The prevalence of *Chlamydia trachomatis* infection in male undergraduates: a postal survey

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**Objectives:** To determine the prevalence of *Chlamydia trachomatis* infection in male undergraduates and to investigate whether prevalence increases with time spent at university. To investigate the feasibility of screening men for *C trachomatis* by self sampling and posting of urine specimens.

**Methods:** The study design was a postal survey undertaken by the Department of Genito-urinary Medicine (GUM) and Student University Health Service (SUHS) in Sheffield. 2607 male undergraduates from the SUHS patient list were invited to participate in the study by providing a first void urine specimen and posting it to the laboratory. The main outcome measure was the detection of *C trachomatis* infection.

**Results:** 758 students participated in the study, a response rate of 29.1%. Nine students (1.2%) tested positive for *C trachomatis*. The prevalence of infection in the first, second, and third year of study was 0.7%, 1.5%, and 1.6% of participants respectively. There was no statistically significant difference in prevalence of infection between first and third year students (χ² test, p = 0.32). However, students with chlamydia had a higher median age (Mann-Whitney U test, p≤<0.05). Contact tracing identified four further cases of *C trachomatis* infection.

**Conclusion:** Screening for *C trachomatis* infection by postal survey is feasible. However, the response rate in this study was poor and the estimated sample size was not reached. Therefore, it has not been possible to determine the true prevalence of infection in this population or to accurately assess changes in prevalence with time spent at university.

**Keywords:** screening; ligase chain reaction; *Chlamydia trachomatis*; postal survey

**Introduction**

The high prevalence of *Chlamydia trachomatis* in the United Kingdom is partly sustained by the asymptomatic nature of the infection resulting in failure to detect it. Few studies have addressed the issue of screening men for *C trachomatis*, but those that have report prevalence rates of 4.1–11.3%. Until recently, invasive procedures have been necessary to provide suitable specimens for *C trachomatis* testing. The introduction of sensitive nucleic acid amplification techniques (NAA) using first void urine specimens increases patient acceptability and gives the potential for “self sampling.”

**Subjects and methods**

Male students reaching the age of 19 or 21 during the academic year of the study (September 1998–August 1999) were identified from SUHS patient lists. It was presumed that a majority of these students would be in the first or third year of study. Addresses were available for 1544 students aged 20–21 and 1063 students aged 18–19. Non-responders to the initial request to participate were not followed up.

**Methods**

Ethical approval was obtained from South Sheffield research ethics committee.

Students were sent an explanatory letter, information sheet, and brief demographic questionnaire together with an appropriately packaged specimen bottle conforming with the International Air Transport Authority 650 Regulations for Transport of Pathological Specimens. The packages were discreet and small enough to be posted through a standard letterbox (3.5 × 5.5 × 12 cm). All were issued with first class return postage and addressed directly to the Public Health Laboratory.

Specimens were tested by LCR using the Abbott LCX system (Abbott Laboratories Diagnostics Division, IL, USA) in pools of five. Pool size was determined as follows:

Proportion of positive pools = \(1 - (1 - p)^n\)

\(p = \) expected prevalence—1–4%, \(p = 0.01\) to 0.04

\(n = \) number of samples in pool.

Urine specimens were refrigerated on receipt in the laboratory and frozen (−20°C) within 48 hours. Specimens were thawed and thoroughly mixed before testing. One ml of each specimen was extracted for LCR and 20 µl aliquots of the extract from each specimen were combined and tested. When a pool gave a positive result, each specimen was tested individually. In each split pool only one constituent tested positive. These specimens were designated confirmed positives having given a positive result by LCR on two occasions.
Participants were informed of their results by post. Those with chlamydia were invited to attend the GUM clinic where a further urine specimen was tested by EIA (Dako IDEIA PCE) and DIF (Syva Micro Trak Chlamydia trachomatis Direct Specimen Kit) to confirm infection. Subsequent management was in accordance with standard clinic protocol.

STATISTICAL METHODS
Sample size was determined assuming a 1% prevalence in first year students, rising to 4% in the third year. Calculations suggested 575 participants would be needed from each year to have a 90% power of detecting a difference at the 5% significance level. Results were analysed by χ² and Mann-Whitney U tests using a statistical package (SPSS).

Results
RESPONSE RATE
A total of 758 students provided urine specimens, a response rate of 29.1%.

RECEIPT OF SPECIMENS
One specimen was unsuitable for analysis; 80.9% (533) of specimens where time of collection was provided were received within one calendar day of collection (table 1).

PREVALENCE
Nine students tested positive for C. trachomatis (1.2%). The prevalence of infection was 0.7%, 1.5%, and 1.6% in first, second, and third year students respectively (table 2). There was no statistically significant difference between first and third year students (χ² test, p = 0.32). However, the median age of infected students was higher than uninfected students (Mann-Whitney U test, p < 0.05).

CONTACT TRACING
The nine males with C. trachomatis infection named 16 contacts. Four tested positive for C. trachomatis, and each was linked to a different index case. Of the remainder, six contacts were confirmed to have been treated and six were untraceable owing to lack of information. Overall, 10 (62.5%) of contacts received treatment.

Discussion
The prevalence of C. trachomatis infection detected in this study is lower than that previously reported in asymptomatic men. However, our response rate was low and the results may not reflect the true prevalence of infection among students. It is possible that students at most risk of C. trachomatis infection may have decided not to participate, so introducing a sample bias. Alternatively, some participants may not have been sexually active or may have been practising safer sex.

Postal surveys have been used successfully by other investigators to screen for C. trachomatis. Unfortunately, low response rates are frequently reported, resulting in failure of screening programmes. This can occur as a result of unwillingness to participate due to fear, stigma or lack of motivation. Inadequate population registers will further reduce response rates. Students are a highly mobile population and do not frequently visit their general practitioner, so registered addresses may be incorrect. Some investigators have gone to great lengths to encourage participation in similar studies, but this approach is unlikely to be practical in a non-study postal screening service.

Six of the male students with C. trachomatis (66.7%) were asymptomatic when they attended clinic and four had only one sexual partner in the past year. It is unlikely that these infections would have been diagnosed without screening. Screening also promotes increased awareness of C. trachomatis.

It was not possible to maintain a strict cold chain with urine samples before LCR testing as recommended by the manufacturers. However, some studies have shown that this is not necessary. Pooling samples could theoretically introduce the risk of an inhibitor from one sample affecting an entire pool, but this method has been shown to be acceptable for screening and the inhibitory effect is reduced by dilution.

The cost of the survey and tests to the NHS was £12 770, which produces a cost per index case detected of £1419. Once clinic attendances, tests, and treatment are included this rises to £1508. These figures are likely to underestimate the cost effectiveness as no account has been taken of subsequent contact tracing or cost savings in future years due to reduced medical complications associated with C. trachomatis infection. Previous studies indicate that cost reductions are possible with contact tracing and by avoiding future disease. However, given the low prevalence of C. trachomatis in this population it is unlikely that they would reduce the cost effectiveness of this screening intervention significantly. Costs incurred in future postal surveys will be even higher if IATA 602 specifications become necessary for the postage of specimens as the packaging is expensive (currently over £5 per package) and too large for a standard letter box, necessitating collection or the use of a courier service.

Table 1  Time from collection of specimen to receipt in laboratory

<table>
<thead>
<tr>
<th>Time from collection of specimen</th>
<th>Number of samples received (n=758)</th>
<th>Percentage of samples received (n=758)</th>
<th>Percentage of samples which tested positive for C.trachomatis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Same day</td>
<td>5</td>
<td>0.7%</td>
<td>0</td>
</tr>
<tr>
<td>One day</td>
<td>528</td>
<td>69.4%</td>
<td>55.6</td>
</tr>
<tr>
<td>Two days</td>
<td>67</td>
<td>8.8%</td>
<td>11.1</td>
</tr>
<tr>
<td>Three days</td>
<td>23</td>
<td>3.0%</td>
<td>11.1</td>
</tr>
<tr>
<td>Four days</td>
<td>13</td>
<td>1.7%</td>
<td>0</td>
</tr>
<tr>
<td>More than 4 days</td>
<td>23</td>
<td>3.0%</td>
<td>0</td>
</tr>
<tr>
<td>Information not specified</td>
<td>99</td>
<td>13.2%</td>
<td>22.2</td>
</tr>
</tbody>
</table>

Table 2  Prevalence of Chlamydia trachomatis infection by year of study

<table>
<thead>
<tr>
<th>Year of study</th>
<th>Number of students participating in study</th>
<th>Number of students with C. trachomatis infection</th>
<th>Prevalence of C. trachomatis infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>First year</td>
<td>284</td>
<td>2</td>
<td>0.7%</td>
</tr>
<tr>
<td>Second year</td>
<td>133</td>
<td>2</td>
<td>1.5%</td>
</tr>
<tr>
<td>Third year</td>
<td>316</td>
<td>5</td>
<td>1.6%</td>
</tr>
<tr>
<td>Fourth or subsequent year</td>
<td>7</td>
<td>0</td>
<td>0%</td>
</tr>
<tr>
<td>Not specified</td>
<td>18</td>
<td>0</td>
<td>0%</td>
</tr>
</tbody>
</table>
Conclusion
In this study we have demonstrated that screening for C trachomatis infection by self sampling and posting of urine specimens is possible. However, there are several practical difficulties associated with postal screening, particularly low response rates, and this approach has not been demonstrated to be cost effective in this low risk population.

Conflict of interest: None.

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Contributors: KR, study design and review of manuscript; SB, study management, data collection and preparation of manuscript; SF, laboratory analysis of specimens, data collection and preparation of manuscript; GK, setting up laboratory protocol for testing of specimens, coordination of specimens and review of manuscript; RP, contact tracing, data collection and preparation of manuscript; MO, provision of patient database, protocol consultation and review of manuscript; SD, economic evaluation and preparation of manuscript.


