Effects of tampon components on growth and dissemination of Neisseria gonorrhoeae

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SUMMARY Six components used in vaginal tampons were tested for their effects on a strain of Neisseria gonorrhoeae isolated from a patient with disseminated infection. Tampon components containing carboxymethyl cellulose or its derivative prolonged the in-vitro survival of gonococci and, when injected with mucin into mice, significantly (p<0.001) increased the dissemination of gonococci from the peritoneal cavity. In contrast, a component extracted from rayon tampons reduced in-vitro survival and appeared to suppress gonococcal dissemination in mice. Since tampons are used by a large number of women at a time when the risk of developing complications from venereal infections are increased, their effects on potential urogenital pathogens warrant further study.

Introduction
The association of toxic shock syndrome (TSS) in menstruating women with vaginal Staphylococcus aureus and the use of tampons has stimulated interest in determining the biological effects of various tampon components on other pathogens associated with the female reproductive tract. The use of intrauterine contraceptive devices (IUCDs) has already been identified as a risk factor for women in developing pelvic inflammatory disease (PID) due to ascending infections with Neisseria gonorrhoeae, Chlamydia trachomatis, and possibly other pathogens. Menstruation also appears to increase the risk of complications of PID and disseminated gonococcal infection (DGI). Whether this is due to physiological changes accompanying menses or to a combination of intrinsic and extrinsic factors, such as IUCDs and tampons, remains to be determined.

This study was undertaken to determine the effects of selected tampon components on in-vitro growth and survival and on in-vivo dissemination of N gonorrhoeae in laboratory animals.

Materials and methods
BACTERIA
A strain of N gonorrhoeae isolated from the joint fluid of a patient with DGI was used for testing tampon materials. Stock cultures of this strain were frozen at -60°C until streaked on to a GC base medium supplemented with 1% IsoVitalex (BBL-Microbiology Systems, Cockeysville, MD). After incubation at 36°C for 20 hours in a candle-extinction jar, the gonococci were harvested with a cotton swab into sterile phosphate-buffered (0.067 mol/l KH2PO4 and Na2HPO4) saline (PBS), pH 7.4. A spectrophotometer set at 540 nm was used to adjust cell suspensions to an optical transmittance of 50%. Serial 10-fold dilutions of the cell suspension were then quantitated in colony-forming units (cfu) by plate counts. The cell suspensions for inoculating animals contained gonococci from 10^5 to 10^8 cfu/ml.

TAMPONS
Two commercial tampons* and six tampon component materials were tested. The Rely tampon (lot 15501A16A, Proctor and Gamble, Cincinnati, Ohio) and the Tampax Super Plus tampon (lot 6T072T, Tampax Inc, Palmer, Mass) were purchased from local stores. Components of the tampons tested included: from Rely tampons (a) cross-linked derivative of carboxymethyl cellulose (CLD-CMC) and (b) polyester foam pieces; from Tampax tampons (a) powdered carboxymethyl cellulose (CMC) 'Aqualon', (b) rayon 8285, (c) 'Fiber-T/-MV/CL/Comber/Aqualon', and (d) 'Fiber-T/-MV/CL/ Comber.'

*Use of trade names and commercial sources is for identification only and does not constitute endorsement by the Public Health Service or by the US Department of Health and Human Services.
IN-VITRO TESTS

Tampon components

Aqueous suspensions containing 1-2% (wt/vol) of tampon material were prepared in PBS pH 7-4 and tested for inhibitory activity to Ngonorrhoeae. Suspensions were tested both before and after filtration through a Millipore membrane with a porosity of 0.22μ (Millipore Co, Bedford, Mass). Non-filtered suspensions were also tested after being boiled for 30 minutes. Drops of 0.05 ml of tampon suspensions were absorbed into areas of about 1·0 cm in diameter on GC base plates or suspensions were pipetted into wells (5·0 mm in diameter) cut into the agar surface. A lawn of 10⁴ cfu of gonococci suspended in 0.05 ml of PBS was then streaked over the prepared plates. After incubation for 20 hours the plates were examined under a stereoscopic microscope to determine the growth pattern in the presence and absence of the tampon suspension.

Whole tampons

The leachability and pH of Rely (CLD-CMC) and Tamap Super Plus (Rayon 8285) tampons were determined after suspending each tampon in 50·0 ml of PBS, pH 7·4. The turbidity imparted by material leached from each tampon was measured at ½-hour, 1½-hour, and three-hour intervals with a spectrophotometer set at 540 nm. Tampons were also compared for their effects on the attachment of mouse peritoneal macrophages and on the survival of gonococcal cells. For these tests, tampons were suspended in glass containers with 100 ml of Hanks balanced salts (HBS) solution (pH 7·0). Each tampon was tested in triplicate along with HBS controls (without tampon). Mouse macrophages, collected by intraperitoneal lavage of mineral oil-primed mice, were washed in HBS and then added (10⁴ cells) to each tampon and control suspension. The number of macrophages remaining 'free' in each suspension was determined by cell counts made at ½-hour, three-hour, six-hour, and 24-hour intervals after inoculation. In addition to the mouse macrophages, 10⁶ cfu of Ngonorrhoeae were added to each suspension. Survival of Ngonorrhoeae was determined by sequential cultures made from 0·05 ml aliquots of each suspension at three-hour, six-hour, and then 24-hour intervals until no growth was obtained.

IN-VIVO TESTS

The ability of tampon materials to enhance or suppress the circulatory dissemination of gonococci was tested in a mouse intraperitoneal infection model.8-10 Granular mucin and bovine haemoglobin were prepared as described11 and included in the challenge inoculum where indicated. The challenge inocula were prepared as follows: (a) 9·3 mg of granular mucin (type 1701-W, Wilson Laboratories, Chicago, Illinois) and 3·3 mg of haemoglobin (Difco Laboratories, Detroit, Michigan) were mixed with gonococci and tampon materials in a final volume of 0·5 ml challenge dose; (b) 16·6 mg of mucin was mixed with gonococci and tampon materials. Immediately after mixing, the inocula were injected intraperitoneally into 3-6-week-old Institute of Cancer Research (ICR) white mice. Twenty-four hours later the challenged mice were narcotised by inhalation of CO₂, and 0·2 ml of heart blood was aspirated from each. Approximately 0·1 ml of each specimen was immediately streaked on to plates of GC base medium. After incubation in a candle jar for 24 hours, each plate was examined and the approximate number of cfu of gonococcal growth noted (tables I and II).

ANALYSIS OF DATA

The infectious dose 50% (ID₅₀) was determined by interpolation between doses of gonococci giving less than and greater than 50% positive blood culture results. Statistical comparisons were made by the x² test.11

### Table 1: Effects of tampon components on dissemination of Neisseria gonorrhoeae in mice challenged intraperitoneally

<table>
<thead>
<tr>
<th>Tampon component* mixed with challenge</th>
<th>Challenge dose of gonococci (cfu)</th>
<th>Infectious dose 50%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10⁶†</td>
<td>10⁷‡</td>
</tr>
<tr>
<td>(A) Aqualon CMC powder</td>
<td>7/30</td>
<td>25/43</td>
</tr>
<tr>
<td>(B) CLD-CMC</td>
<td>0/28</td>
<td>27/46</td>
</tr>
<tr>
<td>(C) Polyester foam</td>
<td>NT</td>
<td>12/27</td>
</tr>
<tr>
<td>(D) Rayon No 8285</td>
<td>0/20</td>
<td>9/36</td>
</tr>
<tr>
<td>(E) Fiber-T-½/MV/CL/Comber/Aqualon</td>
<td>0/10</td>
<td>10/16</td>
</tr>
<tr>
<td>(F) Fiber-T-½/MV/CL/Comber</td>
<td>0/18</td>
<td>6/16</td>
</tr>
<tr>
<td>(G) Phosphate buffered</td>
<td>0/38</td>
<td>19/51</td>
</tr>
</tbody>
</table>

*2·0 mg of tampon material plus 16·6 mg of mucin in 0·5 ml PBS.
†0·1 ml of heart blood cultured from each mouse 24 h after inoculation (number with positive blood culture/number cultured).
‡x² values: A + B + E vs D + F + G = p<0·0001, A + B vs D = p<0·01, and D vs G = 0·2≤p<0·3.
§cfu = colony-forming units; NT = no test.
Effects of tampon components on growth and dissemination of Neisseria gonorrhoeae

TABLE II Effects of tampon components on mortality and gonococcal bacteraemia in mice given intraperitoneal inoculations

<table>
<thead>
<tr>
<th>Inoculum components*</th>
<th>% Mortality†</th>
<th>No cultured</th>
<th>% Mice having indicated cfu of gonococci per ml of blood‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBS diluent control</td>
<td>0</td>
<td>26</td>
<td>100</td>
</tr>
<tr>
<td>CLD-CMC</td>
<td>0</td>
<td>32</td>
<td>69</td>
</tr>
<tr>
<td>Bovine haemoglobin</td>
<td>23</td>
<td>54</td>
<td>50</td>
</tr>
<tr>
<td>plus mucin</td>
<td>25</td>
<td>76</td>
<td>25</td>
</tr>
</tbody>
</table>

*Inoculum (0·5 ml) contained: 10⁷ cfu of N gonorrhoeae, 2·0 mg of cross-linked derivative of carboxymethyl cellulose (CLD-CMC), 3·3 mg of bovine haemoglobin, 9·3 mg gastric mucin suspended in phosphate-buffered saline (PBS), pH 7·4, PBS substituted for components not in each test.
†Mice dying in <24 h and not cultured.
‡Approximately 0·1 ml of heart blood cultured for N gonorrhoeae 24 h after inoculation.

Results

Aqueous extracts of tampons or their components showed significant variation in their effects on the growth, survival, and in-vivo dissemination of N gonorrhoeae. Inocula containing mucin plus tampon components of CLD-CMC or Aqualon CMC produced significantly (P<0·001) higher rates of gonococcal bacteraemia in mice than in mucin controls and in groups not receiving CMC (table I). The CLD-CMC component when combined with mucin and haemoglobin produced a significantly (P<0·001) higher systemic gonococcal infection in mice than did control inocula without CLD-CMC (table II). Rayon extracts appeared to suppress the in-vivo dissemination of N gonorrhoeae in the mouse model. Significantly fewer (P<0·01) mice in group D (rayon 8285) developed bacteraemia than did those in groups A and B receiving CMC or its derivative (table I). Although fewer mice in group D (9/36) became bacteraemic than in group G (PBS-mucin controls, 19/51), this difference was not significant (P=0·02<0·05).

When rayon and CLD-CMC tampons were compared for leachability and pH, and for their effects on gonococci and the attachment of mouse macrophages, (a) CLD-CMC tampons produced substantial levels of turbidity in suspensions with PBS; percentage optical transmittance ranged from 20-23% with CLD-CMC tampons compared with 79-91% for rayon tampons and 100% for PBS controls; (b) suspensions of rayon tampons were slightly more acid, pH 6·7 compared with pH 7·0 for CLD-CMC tampons; (c) by indirect determinations macrophages appeared to attach more rapidly to the rayon tampons (figure, a); and (d) gonococci remained viable for a substantially shorter period in suspension with rayon tampons than in the control HBS or CLD-CMC suspensions (figure, b). In addition, gonococci growing on a translucent medium showed phenotypic changes in morphology from smooth to granular opaque colonies in areas where the rayon extract had been absorbed. A similar predominance of opaque colonies occurred around agar wells containing the rayon extract.

The factor(s) in rayon extracts producing the effects on gonococcal cells appeared to be heat stable, resisting boiling for 30 minutes, and were filterable through a 0·22-μ Millipore membrane. Analysis of these extracts by gas-liquid chromatography failed to detect fatty acids (C W Moss, unpublished data).

Discussion

The purpose of this study was to determine the biological effects and possible mechanisms whereby tampon materials may influence the growth and dissemination of N gonorrhoeae. Several mechanisms potentially affecting the virulence of gonococci became apparent. (a) The CMC components appeared more soluble in aqueous fluids and had a greater tendency to leach out of the tampon into the surrounding fluid, possibly producing conditions favourable to the survival of gonococci. (b) The wrapped CLD-CMC tampons appeared to provide a less effective surface for the attachment of macrophages than did the non-wrapped rayon tampon. This could conceivably leave a greater number of macrophages engorged with gonococci after removal of the tampon from the vagina. The intracellular survival of gonococci in macrophages may thus act as a ‘Trojan horse’ for the dissemination of the infection across otherwise impregnable tissues. (c) Certain tampon components may inhibit or alter the growth of gonococci, rendering cells more susceptible to host defences. Cells from rough or opaque
colonies, similar to those induced by rayon extracts, can be approximately four times less resistant to normal human serum than cells of smooth or transparent colonies.  

The practical implications of these differences in promoting or inhibiting the dissemination of *N. gonorrhoeae* by tampon materials remains to be ascertained. The effects of tampons on pathogenic micro-organisms in the vagina, however, warrants further study because a number of serious disease complications may arise from this anatomical site.

The authors thank D. C. Mackel, R. L. Anderson, and members of the CDC toxic shock task force for helpful discussion; W. O. Schalla for technical help; and S. J. Smith for statistical advice.

References


Figure: Comparison of non-wrapped rayon tampon (○) with one of wrapped cross-linked derivative of carboxymethyl cellulose (×) for effects on (a) attachment of mouse macrophages (WBC) and (b) survival of Neisseria gonorrhoeae in tampon fluid (● = ● diluent control, 100 ml of Hanks balanced salt solution).