Variation within serovars of *Neisseria gonorrhoeae* detected by structural analysis of outer-membrane protein PIB and by pulsed-field gel electrophoresis

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**INTRODUCTION**

The major protein present on the surface of *Neisseria gonorrhoeae* is protein I (PI), an anion-selective porin (Douglas *et al.*, 1981). Unlike the other major surface antigens, pilus, Opa (PII) and LPS, PI does not undergo antigenic shift during infection (Zak *et al.*, 1984). Antibodies directed against PI activate complement-mediated killing, promote phagocytosis by polymorphonuclear leukocytes and inhibit interaction with epithelial cells (Virji *et al.*, 1986, 1987). Although PI is stable within a strain, structural variations occur which generate antigenic differences between strains (Butt *et al.*, 1990b). Thus, in addition to its essential physiological role, PI is of particular significance both as the antigen responsible for serological classification of strains and as a potential vaccine candidate.

Biochemical studies have differentiated PI into two...
major classes, PIA and PIB, which differ in molecular mass and susceptibility to proteolysis (Sandstrom et al., 1982). Each class can be further subdivided into a number of serovars based on patterns of reaction with panels of either PIA- or PIB-specific mAbs (Knapp et al., 1984). This classification into serovars provides an important base for the differentiation between strains in epidemiological studies (Knapp et al., 1987; Ison et al., 1992).

Information on the structural basis of gonococcal serological specificity has been obtained following the sequencing of the por genes, which encode PIA and PIB. Despite the biochemical and antigenic differences between PIA and PIB, considerable similarity exists between the inferred amino acid sequences of the two classes, with about 70% of amino acid residues in common (Carbonetti & Sparling, 1987; Gotschlich et al., 1987). Even greater homology is seen within each class. Comparison of the amino acid sequences of PIB from three different gonococcal strains revealed over 95% homology between the strains, with most differences being clustered in two regions around residues 196 and 237. Sequencing these regions from four further strains of a different serovar confirmed the variability of these two regions, which were designated variable regions 1 and 2 (Var1 and Var2) (Butt et al., 1990b). The importance of these regions for antigenic specificity was confirmed using overlapping synthetic peptides, corresponding to the entire PIB sequence, to map the location of epitopes recognized by PIB-specific mAbs. In each case the epitopes were localized to the Var1 region (Butt et al., 1990b).

Recent studies have revealed that further variations may occur within the Neisseria porins which are not revealed by current serological methods. Comparison of the PIA sequences from two unrelated strains of identical serovar revealed a ‘silent’ amino acid difference between the strains (Mee et al., 1993). Further sequence differences have been demonstrated within sero-subtypes of the related meningococcal class 1 porin, and these permitted much more detailed differentiation between epidemiologically related strains (Brooks et al., 1995). The current method of differentiating gonococcal isolates relies on agglutination reactions with mAbs to define the serovar. This approach has proved very useful for monitoring temporal changes in gonococcal populations, but gives limited information for forensic purposes or for studying sexual networks. In many geographical locations a limited number of serovars account for the majority of the gonococci. Hence, in such a population, isolates known to be from a single source or sexual contact will belong to identical serovars (Ison et al., 1992), but isolates from a different source may also be of that same serovar. We have therefore investigated the use of molecular methods based on PIB sequence analysis and restriction endonuclease digestion of gonococcal DNA to differentiate between gonococcal strains which show similar reactivity in serovar determination, including some strains known to be epidemiologically linked.

**METHODS**

**Source of bacterial strains and growth conditions.** Isolates of Neisseria gonorrhoeae from known sexual contacts attending the Department of Genitourinary Medicine at Basingstoke District Hospital were obtained from the cervix and urethra of females (denoted C and U respectively) and from the urethra of their male partners in a previous study (Zak et al., 1984). Additional isolates, not known to be from linked cases, were from patients attending the same clinic (designated SU strains).

Strains of defined serovar were from the culture collection at St Mary’s Hospital Medical School (IB-2 strains 3039, 3020, 3615 and 3582; IB-3 strains 3682 and 3668). All bacteria were grown on proteose peptone agar at 37 °C in 5% (v/v) CO₂ (Mee et al., 1993).

**Serotyping and monoclonal antibody reactivity.** Serotyping was performed by co-agglutination reaction with the panel of 12 mAbs (Genetic Systems) directed against epitopes on PI, using the nomenclature of Knapp et al. (1984) as described by Gill (1991).

Dot-blot analysis was carried out as described previously (Mee et al., 1993) using the same panel of antibodies and a second panel of mAbs (SM mAbs) directed against gonococcal PIB which have been previously characterized (Virji et al., 1986; Butt et al., 1990b).

**Sequence analysis of PIB.** The sequences of the por genes encoding PIB were determined following selective amplification of the gene by PCR. A few gonococcal colonies were resuspended in water (200 μl) and were lysed by the addition of 0.25 M KOH (400 μl) and heating to 100 °C for 5 min. The lysate was neutralized by the addition of 0.5 M Tris/HCl pH 7.5 (400 μl) and diluted 1:100 for use in PCR. The oligonucleotides used corresponded to the previously determined 5′-CCAAAAAAGGAATACAGC-3′ and 3′-CGACATTAGATTTCGG-3′ of the por gene of gonococcal strain P9 (Butt et al., 1990a). The reaction mixture (100 μl) contained the diluted bacterial lysate in the presence of 50 mM KCl, 10 mM Tris/HCl pH 9.0, 0.1% Triton X-100, 1.5 mM MgCl₂, 0.2 μl of each dNTP (BCL), 400 ng of each primer and 5 U Taq DNA polymerase (Promega). Amplification was performed in a GeneAmp 9600 (Perkin-Elmer Cetus) using 30 cycles of denaturation at 94 °C for 15 s, primer annealing at 48 °C for 20 s and DNA polymerization at 72 °C for 30 s. The final cycle was followed by an additional polymerization step at 72 °C for 3 min. Excess PCR primers were removed and the amplified DNA concentration twofold using the Magic PCR preps DNA purification system (Promega).

The purified DNA was used in sequencing reactions with a Taq DyeDeoxy Terminator cycle sequencing kit (ABI) according to the manufacturer’s instructions, with 200 ng DNA and 15 ng sequencing primer in a final reaction volume of 24.5 μl was used. The cycle sequencing reactions were carried out using 25 cycles of denaturation at 96 °C for 15 s, primer annealing at 52 °C for 10 s and DNA polymerization at 60 °C for 4 min. After removal of unincorporated nucleotides, the resulting products were analysed using a model 373 automated DNA sequenator (ABI). Sequencing of both strands of the por gene was accomplished with additional oligonucleotide primers based on the sequence of the por gene from strain P9 (Butt et al., 1990a).

**Pulsed-field gel electrophoresis (PFGE).** Gonococcal chromosomal DNA was prepared and digested with the restriction endonuclease SpeI, as described by Poh & Lau (1993). Briefly, gonococcal cultures were harvested, washed, resuspended in
Table 1. Comparison of gonococcal isolates determined to belong to serovar IB-2 on initial isolation

Isolates were reacted with mAbs under standard co-agglutination conditions for determination of serovar (Gill, 1991) and also reacted with the same mAbs in dot-blots (Mee et al., 1993). Auxotype was determined as described by Ison et al. (1992). NR, No requirement.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Auxotype</th>
<th>Test</th>
<th>Serotyping PIB mAbs:</th>
<th>Apparent serovar</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td>3C8</td>
<td>1F5</td>
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<tr>
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<td></td>
<td>Co-Agg</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>SU81C</td>
<td>(Cohort 1)</td>
<td>AHU</td>
<td>D-Blot</td>
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<tr>
<td>SU81U</td>
<td></td>
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<tr>
<td>SU82C</td>
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</tr>
<tr>
<td>SU105</td>
<td>AHU</td>
<td>Co-Agg</td>
<td>+</td>
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<td></td>
<td></td>
<td>D-Blot</td>
<td>+</td>
<td>-</td>
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<tr>
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<td>(Cohort 2)</td>
<td>PA</td>
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<td>+</td>
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<td>+</td>
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<tr>
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<td></td>
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<tr>
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<td></td>
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<td>NR</td>
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<tr>
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<td>NR</td>
<td>Co-Agg</td>
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<td>D-Blot</td>
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1 M NaCl in 10 mM Tris/HCl and mixed with an equal volume of 1% low-melting-point agarose at 42°C. After solidifying, the plugs were transferred to lysis solution containing 6 mM HCl, 100 mM EDTA, 0.5% Brij 58, 0.2% sodium deoxycholate, 0.5% sodium lauryl sarcosinate, 10 mg ml⁻¹ lysozyme and 1 U ml⁻¹ RNase, and incubated at 37°C overnight. After further incubation in 0.5 M EDTA (pH 9.9-9.5) containing 1% sodium lauryl sarcosinate and 500 µg proteinase K ml⁻¹, the prepared plugs were digested with 10 U endonuclease in 150 µl restriction buffer (New England Biolabs).

The digested plugs were sealed into slots of a 1% agarose gel (SeaKem GTG agarose, FMC) and subjected to electrophoresis in a contour-clamped homogeneous-electric-field apparatus with a hexagonal electrode array (Bio-Rad) for 20 h at 200 V (Poh & Lau, 1993). Gels were stained with ethidium bromide and photographed under UV transillumination. After photography, the negatives were scanned using a flat-bed scanner (Epson) and the resulting images were analysed using GelManager software (BioSystematica).

Multiple solid-phase peptide synthesis and epitope mapping. Multiple solid-phase peptide synthesis was carried out with the aid of an Epiguide (Labsystems) using pin technology with a commercially available kit (CRB) as described in detail by Butt et al. (1990a) and McGuinness et al. (1993). Immunological reactivity of the peptides with mAbs was assayed by ELISA and the peptides were re-used after bound antibody was dissociated by sonication of the pins for 30 min in 1% SDS, 0.1% 2-mercaptoethanol in 0.1 M phosphate buffer at 60°C. Immunological reactivity was always observed in duplicate peptides and in assays repeated on at least two occasions.

Sequence analysis. Sequence comparisons were carried out using DNAstar (Lasergene) and with the GCG sequence analysis programs using the BBSRC SEQNET facility.

RESULTS

Definition of strains

All strains originally isolated by Zak et al. (1984) which were serotyped as IB-2 at the time of isolation were chosen for detailed study. These included two groups of isolates known to be epidemiologically linked: cohort 1 (four isolates) and cohort 2 (three isolates). Subsequent serotyping showed that the isolates from cohort 1 and one additional strain were serovar IB-3 (Table 1).

In addition to serotyping, the isolates were also subjected to dot-blot analysis, using the same mAbs as used to determine the serovar by co-agglutination. Two distinct patterns of reactivity were observed (Table 1). The serovar IB-2 strains (including cohort 2) gave identical results with both methods, while the IB-3 strains failed to react in dot-blots with mAb 2D6. Since the latter pattern, in co-agglutination, denotes serovar IB-6, the isolates which reacted in this manner in dot-blots were designated IB-3 (d.b.-6). Despite the unexpected differences in immunological reactivity seen between the strains, the isolates from within each cohort always reacted identically, cohort 2 reacting as IB-2 and cohort 1 as IB-3 (d.b.-6).

The isolates were also reacted by dot-blotting with the SM panel of mAbs, which have previously been used in immunochemical analysis of PIB (Virji et al., 1986; Butt...
et al., 1990b). Again, the two groups of strains could be distinguished: the first group reacted with all five mAbs tested, while the second group reacted with mAbs SM24 and SM198, but failed to react with mAbs SM21, SM23 and SM203.

**Comparison of PIB sequences**

In order to investigate the structural basis for serological specificity and the potential variation of PIB within a serovar, the *por* gene was sequenced from all the isolates described above, together with standard strains which reacted unequivocally in both co-agglutination and dot-blot as either serovar IB-3 or IB-6. The corresponding translated protein sequences are shown in Fig. 1. Significant sequence differences were seen between the strains. A dendrogram of the relationships between the sequences revealed that they fell into four distinct groups which precisely corresponded to the four patterns of immunological reactivity demonstrated by the strains, namely IB-2, IB-3 (d.b.-6), IB-3 and IB-6 (Fig. 2).

The four isolates from cohort 1, which belonged to serovar IB-3 (d.b.-6), came from one male and two female contacts and all gave an identical amino acid sequence. This sequence was also identical to that of the one other isolate (SU105) which was also IB-3 (d.b.-6). This isolate, which was not known to be clinically related to cohort 1, was obtained 1 year later at the same location. The three isolates from cohort 2, which belonged to serovar IB-2, had identical amino acid sequences and PFGE patterns, but differed in sequence and PFGE patterns from other IB-2 isolates and from other serovars tested.

In contrast to the sequence identity shown between the epidemiologically linked isolates, comparisons between unlinked isolates of the same serovar showed wide sequence diversity. With the exception of SU105, as discussed above, the only isolates to show sequence identity within a serovar were those from a cohort of partners. The degree of sequence diversity within serovars varied considerably. Among the IB-2 serovar, isolates differed from cohort 2 isolates at between one

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**Fig. 1.** Comparison of predicted amino acid sequences of PIB from gonococcal isolates. Dots indicate positions with identity to the sequence of strain SU50. Boxes indicate the regions predicted to form surface-exposed loops according to the model of Van der Ley et al. (1991). Horizontal lines separate strains according to serovar, and brackets enclose sequences from cohorts 1 and 2. The sequence of PIB from strain R10 is from Gotschlich et al. (1987).
Variation of gonococcal PIB within serovars

SU104

SU92

3039

3020

3682

3668

SU50

SU51C

SU5IU

SU104

SU103

SU62

3682

SU80

SU105

SU81C

SU81U

SU82

SU106

R10

(SU104) and thirteen (3582) positions. The sequence of the two IB-6 isolates differed by two amino acids, while the IB-3 isolates showed sequence differences in a total of nine positions.

Epitope mapping studies

The availability of sequence information facilitates detailed mapping of the epitopes recognized by mAbs. Previous studies with the SM-series mAbs have shown that most recognize linear epitopes in the Var1 region of strain P9 (IB-26) between amino acid residues 190 and 200. Despite their identical reactivity with these mAbs, the IB-2 isolates in the current study showed sequence differences in this region. The previously determined epitope specificities of mAbs SM22, SM21 and SM203 were found to be the closely related sequences 190EYEHQVY197, 192YE(H/Y)Q(V/A)Y197 and 192YEXQX196 (Butt et al., 1990b). Inspection of the different sequences found in the IB-2 group of isolates showed that none contain potentially reactive sequences, while the IB-3 (d.b.-6) isolates all contain the sequence 190EYEHQVY197, which includes an epitope for each of the three mAbs. Thus, the determined sequence, immunological reactivity and epitope specificity are in accord. Similarly, all isolates from both the IB-2 and IB-3 (d.b.-6) groups reacted with mAbs SM24 and SM198 and all contain the sequences 198SIPS201 and 107WESGK111, which are the respective epitopes for these mAbs.

The epitope specificity of the mAbs used for serovar determination had not been previously reported. To investigate the effect of sequence variation on reaction with these mAbs, a series of overlapping decapeptides were synthesized on polyethylene pins, in which adjacent peptides differed by a single amino acid residue. The sequences used corresponded to the Var1 and Var2 regions of gonococcal strains SU80, SU50, R10 and MS11 that were synthesized with adjacent peptides differing by a single amino acid residue. Peptides were reacted in ELISA with the serotyping mAbs. The results shown were obtained with sequences corresponding to SU80 and MS11; strains SU50 and R10 gave essentially identical patterns to those obtained with SU80.

PFGE analysis

PFGE was carried out to investigate possible relationships between strains. Analysis of the PFGE patterns also showed that unrelated isolates within a serovar all gave different profiles. The greatest similarity was seen between the cohort 1 isolates and strain SU105, which had an identical PIB sequence and differed by a single
DISCUSSION

The ability to differentiate reliably between unrelated gonococcal strains is essential for studies of the epidemiology of gonorrhoea. Serotyping with panels of mAbs which differentiate strains on the basis of antigenic differences in PIA or PIB has facilitated such studies, but methods which permit further discrimination are required. The aim of this study was to investigate the potential extent of variations of PIB within a single serovar, which might permit further differentiation of strains. Although all the strains selected were identified as IB-2 on initial isolation, several were subsequently found to react in co-agglutination as IB-3. Previous studies have shown that determination of gonococcal serovar is potentially subject to problems of reproducibility which are influenced by preparation of the reagents used and interpretation of the results (Ison et al., 1992). As found in the present study, consistency in serovar determination over a period of years may therefore be a particular problem. The development of potentially more reliable molecular methods based on knowledge of the structure of the por gene, which encodes PI, are therefore a logical development of the current serotyping schemes.

Comparison of PIA and PIB with the analogous porins from meningococci has led to a model for the organi-
zation of the Neisseria porins within the outer membrane. The model predicts a series of amphipathic transmembrane \( \beta \)-sheets which generate eight surface-exposed loops. The membrane-bound regions are highly conserved between the different porin classes, while the predicted loops show extensive variation in both length and amino acid composition. Within each class so far examined, sequence diversity is far more restricted. The original studies with PIB compared the sequences of the protein from three strains of different serovar and revealed that sequence variation between the three strains was largely located in two regions designated Var1 and Var2 (Butt et al., 1990b), which were subsequently shown to be located at the apices of predicted loops 5 and 6 (van der Ley et al., 1991). In addition, epitope-mapping studies demonstrated that the SM series of mAbs reacted with the Var1 sequences in loop 5, suggesting that this region was the predominant influence on antigenic specificity. The current studies reveal that the situation is more complex than this: sequence variations are not confined to the Var1 and Var2 regions in loops 5 and 6, but occur elsewhere in the protein, particularly in loops 1, 3 and 7.

Epitope-mapping studies with the mAbs used to prepare reagents for serotyping confirmed that the previously designated Var1 and Var2 regions do contribute to serovar specificity: antibody 3C8 reacted with the highly conserved sequence in loop 5 previously shown to be responsible for broad specificity of mAb SM24, while mAb 2D4 reacted with a sequence from the loop 6 region of PIB of strain MS11, but which was not present in any of the strains from the present study. These locations are in accord with studies of Carbonetti et al. (1988), who used artificial hybrid PIA/PIB molecules to demonstrate that the epitopes for 3C8 and 2D4 were located in a central region of PIB between residues 150 and 270, with 3C8 closest to the N-terminus. In the current study, the remaining four mAbs failed to react with any sequences corresponding to the Var1 and Var2 regions of the strains tested. Although this lack of reactivity may result from the epitopes recognized being conformational in nature, other sequences outside these regions must also contribute to serovar specificity. This would be similar to the situation with PIA, where some serovar-specific antibodies react with linear epitopes in loop 6, while others recognize conformational epitopes apparently located in other loops (Mee et al., 1993). In the studies with PIA, it was possible, by comparing sequence differences with mAb reactivity, to identify regions that appeared to contribute to such determinants. In the present study it has not been possible to identify epitopes by sequence comparison, but the occurrence of sequence variations in other predicted loop regions suggests surface location and, hence, exposure to immunological pressure. Certainly, the fact that both loop 5 and loop 6 sequences of IB-2 cohort 2 strains are identical to that of the previously published sequence of strain P9 (IB-26) indicates that other regions of the protein must also contribute to serovar specificity.

The widespread occurrence of sequence variations of PIB within a serovar has potentially important implications. Differentiation between gonococcal strains is important not only for studies of the epidemiology of gonococcal infection, but also in providing information on patterns of sexual mixing used in mathematical models of HIV-transmission dynamics (Anderson et al., 1990). Conventional differentiation of gonococcal strains, based on a combination classification of auxotype and serovar (A/S), suffers from a lack of discriminatory power (O'Rourke et al., 1995). Other methods such as PFGE (Poh & Lau, 1993) and Opa-typing (O'Rourke et al., 1995) are capable of providing much greater resolving power, but since they rely on pattern recognition, they are relatively subjective, particularly when comparing results between laboratories. The occurrence of the 'silent' differences within a serovar shows that, potentially, PIB holds considerably more epidemiological information than can be obtained using current serological reagents. The use of DNA-based methods provides the potential for greater reproducibility and, although methods based on direct \( \text{por} \) sequencing are relatively complex, such studies should provide information for the design of improved reagents for gonococcal differentiation based on gene-probe techniques, as has been proposed for the related \( \text{porA} \) gene, which encodes the class 1 porin responsible for meningococcal subtype specificity (Maiden et al., 1992).

The practical use of porin sequence information in epidemiological studies would clearly be dependent on stability in the \( \text{por} \) gene during transmission over a useful period of time. While it is likely that PI is subject to immunological pressure to undergo variation, the timescale of such variation is unknown. However, evidence for \( \text{por} \) gene stability is provided in the current study, by comparison of cohort 1 isolates with strain SU105. These strains, which were isolated from the same geographical location, but separated by a period of 1 year, were the only unlinked isolates to have an identical PI sequence. In addition, they also share the unusual AHU/IB-3 combination of auxotype and serovar, and differ in PFGE by a single band, indicating a close genetic relationship. This would be consistent with the stability of PIB during transmission within a community over this period. It would also be in accord with studies on the meningococcal \( \text{porA} \) gene which suggest that the gene is stable during epidemic transmission over a period of decades (Suker et al., 1994).

In addition to its importance for serovar specificity, PIB is an important candidate antigen for vaccination against gonorrhoea (Heckels et al., 1990; Elkins et al., 1992). The further diversity of PIB revealed in the current studies strengthens the suggestion that such strategies should focus on antigenic determinants, such as that recognized by mAb SM24, which are known to be widely shared between strains (Heckels et al., 1990).

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