exchange with an antibiotic resistance cassette. Using standard cloning procedures, upstream and downstream flanking regions of the \textit{nmb2132}, \textit{nmb1870}, \textit{nmb1030}, and \textit{nmb2091} genes, (encoding the GNA2132, fHBP, GNA1030 and GNA2091 antigens respectively) were amplified by PCR from the MC58 genome and cloned on either side of an erythromycin resistance (Erm) cassette into the pBluescript (pBS; Pharmacia) cloning vector, generating the knockout plasmids pBSUD287::Erm, pBSUD741::Erm, pBSUD953::Erm, and pBSUD936::Erm, respectively. The upstream regions of the \textit{nmb2132}, \textit{nmb1870}, \textit{nmb1030}, and \textit{nmb2091} genes were amplified using primer pairs U287-F1/U287-R1, U741-F1/U741-R1, U953-F1/U953-R1, and U936-F1/U936-R1 (Table 2), respectively, and the downstream regions using U287-F1/U287-R1, U741-F1/U741-R1, U953-F1/U953-R1, and U936-F1/U936-R1 (Table 2), respectively. For the generation of the NadA knockout construct, the \textit{nmb1994} gene was amplified by PCR from the MC58 genome using the primer pair 961-F3/961-R3 and cloned into the pBluescript cloning vector, and subsequently an internal HincII fragment at the 5’ end of the coding region was substituted with the Erm cassette, generating pBS961::Erm. For generation of the single knockout mutants, the knockout plasmids were linearised and transformed into the MC58 wild type (WT) strain. Erythromycin resistant colonies were selected and checked by PCR for correct insertion due to a double homologous recombination event and colonies with correct PCR profile were further analysed by Western Blot for the lack of expression of the respective antigen. MC58 transformed with plasmids pBSUD287::Erm, pBSUD741::Erm, pBSUD953::Erm, pBSUD936::Erm and pBS961::Erm gave rise to strains ΔGNA2132, ΔfHBP, ΔGNA1030, ΔGNA2091 and ΔnadA respectively. The ΔfHBP (previously called Δgna1870) mutant [14], and the ΔnadA [19] have previously been described.

For generation of a single MC58 derivative strain lacking the expression of all five antigens, a stepwise deletion by allelic replacement strategy was used (Fig. 1A). First, an erythromycin sensitive ΔGNA2132erm’ single null mutant was generated by transformation of the ΔGNA2132 strain with the pBSUD287 plasmid containing contiguous upstream and downstream flanking regions of the \textit{nmb2132} gene (without the Erm cassette) cloned into the pBS vector. Transformants were replica plated on GC agar with and without erythromycin selection, and erythromycin sensitive colonies were selected and checked by PCR for the correct double homologous
recombination event leading to loss of the erythromycin cassette. The ΔGNA2132erm\' strain was then transformed with the pBSUD741::Erm plasmid to generate the knockout of the \textit{nmb1870} gene, and erythromycin resistant transformants were selected and checked by PCR and Western blot for the replacement of the \textit{nmb1870} gene with the erythromycin resistant cassette giving rise to the double knockout strain ΔGNA2132erm\'ΔfHBP\textit{Erm}. The stepwise knockout of the other three antigens was completed by substituting the erythromycin cassette in the pBS961::Erm, pBSUD936::Erm and pBSUD953::Erm knockout plasmid constructs with a chloramphenicol (Cm), a kanamycin (Kan) and a tetracycline (Tet) antibiotic resistance cassette, respectively, generating pBS961:Cm, pBSUD936::Kan and pBSUD953::Tet plasmids, respectively. This permitted the stepwise knockout of the three remaining antigen-expressing genes, via three consecutive transformations selecting for the appropriate antibiotic resistant colonies. Transforming the ΔGNA2132erm\'ΔfHBP\textit{Erm} strain with pBS961::Cm and selection of chloramphenicol resistant colonies lead to the generation of the MC58-3KO strain. Then transformation of the MC58-3KO strain with pBSUD936::Kan and selection of kanamycin resistant colonies lead to the isolation of the 4KO-ΔGNA2091Kan strain that was finally transformed with pBSUD953::Tet and selection on tetracycline lead to the isolation of the MC58-5KO strain. Each strain was checked by PCR for the correct double recombination at the allelic replacement locus and for the lack of expression of the appropriate antigens by Western Blot.

Complementation of the single null mutants was achieved by insertion of the deleted gene under the control of the \textit{P_\text{lac}} promoter, along with the chloramphenicol resistance gene, into a noncoding chromosomal location between the two converging open reading frames (ORFs) NMB1428 and NMB1429, as previously described [13]. To complement the ΔGNA2091 strain, GNA2091 and 64 bp of the non coding upstream region was amplified from MC58 using the primer pair 936compFOR3 and 936compREV3 (Table 2) and cloned as a NdeI-NsiI fragment into the pComPRBS plasmid. The addition of a short upstream nucleotide sequence of GNA2091 in the complemented strain enabled better restoration of expression. For complementation by allelic replacement, the null mutant was transformed with the respective complementation plasmid. Transformants were selected on chloramphenicol and checked by PCR, and complementation of the mutant strain was verified by Western blot analysis.

2.4. \textit{Western blot analysis}

Expression of proteins in whole cell samples was performed essentially as described previously [14]. Briefly, strains were grown at 37°C /5% CO\textsubscript{2} on agar plates overnight, or for time course experiments in broth from a starter inoculum of OD\textsubscript{600} 0.05 to stationary phase. Bacterial cells were resuspended in SDS-PAGE loading buffer in an appropriate calculated volume so as to
normalize cell density to a final OD$_{600}$ of 5. Ponceau staining was performed to ensure equal loadings of the samples. Western blot analysis was performed according to standard procedures using polyclonal antibodies at a 1:1,000 dilution (raised in mice against the recombinant purified proteins, as described previously [14]), followed by a 1:5,000 dilution of HPR-labeled anti-mouse IgG (Sigma-Aldrich).

2.5. **In vitro antimicrobial peptide and environmental stress assays**

Overnight cultures were harvested into GC broth and diluted to approximately $10^4$ CFU/ml. Each assay was started by the addition of 10 µl of a compound to 90 µl of the bacterial suspension. Compounds used are as follows: LL-37 (final concentration: 2-7 µM; Innovagen, Sweden), detergent (0.01-0.1% SDS, Tween, Triton X), reactive oxygen species (2-10 mM H$_2$O$_2$, 0.005-0.02% cumene hydroperoxide), or generators of reactive oxygen species (2.5-10 mM paraquat, xanthine (1mM) /xanthine oxidase (200-400 mU)). Samples were incubated at 37°C /5% CO$_2$ with gentle agitation, and CFU were determined as described above. Sensitivity to osmotic stress was analyzed with respect to the WT by growth in GC broth plus NaCl (0.3-0.5 M) or sucrose (5-15%).

2.6. **Ex vivo human whole blood and human serum models of meningococcal bacteremia**

Cells were harvested into MH broth containing 0.25% glucose and 0.02 mM Cytidine-5'-monophospho-N-acetylneuraminic acid sodium salt (CMP-NANA) to an OD$_{600}$ of 0.05 and grown to mid-log phase (OD$_{600}$ 0.5-0.6) then diluted in MH broth to $10^3$-$10^6$ colony forming units (CFU)/ml depending on the bactericidal activity of the donor. The assay was started by the addition of 240 µl whole human blood, human serum or heat inactivated human serum (56 °C 30 min) to 10 µl of bacterial suspension. Cultures were incubated at 37°C /5% CO$_2$ with gentle agitation, and at various time points an aliquot of the sample was removed and the number of viable CFU were determined by plating serial dilutions onto MH agar and incubating overnight at 37°C/ 5% CO$_2$. Experiments were performed in triplicate on several occasions. The Student's t-test was used to determine the statistical significance of survival of each mutant strain with respect to the isogenic WT strain, with a p-value <0.05 considered to be significant. Whole venous blood, collected from healthy individuals (unimmunized against *N. meningitidis*) and anti-coagulated with heparin (10 U/ml), was used for whole blood experiments as described by Ison et al. [28]. For preparation of human serum, whole blood was coagulated at 25 °C for 30 min, centrifuged at 1,000 xg for 10 min at 4 °C and the supernatant retained.
2.7. *In vivo infant rat model*

The infant rat model was used essentially as previously described [29], modified to include a competitive index (CI) analysis. Bacteria were grown to log phase \( \text{OD}_{600} \) of 0.25 in GC medium, washed, and resuspended at the desired concentration in PBS. Five- to 6-day-old pups from litters of outbred Wistar rats (Charles River) were challenged intraperitoneally with the WT MC58 strain and the MC58-5KO strain together in a 1:1 ratio to establish mixed infections in the animals and determine the CI. Groups of 8 infant rats were used for each infectious dose of \( 1 \times 10^5, 1 \times 10^6 \) and \( 1 \times 10^7 \). A control group of 5 infant rats was injected with PBS. Eighteen hours after the bacterial challenge, blood samples were obtained by cheek puncture, and aliquots (100 \( \mu l \) of undiluted sera and 1:10 and 1:1,000 dilutions) were plated onto columbia agar + 5% horse blood and columbia agar + 5% horse blood plus erythromycin (to select for MC58-5KO bacteria) for viable cell counting. The numbers of CFU/ml of blood were determined after overnight incubation of the plates at 37°C in an atmosphere containing 5% CO\(_2\). Enumeration of WT bacteria and mutant bacteria allowed for the determination of the CI ratio using the following formula: \( \text{CI} = (\text{WT output/mutant output})/(\text{WT input/mutant input}) \). Statistical analysis was performed using the Wilcoxon Signed-Ranks Test.

3. Results

3.1. *Generation of meningococcal strains lacking expression of vaccine antigens*

*N. meningitidis* MC58 derivative strains were generated that lack the expression of all five of the antigens of the MenB vaccine (MC58-5KO), the three main antigens (MC58-3KO; lacking GNA2132, fHBP and NadA), as well as strains lacking the single antigens (\( \Delta fHBP \), \( \Delta \text{NadA} \), \( \Delta \text{GNA2132} \), \( \Delta \text{GNA1030} \), \( \Delta \text{GNA2091} \)). A strategy of stepwise deletion by allelic replacement was used to construct the MC58-3KO and MC58-5KO as described in the Materials and Methods. To allow the generation of the 5KO strain with the four antibiotic resistance cassettes available we started from a \( \Delta \text{GNA2132Erm} \) MC58 strain and generated the \( \Delta \text{GNA2132erm} \) mutant lacking the Erm cassette. The \( \text{nmb1870} \) gene was then replaced with an Erm cassette, followed by replacement of \( \text{nadA} \) with a Cm cassette, replacement of the \( \text{nmb2091} \) gene with a Kan cassette, and finally the \( \text{nmb1030} \) gene was replaced with a Tet cassette. Very few colonies were recovered exhibiting tetracycline resistance from this transformation. The deletion of each gene in the MC58-5KO strain was confirmed by PCR of each of the five loci (Fig. 1B) and Western blot analysis shows the absence of each protein in the intermediate and final knockout strains (Fig. 1C). The successful generation of a 5KO strain demonstrates that none of the genes are essential for *in vitro* growth of MC58 and moreover that the abrogation of the simultaneous expression of all of these proteins in the same strain gives rise to a viable mutant.
Total protein samples taken during a time-course culture of the MC58 WT (Fig. 2A) indicates that each of the five proteins present in the MenB vaccine are expressed throughout the course of growth (Fig. 2B). More specifically, NadA is expressed at a lower level during log phase and increases during stationary phase [as previously seen by 18, 30]; GNA2132 is expressed in higher levels during log-phase than in stationary phase; while fHBP, GNA2091 and GNA1030 are expressed at constant levels throughout the different growth phases with a slight accumulation in stationary phase that is more pronounced for fHBP (Fig. 2B). As shown in Figure 1C, the deletion of the genes in the single knockouts or 3KO strain does not influence the level of expression of the other antigens.

3.2. Characterization of growth in different media: GNA2091 is required for optimal growth

In order to investigate the influence of the vaccine antigens in meningococcal growth and survival, we analysed growth of the MC58 WT and knockout strains in a series of different media. All single and multiple knockout strains generated were viable when grown in culture media including GC, Catlin6 and Mueller-Hinton (MH) broth, as well when plated on GC or MH agar. All knockout strains displayed colony morphology similar to the WT strain when grown on GC agar, and growth in GC broth is equivalent in all strains (~46 min doubling time for WT, 3KO and 5KO; Fig. 3A). Growth in chemically defined Catlin6 broth is also essentially equivalent in all strains, with slightly slower growth seen for the 5KO (~69 min doubling time for WT, 75 min for 5KO; Fig. 3B). However, the 5KO strain has smaller colony morphology when grown on MH agar and decreased growth in MH broth, with respect to the WT and the 3KO strains (doubling times: WT, 88 min; 3KO, 89 min; 5KO 97 min; Fig. 3C). In particular, a similar lag phase was seen, but the growth rate of the 5KO slows during the exponential phase and it does not reach a cell density equivalent to that of the WT, as measured by OD₆₀₀ and CFU count (Fig. 3C-D).

Further investigation revealed the growth phenotype of the 5KO strain to be a result of the lack of GNA2091, with the ΔGNA2091 strain displaying decreased growth in MH that was restored to near WT levels in the complemented strain, MC58Δ2091_C, in which the gene encoding GNA2091 is inserted in trans (doubling times: WT, 80 min; ΔGNA2091, 104 min; ΔGNA2091_C, 83 min; Fig. 3E). The growth phenotype of ΔGNA2091 could be partially rescued by supplementing MH with 0.25% glucose (doubling times: WT, 71 min; Δ2091, 74 min; Fig. 3F). Similarly, the growth phenotype of ΔGNA2091 could be partially rescued by supplementing MH with maltose, a disaccharide compose of two glucose monomers (doubling times: WT, 67 min; ΔGNA2091, 67 min; data not shown). Western blot analysis of the MC58 WT strain grown in MH or MH plus glucose (0.25-1%) showed equivalent expression of GNA2091 under conditions of low and high glucose, indicating that this gene is not regulated by glucose levels (data not shown). Investigation
of this phenotype in ΔGNA2091 knockouts of additional meningococcal strains, including 67/00, NZ98/253 and 2996, revealed that GNA2091 is also required for optimal growth in these strains (data not shown).

3.3. **in vitro survival; fHBP and GNA2091 play a role in various stress responses**

In order to determine whether the vaccine proteins are important for survival under various conditions, we performed several *in vitro* stress assays. Survival of MC58, the 3KO and the 5KO was investigated in the presence of: (1) oxidative stress to mimic the oxidative burst of phagocytic cells; (2) the antimicrobial peptide LL-37; and (3) envelope stress using detergent (SDS and Tween) and osmotic stress (NaCl and sucrose).

In a control experiment, the WT, the 3KO, 5KO and single mutant strains all behave in a similar manner when incubated for the duration of the assays in GC broth, the media used to grow and dilute the cells for these assays (2-3 fold increase in CFU over 2 hours; data not shown). This indicates that all differences observed are not due to growth differences during the assay. No mutant strains displayed increased sensitivity with respect to the WT strain during the *in vitro* oxidative stress assays (including paraquat, hydrogen peroxide and xanthine/xanthine oxidase assays; data not shown).

The 5KO and 3KO strains are sensitive to the antimicrobial peptide LL-37, which is a part of the innate immune system that is constitutively produced by leukocytes (Fig. 4A). As previously reported [13], the ΔfHBP strain is sensitive to LL-37 and the phenotype of the 3KO and 5KO strains can be attributed to the lack of fHBP in these strains (Fig. 4A), rather than to the absence of other proteins since all other single KOs had survival equivalent to that of the WT (data not shown).

The 5KO strain (and to a lesser extent 3KO) showed increased sensitivity to the anionic detergent SDS with respect to the WT, suggesting this strain may have increased membrane permeability (Fig. 4B). Analysis of the ΔGNA2091 single knockout strain revealed that this strain has increased sensitivity to SDS, which could be overcome when the respective knockout strain was complemented by inserting the gene *in trans* (Fig. 4C). The ΔGNA2091 knockout also has increased sensitivity to the non-ionic detergent Tween (data not shown). The 5KO strain did not show increased sensitivity to osmotic stress with respect to the WT when grown in GC broth plus NaCl or sucrose (data not shown).

3.4. **Survival of N. meningitidis in ex vivo human whole blood and human serum models of meningococcal bacteremia**

*N. meningitidis* MC58 WT, the 3KO and 5KO strains, as well as all single knockout strains were analysed in *ex vivo* human whole blood and human serum models of meningococcal
bacteremia in order to determine the influence of the antigens on meningococcal survival and to ensure that the mutant strains do not have increased fitness in these models of infection. The human whole blood assay was used to assess cellular and humoral mechanisms of killing (i.e., complement activity, antibody-mediated serum bactericidal activity (SBA) and opsonophagocytosis (OP), as well as neutrophil, macrophage and antimicrobial peptide killing), while the serum assay was used to assess bacterial killing mediated by the humoral immune response. The 3KO, 5KO and the \( \Delta fHBP \) knockout strains are highly sensitive to killing by both human whole blood (Fig. 5A) and serum (Fig. 6A) with 10-100 fold less survival for the mutant strains with respect to WT over two hours. The sensitivity of the \( \Delta fHBP \) knockout strain is consistent with its reported ability to bind factor H (fH), a key negative regulator of the alternative complement pathway [13, 31]. On the other hand, the remaining four single knockout strains (\( \Delta \text{nadA} \), \( \Delta \text{GNA2132} \), \( \Delta \text{GNA2091} \) and \( \Delta \text{GNA1030} \)) all display equivalent or slightly reduced (typically 2-5 fold lower) levels of survival with respect to the WT in both human whole blood (Fig. 5B) and serum (Fig. 6B), with differences seen between different donors. For example, \( \Delta \text{GNA2091} \) had decreased survival in the blood of donors 1 and 4, and the serum of donors 3 and 4. \( \Delta \text{nadA} \) had decreased survival in the blood of donor 4, and the serum of donors 1 and 3. \( \Delta \text{GNA2132} \) and \( \Delta \text{GNA1030} \) both had decreased survival in the serum of donor 2. The WT, all single mutants, as well as the 3KO and 5KO strains all behave in a similar manner when incubated for the duration of the assay in heat-inactivated serum (no growth or killing seen; data not shown).

3.5. In vivo infant rat competition assay

In order to examine the fitness of the MC58-5KO strain in vivo, a competitive index (CI) assay was performed in infant rats to determine if the WT bacteria could out-compete the 5KO bacteria. WT MC58 and 5KO bacteria were injected i.p. at a ratio of 1:1, at final concentrations of \( 1 \times 10^5, 1 \times 10^6 \) or \( 1 \times 10^7 \) CFU. Bacterial CFU in the blood were determined by replicate plating of serial dilutions on columbia agar and columbia agar plus erythromycin (to select for the 5KO strain). Figure 7 shows the CI of each infant rat for each infectious dose used. A CI of greater than 1 was seen for all rats in each group (median/average CI of 10.5/18.8, 4.2/6.8 and 3.6/10 for \( 1 \times 10^5, 1 \times 10^6 \) and \( 1 \times 10^7 \) respectively) indicating that more WT bacteria survived in the infant rat model of infection than did mutant 5KO bacteria. This suggests that the lack of expression of the vaccine antigens in the 5KO strain significantly affects its ability to survive in vivo.

4. Discussion

In this study we evaluated the influence of the investigational Novartis multivalent MenB vaccine antigens in meningococcal growth and survival. The successful generation of a MC58-5KO
strain, in which each of the antigen encoding genes had been genetically removed, demonstrates that none of the genes are essential for growth of the MC58 strain in vitro nor do they give synergistically negative effects on viability as the abrogation of expression of all of these proteins in the same strain gives rise to a viable mutant. The ability of meningococci to colonize the nasopharyngeal mucosa as well as to survive and multiply within human blood are key factors in the development of fulminant meningococcal disease. Hence we endeavored to evaluate the role of the vaccine antigens in growth and survival in various assays that mimic environmental stresses that the meningococcus may encounter during colonization and infection.

*N. meningitidis* MC58-5KO and ΔGNA20291 mutant strains have decreased growth capabilities *in vitro* with respect to the WT strain in MH broth and agar, which can be rescued by the addition of glucose and maltose to the media. The decreased growth of these strains may be relevant in the context of *in vivo* colonization of the nasopharynx, since saliva and nasal and airway secretions of healthy adults contain very low levels of glucose (10-100 fold less than blood) [32]. Furthermore, the MC58-5KO and ΔGNA2091 mutant strains are susceptible to detergent stress. The detergents SDS and Tween were used to mimic membrane stress, which the meningococcus may encounter in the host during colonization or invasive disease. For example, innate host defense of the nasal/nasopharyngeal mucosa include many compounds that disrupt the integrity of bacterial membranes [33], including surfactant protein D, which is known to play a role in inhibiting nasopharyngeal colonization of the pneumococcus [34].

The MC58-5KO and -3KO strains showed a dramatically increased susceptibility to killing by factors of the human immune response, as displayed in the in vitro LL-37 antimicrobial peptide killing assays, and the *ex vivo* human serum and whole blood models of meningococcal bacteremia. This sensitivity can primarily be attributed to the lack of fHBP in these strains, as the ΔfHBP strain has a similar phenotype. *N. meningitidis* has developed many mechanisms to evade the human immune response [35] and fHBP is emerging as a key player in mediating resistance to host defenses, due to its ability to bind fH, an inhibitory regulator of the alternative complement pathway. The importance of the complement pathway in preventing development of meningococcal disease is evidenced by the increased incidence and recurrence of infection and disease in people with immune disorders related to fH [36-39]. fHBP elicits a strong direct bactericidal response through SBA and OP [12, 14], and anti-fHBP antibodies are able to block binding of fH to the meningococcal surface, increasing susceptibility to killing [10, 40]. fHBP is expressed by almost all *N. meningitidis* strains studied to date, however the level of expression varies between strains (high, intermediate, or low expressors) [14, 41]. MC58 is a ST-32 strain that has been characterized as a high expressor of fHBP and is a strain that is dependent on fHBP for survival in human blood and serum [10, 12, 13] and such high levels of fHBP may mask subtle roles played by other proteins. It
is important to note that some of the other single knockout strains also displayed a small increase in sensitivity in the blood and serum assay, suggesting that they may play a role in survival \textit{in vivo} in the human host. This may be particularly relevant in strains that express less FHBP, and that may be more dependent on other antigens for survival. Furthermore, in the \textit{in vivo} infant rat model used to investigate the competitive index of the WT and the 5KO strains, the WT was seen to out-compete the mutant strain in all infected animals when an infectious dose of $1 \times 10^5$ or $1 \times 10^6$ CFU was used. Since fHBP binds specifically to human, but not rat factor H [42], this decreased survival of the 5KO \textit{in vivo} cannot be attributed to the lack of fHBP. It however strongly suggests that the expression of the one or more of the vaccine antigens plays an important role in the growth, fitness and survival of the meningococcus \textit{in vivo} in this model of infection. Further experiments will be needed to elucidate the mechanism by which these antigens contribute to survival in this model of infection.

We have shown that the proteins present in the MenB vaccine are expressed throughout the \textit{in vitro} growth cycle. It has also recently been shown that the level of antigen expression, and sensitivity to SBA, does not change after up to 5 rounds of selection in the presence of bactericidal antibodies generated in mice against the MenB antigens [43]. Since these proteins are important for growth and survival, they are likely to be expressed \textit{in vivo} during colonization and/or infection, and indeed antibodies against NadA, fHBP and GNA2132 are present children convalescing after meningococcal disease, and antibody levels tend to increase with age [24, 44, 45]. Furthermore, the level of expression of these antigens may be higher \textit{in vivo} than \textit{in vitro}. For example, transcription of fHBP is increased during oxygen limitation in an FNR-dependent manner that may be relevant for certain microenvironments in the host, such as the submucosa or intracellular niches [46]. NadA is repressed by NadR [30, 47], but is derepressed in the presence of 4-hydroxyphenylacetic acid, a metabolite of aromatic amino acid catabolism that is secreted in saliva [30]. As such, NadA may be induced during colonization of the nasopharynx enabling it to mediate adhesion and invasion of the mucosal epithelium. Furthermore, it has been proposed that anti-NadA antibodies may be able to decrease adherence to and invasion of host epithelial cells, since a NadA mutant strain has decreased association and invasion of epithelial cells [19]. Likewise, there is a possibility that anti-GNA2132 antibodies may interfere with its role in heparin binding, rendering it more susceptible to killing. It could also be speculated that anti-GNA2091 antibodies may play a role in decreasing growth of the bacteria in low glucose environments in the host. Unfortunately, no appropriate model is available to evaluate meningococcal colonization.

In summary, the abrogation of expression of the MenB vaccine antigens results in reduced growth \textit{in vitro}, and significantly affects sensitivity of the meningococcus to stresses it may encounter in the host, as well as reducing its fitness in \textit{ex vivo} models of infection. These findings
support the multivalent vaccine approach that has been used, and supports the idea that the potential for selection of vaccine escape mutants should be limited by using this combination of antigens.

Acknowledgements

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References


<table>
<thead>
<tr>
<th>Strain</th>
<th>Description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC58</td>
<td>Clinical isolate from the United Kingdom (1985) for which the genome sequence is available. Classification: B:15:P1.7, 16; ETS; ST-74.</td>
<td>[48]</td>
</tr>
<tr>
<td>MC58-3KO</td>
<td>Mutant derivative of MC58 in which the NMB2132 gene has been substituted by Erm; Erm removed from the NMB2132 gene; the NMB1870 gene has been substituted by Erm; and the NMB1994 gene has been substituted by Cm.</td>
<td>This study see Fig. 1.</td>
</tr>
<tr>
<td>MC58-5KO</td>
<td>Mutant derivative of the MC58 3KO strain in which the NMB2091 gene has been substituted by Kan; and the NMB1030 gene has been substituted by Tet.</td>
<td>This study see Fig. 1.</td>
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<tr>
<td>ΔGNA2132</td>
<td>GNA2132 mutant derivative of MC58 in which the NMB2132 gene has been substituted by Erm.</td>
<td>[24]</td>
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<tr>
<td>ΔfHBP</td>
<td>fHBP mutant derivative of MC58 in which the NMB1870 gene is replaced by Erm.</td>
<td>[14]</td>
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<td>ΔnadA</td>
<td>NadA mutant derivative of MC58 in which the NMB1994 gene has been interrupted with Erm.</td>
<td>[19]</td>
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<td>GNA2091 mutant derivative of MC58 in which the NMB2091 gene has been interrupted with Erm.</td>
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<td>ΔGNA2091_C</td>
<td>Complemented GNA2091 strain; derivative of ΔGNA2091 in which the NMB2091 gene under the control of the heterologous P_{lac} promoter are inserted, with Cm into the noncoding region between NMB1428 and NMB1429.</td>
<td>This study</td>
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3KO, 3 knockout strain; 5KO, 5 knockout strain; Erm, erythromycin resistance cassette; Cm, erythromycin resistance cassette; Kan, kanamycin resistance cassette; Tet, tetracycline resistance cassette.
### Table 2. Oligonucleotides used in this study

<table>
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<th>Name</th>
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<sup>a</sup> Capital letters correspond to nucleotides of the MC58 genome sequence and small letters to either nucleotides of *E. coli* genome or plasmid sequence origin or nucleotides added for cloning reasons, underlined letters indicate sequences of restriction enzyme sites used for cloning PCR fragments.

<sup>b</sup> Enzymes for which the restriction sites are present in the sequence of the primer, added for cloning reasons.
Figure 1. (A) Schematic representation of the stepwise generation of the three (3KO) and five (5KO) protein knockout strains of MC58. (B) PCR verification of the deletion of each gene in the MC58 5KO strain. The expected sizes of the PCR products from the parental MC58 strain and each of the five loci are shown in panel A. (C) Western Blot verification of the absence of each protein in the single, intermediate and final knockout strains.
Figure 2. Expression of MenB vaccine antigens in MC58. (A) Time course of growth of the MC58 WT strain. (B) Western blot analysis of total cell extracts of meningococcal cultures, taken at points shown in A along the time course, showing differential growth-phase expression of the MenB vaccine antigens. The approximate molecular weights of the proteins are indicated on the left of each panel.
Figure 3. Growth rates of MC58 WT, 3KO, 5KO and individual mutant strains in various media. Graphs show growth of MC58, 3KO and 5KO strains grown in (A) GC, (B) Catlin6, and (C-D) Mueller Hinton (MH) broth. MC58 WT, ΔGNA2091, and the complemented strain ΔGNA2091_C in (E) MH broth and (F) MH broth supplemented with 0.25% glucose. Growth rates were assessed by optical density (OD) at 600 nm, or colony forming units per ml (CFU/ml) in panel D, over time.
Figure 4. Survival of MC58 WT, 3KO, 5KO and individual mutant strains during *in vitro* stress assays. (A) Survival of MC58, 3KO, 5KO and ΔfHBP strains in the presence of 3.5 μM LL-37. (B-C) Survival of MC58, 3KO, 5KO, ΔGNA2091, and ΔGNA2091_C strains in the presence of 0.01% SDS. Experiments were performed in triplicate on several occasions and representative results are shown. Error bars indicate ±1 standard deviation of the mean. P values using Student’s T-test for the survival of the mutant strain with respect to the isogenic WT parent strain at 60 min are: panel (A) ≤ 0.01 for all strains; (B) 0.01 for 5KO, 0.045 for 3KO; (C) 0.01 for ΔGNA2091, 0.023 for ΔGNA2091_C.
Figure 5. Survival of MC58 WT, 3KO, 5KO, and individual mutant strains in an ex vivo human whole blood model of meningococcal bacteremia. Survival of (A) MC58, 3KO, 5KO and ΔfHBP in blood from two different donors; and (B) MC58, ΔGNA2091, ΔGNA2132, ΔnadA, and ΔGNA1030 in blood from four different donors. Experiments were performed in triplicate on several occasions and representative results are shown. Error bars indicate ±1 standard deviation of the mean. P values using Student’s T-test for the survival of the mutant strain with respect to the isogenic WT parent strain at 120 min are: panel (A) ≤ 0.001 for strains; (B) ≥ 0.05 for all strains except donor 1 - ΔGNA2091 (0.017) and donor 4 -ΔnadA and ΔGNA2091 (0.001).
Figure 6. Survival of MC58 WT, 3KO, 5KO, and individual mutant strains in an *ex vivo* human serum model of meningococcal bacteremia. Survival of (A) MC58, 3KO, 5KO and ΔfHBP in serum from two different donors; and (B) MC58, ΔGNA2091, ΔGNA2132, ΔnadA, and ΔGNA1030 in serum from four different donors. Experiments were performed in triplicate on several occasions and representative results are shown. Error bars indicate ±1 standard deviation of the mean. P values using Student’s T-test for the survival of the mutant strain with respect to the isogenic WT parent strain at 120 min are: panel (A) ≤ 0.01 for strains; (B) ≥ 0.06 for all strains except donor 2 - ΔnadA (0.011), ΔGNA2132 (0.042), ΔGNA1030 (0.026), donor 3 all strains ≤ 0.07, and donor 4 - ΔGNA2091 (0.02).
Figure 7. Eight 6 day-old infant rats were infected intraperitoneally with $1 \times 10^5$, $1 \times 10^6$ or $1 \times 10^7$ CFU *N. meningitidis* MC58 WT and 5KO strains at a 1:1 ratio. At 18 h post-injection blood was collected and serial dilutions were plated and bacterial colonies were counted. Individual CIs are shown for 8 individual animals for the dose of $1 \times 10^5$ and $1 \times 10^6$, and 7 animals for $1 \times 10^7$ (1 animal died during the experiment), and the mean and median are plotted for each infectious dose. Bold circles indicate underestimated values (0 erythromycin-resistant colonies were isolated, and given a number of 1 in order to calculate the CI). P values using the Wilcoxon Signed-Ranks Test for the survival of the mutant strain with respect to the isogenic WT parent strain are 0.01, 0.01 and 0.02 for the dose of $1 \times 10^5$, $1 \times 10^6$ and $1 \times 10^7$, respectively.