Variation and molecular evolution of HmbR, the Neisseria meningitidis haemoglobin receptor

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Meningococcal disease caused by serogroup B Neisseria meningitidis remains an important health problem in many parts of the world, and there are currently no comprehensive vaccines. Poor immunogenicity, combined with immunological identity to human sialic acids, have hindered the development of a serogroup B conjugate vaccine, resulting in the development of alternative vaccine candidates, including many outer-membrane protein (OMP)-based formulations.

However, the design of protein-based meningococcal vaccines is complicated by the high level of genetic and antigenic diversity of the meningococcus. Knowledge of the extent and structuring of this diversity can have implications for the use of particular proteins as potential vaccine candidates. With this in mind, the diversity of the meningococcal OMP HmbR was investigated among N. meningitidis isolates representative of major hyper-invasive lineages. In common with other meningococcal antigens, the genetic diversity of hmbR resulted from a combination of intraspecies horizontal genetic exchange and de novo mutation. Furthermore, genealogical analysis showed an association of hmbR genes with clonal complexes and the occurrence of two hmbR families, A and B. Three variable regions (VR1–VR3), located in loops 2, 3 and 4, were observed with clonal complex structuring of VR types. A minority of codons (3.9 %), located within putative surface-exposed loop regions of a 2D model, were under diversifying selection, indicating regions of the protein likely to be subject to immune attack.

INTRODUCTION

Neisseria meningitidis, a Gram-negative diplococcus, is a major cause of meningitis and septicaemia worldwide (Tzeng & Stephens, 2000). Successful vaccines, based on conjugate-capsular polysaccharides, have been developed against meningococcal serogroups A, C, W-135 and Y (Mitka, 2005); however, poor immunogenicity, combined with the immunological similarity of the meningococcal serogroup B polysaccharide to human glycoproteins, has hampered the development of vaccines against serogroup B meningococci (Finne et al., 1983). Outer-membrane proteins (OMPs) are widely regarded as leading candidates in the search for alternative vaccine components, and various OMPs have been proposed as vaccine components, including PorA, PorB, FetA, TbpB, NspA, NadA and fHbp (Black et al., 1986; Jacobsson et al., 2009; Jodar et al., 2002; Madico et al., 2006; Wedege et al., 1998). As most meningococcal OMPs are diverse, inadequate cross-protective immune responses to heterologous strains have been attained by most vaccines produced so far (Tapper et al., 1999; Urwin et al., 2004). In several instances, however, OMP-containing outer membrane vesicle (OMV) vaccines have been effective against the particular epidemic strain from which they were derived (Bjune et al., 1991; Sierra et al., 1991; Thornton et al., 2006).

The haemoglobin receptor protein (HmbR) is a meningococcal OMP which has not been included in vaccine
formulations to date. It is a TonB-dependent receptor that binds haemoglobin, extracts the haem and transports it into the periplasm (Perkins-Balding et al., 2003; Stojiljkovic et al., 1995). It has a molecular mass of 89 kDa and, like other pore-forming OMPs and TonB-dependent receptors, it is predicted to adopt a transmembrane β-barrel structure consisting of 22 putative transmembrane regions with 11 extracellular loops (Perkins-Balding et al., 2003). Evidence for the importance of the haemoglobin receptor gene (hmbR) in meningococcal virulence has been described (Harrison et al., 2009; Stojiljkovic et al., 1995), with expression of the gene controlled at the level of transcription by the Fur repressor, and at the level of translation by slipped-strand mispairing at a polyguanosine (G) tract located within the coding region (Lewis et al., 1999).

The diversity of the hmbR gene has been compared for a limited number of meningococcal isolates, one for each of the serogroups A, B and C, as well as for the gene in the related organism Neisseria gonorrhoeae (Stojiljkovic et al., 1996). Sequences are highly conserved within the three meningococcal isolates, whilst the gonococcal hmbR contains a premature stop codon, leading to the expression of a non-functional protein. The extent of the diversity of the gene in large meningococcal populations is poorly defined to date. Here, a comprehensive analysis of N. meningitidis hmbR diversity was undertaken using a panel of meningococcal isolates, which included representatives of the principal hyper-invasive lineages (Maiden et al., 1998). Genealogical analysis identified two hmbR families with clonal complex structuring among both. Analysis of selection and recombination on hmbR sequences revealed that a number of residues, located in regions corresponding to putative surface-exposed loops, were subject to strong positive selection and recombination.

METHODS

N. meningitidis strains. The 107 N. meningitidis isolates employed in the evaluation of the multilocus sequence typing (MLST) typing method were investigated (Maiden et al., 1998). The collection, which was assembled to be representative of organisms causing endemic and epidemic disease in the latter part of the 20th century, included several isolates from each of seven recognized hyper-invasive clonal complexes: sequence type (ST)-1, ST-5, ST-4, ST-11, ST-32, ST-8 and ST-41/44. All meningococcal isolates were grown on Mueller–Hinton agar, supplemented with 5 % (v/v) defibrinated sterile horse blood, for 15 h at 37 °C in a 5 % (v/v) CO2 atmosphere. DNA was extracted from all strains using an IsoQuick Nucleic Acid Extraction kit (ORCA Research) in accordance with the manufacturer’s instructions.

Nucleotide sequence determination. The hmbR genes were amplified using Taq DNA polymerase (AmpliTaQ, Perkin–Elmer) with PCR primers PN1, 5’-CCGGAGGATGATGCCGACAGG-3’, and NE3, 5’-TTAAAACTCCATTCACCGG-3’. For optimal PCR amplification of the hmbR gene, reaction mixes were incubated for 30 cycles; each cycle consisted of 94 °C for 1 min, 60 °C for 2 min and 72 °C for 4 min. PCR products were purified using a PEG precipitation method (Embley, 1991). Sequence termination reactions of the purified PCR products were used the BigDye Ready Reaction mix according to the manufacturer’s instructions (Applied Biosystems), with the sequencing primers listed in Supplementary Table S1. Sequence termination reaction products were separated and the sequence data collected using an ABI Prism 377 automated DNA sequencer (Applied Biosystems).

Gene assembly and alignment. Sequence data were assembled using the STADEN sequence analysis package, with both strands of DNA sequenced for each strain (Staden, 1996). Initial sequence alignment and manipulation employed SEQUA from the GCG package (Wobble, 2000). The upstream non-protein-coding region together with the first 23 codons, encoding the signal peptide and the stop codon, were removed to leave only the mature peptide-coding sequence. In addition, genetic diversity arising from the slipped-strand mispairing region encoded by the polyG tract was removed. The sequences were aligned to maintain maximum positional homology of the nucleotide sequence with consideration for codon constraints. A database was established containing hmbR sequences obtained in this study (http://neisseria.org/nm/typing/hmbR).

Analysis of sequence data. MEGA version 4 was used to calculate overall p-distances of hmbR nucleotide sequences as well as the within- and between-family and clonal-complex-associated p-distances (Tamura et al., 2007). Genealogical trees were constructed using CLONALFRAME version 1.1 (Didelot & Falush, 2007). Inference was performed in a Bayesian framework and a neutral coalescent model was assumed, based on the hypothesis that the bacteria in the sample came from a constant-sized population in which each bacterium was equally likely to reproduce irrespective of its previous history. In the present study, five independent runs each consisting of 100 000 iterations and 100 000 burn-ins were performed with every hundredth tree sampled. These were then assessed for convergence and combined, after which a 75 % majority rule consensus tree was derived. Annotation was then undertaken by importing the tree into MEGA version 4.

Analysis of selection pressure. The START2 program was used for tests of selection using the ratio of non-synonymous to synonymous nucleotide substitutions (dN/dS ratio) (Jolley et al., 2001). A phylogenetic tree representative of the hmbR alleles was constructed using the TREEPUZZLE program (Schmidt et al., 2002). The HKY nucleotide substitution model (Hasegawa et al., 1985) and a gamma distribution of rate variation among sites (with eight rate categories and shape parameter x) were used for tree reconstruction. Values for transition : transversion ratio, gamma distribution shape parameter (x) and base frequencies were estimated by the program from the dataset.

Selection pressures acting on meningococcal hmbR were investigated with a maximum-likelihood approach (Yang, 1997; Yang et al., 2000). Current recommended models used in this analysis were: M0 (one ratio), M1a (nearly neutral), M2a (positive selection), M7 (beta) and M8 (beta and ω) (Wong et al., 2004; Yang et al., 2005). Nested models were compared using a likelihood ratio test (LRT). When parameters for each positive selection model were estimated, a naive empirical Bayesian (NEB) approach was used to infer the most likely category for each codon site (Nielsen & Yang, 1998; Yang, 1997). It has been shown that detecting positively selected sites using NEB under certain conditions can give rise to a large number of false positives (Suzuki & Nei, 2004; Zhang, 2004). These anomalies are due to the failure of the NEB approach to account for maximum-likelihood parameter estimate sampling errors. A subsequent Bayes empirical Bayes (BEB) analysis was used to accommodate uncertainties in maximum-likelihood estimation of parameters in the ω distribution using numerical integration (Yang et al., 2005). All the analyses described above were implemented using the CODEML program from the PAML package version 4b (Yang, 1997; Yang et al., 2005).
Characterization of selection in the presence of recombination was carried out using the OMEGAMAP software package (Wilson & McVean, 2006). This method employed a Bayesian approach to parameter estimation that was independent of phylogeny and therefore less likely to falsely identify sites subject to diversifying selection in sequences displaying clear evidence of recombination. The signature of natural selection was detected using the \( d_\alpha/d_\beta \) ratio and the signature of recombination was detected from the patterns of linkage disequilibrium. In the present study, three runs composed of 100 000 iterations and 100 000 burn-ins each were undertaken, compared to assess convergence and combined. Output from the OMEGAMAP runs was used to visualize possible selection acting on the sequence by means of a graph indicating the posterior probability of positive selection along the sequence. The consensus from all runs was also imported into Microsoft Excel and manipulated to generate a graph showing the levels of positive selection acting on each codon (data not shown).

**Table 1.** Sequence diversity of hmbR

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value for hmbR</th>
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<tbody>
<tr>
<td>Number of alleles</td>
<td>52</td>
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<tr>
<td>Number of isolates of which alleles are representative</td>
<td>89</td>
</tr>
<tr>
<td>Length of sequence</td>
<td>2295–2307</td>
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<tr>
<td>Number of variable sites</td>
<td>426</td>
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<tr>
<td>Sequence divergence: Mean ( p )-distance</td>
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<tr>
<td>Range</td>
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<td>( d_\alpha/d_\beta )</td>
<td>0.164</td>
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<td>Number of amino acid sequences</td>
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<tr>
<td>Number of variable sites</td>
<td>121</td>
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<tr>
<td>Mean ( p )-distance:</td>
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<td>Loop L1</td>
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<tr>
<td>Loop L2</td>
<td>0.135</td>
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<tr>
<td>Loop L3</td>
<td>0.095</td>
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<tr>
<td>Loop L4</td>
<td>0.184</td>
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<td>Loop L5</td>
<td>0.109</td>
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<td>Loop L6</td>
<td>0.051</td>
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<tr>
<td>Loop L7</td>
<td>0.056</td>
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<tr>
<td>Loop L8</td>
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<td>Loop L9</td>
<td>0.011</td>
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<td>Loop L10</td>
<td>0.062</td>
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<tr>
<td>Loop L11</td>
<td>0.036</td>
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</table>

**Results**

**Patterns of diversity**

PCR hmbR amplicons were obtained from 89 of the 107 N. meningitidis isolates, resulting in 52 unique sequences ranging from 2295 to 2307 bp, with allele hmbR01 the most frequent (Table 1 and Supplementary Table S2). The remaining 18 hmbR-negative isolates contained exl2, exl3 or exl4 ORFs and have been described elsewhere (Harrison et al., 2009). The level of sequence diversity was assessed by calculating \( p \)-distances, revealing the proportion (\( p \)) of nucleotide sites at which sequences differed. Nucleotide alleles had average \( p \)-distances varying from 0.012 to 0.116, with a total of 426 polymorphic nucleotide sites. Many of the polymorphisms occurred within the putative surface-exposed loops 2, 3, 4, 5 and 10. The 52 hmbR alleles encoded 44 unique deduced amino acid sequences. Among putative HmbR proteins, there were a total of 121 variable residues, including one indel at codon 323 and two simultaneous indel pairs at codons 676–677 and 680–681.

Two families of hmbR genes were evident following CLONALFRAME analysis (Fig. 1, Supplementary Table S2). A total of 56% of sequences belonged to family A and included hmbR genes from isolates in clonal complexes ST-4, ST-5, ST-18, ST-41/44 and ST-269, as well as three ST-32, one ST-8 and two ST-1 isolates, whilst 43% of isolates occurred in family B and included sequences from the clonal complexes ST-11, the remaining ST-32, ST-8 and ST-1 isolates, as well as many other unique STs. One serogroup Z ST-28 isolate contained a hybrid hmbR gene. Mean \( p \)-distances within families A and B were 0.027 and 0.049, respectively, with a mean \( p \)-distance of 0.076 between both families. The indel observed at codon 323 was predominantly found among HmbR peptides associated with family B, although this was not exclusive, while indels 676–677 and 680–681 were not related to either family. The division of the hmbR sequences into two families was apparent in the deduced peptide alignment of loops 2, 3 and 4 beginning at residue 197 through to site 340 (Supplementary Fig. S1). Allelic variants of the deduced peptides from loops 2, 3 and 4 [denoted variable region (VR)1, VR2 and VR3, respectively] were determined, from which a correlation between hmbR family and VR combination was observed (Supplementary Table S2), with family A hmbR sequences predominantly containing VR types 1, 1, 1 or 2, 1, 2, and family B including 3, 2, 3 or 1, 3, 5 VR combinations. A web-accessible database was established to enable future querying of hmbR sequences and submission of new allele sequences (http://neisseria.org/nm/typing/hmbR). Using this database, the VR type of each hmbR sequence can also be determined.

Genealogical examination of the hmbR genes using CLONALFRAME also provided evidence for antigenic structuring, with hmbR sequences from isolates belonging to the same clonal complex tending to cluster together. A total of nine isolates belonging to the ST-41/44 complex contained the hmbR08 allele and were well conserved (mean \( p \)-
Diversity of the N. meningitidis hmbR gene

Evidence of recombination and positive selection

Genealogical analysis indicated very little horizontal genetic exchange between families A and B, although recombination events were apparent among isolates within each family (Fig. 2). Among family A isolates, horizontal genetic exchange was apparent in one short region before and after nucleotide site 1000 (Fig. 2a, node A). Numerous recombination events were observed among family B isolates, particularly in the N-terminal region of the sequence (Fig. 2b, node B).

The hmbR gene had an average $d_s/d_{ts}$ ratio of 0.16, indicating a level of purifying selection against amino acid change. A codon-by-codon examination of $d_{ks}$ and $d_{ks}$ ratios (parameter $\omega$) using models of codon substitution which differed in the extent to which $\omega$ varied along sequences was employed to detect evidence of selection at individual codons. Phylogenetic relationships obtained from the sequences were incorporated into the models, resulting in independent comparisons (Yang et al., 2000). Models used were: M0, which assumed a single $\omega$ parameter for all sites; M1a, which divided codons into conserved sites ($p_0$), with $\omega_0 < 1$ and neutral sites ($p_1$), with $\omega_1$ set to 1; M2a, which accounted for positive selection through a third category of sites, $p_2$, with parameter $\omega_2 \geq 1$; and models M7 and M8, which both used a discrete $\beta$ distribution (with 10 categories and described by parameters $p$ and $q$) to model $\omega$ ratios among the sites, although M8, unlike M7, considered an extra class of sites for which $\omega$ could be $>1$.

Nested models [differing by extra parameter(s)] were compared using an LRT in which twice the difference in log-likelihood between models was compared with the value obtained under a $\chi^2$ distribution. The degrees of freedom were equal to the difference in the number of
parameters between models. When parameters for each positive selection model were estimated, an empirical Bayesian approach was used to infer to which category each codon site belonged. The posterior probability that a site with a particular dataset belonged to category $k$ (with ratio $\omega_k$) was calculated. The category $k$ that maximized the posterior probability was the most likely category for the site. Positively selected sites (i.e. sites belonging to the third category with $\omega > 1$) were then identified and BEB was used to correct for uncertainties in maximum-likelihood estimation.

In each comparison, the LRT statistics were significantly ($P < 0.0001$) in favour of models that allowed positive selection (M2a and M8) (Table 2). Inference of positive selection using the M2a model and BEB suggested that 14 residues (1.8%) were under diversifying selection ($\omega = 3.604$), whereas 86.8% of codons were under strong purifying selection ($\omega = 0.008$). The M8 model using BEB predicted 30 positively selected sites (3.9%), including those predicted using BEB for the M2a model. Similar results were obtained using the OMEGAMAP program (data not shown), indicating that these sites were not false positives resulting from high levels of recombination occurring in the meningococcus.

### Construction of a structural topology model

A hydrophobicity plot was employed to predict $\beta$-strands for HmbR (Supplementary Fig. S2). By analogy with the crystal structures of other TonB-dependent receptors, HmbR would be expected to adopt a 22-stranded $\beta$-barrel structure, with an N-terminal ‘cork’ region within the barrel. The presence of a ‘cork’ domain was apparent in the hydrophobicity plot through the absence of transmembrane segments (hydrophobic peaks) before residue 152. The poor signal, short length and singularity of the predicted $\beta$-strand at positions 561–568 caused it to be allocated to part of a loop region. The signal from residues 325 to 382 was also ambiguous. Visual inspection of sequence alignments against structural homologues was used to predict this region as consisting of two $\beta$-strands, 341–362 and 377–382. The final topology model for HmbR comprised 22 $\beta$-strands with 10 smaller periplasmic loops and 11 larger external loops (Fig. 3). The largest loop regions were L8 and L11, consisting of 46 and 38 amino acids, respectively. Extracellular loops L6 and L8 both contained pairs of cysteines (residues 447/450 and 560/570, respectively), which could potentially form disulfide bonds in each putative loop region. To complement the 2D topology model, HmbR was compared with known TonB-dependent siderophore receptors, the cobalamin receptor BtuB and the *S. marcescens* haem receptor HasR, whose crystallographic structures have been determined (Buchanan et al., 1999; Chimento et al., 2003, 2005; Ferguson et al., 1998, 2002; Krieg et al., 2009; Locher et al., 1998). A structural alignment was generated, revealing that all receptors shared the same structure, with each contain-
Table 2. Likelihood values, selection pressure parameter estimates and positively selected sites of the \textit{hmbR} gene

<table>
<thead>
<tr>
<th>Model code</th>
<th>( p )</th>
<th>( \ell )</th>
<th>( d_N/d_S )</th>
<th>Estimates of parameters</th>
<th>Likelihood test</th>
<th>( \chi^2 )</th>
<th>( P )</th>
<th>Positively selected sites†</th>
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<td>M0 (one-ratio)</td>
<td>1</td>
<td>-15 011.4</td>
<td>0.138</td>
<td>( \omega=0.138 )</td>
<td>M0 vs M1a</td>
<td>2669</td>
<td>&lt;0.0001</td>
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<td>225P*, 268T*, 299M*, 300V*,</td>
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<td></td>
<td>2</td>
<td>-13 676.9</td>
<td>0.137</td>
<td>( p_0=0.870 ) ( p_1=0.130 ) ( \omega_0=0.007 ) ( \omega_1=1 )</td>
<td>M0 vs M1a</td>
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<td>M2a (positive selection)</td>
<td>4</td>
<td>-13 573.5</td>
<td>0.221</td>
<td>( p_0=0.868 ) ( p_1=0.101 ) ( \omega_1=0.008 ) ( \omega_1=3.604 )</td>
<td>M0 vs M2a M1</td>
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<td>M7 (beta)</td>
<td>2</td>
<td>-13 636.1</td>
<td>0.117</td>
<td>( p=0.011 ) ( q=0.069 )</td>
<td>M7 vs M8</td>
<td>190</td>
<td>&lt;0.0001</td>
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<td>M8 (beta and ( \omega ))</td>
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<td>-13 541.1</td>
<td>0.212</td>
<td>( p=0.008 ) ( q=0.048 ) ( \omega=3.592 )</td>
<td>M7 vs M8</td>
<td>190</td>
<td>&lt;0.0001</td>
<td>150P*, 205P*, 209A*, 211K*,</td>
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</tr>
</tbody>
</table>

†Positively selected sites identified using BEB.

*Represents above 99 % confidence level, whilst other sites listed are above 95 % confidence level.

Fig. 3. Location of positively selected residues on a structural topology model of HmbR. Positively selected codon sites (black background with white lettering) inferred using the maximum-likelihood method with BEB and the M8 model are identified on a structural topology model of HmbR. Amino acids that correspond to inserted/deleted codons are indicated by asterisks. The glycines encoded by the polyG tract are indicated by hashes.
ing 22 \( \beta \)-strands, 10 periplasmic loops and 11 surface-exposed loops. The larger extracellular loops 2, 4, 8 and 11 in HmbR were noticeable, and conserved sequence motifs described previously were observed (Supplementary Fig. S3) (Chimento et al., 2005).

Positively selected sites, identified from the BEB and the M8 selection models were mapped onto the HmbR receptor topology model (Fig. 3). The majority (21 out of 30) were situated within predicted extracellular loop regions, with loops 2, 3, 4, 5, 8, 10 and 11 containing positively selected sites. Of the 10 positively selected selected codons located outside the extracellular loops, seven were situated within the barrel \( \beta \)-strands, two on a periplasmic loop and one within the globular N-terminal domain. The glycine residues encoded by the polyG tract were on periplasmic loop T4. A single codon indel occurred on extracellular loop 4 and two codon indel gap pairs on loop 10.

**DISCUSSION**

Iron-regulated OMPs are among the numerous potential vaccine candidates present in the cell envelope. This class of proteins includes receptors that are involved in the uptake of iron from transferrin (Legrain et al., 1993), lactoferrin (Prinz et al., 1999), haemoglobin (Stojilkovic et al., 1995) and haptoglobin (Lewis et al., 1997). As the concentration of free iron in the human body is insufficient to support microbial growth, the ability to exploit multiple iron sources is believed to play an important role in colonization and dissemination of meningococci in the human body. Additionally, these proteins display conserved functional domains (i.e., the ligand-binding sites) at the cell surface to which the immune response may be directed. Therefore, the iron-limitation-inducible OMPs constitute a group of highly promising vaccine candidates. The vaccine potential of both transferrin and lactoferrin receptors has been evaluated, revealing that both receptors are immunogenic and able to induce bactericidal antibodies, but that the cross-reactivity of these antibodies is limited (Pettersson et al., 2006; Rokbi et al., 1997). The suitability of the HmbR receptor as a vaccine component has not been assessed.

The levels of diversity of \( hmbR \) (52 alleles encoding 44 unique amino acid sequences) were similar to a number of other vaccine candidate OMPs investigated with this same panel of isolates: PorA (33 alleles encoding 33 peptides); PorB (31 alleles encoding 28 peptides); FetA (33 alleles encoding 33 peptides); and Opa (90 alleles encoding 83 peptides) (Callaghan et al., 2006; Urwin et al., 2004). Furthermore, \( hmbR \) alleles clustered with clonal complexes in a similar way to other immunogenic OMPs, including the \( porA, porB, fetA \) and \( opa \) genes (Fig. 1, Supplementary Table S2) (Callaghan et al., 2006; Urwin et al., 2004). The presence of two families of \( hmbR \) gene was reminiscent of the variation observed with meningococcal \( porB \) (Derrick et al., 1999). PorB proteins can be divided into two classes, class 2 and class 3, a distinction originally based on their size as measured by SDS-PAGE, but PorB2 and PorB3 are also homology groups on the basis of nucleotide and amino acid sequences, as observed currently with HmbR (Derrick et al., 1999; Frasch et al., 1986). Sequencing of \( porB \) genes has further identified four regions denoted VR1–VR4 located in loops I, V, VI and VII, respectively, enabling \( porB \) typing in a similar to that of \( porA \), which contains two VRs located in surface-exposed loops I (VR1) and IV (VR2) of PorA (Bash et al., 1995; Feavers et al., 1992; McGuinness et al., 1993; Suker et al., 1994; Zapata et al., 1992). Three VRs (VR1–VR3) located in loops 2, 3 and 4, respectively, were also observed in \( hmbR \) (Supplementary Fig. S1). As with \( porA \) and \( porB \), typing of the \( hmbR \) VRs revealed that combinations of VR types were found with both \( hmbR \) families and thus with clonal complexes (Supplementary Table S2) (Sacchetti et al., 1998). It has been documented that immune selection by the host can cause populations of meningococci to self-organize into stable collections of strains with non-overlapping repertoires of dominant polymorphic determinants, despite frequent horizontal genetic exchange (Gupta et al., 1996). The non-overlapping association observed among the putative surface-exposed VR peptides identified in HmbR indicates that the immune response directed against these putative epitopes is strong enough to structure the population into discrete combinations of VR types, as evidenced with PorA (Gupta & Maiden, 2001; Gupta et al., 1996).

A structural topology model for the HmbR protein has been described, although it is based on an arbitrary \( \beta \)-strand length of 10 residues (Perkins-Balding et al., 2003). Hidden Markov models and neural networking methods were used to predict the \( \beta \)-strands of HmbR more accurately (Martelli et al., 2002). As a result, the topology model proposed here, although similar to that previously described, has different-length \( \beta \)-strands (Fig. 3). The deduced HmbR peptide was included in a structural alignment containing the \( S. marcescens \) haemoglobin receptor HasR, as well as the TonB-dependent receptors FhuA, FepA, BtuB, FecA and FptA, whose crystallographic structures have been determined (Buchanan et al., 1999; Chimento et al., 2003; Ferguson et al., 1998, 2002; Krieg et al., 2009; Locher et al., 1998) (Supplementary Fig. S3), revealing that all receptors shared the same structure, with each containing 22 \( \beta \)-strands, 10 periplasmic loops and 11 surface-exposed loops. Conserved structure motifs previously described in the N-terminal ‘cork’ domain (also known as ‘plug’ or ‘hatch’) were found as well as the NLFN (also occurring as NLFT or NVFT) motif in loop 11. The cork domain has an essential role in haem uptake, such that combinations of residues from the three apices A, B and C located in the cork domain contribute to ligand binding in these receptors, although individual receptors use different combinations to bind their respective ligand (Krieg et al., 2009). The exact role played by the apices in the cork domain of HmbR has not been determined;
however, the presence of the conserved motifs, in addition to the description of HmbR mutants devoid of the cork domain unable to use haemoglobin, is indicative of an essential function (Perkins-Balding et al., 2003). The NPNL conserved motif in loop 7 found in all receptors transporting haem into the periplasm was also apparent, although, in agreement with a previous study, this was not as conserved among the siderophore receptors (Bracken et al., 1999). Histidine residues often serve as axial ligands for haem binding, with two histidine residues in HasR, His-603 in loop 7 and His-189 in apex C, conserved among haemophore receptors (Izadi-Pruneyre et al., 2006). HmbR shared one conserved histidine residue (His-500) with HasR (His-603) in loop 7; however, it has previously been shown that mutagenesis of histidine residues in loop 7 does not affect haemoglobin binding in HmbR, indicating that unlike HasR, these residues do not interact with haem (Perkins-Balding et al., 2003).

Functional studies of the HmbR protein have indicated that loops 2 and 3 are necessary for haemoglobin binding, with deletion mutation resulting in 33 and 9% haemoglobin-binding activity, respectively, compared with the wild-type strain (100%) (Perkins-Balding et al., 2003). Clonal complex structured combinations of VR1 and VR2, which correspond to loops 2 and 3, were observed (Supplementary Table S2). Thus, despite the fact that the surface-exposed ligand-binding site was under immune selection, as observed by the identification of positively selected residues (Fig. 3), the clonal complex structuring of loops 2 and 3 indicated lineage conservation of the haemoglobin-binding site. The non-overlapping structured combinations of hmbR VR peptides including loops 2 and 3 combined with the detection of positively selected residues within these regions is consistent with their being putative epitopes towards which an immune response may be directed.

In contrast, extracellular loops 6 and 7 did not contain residues under strong diversifying selection. These loops are putatively implicated in haemoglobin utilization, with mutation resulting in reduced fitness of the organism (Perkins-Balding et al., 2003). The highly conserved amino acid sequence NPNL identified in haem receptors belonging to bacteria including Yersinia enterocolitica and Haemophilus influenzae was also observed in loop 7 of the meningococcal HmbR proteins in this study (Bracken et al., 1999). The lack of recombination events evidenced by CLONALFRAME analyses (Fig. 2, nodes A and B) is also indicative of the conserved nature of loops 6 and 7. Positively selected sites were also located in surface-exposed loops 4, 5, 8 and 10, which are predicted to be large extracellular regions and have not been implicated in haemoglobin binding or utilization (Perkins-Balding et al., 2003). Correspondingly, these loops were found to contain numerous polymorphic sites and displayed some of the highest p-distances in this study as well as many instances of horizontal genetic exchange (Table 2) (Fig. 2, node B). The putative redundant role of these loops may therefore allow greater diversity. Indeed, VR typing of these regions was not as structured (data not shown).

Although most residues within the transmembrane b-strands of HmbR were highly conserved, 10 positively selected sites were located among these structural regions (Fig. 3), although combined with the data obtained using OMEGAMAP, individual dN/dS ratios for these sites were low, indicating a lower probability of positive selection. Many of these were found to be connected mutations such that a proline residue at site 150 was invariably associated with a glutamic acid residue at site 151, whereas an aspartic acid residue at site 150 was always linked to another aspartic acid residue at site 151. Similarly, a methionine residue at site 532 was consistently linked with serine and glutamic acid residues at sites 528 and 529, respectively, whereas a threonine residue at site 532 was accompanied by glycine and aspartic acid residues at sites 528 and 529. According to the topology model, four positively selected sites (positions 291, 296, 299 and 300 in Fig. 3) were located close to one another on opposite sides of a b-turn. A methionine residue at position 291 was invariably accompanied by tryptophan, serine and leucine residues at positions 296, 299 and 300, whereas a tryptophan residue at position 291 was accompanied by arginine, methionine and valine residues at positions 296, 299 and 300. Within the limitations of the homology model, the side chains of the two residues may be in contact, suggesting that these are compensatory mutations. Elsewhere, the reasons for positive selection within the b-barrel region were more difficult to discern.

Molecular epidemiology has played an important role in the development, implementation and study of meningococcal vaccines. For candidate protein components it is essential to identify the diversity of the protein in suitable bacterial populations, ideally before a vaccine formulation is tested in humans. Thus, although the suitability of the HmbR protein as a vaccine component through the use of functional assays has not been investigated, comprehensive analysis of N. meningitidis hmbR diversity provides the first step in revealing the potential of the gene. Data presented here support the conclusion that HmbR is an OMP whose diversity bears clear signatures of a dynamic interplay with the host immune system. A web-accessible database has been created (http://neisseria.org/nm/typing/hmbR), enabling current and future hmbR sequences to be queried or deposited whilst also allowing the VR type of each hmbR sequence to be determined. It is hoped that as this database grows, it will be constructive in establishing the molecular epidemiology of the hmbR gene in Neisseria populations.

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