Temporal changes in the gonococcal serovar patterns in Stockholm during two years with special reference to PPNG strains

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Abstract
Objective—To analyse temporal changes in gonococcal serovar patterns in Stockholm during a two year study period (1987–1989) to elucidate the dynamics of gonorrhoea epidemiology.

Design—The study population comprised 857 patients with culture proven gonorrhoea and with serotyped gonococcal isolates. The probable geographical origin of the infection was determined in 690 of the patients.

Results—A total of 108 Ph/GS-serovars were identified. Most (73%) of the serovars were recognised only during one or two quarters of the study period and comprised 16% of the isolates. Seven serovars were encountered during all eight quarters. Three of these serovars i.e. Arost/Aedgklih (IA-1, IA-2), Bropt/Bajk (IB-3, IB-6), Brpyust/Bacejk (IB-1, IB-2) were the most prevalent in the overall study, accounting for 60% of the isolates during the first quarter of the study and 36% of the isolates during the last quarter. Fifty-seven percent of the patients were infected in Stockholm (endemic infection). The proportion of endemic isolates among the three most common serovars declined throughout the study period (77% during the first quarter; 47% during the last quarter). A total of 32 Ph/GS-serovars were recognised among 80 PPNG strains. Only four of these 32 serovars were encountered during more than two quarters. Of 57 patients with PPNG strains and with geographical origin of the infection known, only seven (12%), all infected in Sweden, might have transmitted their infection further into the society.

Conclusions—The decline in the total number of gonorrhoea cases seen in Stockholm during the study period, was due mainly to a decline of endemic isolates of the three most prevalent serovars. Results from contact tracing and serotyping indicated that PPNG infections acquired abroad seldom become established in the community. Serovar determination seems valuable mainly as a tool for surveillance whereby the introduction and circulation of gonococcal strains in the community can be pursued.

Introduction
The development of a reliable and well functioning classification system for Neisseria gonorrhoeae by using monoclonal antibodies against protein I, the major outer membrane protein, has provided us with a tool for interpreting gonorrhoea epidemiology at the community level.1-4

In 1987, we started a two year project with the aim of “eradicating” endemic gonorrhoea in Stockholm. At that time the incidence of gonorrhoea in Sweden had been continuously decreasing since 1970, except for a minor peak in 1976. The Stockholm area accounted for about one third of all Swedish gonorrhoea cases, 5.4% of which were caused by beta-lactamase producing Neisseria gonorrhoeae (PPNG) strains.

One object of the project was to characterise endemic versus non-endemic gonorrhoea in order to identify groups at risk for transmission of the infection. This would also include an evaluation of the effects of intensified contact tracing performed by specially trained social workers.7 Another object was to correlate antibiotic susceptibility, serovars, and auxotypes of the gonococcal isolates to geographical origin of the infection.8

The aim of the present study was to describe and analyse temporal changes in serovar patterns in the community to show the dynamics in the spread of Neisseria gonorrhoeae strains with special reference to PPNG strains.

Methods
The aim of the whole project was to eradicate endemic gonorrhoea from Stockholm. Endemic infection is therefore defined as infection acquired in Stockholm and non-endemic infection as infection acquired abroad (imported) or from other parts of Sweden.

Study design The study group comprised all patients in Stockholm with culture proven gonorrhoea during the two year study period (October 1987—September 1989), diagnosed at any of the six bacteriological laboratories and with serologically classified gonococcal isolates. Strains from four patients were not serotyped and these patients were excluded. A patient returning within six weeks with a gonococcal isolate of identical serovar to that isolated on the first occasion and with no negative culture in between was only counted once. Our aim was to see all patients at one or other of the four venereal outpatient clinics in
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Stockholm. Doctors working outside the venereal clinics were asked to refer patients (with the patient's consent) to one of the venereal clinics for treatment, test-of-cure culture and contact-tracing interview performed by a social worker.

The two year study period was divided into quarters with October—December 1987 defined as quarter number 1, January—March 1988 as quarter number 2 etc. For a more detailed description of the principles of design of the study see reference number 7.

Study population Out of 857 patients with gonorrhoea, included in the study, 671 (78%) attended a venereal clinic for a follow-up visit with a contact-tracing interview. The remaining 186 patients did not attend any venereal clinic.

Contact tracing All 671 patients were subjected to a semi-structured interview, performed by social workers. The questions included the number and identity of sexual partners during the previous six months, the probable source of infection, the likely geographical origin of the infection and the existence of a steady partner with whom the patient had an ongoing sexual relationship.

Microbiological methods Specimens were obtained from the urethra in men and from the urethra and cervix in women. Rectal and pharyngeal specimens were obtained when appropriate. The strains were cultured and identified as described earlier.8 All strains were classified serologically with co-agglutination into serogroups (WI and WII/III) and serovars as previously described.4 Two different sets of monoclonal antibody reagents were used: the Ph-panel9,10 (Pharmacia Diagnostics AB, Uppsala, Sweden) and the GS-panel1 (Syva, Palo Alto, USA). For the choice of monoclonal antibodies, nomenclature used for the designation of serovars and correlations between the Ph- and the GS-serovars see Bygden et al.7 The corresponding serovar nomenclature according to Knapp et al2 is given in brackets. A serovar combination of a Ph-serovar and a GS-serovar is in this paper referred to as a Ph/GS-serovar or just a serovar.

Before the start of the study laboratory technicians from all six bacteriological laboratories in Stockholm were educated and trained in the serological classification technique although three of the six laboratories already had used the technique routinely for several years. If the laboratory technician had any problem with interpretation of the co-agglutination results, one laboratory (Huddinge Hospital) had a supervising role. Every new or unusual serovar or serovar combination of the two panels was double-checked at this laboratory.

All isolates were typable. If a patient had strains of the same serovar, isolated at the same time from several sites, only one strain was included in the study.

Statistical analysis Chi square test with Yates' correction was used.

Results General Among the 857 patients (332 women and 525 men) comprising the study population a total of 859 gonococcal strains were isolated. Two patients had a double infection with two strains of different Ph/GS-serovars isolated from the urethra. A probable geographical origin of the infection was determined for 690 (81%) of the patients. Of these 690 patients, 57% were considered to have an endemic infection and 43% to have a non-endemic infection.

Out of the 859 isolates, 231 (27%) belonged to serogroup WI and 626 (73%) to serogroup WII/III. Two isolates from two women reacted with both WI and WII/III reagents (WI-WII/III) and belonged to the same serovar.

Serovar determination A total of 108 Ph/GS-serovars was encountered. Table 1 shows the distribution of these serovars by quarter through the study period. The majority (86%) of the Ph/GS-serovars belonged to serogroup WII/III (table 2). Forty-three percent (6/14) of the WI serovars and 61% (57/93) of those of serogroup WII/III were confined to only one gender (p = 0·3). Forty-one percent of all serovars were recognised in isolates from both

Table 1  Distribution of 859 isolates into 108 Ph/GS-serovars during the 8-quarter study period

<table>
<thead>
<tr>
<th>Serovars</th>
<th>Quarter number</th>
<th>Total number</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Arost/Adgdkih (IA-1, IA-2)</td>
<td>37</td>
<td>14</td>
</tr>
<tr>
<td>Bropt/Baft (IB-3, IB-6)</td>
<td>16</td>
<td>30</td>
</tr>
<tr>
<td>Bppysa/Bacejk (IB-1, IB-2)</td>
<td>29</td>
<td>17</td>
</tr>
<tr>
<td>Arst/Aedh (IA-6)</td>
<td>6</td>
<td>2</td>
</tr>
<tr>
<td>Bppysa/Beygik (IB-1, IB-11)</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Bropt/Baft (IB-1, IB-2)</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Bropt/Bacejk (IB-1, IB-2)</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Bopt/Baft (IB-3, IB-6)</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Bropt/Bacejk (IB-5, IB-7)</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Boppt/Baft (IB-3, IB-6)</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Boppt/Bacejk (IB-5, IB-7)</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Bppysa/Bacejk (IB-1, IB-2)</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Additional 94 Ph/GS-serovars'</td>
<td>26</td>
<td>25</td>
</tr>
<tr>
<td>Total</td>
<td>136</td>
<td>107</td>
</tr>
</tbody>
</table>

* Serovars with less than 10 isolates each.
Table 2  Distribution of 108 Ph/GS-serovars among 859 gonococcal isolates in relation to serogroup and sex

<table>
<thead>
<tr>
<th>Serogroup</th>
<th>No of serovars in</th>
<th>W1</th>
<th>W1/III</th>
<th>W1-W1/III</th>
</tr>
</thead>
<tbody>
<tr>
<td>female isolates only</td>
<td>1</td>
<td>11</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>male isolates only</td>
<td>5</td>
<td>46</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>isolates from both sexes</td>
<td>8</td>
<td>36</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>14</td>
<td>93</td>
<td>1</td>
<td></td>
</tr>
</tbody>
</table>

Most (73%) of the serovars were encountered only during one or two quarters of the study. They accounted for 16% of the isolates. Thus, a minority, 27% of the serovars, represented 84% of the isolates and were seen during 3 quarters or more (table 3).

Seven serovars (6%) with 530 isolates, that is 62% of all isolates were recognised during all 8 quarters (table 3). Three of these serovars (Arost/Aedgkih (IA-1, IA-2), Bropt/Bajk (IB-3, IB-6) and Bpyust/Bacejk (IB-1, IB-2)) accounted for 49% (423/859) of all isolates in the study. During the first quarter of the study period, 60% (82/136) and during the last quarter, 36% (27/75) of all isolates belonged to these serovars (p < 0.005). The decline in total number of isolates from the first quarter to the last (table 4) was due mainly to a decline in number of isolates of these three serovars (fig 1). The proportion of endemic isolates among these serovars was 65% (229/355) of those with known geographical origin. During the first quarter of the study period, 77% (54/70) of the isolates were of endemic origin with a decrease to 47% (9/19) during the last quarter (fig 2). Thus, the decrease in number of isolates of these three serovars was due mainly to a lower number of endemic isolates.

The remaining four serovars (Arst/Aedih (IA-6), Bpyust/Beghjk (IB-4, IB-11), Bpyust/Bejgjk (IB-5, IB-7)) of those identified during all eight quarters, accounted for 13% (108/859) of all isolates; 10% (13/136) of the strains isolated during the first quarter and 12% (9/75) during the last quarter. Each of these serovars was represented mainly by non-endemic isolates. This inflow of non-endemic strains showed a relatively stable pattern during the study period. Among the remaining 101 serovars none became dominant over the study period.

An increasing diversity was identified during the study period between number of isolates and number of isolates, caused by a minor increase in the number of serovars recognised per quarter during the study and a major decrease in the number of isolates (table 4).

Of all patients in the study group, 80 (9%) (14 women and 66 men) had an infection with a PPNG strain. Among the 80 PPNG strains 32 Ph/GS-serovars were seen; 4 W1, 27 W1/III and 1 W1-W1/III. Eighty-four percent of the PPNG strains belonged to serogroup W1/III. There was no dominating serovar but Arst/Aedih (IA-6) was the most commonly encountered with 13 (16%) isolates from 13 patients. PPNG strains of this serovar were identified during five quarters of the study period. In the overall study, Arst/Aedih (IA-6) accounted for 33 strains. Thus, 39% of all were PPNG strains. PPNG strains of the second most common serovar Bpyust/Beghjk (IB-4, IB-11) with 11 isolates (14%) was represented during six quarters. Most of the serovars, 28 (87.5%) out of 32, were found only during one or two quarters.

The ratio between the number of serovars

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The ratio between the number of serovars
and number of PPNG isolates per quarter were relatively stable during the study period.

Use of serovar determination in clinical work with patients, harbouring PPNG strains
Contact tracing was performed in 57 patients with PPNG infection examined at one of the venereal clinics. Forty-three (75%) of these 57 patients reported that they had been infected abroad. The majority (33 or 77%) of those who were infected abroad denied sexual intercourse after returning to Stockholm. Another six patients had had sexual contacts after returning home but examination of their partners resulted in negative gonococcal cultures. The remaining four men had infected their steady partners in Stockholm but with no further transmission.

Fourteen patients stated that they had been infected in Sweden. Five of these patients had a second generation imported infection; four had acquired their infection in Stockholm and were infected by the four men mentioned above; one had a second generation imported infection, acquired in another part of Sweden. None of these five patients, all women, admitted to further contacts.

One man, infected in Stockholm, had two female partners with negative cultures. He had a strain belonging to the most common serovar among PPNG strains, Arst/Aedghk (IA-6). Only one woman had a PPNG strain of this serovar. She had a second generation imported infection and could not be connected to this man.

Another six patients had only unknown partners. Two of these patients were infected in other parts of Sweden and four in Stockholm. Out of the four patients infected in Stockholm, three had unique serovars, not seen in any other PPNG isolates. The fourth patient, a man, had a strain with the serovar Arost/Aedgkh (IA-1, IA-2) and reported having had sex with two unknown female partners. During the study period only four PPNG strains of this serovar were recognized, all isolated from men.

One woman infected in Stockholm admitted to sexual contacts with three partners. She reported that she had informed two of them of the suspicion of gonorrhoea but refused to identify them. The remaining partner was examined and was found to be infected. This man denied additional sexual partners. The PPNG strains involved belonged to the second most common serovar among these strains, Byvuyt/Beghjk (IB-4, IB-11). Of the additional nine patients, all men, with PPNG strains of this serovar, four had acquired their infection abroad and denied sexual contact after returning to Sweden. The remaining five patients were not subjected to contact tracing at a venereal clinic.

Discussion
The aim of the present study was to describe and analyse changes over time in serovar patterns in the community. Two different sets of monoclonal antibodies (Ph- and GS-panels) were used to improve the resolution. Earlier studies have shown that the resolution increases when combining these two sets of monoclonal reagents.5,11

In two previous studies in Stockholm in 1979–8012 and in 1982–83,13 42% and 33% of the strains, respectively, belonged to serogroup WI compared with 27% in the present study. Thus, a progress towards a greater dominance of WII/III strains was noted. This is unfortunate since WII/III strains more often show decreased susceptibility to different antibiotics compared with WI strains.5,14 Bygdeman et al14 have shown that the high sensitivity to antibiotics among WI strains is correlated with one WI serovar, Aedghkh, which in several studies have accounted for the majority of WI strains.3 In the present study, strains of this serovar markedly decreased in number during the study period.

A greater heterogeneity of serovars was seen in serogroup WII/III compared with serogroup WI (table 2). This is in accordance with earlier studies.3,11

Almost half (47%) of the 108 serovars observed were encountered in isolates from men only (table 2). The corresponding figure for women was 12%. In the study in Stockholm 1982–83 mentioned above,13 an almost identical picture was noted with 47% of the serovars found in isolates from men only and 11% in isolates from women. We have shown earlier that men more often acquired their infection outside Stockholm compared with women1 and furthermore that imported strains frequently belong to serovars that are unusual in Stockholm.13

Sixteen percent of the strains showed a transient pattern that was encountered only during one or two quarters of the study period (table 3). These isolates represented the major part (73%) of the 108 serovars.

Only seven serovars were persistent

Figure 2  Distribution by quarter of isolates belonging to the three dominating Ph/GS- serovars in relation to endemic infection, non-endemic infection and infection of unknown origin.
throughout all eight quarters (table 3). Three of these serovars accounted for almost half of the isolates. This is in accordance with previous studies from other geographical areas using the Ph/GS-panels. A majority (65%) of the isolates of these three serovars was acquired in Stockholm and constituted the basis for endemic infections. From the first to the last quarter of the study, the proportion of isolates belonging to these three most prevalent serovars, markedly declined from 60% to 36% without a corresponding decline in the proportion of isolates of other serovars (fig 1). In a five year study in Edinburgh, Scotland, a similar decline was seen in the total number of gonorrhoea cases. Fifty-eight percent of the infections were acquired in Edinburgh, 33% in the rest of the United Kingdom and 5% abroad. The same three GS-serovars dominated as in our study. Isolates of two of these serovars, Aedgkih (IA-1, IA-2), Bacejk (IB-3, IB-6) decreased in frequency in parallel with the overall fall in the prevalence of gonorrhoea. However, one serovar, Bacejk (IB-1, IB-2), was persistent at a constant level. No information was given about the proportion of endemic infections in these serovars. The decrease in the number of gonococcal isolates identified throughout the study period in the present study was mainly due to a decrease among endemic isolates of the three dominating serovars (fig 2).

In a previous study, enrolling 671 of the 859 patients in the present study, we showed by using contact tracing that strains acquired outside Stockholm mostly did not establish themselves in the community. Furthermore, a decrease from 75% to 40% in the proportion of endemic infection was noted during the two year study period. Usually, patients with non-endemic infection did not transmit their infection after their return to Stockholm. Alternatively the infection ended as a second generation imported infection with no further transmission. Thus, we have a considerable inflow of strains that do not become established as endemic strains. The three most prevalent serovars with a dominance of endemic isolates were persistent throughout the study period. The major part of the serovars, however, showed a transient pattern. The reappearance of a serovar could depend on reintroduction either from an external source or from an earlier undetected case, probably with asymptomatic infection. Furthermore, serovar identity of two strains does not ensure that they have the same clonal origin, especially if they belong to the most prevalent serovars or are of different geographical origin.

The proportion of PPNG strains in the study was 9%, with a significant difference between the genders. Out of the 108 Ph/GS serovars seen in the whole study of 859 isolates, 32 (30%) were identified among the 80 PPNG isolates. Only four of these 32 serovars were recognised during more than two quarters suggesting that the major part of the strains were imported. This conclusion was supported by contact tracing data revealing that 75% of the 57 patients with probable geographical origin of the infection determined, had acquired their infection abroad.

The most common serovar combination among PPNG strains in the present study was Arst/Aedih (IA-6) representing 16% of all PPNG strains. In a study in 1982–1983 in Sweden comprising 253 PPNG strains, Arst/Aedih (IA–6) was one of the most prevalent serovars accounting for 20% of the isolates. In a study from Amsterdam during two time periods, 1981–1982 and 1985, Arst/Aedih (IA–6) was the only common serovar among PPNG strains on both occasions.

In the present study we studied the dynamics and transmission of PPNG strains in the society by analysing results of serovar determination and contact tracing in conjunction. The 43 patients with PPNG strains, acquired abroad and who were contact traced at one of the venereal clinics, either did not transmit their infection into the society at all or did only transmit the infection to their steady partner where the transmission ended. Seven of the 14 patients infected with PPNG strains in Sweden, had unknown partners only. Why did we not find possible partners with isolates of the same serovars to these patients infected in Stockholm and with unknown partners? Alternative explanations are possible. The partner was examined outside Stockholm, or outside the study period. The partner had an asymptomatic infection or had typical symptoms and was treated for gonorrhoea without a preceding culture taken. He or she had practised self-medication or had spontaneously healed. Even though every male patient was asked about sexual orientation both by a social worker and a physician, independently, there might have been an incorrect answer. Furthermore, the laboratory might have produced a false negative result. Although the sensitivity of gonococcal cultures is high, false negative results exist. Another possibility would be that the PPNG strain had lost its plasmid encoding for beta-lactamase production. A false interpretation of the co-agglutination results might also have been made in the laboratory. However, every new or unusual serovar was double-checked.

With serovar analysis and contact tracing data we have shown that the number of strains belonging to endemic serovars has continuously declined and endemic infections are becoming uncommon in Stockholm.

4 Knapp JS, Holmes KK, Bonin P, Hook III, EW. Epidemiology of gonorrhoea: distribution and temporal
changes in auxotype/serovar classes of Neisseria gonorrhoeae. Sex Transm Dis 1987;14:26-32.