Virulence of transparent and opaque colony types of *Neisseria gonorrhoeae* for the genital tract of mice

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Summary. The virulence of transparent (Tr) and opaque (Op) colony types of *Neisseria gonorrhoeae* in the genital tract of female mice was evaluated at two stages of oestrous. Isogenic pairs of Tr and Op variants were isolated from *N. gonorrhoeae* strain 57–120. Both variants exhibited a T2 morphology, but only the Op variant possessed protein II (P. II) in outer-membrane fractions. When administered by intravaginal inoculation Op gonococci were highly infective only for mice in late pro-oestrous, whereas Tr gonococci were virulent for mice at both late pro-oestrous and dioestrous. Gonococci recovered from the uterus were of both Tr and Op phenotypes in equal proportions when mice were infected at dioestrous with Tr cells. In contrast, >90% of recovered colonies were of Op phenotype when mice were infected at late pro-oestrous with either Op or Tr cells. These results indicate that the virulence of gonococci for the genital tract of female mice differs from that for the chicken embryo. Furthermore, gonococcal survival in the female genital tract might be attributable to phase variation from Tr to Op phenotypes.

Introduction

Recent investigations of the virulence of *Neisseria gonorrhoeae* have focused on several properties of the macromolecular components on the cell surface. Amongst surface structures implicated in the pathogenesis of gonococcal infection, pili and protein II (P. II) have received particular attention because of their ability to undergo dramatic alterations in their structural characteristics.1–6

Pili are thought to promote adhesion of gonococci to epithelial cells7 as a primary step in infection, thus anchoring the bacteria to the mucosa.8,9 P. II is also considered to contribute10–14 to direct contact between the epithelium and bacterial surface15 as well as to inter-gonococcal adhesion.16,17 Moreover, several studies have revealed that alterations in pili and P. II molecules substantially affect the adhesive properties of the bacteria for different cell types.18–20

Clinical isolates of gonococci can be classified not only into pilate and non-pilate phenotypes, but also into opacity variants on the basis of colonial morphology.21 Opaque-colony (Op) gonococci have one or more P. II proteins,22,23 whereas colourless, transparent (Tr) colonies generally contain bacteria whose outer membranes are devoid of these proteins. Gonococci possessing certain P. II proteins show not only increased colonial opacity but also decreased sensitivity to steroid sex hormones; in particular, oestrogen enhances the growth of Op gonococci in vitro and progesterone inhibits their growth.24

Colonies of gonococci isolated from cervical cultures range from fully opaque to transparent, with some intermediate degrees of opacity, and the colonial phase variation is associated with the menstrual cycle.25 Gonococci expressing P. II are associated only with infections localised to the urogenital mucosa. This strongly suggests that the expression of P. II in vivo may be associated with the pathogenesis of gonococcal infection in the female genital tract.

In the present study we compared the virulence of P. II-expressing (Op) and P. II-non-expressing (Tr) variants of the same strain (isogenic pairs) in our mouse model of gonococcal genital infection.

Materials and methods

Organisms

Isogenic Op and Tr variants were isolated from *N. gonorrhoeae* strain 57–12027 following the criteria of Swanson.21 Kellogg's colonial type28 was determined by the method of Juni and Heym,29 and type 2 colony morphology for both Op and Tr variants was selected. Both variants were passaged sequentially daily by single-colony transfer on clear phenotyping agar,10 until a high degree of colony stability of each variant was ensured. P. II proteins were identified by sodium dodecyl sulphate-polyacrylamide gel electrophoresis.
of whole-cell lysates and by apparent change in molecular weight when heated. The degree of pilation was confirmed by transmission electronmicroscopy. Both variants were stored at -80°C in gelatin disks, so that the same batch of each variant could be used throughout the study.

For each experiment, one of the disks was dissolved in 1 ml of Mueller-Hinton Broth (Difco) and cultured on GC Agar Base (Difco) enriched with IsoVitalex (Baltimore Biological Laboratories) 2% v/v at 37°C for 18 h in air + CO₂ 5%. Organisms grown on GC agar were transferred to the liquid medium and incubated overnight at 37°C with continuous rotation as described previously. After incubation for 24 h, bacteria were harvested and washed several times with phenol red-free Eagles’ Minimal Essential Medium (MEM; Flow Laboratories, McLean, VA, USA) supplemented with 20 mM HEPES (HEPES-MEM); washed bacteria were suspended in HEPES-MEM. Subculture on GC agar of each variant grown in the liquid medium showed that each phenotype was preserved (>98% viability) during incubation for 24 h in liquid culture. The concentration of the gonococcal inoculum was quantified spectrophotometrically, and confirmed by plate counts.

Animals

Female virgin mice of the ddY strain, 10–12 weeks old, were used throughout the investigation. They were fed a commercial laboratory animal diet and water ad libitum. The stages of their oestrous cycle were determined by the vaginal smear technique as described previously.

Inoculation procedure

Intravaginal inoculation of gonococci was performed as described previously. The vaginal openings were disinfected with alcohol before inoculation. Gonococci were suspended in pre-warmed HEPES-MEM containing gelatin 1% w/v at a concentration of 10⁶ cfu/ml, and 20 μl of the suspension were inoculated into the vagina with a sterile tip on a micropipette (Pipetten).

Recovery of gonococci from the genital tract

At regular intervals, mice were killed by cervical dislocation. The vagina was washed twice with 50 μl of pre-warmed HEPES-MEM by means of a micropipette; washings were pooled and 20 μl of the pooled fluid was cultured on GC agar with vancomycin 3 mg/L, colistin 7.5 mg/L, and nystatin 12 500 U/L (TM agar) at 37°C in air + CO₂ 5% for 48 h. The washed tissues were then homogenised in 1-0 ml of HEPES-MEM, supplemented with saponin 0-5% w/v, with a teflon homogeniser. Dilutions of the homogenate (20 μl) were cultured on TM agar at 37°C in air + CO₂ 5% for 48 h to quantitate the numbers of viable gonococci.

The contralateral uterine horn and body were fixed with formaldehyde 3-5% v/v, embedded in paraffin, and sectioned for haematoxylin and eosin (HE) staining. For scanning transmission electronmicroscopy (SEM), tissues were fixed with glutaraldehyde 1% v/v in 0.15 M sodium cacodylate buffer (pH 7.2) immediately after washing with HEPES-MEM. The critical point method was employed for drying the specimens which were then examined with a JEOL scanning electronmicroscope (JSM-T20) at 20 kV after coating the specimens with gold-palladium (Au 60%, Pd 40%) while rotating them in a vacuum evaporator.

Direct immunofluorescence

Antiserum to N. gonorrhoeae strain 57–120 was raised in rabbits by injecting formalin-killed organisms emulsified in Freund’s complete adjuvant subcutaneously three times at 12-day intervals. The rabbits were bled 7 days after the final injection. Fluorescein-labelled rabbit anti-N. gonorrhoeae strain 57–120 immunoglobulin was prepared by the method of Johnson et al. The labelled immunoglobulin was pre-absorbed with normal mouse uterine homogenates. Slides of uterine sections were stained with this pre-absorbed fluorescein-labelled antibody and then observed with an Olympus epifluorescence ultraviolet microscope. Control slides of sections of uninfected mouse uterus showed no specific immunofluorescence.

Statistical analysis

The levels of significance for the observed frequencies were determined by Student’s t test.

Results

Comparison of infectivity of Op and Tr variants

The infectivity of both variants for the mouse uterus was compared in mice at late pro-oestrous and at dioestrous stages. The results are summarised in table 1. Both phenotypes were highly infective for mice at late pro-oestrous; Op variants showed greater infectivity than Tr variants. In contrast, at dioestrous, Op variants showed low infectivity, whereas Tr variants exhibited significantly greater infectivity.

Enumeration of gonococci in the uterus

The number of gonococci recovered from uterine homogenates was determined (fig. 1). In mice that had
Table I. Relationship between oestrous cycle of mice and infectivity of colonial variants of N. gonorrhoeae 57–120

<table>
<thead>
<tr>
<th>Experiment no.</th>
<th>Op variant at late pro-oestrous</th>
<th>Op variant at dioestrous</th>
<th>Tr variant at late pro-oestrous</th>
<th>Tr variant at dioestrous</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>10</td>
<td>1</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>II</td>
<td>9</td>
<td>0</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>III</td>
<td>10</td>
<td>0</td>
<td>8</td>
<td>5</td>
</tr>
</tbody>
</table>

been infected at late pro-oestrous with either variant, the number of gonococci recovered from homogenates increased slowly, but steadily, from 12 h to 7 days after inoculation, although the number of bacteria in the mice injected with the Op variant remained about 1-0 log_{10} higher than in the mice injected with the Tr variant at each time point. However, in mice that had been infected at dioestrous with either Op or Tr variants, gonococci were not recovered 12 h after inoculation. After 96 h, gonococci were recovered only from mice that had been infected with the Tr variant and the number of bacteria remained, at most, 100 cfu per uterus. Gonococci were not recovered from mice that had been infected at dioestrous with the Op variant.

The number of gonococci recovered from the uterine washings was greatest at 24 h after inoculation in mice infected at late pro-oestrous with either variant, and in those infected at dioestrous with the Tr variant (table II). Thereafter, the number of gonococci in washings decreased with time. Furthermore, no significant colonial phase variation occurred over a 7-day period. In mice infected at dioestrous with the Op variant, cultures were negative over a 7-day period.

Pathological changes in vaginal discharge

Smears of the vaginal discharge from mice infected at dioestrous with either variant demonstrated the presence of gonococci attached to or engulfed by polymorphonuclear leucocytes (PMNL) until 48 h after infection (fig. 2a). Thereafter, the number of gonococci decreased rapidly with time, and PMNL infiltration continued over a 7-day period (fig. 2b). These changes were common to both colonial types. However, when mice at late pro-oestrous were infected with either variant, PMNL infiltration was not discernable throughout the 7-day experiment, but a large number of gonococci were seen attached to epithelial cells (fig. 2c). Furthermore, gonococci were recovered from vaginal discharges over a 7-day period from mice infected at late pro-oestrous with either variant, and from mice infected at dioestrous with the Tr variant, whereas cultures were positive only for 2 days in mice infected at dioestrous with the Op variant (table III). The predominant colonial type of gonococci recovered from the vagina was the same as that of gonococci recovered from the uterine washings.

Table II. Recovery of N. gonorrhoeae from uterine washings after infection*

<table>
<thead>
<tr>
<th>Colony type of inoculum</th>
<th>Stage of oestrous cycle at inoculation</th>
<th>Mean (SD) number of gonococci (log_{10} cfu) in washings on day (after inoculation)</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Op</td>
<td>Late pro-oestrous</td>
<td>3.8 SD 0.5</td>
</tr>
<tr>
<td></td>
<td>Dioestrous</td>
<td>—</td>
</tr>
<tr>
<td>Tr</td>
<td>Late pro-oestrous</td>
<td>2.4 SD 0.4</td>
</tr>
<tr>
<td></td>
<td>Dioestrous</td>
<td>2.0 SD 0.2</td>
</tr>
</tbody>
</table>

—, None recovered.

*Data from three experiments with 10 mice in each.

†Numbers in parenthesis indicate percentage of the predominant phenotype.
However, in mice infected at dioestrous with the Op variant, Op was the predominant type (about 64% of total colonies) only on day 1 and >66% of colonies had become Tr variants on day 2.

**Histological study**

No remarkable histological changes were observed by light microscopy in sections of uterus from mice infected with either variant until 24 h after infection, regardless of the stage of the oestrous cycles at which the mice were infected. The appearance was no different from that of sections of uterus from mice matched for oestrous-cycle stage.

In mice infected at dioestrous with the Tr variant, PMNL infiltration of the mucosal epithelium and subepithelium in the uterus was significantly increased 48 h after infection (fig. 3a), compared with the normal changes in PMNL numbers associated with changes in the oestrous cycle (fig. 3b). After 72 h, only slight oedema was observed in the subepithelial stroma with minimal infiltration, and the epithelial cell layer was not affected (fig. 3c). No histopathological changes were detected in the uterus of mice infected with the Op variant when animals were infected at dioestrous.

In contrast, when mice were infected at late pro-oestrous with either variant, there was minimal acute inflammatory reaction within 48 h and only slight oedema was seen in the subepithelial stroma (fig. 3d). Gonococci were detected by direct fluorescent staining in the epithelium and submucosal stroma (fig. 3e). The most striking change in mice infected at this stage was detected 72 h after inoculation; it was characterised by the presence of multiple gonococcal clusters in the subepithelial stroma (fig. 3f). This change was more evident in the mice infected with the Op variant than in those infected with the Tr variant.

Uterine tissues from all infected mice were also examined by SEM. Within 4 h of inoculation, gonococci were seen to be attached to the surface of the epithelium of the uterine body (fig. 4a), regardless of colonial phenotype of the inoculum, when mice were infected at late pro-oestrous. Normal bacterial flora were visible on the surface of the uterine body, but the

![Fig. 2. Cytosmears of vaginal discharge: (a) 48 h and (b) 7 days after mice at dioestrous were infected with Tr variant; (c) 48 h after mice at late pro-oestrous were infected with Op variant (bar, 10 \( \mu \)m).](image)

<table>
<thead>
<tr>
<th>Table III. Recovery of <em>N. gonorrhoeae</em> from vaginal washings*</th>
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</thead>
<tbody>
<tr>
<td>Colony type of inoculum</td>
</tr>
<tr>
<td>-------------------------</td>
</tr>
<tr>
<td>Op</td>
</tr>
<tr>
<td>Dioestrous</td>
</tr>
<tr>
<td>Tr</td>
</tr>
<tr>
<td>Dioestrous</td>
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</table>

*See footnotes to table II.*
surface was not affected by the attachment of gonococci. Gonococci were seen on the surface of the uterine horn from as early as 12 h after inoculation. The surface to which gonococci were adherent was devoid of microvilli, and appeared rough compared to uninfected sites (fig. 4b). Phagocytic cells were not detected on the surface of the uterine horn of mice infected at late pro-oestrus, and gonococci seemed to be sinking into the mucosal surface 24 h after inoculation (fig. 4c). This change was common to infections with both variants.

However, when mice were infected at dioestrus, gonococci were not detected on the surface of the uterine body until 24 h after inoculation, regardless of the colonial phenotype of the inoculum. Thereafter, gonococci were seen on the surface of the uterine horn with leucocytes migrating towards the bacteria (fig. 4d), but only in mice infected with the Tr variant.

Colonial phenotypes of recovered gonococci

The number of gonococci in the uterus 7 days after inoculation was about 2 log\(_{10}\) higher in mice infected with the Op variant than in those infected with the Tr variant when animals were infected at late pro-oestrus (table IV). Furthermore, >90% of gonococci recovered from mice infected at late pro-oestrus were of Op phenotype and <5% of colonies represented the Tr phenotype, regardless of the colonial phenotype of the gonococci inoculated. In contrast, when mice
were infected at dioestrous with the Tr variant, both phenotypes were recovered in almost equal amounts.

**Discussion**

It has been reported\(^{25}\) that strains of gonococci isolated from infected women at the time of menses give rise primarily to Tr colonies, whereas isolates obtained in mid-cycle yield predominantly Op colonies. Moreover, a high frequency of complications such as pelvic inflammatory disease is observed at or near menses.\(^{33}\) These clinical observations indicate that the pathogenicity of gonococci in the female genital tract is closely related to the menstrual cycle. Although the colonial phase variation of isolates from cervical cultures has been studied at various stages of the menstrual cycle, little is known about the relationship between the infectivity of gonococcal variants and the menstrual cycle. Therefore, the present study was undertaken to clarify this relationship in studies with our mouse model of gonococcal genital infection.

The results obtained have demonstrated that Op variants are infective in the genital tract of female mice only at late pro-oestrous, whereas the Tr variant was pathogenic at dioestrous as well as late pro-oestrous. Furthermore, Op variant colonies were predominant amongst isolates recovered from the uterus of mice infected at late pro-oestrous with either variant, whereas both phenotypes were recovered with equal frequency from animals infected at dioestrous with the Tr variant. These findings indicate that although Tr variants are infective at late pro-oestrous and dioestrous, intrauterine survival of the gonococci could be augmented by the phase variation from Tr to Op phenotype.

Gonococci of both phenotypes, loosely attached to the uterine surface, did not multiply there as much as in the uterine tissue, and transition from Tr to Op phase in uterine washes did not occur at the same rate.
Table IV. Recovery of N. gonorrhoeae 57–120 Op and Tr variants from the uterus of mice infected at late pro-oestrous or dioestrous*

<table>
<thead>
<tr>
<th>Colonial type of inoculum</th>
<th>Stage of oestrous cycle at inoculation</th>
<th>Log10 cfu/uterus at day 7</th>
<th>Percentage of total colonies that were</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Op</td>
</tr>
<tr>
<td>Op</td>
<td>Late pro-oestrous</td>
<td>5.43 SD 0.86†</td>
<td>92.8 SD 10.2</td>
</tr>
<tr>
<td></td>
<td>Dioestrous</td>
<td>Not recovered</td>
<td>4.8 SD 9.6†</td>
</tr>
<tr>
<td>Tr</td>
<td>Late pro-oestrous</td>
<td>3.56 SD 0.53†</td>
<td>96.3 SD 11.2</td>
</tr>
<tr>
<td></td>
<td>Dioestrous</td>
<td>1.25 SD 0.24</td>
<td>49.4 SD 9.8</td>
</tr>
</tbody>
</table>

*Data were obtained from three separate experiments, and expressed as the mean (SD) for 10 mice.
†P < 0.005.
‡P < 0.001.

as transition from Tr to Op phase in uterine homogenates. Furthermore, the number of gonococci in uterine washes decreased with time, compared with viable counts in uterine homogenates. Therefore, Op variants seemed to be heterogeneous, as some gonococci of the Op variant could invade the uterine epithelium but some were only attached to the surface of the uterus. This discrepancy between invasive and non-invasive types of Op variants could be explained by the differences in P. I1 proteins or other surface appendages such as pili.

Tr variants are known to be more virulent in the chicken embryo model and more resistant to human serum than Op variants. Furthermore, Rest et al. have suggested that lectin-like components of gonococci expressing P. I1 can stimulate the oxidative burst of human phagocytes by which organisms are killed intracellularly. The present study has shown that PMNL infiltration did not occur in the genital tract after mice at late pre-oestrous were infected with either variant. Thus, it is unlikely that Op gonococci are killed by phagocytes in the genital tract of mice during late pro-oestrous. Moreover, Tr gonococci converted to Op within the uterus; phase variation rates were > 90% in mice infected at late pro-oestrous, and c. 50% in mice infected at dioestrous. This result contrasts with the report by Salt and Gotschlich that Op gonococci inoculated into the chicken embryo are replaced by Tr variants. We have isolated isogenic variants from seven clinical isolates of N. gonorrhoeae, and tested them for infectivity and survival in the uterus. By comparing their behaviour in the genital tract, we have found that they are similar to isogenic variants of strain 57–120 (unpublished observation).

Histological studies showed clusters of Op gonococci in the subepithelial stroma to which phagocytic cells did not respond. P. II proteins were reported to increase inter-gonococcal adherence as well as attachment to epithelial cells. Therefore, clustering of Op gonococci may be attributable to P. II proteins. Additionally, cluster formation might be associated with increasing oestrogenic activity in the genital tract, because oestrogen promotes the growth of Op gonococci and is more active at late pro-oestrous than at other stages of oestrous. However, P. II has proved to be a most intriguing group of highly variable and diverse proteins, and gonococcal variants, within a strain, may express none, one, or several different protein(s). Moreover, P. II proteins may differ within a strain in both molecular weight and antigenicity. This may explain why Op variants exhibit differing abilities to invade uterine tissue. Thus, it is important to determine which of the P. II proteins can be induced or altered by oestrogen, and whether this P. II protein can contribute to cluster formation. Tentatively P. II could be involved in the intra-uterine survival of gonococci.

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References


