Taking cell cultures to the patient in an attempt to improve chlamydial isolation

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SUMMARY McCoy cell cultures were inoculated with 121 urethral and cervical specimens taken from patients attending one of two sexually transmitted disease clinics. The mean number of Chlamydia trachomatis inclusions was greater when the cultures were inoculated with the specimens and centrifuged in the clinic than when the specimens were first stored in liquid nitrogen. Furthermore, 18 of the 29 chlamydia-positive specimens produced larger numbers of inclusions when inoculated immediately. Despite this, the isolation rate from specimens inoculated directly (22%) was about the same as from specimens which had been frozen (21%). Of the 30 occasions on which the cell monolayers were disrupted, 29 followed immediate inoculation. This may possibly have been due to some toxic factor in some of the specimens.

Introduction

The results of various studies indicate that Chlamydia trachomatis organisms are isolated from 40-50% of men with nongonococcal urethritis and from 20-30% of women with non-specific genital infections.1 These figures are derived from centrifuge-assisted isolation of chlamydia in McCoy cells which have been treated in various ways to enhance detection of inclusions and after specimens have been at 4°C or room temperature for a few hours or after storage at −70°C or in liquid nitrogen. Some viruses, such as respiratory syncytial virus, are isolated more efficiently by taking susceptible cell cultures to the patient and inoculating the cultures immediately the specimens have been taken. This procedure should provide the maximum opportunity for recovery of most micro-organisms. We therefore inoculated McCoy cell cultures with specimens immediately after obtaining them and centrifuged the cultures in the clinics to determine whether this would enhance the isolation of C trachomatis.

Patients and methods

STUDY POPULATION
Sixty-two consecutive male patients attending the special clinics at St Mary’s Hospital or the Central Middlesex Hospital, London, with gonococcal or nongonococcal urethritis and 59 female patients with non-specific genital infections were studied. The investigation formed part of a larger study, for which ethical approval had been obtained from the ethics committee.

CULTURE TECHNIQUE

McCoy cells These were obtained originally from Dr J H Pearce (Microbiology Department, Birmingham University) and are maintained routinely for no more than 30 passages after removal from storage in liquid nitrogen. Complete medium with antibiotics (CMA) for cell maintenance and the same medium with additional glucose and serum (CMGA) for chlamydial growth, have been described.2

Procedure

Urethral swabs from men and cervical swabs from women were agitated in 0-8 ml of 2SP medium. Half of this was then stored in liquid nitrogen and the other half was diluted in CMGA medium and put directly on a monolayer of McCoy cells on a coverslip contained in a plastic flat-bottomed tube. These inoculated cultures were centrifuged in the clinic at 2800 × g for one hour at room temperature in an MSE Super Minor centrifuge. They were then transported to the laboratory and incubated at 37°C for two hours, after which the medium was replaced with 2 ml of CMGA medium containing 1 µg/ml of cycloheximide. After incubation for a further 48 hours, the medium was removed and the cells were fixed with methanol for 10 minutes, stained with
Giemsa, and examined by dark-ground microscopy. Specimens which had been put in liquid nitrogen were transported to the laboratory in the frozen state and after rapid thawing processed in the same way.

Results

*Chlamydia trachomatis* was isolated from 30.6% of 62 male urethral and from 16.9% of 59 cervical specimens. The mean number of inclusions detected in monolayers of cells after immediate inoculation and centrifugation in the clinic was three to four times greater than the mean number detected after storage of an equal portion of the same specimens in liquid nitrogen (table I). Moreover, 18 of the 29 specimens produced a larger number of inclusions after immediate inoculation than after storage. Three specimens produced inclusions after immediate inoculation but not after storage. Two others produced inclusions after storage only, so that the chlamydial isolation rates resulting from use of the two procedures remained similar—82% for samples dealt with immediately and 81% for samples which had been stored.

In the case of the two specimens which produced inclusions after storage unstored aliquots caused complete disintegration of the cell monolayers so that inclusions could not be sought. Indeed, complete or incomplete removal of cells from the coverslips or contamination of the cultures by other microorganisms rendered evaluation impossible with 30 specimens, 29 of which had been inoculated immediately after collection. The occurrence of bacteria and other micro-organisms in these specimens, which may have accounted for the disintegration of the McCoy cell monolayers, is shown in table II. It seems clear that the disruption was not due to overwhelming chlamydial infection because chlamydial inclusions were not detected in cell monolayers which remained intact after they had been inoculated with the frozen portions of the same specimens.

Discussion

We found that there was no advantage in inoculating cell cultures with specimens immediately they had been obtained in the clinic because it did not enhance the chlamydial isolation rate. This finding differs from that of Reeve et al., who reported that inoculation of specimens before freezing produced a 20% higher isolation rate than after freezing and storage. This difference is difficult to reconcile but might be due to the fact that their specimens were frozen at −70°C and possibly incurred a greater chlamydial loss than ours, which were put into liquid nitrogen. Under the right conditions few chlamydial organisms are lost by freezing in liquid nitrogen. Nonetheless, in the experiment of Reeve et al. and also in ours, freezing of specimens reduced the number of inclusions which could be detected, so that immediate inoculation might be valuable when a small number of organisms is being sought, for example after antibiotic therapy.

**TABLE I** Effect of immediate inoculation and of storage on the number of inclusions detected in *Chlamydia trachomatis*-positive specimens

<table>
<thead>
<tr>
<th>Specimens</th>
<th>No chlamydiosis-positive*</th>
<th>Mean No of inclusions after</th>
<th>No of specimens</th>
<th>Disrupting cell monolayers†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Immediate inoculation (A)</td>
<td>Storage (B)</td>
<td>Producing more inclusions by A than after B</td>
</tr>
<tr>
<td>Urethral (men)</td>
<td>19</td>
<td>6851</td>
<td>1783</td>
<td>13 (9 at least two-fold)</td>
</tr>
<tr>
<td>Cervical (women)</td>
<td>10</td>
<td>20 549</td>
<td>7452</td>
<td>2 (completely)</td>
</tr>
</tbody>
</table>

*Includes three specimens in which bacterial contamination made counting inaccurate.
†Cell monolayer disruption occurred only with specimens inoculated immediately.
‡Three of the 19 specimens produced at least two-fold more inclusions after B than by A.

**TABLE II** Micro-organisms contaminating McCoy cell monolayers which were disrupted after immediate inoculation of specimens*

<table>
<thead>
<tr>
<th>Disruption of monolayer</th>
<th>Bacteria</th>
<th>Candida spp</th>
<th>Trichomonas vaginalis</th>
<th>No organisms</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No</td>
<td>%</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Complete</td>
<td>8</td>
<td>2</td>
<td>3</td>
<td>7</td>
<td>20</td>
</tr>
<tr>
<td>Partial</td>
<td>3</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>Total</td>
<td>11</td>
<td>5</td>
<td>3</td>
<td>10</td>
<td>29</td>
</tr>
</tbody>
</table>

*Only one stored specimen, which contained *T vaginalis*, produced disruption and this was partial.
However, we found that the disruption of cell monolayers which occurred after immediate inoculation was a problem. It is interesting that freezing of specimens generally destroyed whatever was responsible for the cell disruption. Often there was an obvious microbial contaminant which may have accounted for it, but sometimes no contaminating micro-organism was detected. Furthermore, it was clear that the cell monolayers had not been disrupted by an overwhelming chlamydial infection. In such circumstances, the possibility that the specimens may contain a toxin, in the same way that faecal specimens from patients with pseudo-membranous colitis contain the toxin liberated by Clostridium difficile, should not be ignored.

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References