TITLE: Study of Chlamydia trachomatis in Military Women; Prevalence, Risk Factors, and a Cost Benefit Analysis of Early Diagnosis and Treatment

PRINCIPAL INVESTIGATOR: Charlotte A. Gaydos, Ph.D.

CONTRACTING ORGANIZATION: The Johns Hopkins University
Baltimore, Maryland 21205-2196

REPORT DATE: September 1998

TYPE OF REPORT: Annual

PREPARED FOR: Commander
U.S. Army Medical Research and Materiel Command
Fort Detrick, Frederick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for public release; distribution unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.
In the third project year, 5,657 women and 1,203 male military recruits were screened at Fort Jackson, SC. Over the course of the project, there have been 16,727 women and 1,203 men screened. Those that tested positive received treatment at the Troop Medical Clinic (TMC). **Methods:** After receiving an hour of instruction on chlamydia including symptoms and sequelae, prevention, and screening and treatment, potential subjects were asked for their informed consent. All potential subjects answered a chlamydia risk history questionnaire, however, some who chose each month not to participate submitted the questionnaire anonymously. Study participants submitted a first catch urine specimen for testing at Johns Hopkins by ligase chain reaction (LCR). **Results:** 583 women (10.3%) and 59 men (4.9%) tested positive and were subsequently treated. If a questionnaire could be avoided and young age (< 25 years) used alone as the screening criterion, 87.9% (11,603/13,204) of the population would need to be tested, and 95.3% (1,162/1,219) of the positives would be identified. Young age was not a risk factor for men, as it was for women. **Conclusion:** We recommend screening women < 25 years of age in this population.
Opinions, interpretations, conclusions and recommendations are those of the author and are not necessarily endorsed by the U.S. Army.

Where copyrighted material is quoted, permission has been obtained to use such material.

Where material from documents designated for limited distribution is quoted, permission has been obtained to use the material.

Citations of commercial organizations and trade names in this report do not constitute an official Department of Army endorsement or approval of the products or services of these organizations.

In conducting research using animals, the investigator(s) adhered to the "Guide for the Care and Use of Laboratory Animals," prepared by the Committee on Care and Use of Laboratory Animals of the Institute of Laboratory Resources, National Research Council (NIH Publication No. 86-23, Revised 1985).

For the protection of human subjects, the investigator(s) adhered to policies of applicable Federal Law 45 CFR 46.

In conducting research utilizing recombinant DNA technology, the investigator(s) adhered to current guidelines promulgated by the National Institutes of Health.

In the conduct of research utilizing recombinant DNA, the investigator(s) adhered to the NIH Guidelines for Research Involving Recombinant DNA Molecules.

In the conduct of research involving hazardous organisms, the investigator(s) adhered to the CDC-NIH Guide for Biosafety in Microbiological and Biomedical Laboratories.

Signature
Charlotte Gaydos, D.P.H.  7-27-98
Date
PI - Signature  Date
## Table of Contents

1 Front Cover ................................................................. 1

2 Form 298 ...................................................................... 2

3 Foreword .................................................................... 3

4 Table of Contents ........................................................... 4

5 Introduction ................................................................. 7

6 Body ........................................................................... 9
   6.1 Methods ............................................................. 9
   6.2 Results ............................................................... 10

7 Conclusions and Future Plans ....................................... 16

8 References .................................................................. 20

9 Appendices ................................................................. 23
   9.1 Articles published during the project’s third year ................. 24


9.2 Abstracts presented at conferences during the project’s third year

9.2.1 International Symposium of Sexually Transmitted Disease Research, Seville 1997.


9.2.1.2 Stability of *C. trachomatis* and *N. gonorrhoeae* in urine at room temperature for LCR and PCR tests.


9.3 Abstracts submitted during the project’s third year for conferences


9.4 Maps

9.4.1 Percentage Chlamydia Positive Females by Geographic Origin 3rd Year
9.4.2 Percentage Chlamydia Positive Males by Geographic Origin Total
9.4.3 Percentage Chlamydia Positive Females by Geographic Origin for Project Years 1-3

9.5 Armed Forces Epidemiology Board and the Institutional Review Boards of Johns Hopkins University and Ft. Jackson (Eisenhower, Ft. Gordon)
5 Introduction

Each year in the United States, there are approximately 4.5 million cases of *Chlamydia trachomatis* infection. Although many *C. trachomatis* infections are asymptomatic, the symptoms and sequelae from infection, including cervicitis, urethritis, chronic pelvic pain, pelvic inflammatory disease (PID), ectopic pregnancy, and infertility represent a large disease burden. Infection rates for young, sexually active women range from 5-20%. Infants infected perinatally can develop pneumonia, otitis media, and conjunctivitis (if not given ocular prophylaxis). In men, chlamydia infections can cause urethritis, epididymitis, and proctitis.

*C. trachomatis* can be transmitted through vaginal, anal, and oral sex as well as mother-to-infant. Risk factors in women for infection include having cervical ectopy, being a young female, having multiple sex partners, inconsistent condom use. In men risk factors include having multiple sex partners, engaging in unprotected sex, and having multiple sex partners.

Detection of genital *C. trachomatis* infection by ligase chain reaction (LCR) using first void urine is a non-invasive, highly sensitive, and highly specific procedure. Although the cost of LCR is higher than other tests such as direct fluorescent antibody (DFA), antigen detection by enzyme immunoassay (EIA), and nucleic acid probe tests, it is more sensitive and more specific. Culture has been considered to be the gold standard in the past, but costs more and is less sensitive than either LCR or PCR.

In the U.S., the annual costs of chlamydia infections and their sequelae are estimated at $5 billion. Screening with the urine LCR for the detection of *C. trachomatis* and treatment of those persons testing positive with a single dose therapy of azithromycin could prevent a large burden of disease caused by *C. trachomatis* infection.

This grant has been in progress for three years. Based upon findings of the study, recommendations will be made for an effective, cost-effective chlamydia control program designed to reduce morbidity due to *C. trachomatis*.
The objectives of this study are to:

1. Determine the prevalence of infection in several military populations at risk of *C. trachomatis* infection including: female military recruits, and women attending the Troup Medical Clinic (TMC) with urogenital symptoms or the Pap Smear Clinic,
2. Determine risk factors predictive of infection;
3. Conduct a cost-effectiveness analysis comparing universal screening versus selective screening utilizing risk factor criteria;
4. Recommend a chlamydial control program: selective screening and treatment, universal screening and treatment, or mass therapy for all female basic recruits; and
5. Monitor PID and ectopic pregnancy rates, over the 3 year period of chlamydia screening.

Determination of the prevalence of chlamydia in the TMC and Pap Clinics were completed in previous study years. Data regarding these portions of the grant were analyzed for and reported in previous annual reports.

This year, a number of study objectives were achieved. Incoming female recruits were screened and the data were analyzed for a third year. Finally, cost-effectiveness models were finalized to determine how best to prevent the sequelae from *C. trachomatis* infections in the upcoming projects' fourth year.

In addition to the aforementioned study objectives, we screened over one thousand incoming male recruits to estimate prevalence in men. Risk factors for infection in male recruits were also studied. This project was undertaken to determine, in part, reinfection potential of women undergoing the screening program.

From the previous three complete years of research, we make the recommendation that young age (<25 years) would be a valid risk factor upon which to screen female recruits joining the U.S. Army.
6 Body
6.1 Methods

STUDY SUBJECTS

Every Sunday, recruits undergoing in processing at the Reception Battalion, Ft. Jackson were directed to a classroom to be taught about the transmission, symptoms, and sequelae of chlamydia by a research nurse. Potential subjects were then asked to participate in the study by filling out a demographic and risk behavior questionnaire and submit a urine specimen. Only female recruits were screened with the exception of the period from May 24, 1998 to June 14, 1998 when only male (n=1,203) recruits from the Physical Exam Section (PES) were screened.

After giving their informed consent, each subject completed a questionnaire for demographic information and sexual risk factor history and provided a urine sample. The questionnaire was a one page, two-sided scanning form (Scantron Corporation, Tustin, CA). All urine specimens, consent forms, and questionnaires were shipped to Johns Hopkins University Chlamydia Laboratory, under appropriate environmental conditions (4°C for urine specimens).

To determine comparability of the volunteer and non-volunteer recruits at the Reception Battalion, with regard to demographics and risk history, a sub-sample of those non-volunteering recruits were invited to anonymously fill out a questionnaire. This sub-sample was collected on the first Sunday of each month from female recruits and during all four weeks of male recruit screening.

CHLAMYDIA URINE LCR TESTS.

Urices were processed and tested by ligase chain reaction (LCR) (Abbott Laboratories, Abbott Park, IL) for chlamydial DNA, according to manufacture’s directions. The LCR test is cleared by the FDA for use with female and male urine specimens.

DATA MANAGEMENT AND ANALYSIS

The scan forms were scanned into an Ascii text format and then appended to the main data base stored as an Access97 (Microsoft Corporation, Seattle, WA) file. The LCR results, demographics, and risk factor information were analyzed using chi-squared tests, Fisher’s exact tests
and logistic regression analysis (Stata 4.0, College Station, TX). Data for the multivariate models were recorded as dichotomous variables (presence of risk vs. no risk or unknown risk) according to the findings of the univariate analysis.

6.2 Results

FEMALE RECRUITS, FT JACKSON, 1997-8 (3rd Year): Of 6,812 recruits presenting at the Physical Exam Station, 5,657 (83.0%) volunteered from August 31, 1997 to August 30, 1998. From the 5,567 participating female recruits, 88.5% (5007/5655) were age 25 or younger, 49.2% (2774/5640) were Caucasian, 35.1% (1979/5640) were African American, and 15.7% (887/5640) were other races. The prevalence for *C. trachomatis* by urine LCR for the population was 10.3% (583/5657).

FEMALE RECRUITS, FT JACKSON, 1996-7 (2 project years): A thorough analysis of this population from January 1996 to December 1997 was completed on 13,223 female subjects presenting at the Physical Examination Section (See Appendix 9.1.3 for the full article). The infection status of nineteen individuals could not be determined due to missing data items or insufficient urine. Of the remaining 13,204 female recruits, the median age was 21 years (range 17 to 39) and 87.9% (11,603/13,204) were age 25 or younger. Half of the women were Caucasian, 35.9% were African American, and 13.1% were other races. The prevalence for *C. trachomatis* by urine ligase chain reaction for the entire population was 9.2%.

By questionnaire, 93.1% reported having had vaginal sex, 26.7% had more than 1 sex partner in the previous 90 days, 31.4% had a new sex partner in the previous 90 days, and only 16.9% always used condoms. A prior history of chlamydia infection was reported in 9.1%, gonorrhea in 3.3%, syphilis in 0.6%, and trichomonas infection in 4.6%. Of those volunteers who denied vaginal sex, 1.4% (13/914) were chlamydia positive, and of those who reported always using condoms, 8.4% (177/2,115) were chlamydia positive.

Of the 823 non-volunteer recruits who filled out a questionnaire anonymously, 203 (24.7%) did not provide age data and were dropped from the analysis. Their mean age was 21 (range 17 to 36), 51.3% were Caucasian and 31.9% were African American and not significantly different from
volunteers. Only 66.9% reported having vaginal sex compared to 93.1% of the volunteers ($p < 0.001$). In the non-volunteers, four variables were significantly different from those of the volunteers, even when vaginal sex was controlled for: only 4.0% reported prior chlamydia infections ($p = 0.013$), 18.2% had a new sex partner ($p = 0.002$), 20.1% consistently used condoms ($p < 0.001$), and 90.7% reported no previous STD diagnosis ($p = 0.001$). Of the non-volunteers, 17.7% had more than one sex partner in the prior 90 days, similar to the volunteers when controlling for vaginal sex ($p = 0.189$).

The age-specific prevalence of \textit{C. trachomatis} infection for the 13,204 volunteers is shown in Fig. 1. The highest prevalence of chlamydia was in females aged 17 (12.2%) years. Prevalences declined sharply with increasing age to below 5% for ages older than 25 years. For further analysis, the youngest age categories (17 to 25 yr.) were combined into a variable called “young” (prevalence 10.0%, 1162/11,603). In the “older” category (ages 26 to 39 yr.) the prevalence was 3.6% (57/1,601). By race, prevalences were 5.5% (369/6,715) for Caucasian, 14.9% (707/4,733) for African-American, and 8.1% (143/1,756) for other races.

**Univariate Analysis.**

Univariate analysis identified 10 significant variables associated with chlamydia infection: young age (17 to 25 years), African-American race, ethnicity other than Caucasian or African American, ever having vaginal sex, > 1 sex partner in the previous 90 days, new sex partner in the previous 90 days, having inconsistently used condoms last 90 days, a previous diagnosis of gonorrhea, a previous diagnosis of trichomonas, and history of any sexually transmitted disease (Table 2). Prior diagnosis of chlamydia or syphilis were not significantly associated with being chlamydia positive.

**Multivariate Analysis.**

In the complete multivariate model, vaginal sex (odds ratio 5.9, 95% confidence interval 3.2, 10.6), young age (odds ratio 3.0, 95% confidence interval 2.3, 4.0), African American race (odds ratio 3.4, 95% confidence interval 2.9, 3.8), more than 1 sex partner (odds ratio 1.4, 95% confidence interval 1.2, 1.7), having a new sex partner (odds ratio 1.3, 95% confidence interval 1.1, 1.6), having inconsistently used condoms (odds ratio 1.4, 95% confidence interval 1.1, 1.6), and history of any
sexually transmitted diseases (odds ratio 1.2, 95% confidence interval 1.0, 1.4) were independent predictors for chlamydia infection (Table 3).

**Strategies for Selective Screening Criteria for Identification of Chlamydia Infections.**

A screening strategy using all variables identified as independent predictors would require testing 100% of the population, and would detect 100% of the positive individuals. In this model, the magnitude of risk associated with having a new sex partner may vary across race categories. For the purpose of a screening program, this would not alter the proportion of the population tested or the percent of positives detected using this model. Because screening based on race would likely be viewed as inequitable, a strategy excluding race was examined. This strategy would necessitate a questionnaire, and include those reporting high risk behaviors, such as more than one or a new sex partner in the previous 90 days, lack of condom use, prior sexually transmitted disease, or young age. Screening on these criteria would still require testing 100% of the population. If a questionnaire could be avoided and young age (≤ 25 years) used alone as the screening criterion, 87.9% (11,603/13,204) of the population would need to be tested, and 95.3% (1,162/1,219) of the positives would be identified. We therefore recommend screening women ≤ 25 years of age only for the last year of the study.

**MALE RECRUITS, FT JACKSON (1998):** There were 1,203 men screened, the prevalence of infection was 4.9%. The mean age was 19.8 (±2.61) years, 61.6% were Caucasian, 33.8% had more than one sex partner, and 36.7% had a new sex partner in the last 90 days. Only 21.2% of men reported using condoms regularly, and 2.6% reported having prior chlamydial infections. Of the men infected with *C. trachomatis*, only 13.6% reported having symptoms. Risk factors that proved useful for predicting chlamydial positivity included: African-American race, more than one sex partner, and a new sex partner. Young age was not a risk factor, as it was for women. This information was submitted to two conferences in abstract form (See Appendices 9.3.2 and 9.3.3).

**CHLAMYDIA PREVALENCE BY PUBLIC HEALTH REGION ORIGIN OF RECRUITS:** The following table, Table 1, describes the geographic variability by the public health region origin
of the recruits for 1) females screened this third project year, 2) males screened this third project year, and 3) females from years one to three of the project. Figures in Appendices 9.4.1, 9.4.2, and 9.4.3.
Table 1. Chlamydia prevalence by public health region in study recruits; 1) female 3rd year, 2) females for 3 project years, and 3) males 3rd year.

<table>
<thead>
<tr>
<th>Public Health Region</th>
<th>States within the Region</th>
<th>Chlamydia Prevalence (%) Females (3rd year)</th>
<th>Chlamydia Prevalence (%) Females (Total)</th>
<th>Chlamydia Prevalence (%) Males (3rd Year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region 1</td>
<td>Massachusetts, Connecticut, Rhode Island, Vermont, New Hampshire, Maine</td>
<td>7/154 (4.5)</td>
<td>19/486 (3.9)</td>
<td>1/20 (5.0)</td>
</tr>
<tr>
<td>Region 2</td>
<td>New York, New Jersey, Puerto Rico, Virgin Islands</td>
<td>61/613 (10.0)</td>
<td>153/1713 (8.9)</td>
<td>3/70 (4.3)</td>
</tr>
<tr>
<td>Region 3</td>
<td>Delaware, Maryland, Pennsylvania, Virginia, West Virginia, District of Columbia</td>
<td>62/664 (9.3)</td>
<td>150/1926 (7.8)</td>
<td>4/124 (3.2)</td>
</tr>
<tr>
<td>Region 4</td>
<td>Georgia, North Carolina, South Carolina, Florida, Alabama, Mississippi, Tennessee, Kentucky</td>
<td>214/1391 (15.4)</td>
<td>564/4328 (13.0)</td>
<td>22/340 (6.5)</td>
</tr>
<tr>
<td>Region 5</td>
<td>Illinois, Minnesota, Michigan, Indiana, Ohio, Wisconsin</td>
<td>54/667 (8.1)</td>
<td>153/2153 (7.1)</td>
<td>4/135 (3.0)</td>
</tr>
<tr>
<td>Region 6</td>
<td>Arkansas, Louisiana, New Mexico, Oklahoma, Texas</td>
<td>92/798 (11.5)</td>
<td>282/2350 (12.0)</td>
<td>12/166 (7.2)</td>
</tr>
<tr>
<td>Region 7</td>
<td>Nebraska, Missouri, Iowa, Kansas</td>
<td>15/251 (6.0)</td>
<td>44/684 (6.4)</td>
<td>3/83 (3.6)</td>
</tr>
<tr>
<td>Region 8</td>
<td>Colorado, Montana, North Dakota, South Dakota, Utah, Wyoming</td>
<td>7/138 (5.1)</td>
<td>17/469 (3.6)</td>
<td>1/31 (3.2)</td>
</tr>
<tr>
<td>Region 9</td>
<td>Arizona, California, Hawaii, Nevada, American Samoa, Guam, Mariana Islands, Marshall Islands, Micronesia, Palau</td>
<td>47/627 (7.5)</td>
<td>130/1674 (7.8)</td>
<td>4/92 (4.3)</td>
</tr>
<tr>
<td>Region 10</td>
<td>Oregon, Washington, Idaho, Alsaka</td>
<td>11/221 (5.0)</td>
<td>24/647 (3.7)</td>
<td>1/22 (4.5)</td>
</tr>
</tbody>
</table>
DISCUSSION IN RELATION TO STATEMENT OF WORK AND PROBLEMS.

For Year Three, the Statement of Work as stated in the grant proposal included 2 tasks.

**#8. Proposed in original grant: Control Program.** Based on the results from the cost analysis results computed in the second year, we will institute a chlamydia control program for incoming basic recruits. If the mass therapy option is used, the effectiveness of it will be compared to a control group, which will consist of universally screened women with treatment of infected individuals, and which will be built into the study design.

**Performed:** The screening and treatment of female military recruits was continued.

**Issues:** Mass treatment was computed to be the most cost-effective intervention if the follow-up time was 5 years; however, the Fort Jackson IRB (Eisenhower, Ft. Gordon) denied approval to test such an intervention. The Fort Jackson IRB response is attached in Appendix 9.5. For this reason, continued screening of female recruits and treatment of positives was performed. Permission to have a mass treatment arm of the study was granted, however, by the Institutional Review Board (IRB) at Johns Hopkins. We plan to screen based on young age (<25 years) the last year of the study.

**#9. Proposed in original grant: Screening.** We will continue to screen approximately 15,000 female recruits by urine LCR at Ft. Jackson to monitor prevalence. Treatment with azithromycin, 1.0 gram dose orally, will be offered to those who are infected with chlamydia.

**Performed:** There were 5657 females and 1203 males who were screened during the year.

**Issues:** Inability to schedule troops, due to troop commitment to other duties decreased the total number of female recruits screened over last year by 13.1%. We will start screening based on age in the fourth study year.
7 Conclusions and Future Plans

In view of results from the analyses described above and events over the past year, we are negotiating the following activities for the final year of this study. Each of these six point will be discussed further below.

1) continuation of the retrospective analysis of care obtained for PID in the women screened and those women not screened from January 1996 to September 1998 under the protocol described above;
2) re-specification of the CEA from a one-year analytic horizon;
3) continued screening of male recruits;
4) implementation of a screening protocol based on age equal to 25 or less for women in processing at Fort Jackson;
5) assessment of the impact of screening based on age on out-patient care obtained for PID in women offered screening and those not offered screening;
6) implementation of a questionnaire to obtain information on patient preferences for the use of self-administered swabs and possible use of the self-administered swab by a sub-set of volunteers at baseline.

Consistent with the protocol described and discussed above, we will continue to retrospectively document in-patient PID in women. Discussions amongst the investigators have recently determined that in the absence of data on sex mixing patterns after basic training, reinfection, and lack of longitudinal screening programs, conduct of the cost-effectiveness analyses from primarily a one-year analytic horizon, is most relevant to the goals of this study due to potential for re-infection. Consequently, the cost-effectiveness analysis will be conducted from a one-year analytic horizon which considers only PID prevented, as well as a multi-year horizon which considers PID, chronic pelvic pain, and ectopic pregnancies prevented, as has been done in the past.

Preliminary screening efforts for male recruits with internal review board (IRB) approval was conducted at Fort Jackson, SC from May 1998 to June 1998 identified a C. trachomatis prevalence
of 4.9%. Chlamydial infections in men directly impact re-infection rates in female recruits, limiting the ability of only one screening event to prevent sequelae in females. Documentation of the extent of infection in male recruits provides preliminary data to determine this potential of re-infection in women. The pilot data in men were based on 1,203 total men screened. To strengthen confidence in the prevalence estimates in male recruits, we will again screen for chlamydia in approximately 1,000 male recruits at Fort Jackson consistent with the protocol implemented during the spring of 1998. This project will be assisted by Dr. Jane Cecil, an infectious disease fellow at the Johns Hopkins University who will travel to Fort Jackson, SC. Arrangements for submission of a modified IRB application are in progress.

Consistent with the findings of the risk factor analysis noted above and documented in the recent New England Journal of Medicine publication, young age (i.e., age 25 or less) was the only factor available for practical use (i.e., facilitated access to data and high political feasibility) which was significantly associated (O.R. 3.0) with chlamydial infection in the female recruits screened at Fort Jackson, SC. Additionally, this screening strategy provides a cost-savings over no screening. Consequently, commencing in October 1998 we will alter the screening protocol to be consistent with the results of the analysis and screening in the PES will be limited to young women age 25 years or less. This strategy, which will provide screening for an estimated 87.9% of the incoming recruits, is expected to offer the potential to identify 95.3% of the infections in new female recruits.

Given that approximately 4.7% of all infections in new female recruits would not be identified by an age based screening protocol, we will conduct chart reviews to assess the magnitude of out-patient care for PID at Fort-Jackson in women who are screened and those not screened for the last year of the study. While this review is not intended to provide data for an exhaustive analysis of the impact of screening on PID, it will indicate potential impact on care sought within the first eight weeks of service. An addendum to the IRB to obtain informed consent to review records will be submitted for approval.

The impact of screening based on age on out-patient care obtained for PID in women offered screening and those not offered screening is currently being assessed. Hospitalizations (PASBA) for PID (ICD9 codes 614 and 615), infertility (ICD9 code 628), and ectopic pregnancy (ICD9 code 633) in Army enlisted females with less than or equal to one year of service are being examined per
person year for 1995, 1996, and 1997. Denominator data is being collected from the Defense Manpower Data Center, Monterey, CA. The Army recruits screened are being followed for hospitalization. Cases are being comprised of only those subjects that entered full-time active duty and exclude those going from basic training at Fort Jackson into the reserves who would not be hospitalized in the military healthcare system unless on temporary active duty. The remaining females entering the Army as enlisted soldiers during the same time period in which the cases were gained (as per DMDC) comprise the remainder of the cohort. The cohort is being followed for hospitalization for chlamydia sequelae through their first year of service.

Another Women's Defense Grant (P.I. Anne Rompalo, M.D.) being conducted at Fort Bragg, also a collaborative effort between Johns Hopkins and the Army, has found self-administered vaginal swabs (SAS) to be a potentially convenient alternative to the use of cervical and urine specimens with high sensitivity and specificity. Others have found the use of SAS to provide a sensitive and specific way to identify \textit{C. trachomatis} in women which is highly acceptable \textsuperscript{19}. To assess the potential feasibility of future use of self-administered swabs we will implement a questionnaire to approximately 100 volunteers to determine the preferences of new female recruits with regard to screening test procedure, as well as ask them to provide a self-administered swab. We will compare prevalence obtained by urine to that from vaginal swabs in this sub-set of volunteers. An application for IRB approval is being drafted.

Urine-based screening for \textit{C. trachomatis} by Ligase Chain Reaction was effective in a female military recruit population, as well as in a symptomatic Troop Medical Clinic population and an asymptomatic PAP clinic population. Acceptance was high, the urine specimens were readily obtained, and the assays were able to be performed quickly and efficiently.

The study has demonstrated a high prevalence (9.0\%) for female recruits from a geographically and demographically diverse group, a substantial prevalence (11.9\%) from a symptomatic Troop Medical Clinic population, and a higher than expected prevalence (7.3\%) from an asymptomatic PAP clinic population. These results have indicated the need for an ongoing chlamydial control program in such female military groups.

Among recruits, risk factor analysis by multivariate logistic regression identified five independent, statistically significant, predictors for being infected with chlamydia: young age,
African American, vaginal intercourse, more than one new sex partner, and a new sex partner in the prior 90 days. Women that volunteered from the recruit population appeared to have behavioral characteristics that put them at high risk for chlamydial infections. Women, who were non-volunteers appeared to be similar demographically and many also practiced high risk behavior, but were significantly less likely to have these risk factors (prior chlamydia infection, vaginal sex, new sex partner, more than one sex partner, and inconsistent condom use) than were the volunteers.

Although the women from the symptomatic (TMC) and asymptomatic (PAP) groups had prevalences that were higher than the recruit population, their demographic and risk factor profiles were similar. The numbers of women enrolled at the present time are insufficient to perform univariate or multivariate regression analyses.

A chlamydial screening program that focused on screening all young female recruits (age 25 or less) would require that 87.2% of this population would be screened and 95.8% of all positive infections would be identified. Such a screening program as this, which employed urine LCR testing, has the potential to prevent pelvic inflammatory disease and ectopic pregnancy in Army women.
8 References


9 Appendices
9.1 Articles published during the project’s third year


Use of Ligase Chain Reaction with Urine versus Cervical Culture for Detection of \textit{Chlamydia trachomatis} in an Asymptomatic Military Population of Pregnant and Nonpregnant Females Attending Papanicolaou Smear Clinics

CHARLOTTE A. GAYDOS, M. RENE HOWELL, THOMAS C. QUINN, JOEL C. GAYDOS, AND KELLY T. MCKEE, JR.

Infectious Disease Division, The Johns Hopkins University, Baltimore, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, and U.S. Army Center for Health Promotion and Preventive Medicine, Aberdeen Proving Ground, Maryland, and Preventive Medicine Service, Womack Army Medical Center, Fort Bragg, North Carolina

Received 13 November 1997/Returned for modification 27 January 1998/Accepted 16 February 1998

Ligase chain reaction (LCR) (Abbott Laboratories, Abbott Park, Ill.) with first-catch urine specimens was used to detect \textit{Chlamydia trachomatis} infections in 465 asymptomatic military women attending clinics for routine Papanicolaou smear tests. Results were compared to results of cervical culture to determine the sensitivity of the urine LCR and the possible presence of inhibitors of amplification in pregnant and nonpregnant women. Discrepant results for LCR and culture were resolved by direct fluorescent antibody staining of culture sediments, two different PCR assays, and LCR for the outer membrane protein 1 gene. The prevalence of \textit{Chlamydia} in specimens by urine LCR was 7.3% compared to 5% by culture. For 434 women with matching specimens, there were 11 more specimens positive by LCR than were positive by culture, of which all but one were determined to be true positives. There were four culture-positive, LCR-negative specimens, all from nonpregnant women. The sensitivity, specificity, and positive and negative predictive values of urine LCR after discrepant results were resolved were 88.6, 99.7, 96.9, and 99.0%, respectively. The sensitivity of culture was 71.4%. From the 148 pregnant women (prevalence by LCR, 6.8%), there were no patients who were cervical culture positive and urine LCR negative to indicate the presence in pregnant women of inhibitors of LCR. Additionally, a subset of 55 of the LCR-negative frozen urine specimens from pregnant women that had been previously processed in LCR buffer were inoculated with 5 cell culture inclusion forming units of \textit{C. trachomatis} and retested by LCR; all tested positive, indicating the absence of inhibitors of LCR in urine from these pregnant women. The use of LCR testing of urine specimens from asymptomatic women, whether pregnant or not, offers a sensitive and easy method to detect \textit{C. trachomatis} infection in women.

Approximately 4 million \textit{Chlamydia trachomatis} urogenital infections occur in the United States annually, and more than 50 million cases occur worldwide (7, 28). Unfortunately, symptoms are often mild or absent among infected men and women, leaving a large reservoir of infected persons to continue transmission to new sex partners (29). Chlamydial infections occur primarily among young sexually active persons. A high prevalence is common to all socioeconomic groups and may range from 5 to 20% in various groups of young adults (32, 33). Because of the high probability of progression of asymptomatic disease to serious sequelae, it has been recommended that individuals at risk for chlamydial infections be screened, especially women who are vulnerable to the serious consequences of genital infections, such as pelvic inflammatory disease, ectopic pregnancy, and tubal infertility (7, 11). Urine can now be used to detect chlamydial infections in women by ligase chain reaction (LCR) (2, 8, 14, 20, 31, 34), which with its easily obtainable specimen is a cost-effective method for screening programs for asymptomatic women (16). Because asymptomatic military populations have not been studied widely with regard to chlamydial infections (4, 6, 10, 21, 26, 27) and because the sensitivity of the urine LCR assay has been reported to be low for samples from pregnant women due to the presence of inhibitors to amplification (18), we compared urine LCR to cervical culture for the detection of \textit{C. trachomatis} in asymptomatic women attending clinics for routine Papanicolaou (PAP) smear tests.

**MATERIALS AND METHODS**

**Populations and specimens.** Military women (n = 480) attending clinics for a routine PAP smear test volunteered for a study to compare urine LCR tests to cervical cultures for the detection of \textit{C. trachomatis} infections. The volunteer rate of the women approached by the civilian research nurse was 71%. The study was approved by the Institutional Review Boards of The Johns Hopkins University, the U.S. Army Medical Research Material Command, Fort Detrick, Frederick, Md., and Womack Army Medical Center, Fort Bragg, N.C. Of 480 women enrolled, 465 provided a urine specimen. All subjects completed a questionnaire for demographic information and behavioral risk factors for sexually transmitted diseases. The data instrument was a one-page, two-sided scannable bubble form (Scanntron Corporation, Tustin, Calif.). During the pelvic examination, an endocervical swab was obtained by the attending clinician at the PAP smear clinic, who recorded clinical signs and symptoms on the data form. Culture swabs were placed into 2-sucrose-phosphate chlamydia transport medium. Commercial transport medium was replaced with in-house transport medium after 1 month of the study due to some toxicity of the former to tissue culture cells. Specimens were stored appropriately (4°C for urine specimens and −70°C for cultures) until shipping of the urine specimens at 4°C and cultures at −70°C. Shipment were made to ensure arrival at the laboratory within 4 days of collection. All specimens, consent forms, and data forms were shipped to Johns Hopkins Chlamydia Research Laboratory.
Laboratory procedures. Urine specimens were processed and tested by LCR (Abbott Laboratories, Abbott Park, Ill.) according to the manufacturer's instructions. Briefly, 1 ml of urine was centrifuged at 13,000 x g for 15 min. After the supernatant was removed, 1 ml of urce buffer was added to the pellet and the mixture was vortexed. After being heated at 97°C for 15 min, specimens were cooled and 100 μl of each specimen was added to an LCR unit dose tube. An appropriate chlamydia-positive control was included for the processing steps for each group of specimens. Additionally, two negative controls and two positive calibrator controls supplied by the manufacturer were used for each LCR assay run. After the amplification step in the automated thermocycler, unit dose tubes containing the specimens and controls were transferred to the automated enzyme immunoassay machine for the detection of amplified products. Tubes containing the amplified products were then opened: the automated enzyme immunoassay process sampled tubes by piercing the tops of the unit dose tubes, which prevented amplicon contamination. In order to prevent other sources of contamination, specimens were processed in a designated room separate from the room used to amplify and detect specimens. Gloves were frequently changed and aerosol-barrier pipe tips and dedicated pipettes were used. Strict quality-control measures such as machine maintenance checks, daily cleaning of laboratory areas and equipment with bleach, and area swipe tests to monitor ampli-

Comparison of urine LCR to cervical culture. Of the 465 women, 31 women did not have matched culture specimen results. Ten specimens were toxic for tissue culture and no cervical cultures were collected from 21 women, leaving 434 matched specimens for comparison. After the use of the commercial chlamydia transport buffer was stopped and the in-

mydia infection by LCR was 7.3%. The prevalences of infec-
tion for other categories based on LCR included 11.0% for women ≤25 years of age, 8.9% for African-American women, and 6.8% for pregnant women. By risk category the preval-
ences were 15.1% for those with a new sex partner in the previous 90 days, 10.3% for those with more than one sex partner in the previous 90 days, 7.5% for those with inconsistent condom use, 7.4% for those reporting vaginal sex, and 3.6% for those with a prior chlamydial infection.

In univariate analysis only young age (≤25 years) (odds ratio [OR], 4.23; 95% confidence interval [CI], 1.72 to 10.43) and a new sex partner (OR, 2.61; 95% CI, 1.11 to 6.1) were predic-
tors of chlamydial infection (Table 2). However, when we con-
trolled for age, a new sex partner was no longer significant.

Comparison of urine LCR to cervical culture. Of the 465 women, 31 women did not have matched culture specimen results. Ten specimens were toxic for tissue culture and no cervical cultures were collected from 21 women, leaving 434 matched specimens for comparison. After the use of the commercial chlamydia transport buffer was stopped and the in-

mydia infection by LCR was 7.3%. The prevalences of infec-
tion for other categories based on LCR included 11.0% for women ≤25 years of age, 8.9% for African-American women, and 6.8% for pregnant women. By risk category the preval-
ces were 15.1% for those with a new sex partner in the previous 90 days, 10.3% for those with more than one sex partner in the previous 90 days, 7.5% for those with inconsistent condom use, 7.4% for those reporting vaginal sex, and 3.6% for those with a prior chlamydial infection.
house 2-sucrose-phosphate medium was used, no further specimens toxic to tissue culture were observed. Among the 31 specimens without matched results, there were two LCR-positive urine specimens for which a matching cervical culture was not collected.

From the 434 matched specimens, 32 (7.4%) were LCR positive, of which 31 (7.3%) were confirmed as true positives (Table 3). There were 21 LCR-positive, culture-positive specimens. Four patients had urine-LCR-negative, cervical-culture-positive specimens. Discrepancy analysis of these LCR-negative, culture-positive specimens demonstrated that one was positive in the repeat LCR assay and was OMP-1 LCR positive, one had a negative value which was close to the cutoff value for a positive result and was PCR positive when the archived frozen urine was tested, one had a culture transport specimen that was PCR positive, and the results of one could not be confirmed by any of the ancillary tests, including repeat culture. The initial LCR-negative results from these four urine specimens were all considered to be false negatives.

There were 11 specimens that were LCR positive and culture negative, 10 of which could be confirmed as true-positive specimens (Table 4). Five were DFA positive, six were urine PCR positive, seven were culture PCR positive, and eight were OMP-1 LCR positive. Thus, all but one of these LCR-positive specimens were confirmed as true positives by at least one or more additional assays. After resolution of the discrepant results, the sensitivity, specificity, and positive and negative predictive values of urine LCR were 88.6, 99.7, 96.9, and 99.0%, respectively (Table 3), and the sensitivity of culture was 71.4%.

### TABLE 2. Univariate analysis of results relative to factors associated with positive urine LC Rs for military women attending PAP smear clinics

<table>
<thead>
<tr>
<th>Factor*</th>
<th>% with a positive LCR Factor absent Factor present</th>
<th>OR (95% CI)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age &lt;25 yr (254)</td>
<td>2.8 11.0</td>
<td>4.2 (1.72, 10.43)</td>
<td>0.002</td>
</tr>
<tr>
<td>African-American (233)</td>
<td>6.8 8.6</td>
<td>1.3 (0.61, 2.71)</td>
<td>0.501</td>
</tr>
<tr>
<td>Pregnant (142)</td>
<td>7.4 7.0</td>
<td>0.94 (0.44, 2.03)</td>
<td>0.882</td>
</tr>
<tr>
<td>Normal pelvis exam (275)</td>
<td>12.5 6.6</td>
<td>0.49 (0.18, 1.31)</td>
<td>0.154</td>
</tr>
<tr>
<td>Prior diagnosis of STD* (127)</td>
<td>8.3 3.9</td>
<td>0.45 (0.17, 1.2)</td>
<td>0.110</td>
</tr>
<tr>
<td>Having had more than one sex partner in last 90 days (68)</td>
<td>6.8 10.3</td>
<td>1.6 (1.53, 3.73)</td>
<td>0.316</td>
</tr>
<tr>
<td>Having had a new sex partner in last 90 days (53)</td>
<td>6.4 15.1</td>
<td>2.6 (1.11, 6.10)</td>
<td>0.027*</td>
</tr>
</tbody>
</table>

* Numbers in parentheses represent numbers of individuals with the factor present (n = 465).

### LCR of urine of pregnant women

There were 148 urine specimens from pregnant women. The prevalence of chlamydia infection by LCR for the pregnant women was 6.8%, and that for the nonpregnant women was 7.8%. There were no culture-positive, LCR-negative results from pregnant women which could have indicated the presence of LCR inhibitors. All four of the culture-positive, LCR-negative specimens were from women who were not pregnant. In addition, a subset of 55 LCR-negative urine specimens, previously processed in LCR buffer and frozen, which were from pregnant women and were inoculated with chlamydia and retested by LCR were all LCR positive, indicating the lack of inhibitors. Of the 65 available archived urine specimens from pregnant women which were LCR negative and tested in the internal control assay, there were 3 (4.6%) that exhibited inhibition based on a negative value for amplification of the internal control.

### DISCUSSION

Chlamydia infections were of a higher prevalence than expected from these asymptomatic military women attending a clinic for a routine PAP smear test. An LCR prevalence of 7.3% underscores the necessity for the recommendation to screen all sexually active young women when they are attending a routine health care clinic (7). The high prevalence of 11.0% for those <25 years of age confirm the result of studies of others that young age is a significant risk factor for chlamydial infections (13, 17, 22). These results indicate the need to screen all sexually active young women when they are attending a routine health care clinic (7). The high prevalence of 11.0% for those <25 years of age confirm the result of studies of others that young age is a significant risk factor for chlamydial infections (13, 17, 22). These results indicate the need

### TABLE 3. Resolution of urine-LCR-positive and cervical-culture-negative discrepant results for C. trachomatis in military women attending PAP smear clinics (n = 11)

<table>
<thead>
<tr>
<th>Test result</th>
<th>LCR (urine)</th>
<th>DFA*</th>
<th>PCR (urine)</th>
<th>PCR (cervix)</th>
<th>LCR for OMP-1 (urine)</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory no.</td>
<td>1264</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>2407</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>3197</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>3659</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>3891</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Confirmed</td>
</tr>
<tr>
<td>5560</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>5570</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>Unconfirmed</td>
</tr>
<tr>
<td>6082</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>6280</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>6966</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
<tr>
<td>8016</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Confirmed</td>
</tr>
</tbody>
</table>

* DFA staining of culture transport vial specimen.

**Table 3. Comparison of urine LCR to cervical culture for C. trachomatis in military women attending PAP smear clinics**

<table>
<thead>
<tr>
<th>Test</th>
<th>Result</th>
<th>No. of % with test result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cervical culture</td>
<td>Positive</td>
<td>25 (5.8)</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
<td>409 (94.2)</td>
</tr>
<tr>
<td>Urine LCR</td>
<td>Positive</td>
<td>32 (7.4)</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
<td>402 (92.6)</td>
</tr>
</tbody>
</table>

* Four hundred eighty women enrolled; 434 had matching specimens.

**Table 4. Resolution of urine-LCR-positive and cervical-culture-negative discrepant results for C. trachomatis in military women attending PAP smear clinics (n = 11)**
for an ongoing chlamydial control program for such female military personnel as those enrolled in this study. This population demonstrated a high degree of sexual behaviors placing them at risk for sexually transmitted diseases, with 98% being sexually active, 15% having more than one partner, 11% having a new partner in the last 90 days, and 88% using condoms inconsistently. All of these behaviors have been shown by others to be predictive of chlamydial infection (1, 22-24, 36). In the univariate analysis for this study, both young age (prevalence, 11.0%) and having a new partner (prevalence, 15.1%) reached statistical significance. However, when we controlled for age, a new sex partner was not significant. Young age (≥25 years), which is an easily determined risk factor and which is a nonthreatening question for those women who may be reticent to answer questions about their sexual behavior, appears to be an excellent predictor of chlamydia infection and can be recommended for deciding who should be screened in clinical or outreach situations (13, 17).

Urine LCR performed well in this study of asymptomatic women, with a sensitivity of 88.6%, which is similar to that demonstrated by others for asymptomatic women (87.5%) (2). Compared to cervical culture, which had a sensitivity of 71.4%, LCR detected more infected women. Many reasons can account for the lower culture sensitivity. Not only can the cold chain of transport be interrupted, but the quality of the transport medium is important as well. Initially, a commercially available transport medium was used in this study, which resulted in many (10) toxic tissue culture results. Quality-control assays of the remaining lot of uninoculated transport medium demonstrated that it was toxic to cells in tissue culture. After switching to the use of our own transport medium, which is quality controlled in tissue culture, we observed no further toxicity.

Additionally, the quality of the endocervical specimen, as measured by the presence of columnar epithelial cells, has been shown to play a significant role in the numbers of positive specimens (19, 37). In another study of family-planning clinics in Baltimore, Md., clinicians obtained adequate specimens only 72.3% of the time (37). Thus, inadequate cervical swab specimens could have contributed to the lower sensitivity of culture in our study. Other studies have demonstrated higher sensitivities for urine LCR than cervical culture (2, 5, 8, 20, 31, 34). Sensitivities for cervical culture in these studies has ranged from 45.5 to 46.9% to 55.6 to 65.0% (2, 5, 8, 34). Schachter et al. have demonstrated that the sensitivity of culture for C. trachomatis may be increased from 67.1% to 74% by adding a urethral swab culture, which could be indicative that some women may be infected only in the urethra and not the cervix (31). This could help explain the higher number of positives found by urine LCR, presumably reflecting infections from both the cervix and the urethra. Because urine is an easy-to-obtain, noninvasive specimen giving accurate results with LCR, it is ideal for screening asymptomatic individuals who may not be presenting for a pelvic exam or for outreach screening programs.

Although our study enrolled only 148 women who were pregnant, we did not observe any indication of inhibitors in urine specimens, as evidenced by the lack of urine-LCR-negative results when the cervical culture was positive. Although there were four such specimens in this study, they were all from nonpregnant women. Another study has reported a significant problem with inhibitors in urine with use of the LCR test; however, the urine specimens were transported at ambient temperatures, which may have influenced the LCR results (18, 25, 30). The spiking experiment in our study did not demonstrate any inhibitors in the SS LCR-negative, previously frozen urine specimens from pregnant women. It is possible that freezing and thawing of these processed urine specimens reduced or destroyed some LCR inhibitors. Freezing and thawing reduced the inhibition from 19 to 16% in one study (35). Additionally, the experiment which tested the archived urine of 65 pregnant women demonstrated only three (4.6%) inhibited specimens. This value is of the same order of magnitude as that reported by others for inhibition in urine specimens (2.6 and 1.8%) for amplified testing (3, 15). Most investigators now believe that inhibitors to amplification exist for both urine and cervical specimens (3, 15, 35). A combination of heat treatment (95°C for 10 min) and 10-fold dilution of the processed specimen reduced inhibition of PCR from 19 to 4% in one study (35). The pH of the cervical mucosa was partly correlated with inhibitors (35). Decreased inhibition was found at pH values of ≥7.5. The degree to which inhibitors to amplification influence the prevalence detected by LCR and PCR needs to be further studied. Roche Molecular Systems has addressed this problem by incorporating an internal DNA control amplification and detection assay into their new combination PCR assay for C. trachomatis and Neisseria gonorrhoeae, which will prove to be a great advance in the diagnostic capability of amplification assays. Specimens exhibiting inhibitors can be diluted or heated and their DNA can be extracted, and tests can be repeated. The use of the internal control will give a greater degree of confidence to the validity of a negative amplification result. Consideration of the use of an internal control should be given for amplification tests in the future. The College of American Pathologists now requires examination of a control to assess the presence of inhibitors in all amplification procedures.

In summary, young sexually active women, including those in the military, should be frequently screened for chlamydia infections. Urine LCR offers an easy and sensitive method to accomplish this, especially for women not presenting for a pelvic examination. It is cost-effective in preventing the expensive sequelae of pelvic inflammatory disease, ectopic pregnancy, and tubal infertility (16).

ACKNOWLEDGMENTS

We thank the study coordinator, Barbara Pare; the research nurses Eleanor Howard, Katy Cline, and Bobbi Jones and the staffs of the Fort Bragg clinical sites for obtaining specimens; the laboratory technicians Graciela Jasek, Laura Welsh, Dion Pham, Diana Perkins, Sandy Leister, and Kimberly Crockfied for performance of laboratory tests and data entry; Katherine Clark for statistical assistance; and Pat Buist for assistance in manuscript preparation.

Funding for this study was from Department of the Army grant DAMD 17-95-1-5064.

REFERENCES


Pooling Urine Samples for Ligase Chain Reaction Screening for Genital *Chlamydia trachomatis* Infection in Asymptomatic Women

KATHERINE A. KACENA, SEAN B. QUINN, M. RENÉ HOWELL, GUILLERMO E. MADICO, THOMAS C. QUINN, AND CHARLOTTE A. GAYDOS

Division of Disease Control, International Health, School of Hygiene and Public Health, and The Division of Infectious Diseases, The Johns Hopkins University, Baltimore, and National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland, and Universidad Peruana Cayetano Heredia, Lima, Peru

Received 23 July 1997/Returned for modification 19 September 1997/Accepted 4 November 1997

The accuracy of pooling urine samples for the detection of genital *Chlamydia trachomatis* infection by ligase chain reaction (LCR) was examined. A model was also developed to determine the number of samples to be pooled for optimal cost savings at various population prevalences. Estimated costs included technician time, laboratory consumables, and assay costs of testing pooled samples and retesting individual specimens from presumptive positive pools. Estimation of population prevalence based on the pooled LCR results was also applied. After individual urine specimens were processed, 568 specimens were pooled by 4 into 142 pools and another 520 specimens were pooled by 10 into 52 pools. For comparison, all 1,088 urine specimens were tested individually. The sample-to-cut-off ratio was lowered from 1.0 to 0.2 for pooled samples, after a pilot study which tested 148 samples pooled by 4 was conducted. The pooling algorithm was 100% (48 of 48) sensitive when samples were pooled by 4 and 98.4% (61 of 62) sensitive when samples were pooled by 10. Although 2.0% (2 of 99) of the negative pools of 4 and 7.1% (1 of 14) of the negative pools of 10 tested presumptive positive, all samples in these presumptive-positive pools were negative when retested individually, making the pooling algorithm 100% specific. In a population with 8% genital *C. trachomatis* prevalence, pooling by four would reduce costs by 39%. The model demonstrated that with a lower prevalence of 2%, pooling eight samples would reduce costs by 59%. Pooling urine samples for detection of *C. trachomatis* by LCR is sensitive, specific, and cost saving compared to testing individual samples.

There are 89.9 million cases of genital *Chlamydia trachomatis* infection every year worldwide (13). 4.5 million of which occur in the United States (4). Although many *C. trachomatis* infections are asymptomatic (16), the sequelae from infection, including pelvic inflammatory disease (PID), and infertility, represent a large burden for populations worldwide. Furthermore, inflammatory sexually transmitted diseases, such as those caused by *C. trachomatis*, increase the risk of both human immunodeficiency virus (HIV) transmission and infection (7, 11). Together, the high percentage of asymptomatic infections, the sequelae of infections, and the increased association with HIV transmission underscore the importance of screening as a necessary intervention to reduce the burden of diseases caused by *C. trachomatis*.

Detection of genital *C. trachomatis* infection by ligase chain reaction (LCR) with first-void urine is a noninvasive, highly sensitive, and highly specific procedure (2, 8). Although the cost of LCR is higher than that of other tests such as direct fluorescent antibody, antigen detection by enzyme immunoassay, and nucleic acid probe tests, LCR is more sensitive and more specific (15, 17). Culture has been considered to be the "gold standard" in the past but costs more and is less sensitive than either LCR or PCR (3, 5, 10, 12, 14).

Pooling serum samples for HIV testing was found to be accurate and has been used to reduce the cost of enzyme-linked immunosorbent assays for detection of antibody to HIV (1, 6). Pooling for HIV testing has been used to develop both population estimates and, in a multiple-step procedure, to determine which individual sample is positive. Pooling has also been applied to the PCR detection of *C. trachomatis* in endocervical and urethral scrapes (9), but in that study the sample size was small. The investigators acknowledged the need for subsequent studies to rule out the possibility of reduced sensitivity by diluting out individual specimens in the pool.

The screening of women at risk for *C. trachomatis* has been recommended by the Institute of Medicine as a cost-effective program which would prevent the high cost of untreated infections which lead to PID (4). As a screening and treatment intervention reduces the prevalence of *C. trachomatis* infection over time, the cost per specimen tested with the pooling protocol algorithm would be further decreased. The reduction in price occurs for two reasons: (i) as prevalence decreases, pooling a greater number of samples increases cost savings and (ii) the samples from fewer pools would test presumptive positive such that fewer samples would be retested individually. Therefore, the cost for finding one case does not increase dramatically as prevalence decreases, as is the case when samples are tested individually.

In this study we examined the accuracy and cost-saving ability of pooling urine specimens for the detection of genital *C. trachomatis* infections by LCR. A cost analysis of the pooling protocol was conducted to determine the number of specimens it would be necessary to pool in order to provide the
highest cost savings, taking into account the prevalence of infection in the population screened.

**MATERIALS AND METHODS**

**Sample size and parameters.** As part of an ongoing study to determine chlamydia prevalence in asymptomatic U.S. Army females with a mean age (± standard deviation [SD]) of 22 (±2 years), urine samples were tested by LCR to ascertain genital *C. trachomatis* infection. A sample of 506 processed urine specimens was pooled by 4 into 142 pools, and 525 specimens were pooled by 10 into 52 pools. Pools were formed by order of consecutive laboratory accession number. All 1,088 pooled urine specimens were also tested individually. For all discordant individual and pool results, both the individual samples and the pools were retested to confirm results.

**Urine specimen, collection, preparation and assay setup.** Specimen, collection, preparation, and assay setups were performed according to the manufacturer's instructions for the urine-based chlamydia LCR assay (Abbott Laboratories, Abbott Park, Ill.). Specimens were refrigerated immediately after collection and shipped overnight delivery with wet packs to maintain refrigeration temperature. Specimens were either processed immediately on arrival at the laboratory or refrigerated and processed within 2 days. The total time before processing never exceeded 4 days as per the LCR package insert. Pooled refrigerated specimens were amplified the day after processing. Processed urine can be refrigerated or frozen for up to 30 days before testing. We refrigerated our processed specimens for up to 7 days in case retesting was needed. One milliliter of urine was centrifuged at 10,000 × g for 15 min (±2 min) at room temperature. The supernatant was removed, and the pellet was resuspended into 0.1 ml of LCR urine specimen resuspension buffer and vortexed. Preparations were then boiled at 97°C (±2°C) for 15 min (±1 min) to extract the DNA and stored at 2 to 8°C for up to 7 days until tested. Processed urine specimens were subsequently tested individually and tested pooled.

When specimens were tested individually, a volume of 100 μl of processed urine specimen was placed into its own LCR chlamydia amplification unit (unit dose). For each pool of four, 25 μl of each of the four processed specimens was placed into a single unit dose. For each pool of 10, 10 μl of each of the ten processed specimens was placed into a single unit dose. The total specimen volume was then 100 μl for each unit dose. Two negative controls, two positive calibrators, and a positive processing control were included in every amplification run in accordance with the manufacturer's instructions.

**DNA amplification and detection.** Unit dose tubes containing DNA preparations were amplified under the following conditions: 40 cycles of denaturation at 93°C for 1 s, annealing at 59°C for 1 s, extension at 62°C for 1 s, and soaking at 25°C in an LCR thermocycler (Abbott Laboratories). Amplified DNA was detected in an LCR automated machine which performed a particle-based enzyme immunoassay with a fluorescent signal. For individually tested samples, a sample-to-cutoff ratio (SOCO) of ≥1.0 was considered positive, and borderline negative samples (0.80 to 0.99 S/CO) were retested, according to manufacturer's instructions.

**Pilot study.** Because the volume for each individual urine specimen is decreased in the pooled assay, a pilot study was conducted to determine an appropriate S/CO for the pooled assays. The desired S/CO would detect all positive pools, and not more than 0.1% of all negative pools. The pilot study consisted of 448 processed urine samples from the ongoing study of female U.S. Army recruits. The technician, blinded to the individual test results, pooled and tested these 448 samples. By lowering the S/CO from 1.0 to 0.2, all of the pooled tested positive (100%) (25 of 25) and by lowering the S/CO only 2.7% (1 of 37) of the negative pools tested presumptive positive. Since all pools which test positive are retested, specificity with the pooling algorithm is 100%, i.e., no different than with testing processed specimens individually.

**Cost analysis.** A model was developed to determine the pool size that yielded the highest cost savings. The binomial distribution was used to estimate the number of pools that are likely to be positive given a selected pool size and population disease prevalence. Next, the optimal pooling number for a range of disease prevalences was calculated. For a dichotomous outcome (i.e., positive or negative test result for a genital *C. trachomatis* infection), independence was assumed (i.e., the order of the samples received was random with regard to the distribution of the positive or negative samples in the population). The expected proportion of positive pooled assays was determined using the following equation: $s = \left(1 - \binom{n}{r} \binom{p}{r} \binom{n-r}{n-r} \right) \times 100\%$, where $s$ is the expected number of positive samples tested, $n$ is the number of positive samples tested, $n$ is the total number of samples tested, $r/n$ is the prevalence of disease, and $c$ is the number of specimens pooled. This equation accounted for the probability that from 1 to $c$ samples in the pool were positive.

A baseline total cost of $12.76 per individual sample which included $0.36 for laboratory consumables, $3.56 for technician cost, and $8.84 for the LCR assay was used. Laboratory consumables include gloves and supplies used for handling samples. Technician cost was calculated assuming an average of 10 runs per week (i.e., 380 samples), an annual salary of $30,000 with an additional 28% of salary in benefits, and a 69% laboratory or university overhead (i.e., $30,000 \times 1.25 \times 1.09 \times 0.69$). The cost of the five controls used for each 19 specimens tested was calculated into the LCR assay cost, in which the base cost per unit dose was $5. A sensitivity analysis was also done first, with the base cost per unit dose ranging from $5 to $15, technician cost set at $3.56, and cost for laboratory consumables set at $0.36 per specimen tested. In the second sensitivity test, the annual salary of the technician ranged from $20,000 with 28% overhead to $40,000 with 69% overhead, unit dose cost was set at $7.00, and cost for laboratory consumables set at $0.36 per specimen tested. Low technician and low assay costs as well as high technician and high assay costs were also calculated.

**RESULTS**

**Sensitivity and specificity of the pooled assays.** A comparison of the distribution of S/COs for the individual and pooled samples indicated that lowering the S/CO from 1.0 to 0.20 for determining positive pools resulted in high sensitivity with a low proportion of specimens from negative pools that need to be retested individually (Fig. 1). There were two weakly positive individual specimens (i.e., an S/CO of ≥1 but <2.0) in the pilot study (pooled by 4), six weak positives in the study pooled by 4, and four low positives in the study pooled by 10. These weak positives were the only positive specimens in the pool. These pools all tested between 0.2 and 1.0 and sometimes higher.

The pooling algorithm was 100% (48 of 48) sensitive when pooling by 4 and 98.4% (61 of 62) sensitive when pooling by 10 (Table 1). Although 2.0% (2 of 99) of the negative pools of four and 7.1% (1 of 14) of the negative pools of 10 tested presumptive positive, all of the samples in these pools would be retested individually, according to the pooling algorithm. Re-testing the individual samples in the presumptive-positive pools resulted in no false-positive specimens (100% specificity).

**Cost analysis.** For a population with 8% genital *C. trachomatis* prevalence, which is close to the 8.5% prevalence found in our study population of female U.S. Army recruits, pooling by four provided the highest cost savings. The reduction of total assay costs per specimen, which included technician time, decreased from $12.76 to $7.78, i.e., by 39%. The model demonstrated that with a 2% prevalence, pooling eight samples would reduce the cost per sample by 59%. A population prevalence graph was constructed from the model to determine the number of pooled samples that would achieve the highest cost savings (Fig. 2).

A sensitivity analysis for the cost savings model was conducted with ranges of both technician and LCR unit dose costs. For specimens tested individually, raising the base cost of the LCR unit dose from $7 to $15 resulted in an increase of the total cost per specimen tested from $12.76 to $22.87, whereas lowering the cost of the LCR unit dose to $5 reduced the total cost per specimen tested to $10.24. Similarly, raising the annual salary of the technician from $30,000 to $40,000 and assuming 28% benefits and 69% overhead increased the total cost per specimen tested from $12.76 to $13.95, whereas lowering the technician's annual salary to $20,000 and the overhead to 20% reduced the total cost per specimen tested from $12.76 to $10.89. The cost per specimen tested with the low unit dose and low technician costs was $11.43, which is the highest unit dose and technician costs yielded a cost of $24.06 per specimen tested.

For a population prevalence of 8% and pooling by four, ranging the unit dose cost from $5 to $15 would result in a total...
POOling Urine Samples for C. Trachomatis LCR Assays

VOL. 36, 1998

FIG. 1. Detection of genital C. trachomatis infection by LCR. Graphs A and C show the distribution of individual urine samples from two groups of 568 and 520 women taken from the study population. Graph B shows the distribution of the samples in A pooled by 4 (n = 142), and graph D shows the distribution of the samples in C pooled by 10 (n = 52). The S/CO for individual samples, which was 1.0, was lowered to 0.2 for pooled samples, as indicated.

The cost of $6.43 and $13.17, respectively, per specimen tested. Raising technician cost from low to high would result in a total cost increase of $6.36 and $8.68, respectively, per specimen tested. The overall savings of the pooling algorithm over individual testing ranged from 37 to 42% when low to high unit dose cost was considered. Similarly, the overall savings ranged from 42 to 38% when a low-to-high technician cost was considered. The total cost per specimen with the pooling by four algorithm with both the low unit dose and low technician cost was $5.01 per specimen tested, and that with the high unit dose and high technician cost was $14.07 per specimen tested. In all of these scenarios, pooling provided a cost savings compared with individual testing.

Estimation of population prevalence with pooled data. The observed prevalence for the individual samples in the 142 pools of 4 was 8.5% (48 of 568), and that for the 52 pools of 10 was 11.9% (62 of 520) (Table 1). The estimated population prevalence, back calculated from the number of positive pools, for the 142 pools of 4 was 9.1 (95% CI: 6.5 and 11.6), and for the 52 pools of 10 it was 12.9 (95% CI: 8.8 and 17.0). Each 95% CI included the observed prevalence of the subsample, 10.1% (110 of 1,088). Additionally, each 95% CI included values within 8 to 9%, the overall prevalence measured in a much larger sample (>10,000) of this population.

DISCUSSION

In this study we evaluated pooling of processed urine specimens for LCR detection of C. trachomatis for both accuracy and cost-saving ability. The high sensitivity and specificity of LCR was not affected by pooling up to 10 samples when the S/CO was adjusted from 1.0 to 0.2. Although a small percentage of negative pools tested presumptive positive, no specificity was lost with the pooling algorithm, since all specimens in pools which test presumptive positive are retested individually with the manufacturer's specified S/CO for the individual test. Since retesting negative pools does increase costs, the specificity of pools must be high.

The cost analysis model showed that depending on the prevalence of C. trachomatis, the number of specimens that should be pooled for optimal cost savings varies. As prevalence decreases, the pooling protocol for screening could save more than 59% of the cost per specimen compared to that for testing individual samples only. Also, early studies have shown that C.
trachomatis screening and treatment programs are cost effective; the Centers for Disease Control and Prevention has estimated that for every dollar spent on prevention, $12 is saved in treating sequelae (4). The use of the pooling algorithm for testing samples obtained during screening could further increase savings in health care costs.

Since C. trachomatis prevalence levels have ranged from 4 to 20% in various populations in the United States, pooling three to four samples is likely to provide the highest cost savings. Furthermore, the cost saved does not significantly change the sensitivity or specificity of the assay. In the event that screening is not conducted, pooling can be used to determine population prevalences over time in order to measure the benefits of disease interventions such as mass treatment or behavioral interventions. The population prevalence back calculation, described previously (6), gave an accurate estimate of the observed population prevalence in this study.

Use of the pooling algorithm would benefit investigators and program planners in two ways: (i) money saved from the use of the pooling algorithm could be applied to other areas of disease prevention and/or (ii) the amount of money allocated to screening would allow more specimens to be tested for the same total cost. Pooling samples for the detection of genital C. trachomatis infection in urine samples is cost saving and simple to perform and could be applicable in screening programs in the United States and in population-based research worldwide.

Pooling is a technique which could be immediately used for significant cost savings in high-volume laboratories such as state labs and referral labs. Laboratories which are currently using less sensitive and specific and less costly techniques could introduce both LCR and pooling into their laboratories.

Specific populations or laboratories that might benefit from pooling include any lab in which the combination of turn-around time and volume allows at a minimum a combination of 19 pools and retests per day. With 96 specimens at a population prevalence of about 4%, pooling by six would fill up one full run (38 test unit doses) per day. The run would include, on average, 16 pools of six and 22 retests.

Laboratory managers should consider two points before using pooling. First, processed specimens from presumptive-positive pools need to be amplified and detected individually. This additional step adds a minimum of 3 hours until individual test results for specimens in presumptive-positive pools are known. Second, laboratory managers should estimate the cost savings they expect to gain for their laboratories. This estimate is a combination of both technicians’ salaries and their benefits, institutional overhead, and the prevalence of chlamydia in the populations served by the laboratory. Pooling a greater number than is recommended for certain population prevalences can cost more money than testing specimens individually.

A potential limitation of the pooling algorithm is the possibility of technician error while processed samples are pooled in the LCR run. The use of tray maps simplifies this process. Samples should be organized by skipping a space after each pooled group in the specimen rack. Thus, pooling adds no significant complexity to setting up unit doses. Additional technician error can be avoided when samples from presumptive-positive pools (detected in the previous run) are retested individually before the routine testing of the new pooled groups. Therefore, each run has a combination of samples that are retested individually and new pooled samples from the next batch of specimens.

The study laboratory has met Clinical Laboratory Improvement Act requirements for the modification of a clinical laboratory procedure from a Food and Drug Administration-approved diagnostic kit. Investigators consider performance documentation of the required study adequate for including the pooling protocol in testing clinical specimens in the study laboratory. Each laboratory that wishes to introduce pooling must meet the requirements to modify a Food and Drug Administration-approved package insert. These requirements include meeting the regulations as set forth in the Federal Register (3a).

Use of pooling processed urine samples for LCR testing of C. trachomatis will decrease the cost of screening, providing more evidence that screening programs can and should be implemented. Further applications of pooling include pooling urine specimens for the LCR detection of Neisseria gonorrhoeae.

![Graph showing cost-saving ability of pooling processed urine specimens before the performance of the LCR test for the detection of genital C. trachomatis infections. The graph shows the cost per sample when the pooling algorithm was used, depending on the number of samples pooled and taking into account various prevalences of infection in the population screened. A baseline total cost of $12.76 per individual sample which included laboratory consumables, technician time, and LCR unit dose costs was used.](image-url)
rhihoeae. The cost savings of pooling urine for both N. gonorrhoeae and C. trachomatis should also be considered.

ACKNOWLEDGMENTS

We acknowledge D. Perkins for her collaboration and T. and A. Kacena for their assistance.

REFERENCES


ABSTRACT

Background Asymptomatic genital Chlamydia trachomatis infections in women can lead to pelvic inflammatory disease, infertility, and ectopic pregnancy. To design a chlamydia-control program, we conducted a large survey of women in the U.S. military.

Methods From January 1996 through December 1997, urine samples from 13,204 new female U.S. Army recruits from 50 states were screened by ligase chain reaction for C. trachomatis infection. Information on potential risk factors was obtained by questionnaire. With multivariate analysis, we identified criteria for a screening program.

Results The overall prevalence of chlamydial infection was 9.2 percent, with a peak of 12.2 percent among the 17-year-old recruits. The prevalence was 15 percent or more among the recruits from five southern states. The following risk factors were independently associated with chlamydial infection: having ever had vaginal sex (odds ratio for infection, 5.9), being 25 years of age or less (odds ratio, 3.0), being black (odds ratio, 3.4), having had more than one sex partner in the previous 90 days (odds ratio, 1.4), having had a new partner in the previous 90 days (odds ratio, 1.3), having had a partner in the previous 90 days who did not always use condoms (odds ratio, 1.4), and having ever had a sexually transmitted disease (odds ratio, 1.2). A screening program for subjects 25 years of age or less (87.9 percent of our sample) would have identified 95.3 percent of the infected women.

Conclusions Among female military recruits, the prevalence of chlamydial infection is high. A control program that screens female recruits who are 25 years old or younger with urine DNA-amplification assays has the potential to reduce infection, transmission, and the sequelae of chlamydial infection.

METHODS

Population and Specimens

All female Army recruits who were present on Sundays between January 1996 and December 1997 at the Physical Examination Section, Reception Battalion, Fort Jackson, South Carolina, were invited to participate in this study. The study was approved by the institutional review boards of Johns Hopkins University and Fort Jackson (Eisenhower Army Medical Center, Fort Gordon, Ga.), as well as the Human Subjects Research Review Board of the U.S. Army Surgeon General. Of the 16,593 recruits approached, 13,223 (79.7 percent) volunteered to participate in the study and were given a briefing about the study as well as an educational briefing about chlamydial infections by the civilian research nurse.

All subjects signed an informed-consent form and completed a questionnaire regarding demographic information, home state, medical care, screening of young, sexually active women has been recommended. In the past, screening for C. trachomatis infections in women has been limited by the need for access to a medical clinic and a pelvic examination. However, C. trachomatis infections can now be detected with high sensitivity (85 to 95 percent) and specificity with DNA-amplification assays performed on urine specimens, allowing cost-effective screening of large numbers of women in nonclinic settings.

Few studies of the prevalence of chlamydial infection in U.S. military populations have been published, and there have been no studies using DNA-amplification techniques among women not seeking health care. Because adolescents have the highest prevalence of disease and most military recruits are young, we conducted a large prevalence study and risk-factor analysis of female recruits from throughout the United States who began basic training at Fort Jackson, South Carolina. We performed this study to determine the extent of infection, assess the feasibility of screening female recruits for C. trachomatis by the ligase chain reaction, and assess which epidemiologic correlates would be useful for implementing an effective chlamydia-control program for female recruits.

MORE than 4 million urogenital Chlamydia trachomatis infections occur in the United States annually. They occur in young, sexually active persons from all socioeconomic groups, with prevalence ranging from 5 percent to 20 percent. Women, especially, bear the burden of disease, with consequences of genital infections ranging from pelvic inflammatory disease to ectopic pregnancy and infertility. These sequelae are associated with a large economic burden. Because up to 80 percent of infected women are asymptomatic and therefore do not seek medical care, screening of young, sexually active women has been recommended. In the past, screening for C. trachomatis infections in women has been limited by the need for access to a medical clinic and a pelvic examination. However, C. trachomatis infections can now be detected with high sensitivity (85 to 95 percent) and specificity with DNA-amplification assays performed on urine specimens, allowing cost-effective screening of large numbers of women in nonclinic settings. Few studies of the prevalence of chlamydial infection in U.S. military populations have been published, and there have been no studies using DNA-amplification techniques among women not seeking health care. Because adolescents have the highest prevalence of disease and most military recruits are young, we conducted a large prevalence study and risk-factor analysis of female recruits from throughout the United States who began basic training at Fort Jackson, South Carolina. We performed this study to determine the extent of infection, assess the feasibility of screening female recruits for C. trachomatis by the ligase chain reaction, and assess which epidemiologic correlates would be useful for implementing an effective chlamydia-control program for female recruits.
and sexual history. The data instrument was a two-sided scannable form (Scantron, Tustin, Calif.). To determine the similarity of the study subjects and those who chose not to participate in the study with regard to demographic characteristics and sexual history, 823 of the 3370 women who did not volunteer were invited to fill out an anonymous questionnaire. Nonvolunteers were asked to fill out a questionnaire only during the first week of each month.

Each volunteer was instructed to collect 20 to 30 ml of first-catch urine (the first part of the urine stream). A unique study number was assigned to each volunteer. All urine specimens, consent forms, and questionnaires were shipped to the Johns Hopkins University chlamydia laboratory. Urine specimens were kept at 4°C until processed, within 48 hours.

**Laboratory Procedures and Treatment**

Urine specimens were processed and tested by the ligase chain reaction (Abbott Laboratories, Abbott Park, Ill.) for chlamydial DNA according to the manufacturer's directions. Each week a list of infected subjects was sent to the research nurse. The infected subjects were contacted and treated at the Troop Medical Clinic at Fort Jackson by directly observed therapy with a single 1-g dose of azithromycin. The subjects were also tested for coexisting sexually transmitted diseases. The sensitivity and specificity of the ligase chain reaction in urine specimens as compared with cervical culture for chlamydia had been previously determined to be 88.6 percent and 99.7 percent, respectively, in another military population.14

**Statistical Analysis**

Questionnaire forms were scanned into a data base (dBASE III Plus, Borland International, Spring Valley, Calif.). The results of the ligase chain reaction, demographic information, and risk-factor information were analyzed as dichotomous variables with the chi-square test. Univariate and multivariate logistic-regression analysis for factors associated with chlamydial infection was performed with Intercooled Stata software (version 4.0, Stata, College Station, Tex.). All independent variables were entered into the model, and a two-sided P value of less than 0.05 was considered to indicate statistical significance. The 95 percent confidence interval for the prevalence value for recruits from each state was calculated with Stata software. A one-way analysis of variance was performed to assess the degree of significance of differences in prevalence between states.

**RESULTS**

**Characteristics of the Subjects**

Of 13,223 subjects presenting at the Physical Examination Section on Sundays from January 1996 through December 1997, 19 could not be evaluated because of missing data or insufficient urine. The median age of the 13,204 who could be evaluated was 21 years (range, 17 to 39); 87.9 percent (11,603) were 25 years old or younger (Table 1). Fifty-one percent of the women were white, 35.9 percent were black, and 13.1 percent were of other races. For the entire population, the prevalence of C. trachomatis infection according to the urine ligase chain reaction was 9.2 percent.

On the questionnaire, 93.1 percent of the subjects reported having ever had vaginal sex, 26.7 percent having had more than one sex partner in the previous 90 days, and 31.4 percent having had a new sex partner in the previous 90 days. Only 16.9 percent reported that their partners always used condoms. A history of chlamydial infection was reported by 9.1 percent of the subjects, gonorrhea by 3.3 percent, syphilis by 0.6 percent, and trichomonsa infection by 4.6 percent. Of the volunteers who reported having had no vaginal sex, 1.4 percent (13 of 914) were chlamydia-positive, and of those who reported that their partners always used condoms, 8.4 percent (117 of 2115) were chlamydia-positive.

Of the 823 nonvolunteer recruits who filled out a questionnaire anonymously, 203 (24.7 percent) did not provide their ages and were dropped from the analysis. The mean age of the remaining nonvolunteer recruits was 21 years (range, 17 to 36); 51.3 percent were white, and 31.9 percent were black. The mean age and the racial distribution of these recruits were not significantly different from those of the volunteers. Only 66.9 percent reported having had vaginal sex, as compared with 93.1 percent of the volunteers (P<0.001). This group differed significantly from the volunteers in four variables, even after adjustment for whether the women reported having had vaginal sex: only 4.0 percent reported prior chlamydial infections (P=0.013), 18.2 percent had had a new sex partner in the previous 90 days (P=0.002), 20.1 percent had partners who consistently used condoms (P<0.001), and 90.7 percent reported no previous diagnosis of a sexually transmitted disease (P=0.001).

Of the nonvolunteers, 17.7 percent had had more than one sex partner in the previous 90 days; the proportion of the volunteers who had had more than

<p>| TABLE 1. CHARACTERISTICS OF 13,204 FEMALE ARMY RECRUITS SCREENED FOR CHLAMYDIA TRACHOMATIS. |
|---------------------------------------------|---------------------------------|-----------------|</p>
<table>
<thead>
<tr>
<th>CHARACTERISTIC</th>
<th>VALUE</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Age — yr</td>
<td>21</td>
<td>Median</td>
</tr>
<tr>
<td>Range</td>
<td>17-39</td>
<td></td>
</tr>
<tr>
<td>Race — no. (%)*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>6,715 (51.0)</td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>4,733 (35.9)</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>1,726 (13.1)</td>
<td></td>
</tr>
<tr>
<td>Ever had vaginal sex — no. (%)†</td>
<td>12,281 (93.1)</td>
<td></td>
</tr>
<tr>
<td>Sexual history in previous 90 days — no. (%)‡</td>
<td>3,478 (26.7)</td>
<td></td>
</tr>
<tr>
<td>More than one partner§</td>
<td>4,076 (31.4)</td>
<td></td>
</tr>
<tr>
<td>New partner‖</td>
<td>2,115 (16.9)</td>
<td></td>
</tr>
<tr>
<td>Partner always used condoms¶</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Previous diagnosis of sexually transmitted disease — no. (%)</td>
<td>1,206 (9.1)</td>
<td></td>
</tr>
<tr>
<td>Chlamydia trachomatis</td>
<td>430 (3.3)</td>
<td></td>
</tr>
<tr>
<td>Neisseria gonorrhoeae</td>
<td>74 (0.6)</td>
<td></td>
</tr>
<tr>
<td>Syphilis</td>
<td>61 (0.6)</td>
<td></td>
</tr>
<tr>
<td>Trichomona</td>
<td>11,272 (86.1)</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>1,219 (9.2)</td>
<td></td>
</tr>
</tbody>
</table>

*Data were missing for 30 subjects.  †Data were missing for 9 subjects.  ‡For 225 subjects, data were missing or subject did not know answer.  §For 168 subjects, data were missing or subject did not know answer.  ¶For 684 subjects, data were missing or subject did not know answer.
Prevalence of Infection

The age-specific prevalence of *C. trachomatis* infection among the 13,204 volunteers is shown in Figure 1. The highest prevalence of chlamydial infection (12.2 percent) was among 17-year-olds. The prevalence declined sharply with increasing age, to below 5 percent for women over 25 years of age. For further analysis, the youngest age groups (17 to 25 years) were combined into a category called "young." The prevalence in this group was 10.0 percent (11,622 of 116,031). In the older-age category (26 to 39 years), the prevalence was 3.6 percent (570 of 16,010). The prevalence was 5.5 percent (369 of 6715) for whites, 14.9 percent (707 of 4733) for blacks, and 8.1 percent (143 of 1756) for other races.

Univariate Analysis

Univariate analysis identified 10 variables significantly associated with chlamydial infection: young age (17 to 25 years), black race, race other than white or black, ever having had vaginal sex, having had more than one sex partner in the previous 90 days, having had a new sex partner in the previous 90 days, having had a partner who did not always use condoms in the previous 90 days, and a history of any sexually transmitted disease (Table 2). A prior diagnosis of chlamydia or syphilis was not significantly associated with being positive for chlamydial infection.

Multivariate Analysis

In the complete multivariate model, having had vaginal sex, an age of 25 years or less, black race, having had more than one sex partner in the previous 90 days, having had a new sex partner in the previous 90 days, having had a partner who did not always use condoms in the previous 90 days, and a history of any sexually transmitted disease were independent predictors of chlamydial infection (Table 3).

Strategies for Selective Screening

A screening strategy involving all variables identified as independent predictors would require that 100 percent of the population be tested and would detect 100 percent of the positive subjects. In this model, the magnitude of risk associated with having had a new sex partner might vary according to race. For the purpose of a screening program, this would not alter the proportion of the population tested or the percentage of positive subjects detected with this model. Because screening on the basis of race would probably be viewed as inequitable, a strategy excluding race was examined. According to this strategy,

![Figure 1. Mean (±SE) Age-Specific Prevalence of Chlamydial Infection among 13,204 Female Army Recruits, According to Ligase-Chain-Reaction Assays of Urine Specimens.](image-url)
**Table 3. Multivariate Analysis of Factors Independently Associated with Chlamydial Infection in Female Army Recruits.**

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Odds Ratio (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age ≤25 yr</td>
<td>3.0 (2.3-4.0)</td>
</tr>
<tr>
<td>Black race†</td>
<td>1.4 (2.9-3.8)</td>
</tr>
<tr>
<td>Other (nonwhite, nonblack) race†</td>
<td>1.7 (1.4-2.1)</td>
</tr>
<tr>
<td>Having ever had vaginal sex</td>
<td>5.9 (3.2-10.6)</td>
</tr>
<tr>
<td>Having had &gt;1 partner in previous 90 days‡</td>
<td>1.4 (1.2-1.7)</td>
</tr>
<tr>
<td>Having had a new partner in previous 90 days§</td>
<td>1.3 (1.1-1.6)</td>
</tr>
<tr>
<td>Having had a partner who did not always use condoms in previous 90 days¶</td>
<td>1.4 (1.1-1.6)</td>
</tr>
<tr>
<td>Having ever had a sexually transmitted disease</td>
<td>1.2 (1.0-1.4)</td>
</tr>
</tbody>
</table>

*CI denotes confidence interval.
†The reference group consisted of the white subjects.
‡The reference group consisted of the subjects who did not have >1 partner, who answered that they did not know, or for whom data were missing.
§The reference group consisted of the subjects who did not have a new partner, who answered that they did not know, or for whom data were missing.
¶The reference group consisted of the subjects who did not have a partner who always used condoms.

**DISCUSSION**

Although the diagnosis and treatment of sexually transmitted diseases has always presented a challenge, there has been no routine screening of recruits for chlamydial infections at entry into the U.S. Army. Because most chlamydial infections are asymptomatic in women and because the sequelae of disease present a severe and costly burden, screening women at entry into the Army is an appropriate way to identify infections early and to explore opportunities for a control program.

Most of these control programs have used diagnostic assays that require pelvic examinations and cervical specimens. However, it has recently been shown that testing urine specimens by DNA-amplification techniques is cost effective for screening large numbers of persons in different settings. We used this new technique to determine the prevalence of chlamydia and to identify screening criteria for a program to control chlamydia in the military.

There was considerable variation in the prevalence of chlamydial infection according to the state of origin of the recruits (F<0.001 by one-way analysis of variance). The prevalence was more than 15 percent for recruits from South Carolina, Georgia, Alabama, Louisiana, and Mississippi. For New Jersey, North Carolina, Kentucky, Texas, Oklahoma, and Arkansas, the prevalence was 10 to 15 percent, and for 17 other states and Puerto Rico, it was 5 to 10 percent. For five states (Washington, Oregon, Minnesota, Arizona, and Massachusetts), the prevalence was less than 5 percent. Fewer than 100 recruits were tested from each of 17 states, 3 territories, and the District of Columbia, and prevalence figures from these areas were therefore not included in the analysis. The prevalence for the five states with the highest prevalence and the five states with the lowest prevalence differed significantly, since the 95 percent confidence intervals for prevalence did not overlap.

Using the ligase chain reaction with urine samples, we found a high prevalence of C. trachomatis infection (9.2 percent). This prevalence was higher than that observed in family-planning clinics but not as high as that reported in some adolescent health clinics. Our data agree with those from previous studies of chlamydial infections in Army women, in which prevalence rates ranged from 8.2 percent to 9.8 percent. In one large, community-based screening study, the overall prevalence of chlamydia in young women was 8.6 percent, as detected by the urine ligase chain reaction, a prevalence similar to that found in our study. Because our population was not clinic-based and was not made up of women seeking health care, the finding of such a high prevalence in these women warrants the institution of a control program for the routine identification and treatment of chlamydial infections in order to prevent sequelae and transmission to sex partners.

The study population consisted of a young, sexually active group of female recruits with sexual risk factors known to be associated with chlamydial infection. Although 9.1 percent of the subjects reported having had chlamydial infection in the past, this factor was not associated with the risk of current infection. The highest prevalence was observed among 17-year-olds. This prevalence is similar to comparable age-specific rates in other studies, confirming that young age is associated with chlamydial infection. In our study, young age was associated with being chlamydia-positive in both univariate and multivariate analyses (odds ratio, 3.0). In order to include more positive subjects, we used an age cutoff of 25 years, which allowed the detection of 95.3 per-
per cent of the chlamydial infections. Other studies have supported age-based screening for chlamydia.27,31,32
Thus, for this group of female recruits coming from a civilian background, who were tested within three days of starting basic training, young age alone can be recommended as a single indicator of who should be tested for chlamydial infection. Other models considered in this study offered high sensitivity, but the models were more complex and required valid sexual-risk histories. We documented 13 chlamydial infections (prevalence, 1.4 percent) among 914 recruits who denied being sexually active, as well as chlamydial infections in 8.4 percent of those who reported that their partners consistently used condoms. These figures indicate that self-reported sexual-risk histories are not always valid.33 The lower prevalence of chlamydial infection among recruits for whom the data on condom use were missing, or who indicated on their questionnaires that they did not know whether their partners always used condoms, may be due to lack of sexual activity, because 58.6 percent of the 684 recruits in these categories reported that they had never had vaginal sex. There is a fixed laboratory budget available for population screening in the Army. Young age is the simplest, least expensive, and most easily documented risk factor on which to base a recommendation for a screening program, as well as being highly sensitive. Alternatively, since the use of age as a selective screening criterion would have missed 4.7 percent of the infections, universal screening might be more cost effective from a societal perspective, and future studies of cost effectiveness are warranted.28
This was one of the largest programs for screening young, sexually active subjects that was not clinic-based and whose results were derived from urine DNA-amplification assays. The geographic variation in prevalence was striking. From more than 15 percent in the five states with the highest prevalence to less than 5 percent in the five states with the lowest, these differences may reflect the levels of disease burden in certain states. These regional variations also appear to reflect regional differences in chlamydial disease, as reported by the Centers for Disease Control and Prevention.34,35 For example, the prevalence in North Carolina reportedly varied from 10 percent to 17 percent.36 The prevalence is lower in regions such as Wisconsin and Washington State, where clinic-based chlamydia-control programs are in place and where declining rates of prevalence of chlamydia have been reported.26,27,32,26 In our study, the prevalence was 11.3 percent for North Carolina and 3.8 percent for Washington State. Our data imply that chlamydial infection remains common in young women across the United States. With a volunteer rate of 80 percent among women who were approached and representation from 50 states and 4 territories, our study had a wide geographic sampling.

One limitation of our study is that it is not known whether the prevalence of risk factors for chlamydial infection differs between young women who decide to join the military and those who do not. However, the demographic and sexual risk-factor characteristics of our subjects appear to be similar to those of other regional and clinic-based populations,26 as well as those from a large, community-based study.31 An additional limitation is that the nonvolunteers in our study differed from the volunteers with regard to sexual risk factors for chlamydia. However, the nonvolunteers represented a group who were mostly sexually active, who had had new sex partners in the previous 90 days, and whose partners did not use condoms. Thus, their risk of chlamydial infection may have been as high as that of the subjects in our study.

Although amplified-DNA tests are more expensive than traditional nonculture tests, the savings associated with not having to have a clinician collect specimens from a pelvic examination and the advantages of being able to use urine as a diagnostic specimen may outweigh the extra cost of the test.29 In addition, it has been demonstrated that amplified-DNA testing of urine specimens is cost effective, and treating chlamydial infections prevents serious complications such as pelvic inflammatory disease, ectopic pregnancy, and infertility.15,38,37

In conclusion, our study indicates that with the limited funding available at present, young age (25 years or less) would be the best criterion on which to base a screening program using amplified-DNA testing of urine for female Army recruits and perhaps for other young women. Institution of such a control program has the potential to reduce drastically the burden of chlamydial disease in the U.S. Army and to prevent morbidity due to these infections.37

Supported by a grant (DAMD17-95-1-5064) from the Department of the Army.
The opinions expressed in this article are those of the authors and are not necessarily endorsed by the U.S. Army or the Department of Defense.

We are indebted to Dr. Daniel Scharfstein, Mary Sheppard, Dr. Guillermo Madico, and Dr. Anne Rampel for statistical assistance; to Kathryn Kacena for data entry; to Diane Perkins, Sandra Leiser, Dier Whim, and Graciela Jachek for performance of the laboratory tests; to Lt. Col. Cynthia Dodd, Lt. Col. (ret.) Delois Daniel, and Kimberly Riley for treatment of infected women and processing of specimens; and to Patricia Huit for assistance in the preparation of the manuscript.

REFERENCES
9.2 Abstracts presented at conferences during the project's third year

9.2.1 International Symposium of Sexually Transmitted Disease Research, Seville 1997.


9.2.1.2 Stability of *C. trachomatis* and *N. gonorrhoeae* in urine at room temperature for LCR and PCR tests.


CHLAMYDIA TRACHOMATIS INFECTIONS IN SYMPTOMATIC AND ASYMPTOMATIC COUPLES

Petersen EE, Claud A, Univ. Frauenklinik Freiburg, D-79106 Freiburg, Germany

Objective: Normally, only symptomatic patients are tested for chlamydia infections. Little is known about partner infection in asymptomatic couples. First void urines of symptomatic (STD outpatient clinic) and asymptomatic females and their asymptomatic partners (genital chlamydial prevalence study with LCR on 4,381 persons, Germany) were collected to assess the prevalence of Chlamydia trachomatis in couples.

Methods: Urines and cervical swabs were tested with the ligase chain reaction (LCR, Abbott). In the prevalence study only one urine from each woman and her male partner was collected. In symptomatic women cervical swabs and urine and partner urines were tested and retested at follow-up visits.

Results: Only couples with one or both partners positive in the chlamydial LCR are presented here. Of 53 symptomatic women 37 (69%) were positive. 39 (72%) of the asymptomatic male partners had positive urines. In 21 (38%) couples both partners were positive. In 84 asymptomatic couples 46 (55%) of the women and 67 (80%) of their male partners were positive. In 29 (36%) couples both partners were positive.

Conclusion: Asymptomatic women significantly more chlamydia infections could be detected when also testing their male partners. Testing of one male (urine) and one female sample (urine or cervical swab) by LCR appears to be more cost-efficient than testing several female samples.

COST-EFFECTIVENESS OF SCREENING vs MASS THERAPY FOR C. TRACHOMATIS IN FEMALE ARMY RECRUTS

Howell MD,1 McKee K,2 Ellis D,1 Gaydos JP,1 Hendrix R,1 Quinn TC1, Gaydos CA1
1The Johns Hopkins University, Baltimore, MD; 2Fort Bragg, NC; 3Fort Jackson, SC; 4CHLIM, Aberdeen Proving Ground, MD; 5HAID, NIH, Bethesda, MD.

Objective: In US Army women C. trachomatis (CT) may cause a significant degree of morbidity. We sought to assess the relative cost-effectiveness of three screening and treatment strategies for CT in a military setting.

Methods: We compared universal and targeted screening to mass therapy with azithromycin for CT in female recruits using a cost-effectiveness analysis. At Fort Jackson, SC, 7,191 recruits presenting for basic training from 1/96-3/97 were randomized into one of three groups: (1) no intervention (No Intervention), (2) universal urine LCR screening, and (3) targeted CT screening. The above results are clinically significant, when considering the low prevalence of C. trachomatis IgG & IgA antibodies observed in the three control groups, blood donors (8%, 4.4%), children (3%, 0%) and pregnant women (6%, 0%), respectively.

The high specificity and sensitivity of SeroCT for the specific detection of C. trachomatis antibodies makes this a new generation test, an accurate efficient and cost effective screening tool for the differential diagnosis between C. trachomatis and C. pneumoniae infections.

<table>
<thead>
<tr>
<th>Strategy</th>
<th>Projected Costs (US$1995)</th>
<th>Projected PID (%) (silent &amp; asymptomatic)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Program</td>
<td>Sequelae</td>
</tr>
<tr>
<td>No Intervention</td>
<td>$973,100</td>
<td>316</td>
</tr>
<tr>
<td>Screen _25 yrs &amp; Treat (+)</td>
<td>$120,600</td>
<td>$737,000</td>
</tr>
<tr>
<td>Screen All &amp; Treat (+)</td>
<td>$137,900</td>
<td>$236,400</td>
</tr>
<tr>
<td>Mass Therapy</td>
<td>$207,400</td>
<td>$138,600</td>
</tr>
</tbody>
</table>

Conclusions: The cost-effectiveness of universal screening for CT in female recruits is comparable to mass therapy, with a 10% increase in the cost of universal screening. The use of universal CT screening may be a cost-effective strategy for CT prevention in female recruits.
in the Netherlands, the municipal policy of toleration and decriminalisation towards prostitution has led to a debate over its regulation and legislation. Organised prostitution in the Netherlands is forbidden, while in practice brothels are allowed to operate with varying amounts of freedom. This semi-

In the Netherlands, the municipal policy of toleration and decriminalisation allows local authorities to impose punishment for operating infractions. In anticipation of a change in the national law, the Municipality of Amsterdam developed their own new regulation. Since Jan 96, it is illegal for women from outside the EC to work as prostitute and, brothel owners, can get a license the majority of the time (hygiene, fire, no engagement of minors).

In Amsterdam, in the years 1994, 1995, 1996, transvestites working in windows were mainly Latin American women. As research revealed that they and their clients were at increased risk of both STD and HIV, public health nurses (together with cultural mediators) frequently visited these women in order to give information on safe sex, HIV/STD, birth control and tuberculosis and to inform them about our STD clinic. In 1995 first contacts were made with 969 prostitutes, including 569 Latin American prostitutes, 73 African, 134 Eastern European and 58 Asian women. In 1996 the number of contacts decreased considerably (a =651), despite a similar fieldwork.

Where did the women disappear to? The consequences of this new regulation will be discussed with regard to accessibility of the women and the segregation of prostitution networks, including the increased risk for STD/HIV transmission.

**O174 THE USE OF LIGASE CHAIN REACTION (LCR) IN THE DIAGNOSIS OF GONOCOCCAL AND CHLAMYDIAL EPIDIDYMITIS.**

Feher HG, Raderc F, Dangor Y, Grobelnaa T, Ballard RC.
National Reference Centre for STDs, SA Institute for Medical Research, and University of the Witwatersrand, Johannesburg, South Africa.

**Objective:** A study was conducted to determine the value of the ligase chain reaction (LCR) technique as a diagnostic tool in cases of epididymitis.

**Methods:** Ninety-five consecutive cases of acute epididymitis among migrant mineworkers attending an STD clinic at the Leslie Williams' Memorial Hospital, Carletonville, were included in the study. In each case, endocervical swabs were collected for gonococcal and chlamydial cultures. A first catch urine sample was collected for LCR for both Neisseria gonorrhoeae (GC-LCR) and Chlamydia trachomatis (CT-LCR). In addition, blood specimens were tested for syphilis, chlamydial and HIV antibodies.

**Results:** Overall, either by culture or LCR, 43 patients (45%) had evidence of gonococcal infection, while 25 patients (26%) had proven chlamydial infection. A further 23 cases (24%) had high levels of antisiphilis antibody of ≥ 1:64 using the indirect microimmunofluorescence test (micro-IF). Of the 25 cases with proven chlamydial infection, 13 cases also had concomitant gonococcal infection. Among the cases with endocervical swabs, 38% were detected by GC-LCR, while only 6% were culture-positive. Similarly, using CT-LCR, 88% of chlamydial infections were detected, while only 40% were culture-positive. In 23 cases (34%) no evidence of infection with either organism was obtained. Reactive syphilis serology was found in 12% of cases and 96% were HIV-negative.

**Conclusion:** LCR proved to be more sensitive than culture for the diagnosis of epididymitis. Chlamydial micro-IF serology proved a useful adjunct to the diagnosis of chlamydial infection. In approximately a quarter of all cases no aetiology could be determined.

**O176 CHLAMYDIAL DIAGNOSIS WITH THE LCR, COBAS AMPLICOR, AND TRANSCRIPTION MEDIATED AMPLIFICATION USING VULVAL SAMPLES.**

Study A: Hartmanns B, Schubat E, Kershbeuerm M, Outpatient's Centre for Infectious Venereal-Dermatological Diseases, Vienna, Austria.

**Objective:** To evaluate whether vulval smears may serve as an alternative non-invasive specimen type for chlamydial diagnosis, the performance of all commercially available amplification tests, COBAS AMPLICOR (Giga-Pros) Transcription-Mediated Amplification (TMA) with vulval specimens as well as with urine, and cervical specimens was compared with culture on endocervical and vulval specimens.

**Methods:** The study was performed in female patients attending the Outpatient's Centre mainly because of a suspected genitinal tract infection. Partner control, contact tracing, and health check up. In addition to a first void urine (FVU), samples were obtained from the vulval region and endocervix and tested by culture and amplification tests.

**Results:** The sensitivity (Table) and specificity for all sampling sites and amplification methods was high, calculated on infected women as the goldstandard established by positive culture or different test procedures.

**Conclusion:** The results demonstrate that in contrast to culture, all amplification methods performed with a high sensitivity with vulval smears (89.6%), comparable with urine (86.6%) and cervical specimens (90.6%). The data indicate that vulval smears can be used as alternative non-invasive specimens useful for chlamydial screening programs.

**O177 STABILITY OF C. TRACHOMATIS AND N. GONORRHOEAE IN URINE AT ROOM TEMPERATURE FOR LCR AND PCR TESTS.**

Gayaia CA', Welsh L', Krielman S', Perkins D', Schmidt K', Chou J', Qinbo T,15 The Johns Hopkins Univ., Baltimore, MD; NIAID, NIH, Bethesda, MD.

**Objective:** To determine the stability of concentrations of C. trachomatis and N. gonorrhoeae in urine at room temperature (25°C) for testing by LCR and PCR.

**Methods:** Aliquots of urine concentrations of CT from 100 to 1 IFU/ml and of NG from 100 to 1 CFU/ml were sequentially processed and tested by LCR and PCR at times 0, 24 hr, 48 hr, 72 hr, 7 days, and 14 days.

**Results:** CT/LCR: All concentrations of CT from 100 to 1 IFU were stable in urine, giving positive results, until 14 days at 25°C. CT/PCR: At 0 hr, concentrations from 100 to 1 IFU were positive. One IFU was negative. At 48 hr, 72 hr, and 7 days, concentrations from 100 to 5 IFU were positive. At 7 days, only 10 IFU were positive. NG/LCR: At 0 and 24 hr, 48 hr, and 72 hr, concentrations from 100 to 50 CFU were positive. At 7 days, only 10 and 100 IFU were positive. At 48 and 72 hr, only 10 IFU were positive. At 7 days, only 10 IFU were positive. NG/PCR: At 0 and 24 hr, all samples were positive. At 7 days, only 10 IFU were positive. At 48 and 72 hr, only 10 IFU were positive. At 7 days, only 10 IFU were positive.

**Conclusion:** Stability of C. trachomatis and N. gonorrhoeae in urine at room temperature for LCR and PCR tests.
Single-dose Azithromycin (AZ) for Mass Therapy to Control Chlamydia trachomatis (CT) in Female Army Recruits: a cost-effectiveness model

HOWELL MR, QUINN TC, HENDRIX R, MCKEE K, GAYDOS J, GAYDOS CA

The Johns Hopkins Univ., Baltimore, MD; NIAID, NIH, Bethesda, MD; Fort Jackson, SC; Fort Bragg, NC; Jackson Foundation, Rockville, MD.

Objective: There is a high prevalence of C. trachomatis (CT) in female U.S. Army recruits, which may cause serious morbidity due to PID. We evaluated the cost-effectiveness of treating all recruits with AZ to prevent chlamydial sequelae compared to testing all recruits and treating only those with positive CT tests.

Methods: In a decision analysis, we modeled mass treatment and universal testing to determine the incremental cost-effectiveness of these two strategies. From a military perspective, we calculated the total program costs, the projected costs of lost training, the projected costs of sequelae due to CT, the projected level of prevented disease (PID, chronic pelvic pain, and ectopic pregnancy) and the projected number of military discharges for medical conditions. 9,192 recruits presenting for basic training at Ft Jackson SC 1996-97 were tested by urine LCR to determine CT prevalence. Results were extrapolated to a hypothetical population of 10,000 recruits per annual cohort.

Results: The recruit sample had a CT prevalence of 9.0%. 218 cases of PID would be expected in the absence of a control program. Mass therapy would prevent 187 cases and save $39,500 over universal testing for a 5-yr follow-up period and $357,300 over no program. At CT prevalence > 6.2% and < 14 clinic visits per 100 individuals for moderate to severe side-effects, mass therapy provided a cost-savings over universal testing. Both strategies provided cost-savings over no program at CT prevalences over 3%.

Conclusion: Mass therapy would prevent sequelae, would likely decrease the number of military discharges for medical reasons in the first 6 mo. of service, and would save overall costs relative to universal testing. Mass therapy could be considered as a first line of control for chlamydia in a well-defined cohort of young women with a high prevalence of CT.
dom use was noted in persons who received more recent positive test results than among persons with positive test results from prior years (p<0.001), though this trend was not noted in heterosexual men. The proportion of persons who injected drugs (IDU) increased with more recent years of first positive HIV test (p<0.05), especially among women. There were slight decreases in the proportion of persons who traded sex for money or drugs, and who used cocaine, with more recent years of first positive HIV test. But the proportion using crack increased.

Conclusions: High-risk practices, such as sex without condoms and IDU, have decreased among MSWM and women. Heterosexuals are emerging as more prominent group for HIV infection.

P-12.3 The Structure of a Newly Identified Genetic Subtype in the HIV-1 Epidemic in Africa. J.K.CARR, F.E. MCCUTCHEON, M. EMERSON, D.BIRX. Henry M. Jackson Foundation for the Advancement of Military Medicine, Division of Retrovirology, Walter Reed Army Institute of Research, Rockville, MD.

Objective: Genetic analysis of HIV-1 genomes has led to the description of at least 10 different genetic subtypes, each with its own geographic or demographic distribution. Although subtype A is one of the most common genetic subtypes of HIV-1 in Africa, only three full-length genomes of this subtype have been sequenced: two from Uganda and one from Nigeria. Methods: Two isolates, DJ263 and DJ264, characterized as subtype A by gag and env sequencing, have been cloned and sequenced in full for the first time. The entire genome was sequenced except for a 73 nt region in theLTR. These subtype A’ viruses were from Djibouti, on the coast of East Africa. Phylogenetic analysis of the full-length genome was performed with these new isolates. Results: The Djibouti viruses were found to form a unique cluster in the full-length analysis, grouping with previously sequenced isolate, IbNG, from Nigeria. They were distinct from subtype A and similar to each other from the 5’ to the 3’ end. Detailed analysis of the subtype structure of these isolates in comparison with reference sequences of the main HIV-1 subtypes, revealed them to be complex recombinants between subtypes A, G and an original "IbNG" parental strain. Both Djibouti isolates and the Nigerian isolate had the same complex subtype structure.

Conclusions: Full-length genomes from one of the common subtypes in Africa will be fully characterized in this report. Although related to subtype A in both gag and env, the isolates were found to be a homogeneous viral subtype of their own, with a complex subtype structure recombinant between two known subtypes, subtypes A and G, and one unknown subtype, the putative parental subtype. It is recommended that this viral subtype be given the new subtype designation of "IbNG*". Like the subtype E* viruses, which are E/A recombinants, this subtype has a segment from an original parental virus, a virus which has never been found in its 'pure', or non-recombinant, state.

P-12.4 A Silent Epidemic of Chlamydia trachomatis Genital Disease in Female U.S. Army Recruits. CA GAYDOS, MR HOWELL, KL CLARK, KT MCKEE, RM HENDRICKS, D ELLIS, TC QUINN, JC GAYDOS. Johns Hopkins Univ, Baltimore, MD; Walter Reed Army Inst Res, Washington, DC; Ft. Bragg, NC; Ft Jackson, SC; NAID, NIH, Bethesda, MD; HM Jackson Fdn, Rockville, MD.

Background: C. trachomatis (CT) infections in the U.S. exceed 4 million cases annually, occur mostly in the young and lead to costly sequelae (PID, ectopic pregnancy, and infertility). Most (>80%) infections are asymptomatic in women, leading to epidemic rates in sexually active youth. Our hypothesis was that a large epidemic of treatable CT genital infections, related to the ongoing civilian epidemic, was occurring in female recruits. Methods: Ligase Chain Reaction (LCR) of urine specimens was used to determine the prevalence of CT infections in female military recruits (N =9,192) at induction. Demographic and sexual risk data were collected. Results: Overall prevalence was 9.0%. By questionnaire, 93.6% reported vaginal sex, 26% had >1 sex partner and 30.5% had a new partner. Only 15.5% always used condoms. By age, prevalences were: 11.0% (17-20 yr), 8.0% (21-25 yr), 3.1% (26-30 yr) 1.9% (31-35 yr). By home state, 4 states had a prevalence >15%, 8 states 10-15%, and 12 states 5-10%. By multivariate analysis, significant variables associated with CT positivity were vaginal sex (OR 4.1), age (17-25 yr age (OR 3.4), African American (OR 2.7), >1 partner (OR 1.4), and a new partner (OR 1.5). A model consisting of LCR screening young female recruits (17-25 yr) would test 87.2% of the recruit population and identify 95.8% of CT positives. Conclusions: A high prevalence of CT infection exists in female army recruits, especially those of young age. Differences by state may be representative of national, geographically diverse prevalences. Detection of infection by LCR of urine provided a large, efficient screening program. Data collected are being used to develop a cost-effective treatment program.

P-12.5 Epidemiology of Shigellosis in San Francisco during the HIV Era. J. T. BAER, D. J. VUGIA, A. L. REINGOLD, T. ARAGON, F. J. ANGULO, W. Z. BRADFORD, for the CA Emerging Infections Program, CA Emerging Infections Program, CA Department of Health Services, UC-Berkeley School of Public Health, Berkeley, CA; Centers for Disease Control and Prevention (CDC), Atlanta, GA; San Francisco Department of Public Health, UC-San Francisco.

Objective: The impact of HIV infection on the epidemiology of Shigellosis and its potential role as a risk factor have not been well described. To better understand this relationship and to assess the use of gay sex, a recognized risk factor for Shigellosis, we conducted an investigation of cases of Shigellosis in San Francisco, a city with a high prevalence of both HIV infection and men engaging in gay sex.

Methods: As part of CDC's Emerging Infection Program, active surveillance for infections caused by Shigella species was conducted in SF during 1996. All available medical records were reviewed using a standardized data collection instrument and data previously collected by the SF Department of Public Health (DPH) during routine interviews of cases were obtained. The estimated prevalence of HIV infection and men engaging in gay sex was obtained from the SF DPH. Results: A total of 228 culture-confirmed cases were identified, including 142 and 73 caused by S. sonnei and S. flexneri, respectively. The incidence rate in cases per 100,000 population was 31.5, compared with an active surveillance rate of 10.9 in neighboring Alameda County (AC), and a reported rate of 7.3 in the U.S. The incidence rate in the 25-44 year age group was 46.0 cases per 100,000 population, compared with 6.8 cases per 100,000 population in AC, and the rate in HIV-infected persons was 442 cases per 100,000 population. Adult cases (age > 17 years) comprised 80% (181/228) of the total, and had the following characteristics: male gender (75%), white race (70%), gay male (65%), HIV-infected (52%), and sexually active in the 10 days prior to interview (69%). For one percent (96/190) of infections occurred in gay, adult men. When compared to the non-gay and non-HIV infected population, the incidence rate ratios for the gay and non-HIV infected, the non-gay and HIV-infected, and the gay and HIV-infected populations were 9.03, 4.80, and 8.00, respectively. The incidence rate ratio for the gay and HIV-infected population was 6.9 (95% CI 4.2-11.3). Conclusion: These population-based data demonstrate high overall rates of Shigellosis and dramatically elevated rates in HIV-infected persons. The high prevalence of HIV infection and men engaging in gay sex are likely important determinants of the Shigellosis rates in SF. The relative contribution of behavioral factors as compared to compromised host immunity in HIV-infected individuals warrants future investigation.
9.3 Abstracts submitted during the project’s third year for conferences


North-South-East-West: Geographic, Race and Age Correlates in Non-Health Care Seeking U.S. Female Army Recruits

MR Howell¹, JC Gaydos²⁵, KT McKee⁴, K Clark³, TC Quinn¹⁵, CA Gaydos¹
¹ Johns Hopkins University, Baltimore, MD; ² HM Jackson Foundation, Rockville, MD; ³ Walter Reed Institute of Research, Washington, DC; ⁴ Fort Bragg, NC; ⁵ NIAID, NIH, Bethesda, MD.

Background/Rationale: Regional variations in C. trachomatis prevalence have been documented in clinic populations. Similar regional variations in non-health care seeking women have not been examined to determine the influence of confounding variables such as demographics. It is necessary to understand differences in populations from non-traditional settings from a diverse geographic perspective.

Objectives: To describe and assess the significance of regional variations and determine if these differences remain when controlled for race and age in a non-health care seeking military population.

Methods: From 1/96 through 12/97, we tested 13,204 female recruits by ligase chain reaction of urine for C. trachomatis infection upon entry to the U.S. Army. The impact of regional variations (CDC reporting areas: northeast, south, midwest, west and territories) on chlamydia prevalence among these women was analyzed by ANOVA. The effect of intervening factors (self-reported race and age) was assessed by logistic regression.

Results: An overall prevalence of 9.2% (1,219/13,204) was observed. Home state records were available for 13,152. Women coming from the northeast had a chlamydia prevalence of 6.9% (146/2103); the south 11.9% (760/6381); the midwest 7.1% (167/2351); the west 5.6% (117/2100); and the territories 10.0% (21/211), (F<0.001). Region, African American race and young age (age < 25) were all independent predictors of infection. Women coming from the southern states were more likely (O.R. 1.5, 95% CI: 1.3-1.7) to be positive for chlamydia than women coming from the other four regions. Conversely women in the west were less likely (O.R. 1.4, 95% CI: 1.1-1.7) to have a positive test than other women.

Conclusions: In non-health care seeking U.S. Army recruits regional disparities in chlamydia prevalence exist. Other national studies in non-health care seeking populations are necessary.

Learning Objectives:
1. Using the results of this analysis participants will be able to describe the regional variations in a non-health care seeking population.

2. Using the results of this analysis participants will be able to describe the association between race, age, region and infection with C. trachomatis.
Infectious Diseases Society of America
36th Annual Meeting

Deadline: August 5, 1998

Prevalence and Risk Factors of C. trachomatis Infection in Male Military Recruits Using Urine Ligase Chain Reaction (LCR)

JA CECIL1, R HOWELL1, JC GAYDOS2, KT MCKEE3, D ELLIS4, RM HENDRIX4, TC QUINN5, CA GAYDOS1. Johns Hopkins University1; HM Jackson Foundation, Rockville, MD2; Ft. Bragg, NC3; Ft. Jackson, SC4; NIH, NIAID, Bethesda, MD5.

Chlamydia trachomatis urogenital infections in women are associated with significant morbidity, including PID, ectopic pregnancy and infertility. In a study of female Army recruits using urine ligase chain reaction (LCR, Abbott Laboratories, Abbott Park, IL), C. trachomatis was identified in 9.3% of women screened. The effectiveness of efforts to limit chlamydial infections in women may depend on the prevention of re-infection through simultaneous screening and treatment of men. We tested male recruits for C. trachomatis using urine LCR to determine prevalence and to assess potential screening criteria for C. trachomatis infection in men. Each recruit who volunteered to participate provided a urine sample and answered a questionnaire. Among 1,203 men screened, the prevalence of infection was 4.9%. The mean age was 19.8 (±2.61) years, 61.6% were Caucasian, 33.8% had more than one sex partner, and 36.7% had a new sex partner in the last 90 days. Only 21.2% of men reported using condoms regularly, and 2.6% reported having prior chlamydial infections. Of the men that tested positive, only 13.6% reported having symptoms. Risk factors that proved useful for predicting chlamydial positivity included: African-American race, more than one sex partner, and a new sex partner. Young age was not a risk factor, as it was for women. Urine LCR is a convenient and well-accepted method for screening large numbers of men, and may be useful in developing strategies for limiting the spread of C. trachomatis in the population.
Abstract:

Prevalence and Risk Factors of *C. Trachomatis* Infection in Male Military Recruits: Multivariate Analysis

JA Cecil, MD; MR Howell, Johns Hopkins University; JC Gaydos, MD, MPH, HC Jackson Foundation; KT McKee, MD, MPH, Ft. Bragg, NC; RM Hendrix, D.O., Ft. Jackson, SC; TC Quinn, MD, Johns Hopkins University, NIH, NIAID: CA Gaydos, DrPH, Johns Hopkins University

Objectives: To determine the prevalence and assess potential screening criteria for *Chlamydia trachomatis* infection in male military recruits, to help target future interventions.

Methods: 1,203 male Army recruits volunteered to provide a urine sample and completed a questionnaire addressing demographic and behavioral characteristics, during inprocessing at Ft. Jackson, SC. Urine samples were tested for the presence of *C. trachomatis* using ligase chain reaction. Multivariate logistic regression was performed to access criteria for a screening program.

Results: The volunteer rate was 80.9%. The prevalence of infection was 4.9%. The mean age was 19.8 years and 61.6% were Caucasian. Only 13.6% of infected men had symptoms. Independent risk factors predictive for being chlamydia positive included: African American race, having more than 1 partner in the last 90 days, and the presence of symptoms. Of 3.4% who were symptomatic, 19.5% were infected. Of asymptomatic men, 4.4% were infected. 3% of those reporting condom use were chlamydia positive.

Conclusion: A significant number of male Army recruits are infected with *C. trachomatis*, and the majority of these men do not have associated symptoms to suggest infection. These men may be less likely to seek appropriate medical attention, and may contribute to the transmission of chlamydia infections.

Urine LCR is a convenient, noninvasive method for screening large numbers of men, and will be used to screen more recruits as part of this ongoing study. Additional data from future screening will be presented.
9.4 Maps

9.4.1 Percentage Chlamydia Positive Females by Geographic Origin 3rd Year

9.4.2 Percentage Chlamydia Positive Males by Geographic Origin Total

9.4.3 Percentage Chlamydia Positive Females by Geographic Origin for Project Years 1-3
Chlamydia Prevalence by Public Health Region Origin of Recruits

Region 10 5.0%
Region 8 5.1%
Region 5 8.1%
Region 2 10.0%
Region 3 9.3%
Region 4 15.4%
Region 6 11.5%
Region 7 6.0%

Females only (project third year)
Chlamydia Prevalence by Public Health Region Origin of Recruits

Males only (project third year)
Chlamydia Prevalence by Public Health Region Origin of Recruits

Females only (project years 1 to 3)
9.5 Armed Forces Epidemiology Board and the Institutional Review Boards of Johns Hopkins University and Ft. Jackson (Eisenhower, Ft. Gordon)
Minutes of the Institutional Review Committee Meeting 9 April 1998

The patient was hospitalized. The patient was found to have complete heart block, systolic BP over 200mmHg, potassium level of 7.9mg/dl. She was dialyzed and a pacemaker was placed. The patient became stable and pacemaker was removed. The reporting physician felt that hyperkalemia was related to therapy with Losartan or control. The report dated 26 Jan 98 involved a 68 year old male patient who had a history of asthma and was hospitalized for asthma. The patient recovered and was discharged from the hospital. The reporter felt that the asthma was possibly related to study drug therapy. Four patients have been enrolled in this study at EAMC. There has been an adverse event at this site for ruled-out MI.

Recommendation: Full Committee Acknowledges

The Informed Consent dated 11 Sep 97 was submitted to delete the following statement, "If you are male, you should avoid fathering children by using condoms when having sexual intercourse unless you have had a vasectomy." The legal department of Merck, Inc requested the removal of this statement.

Recommendation: Approve

Human Use Committee Vote: 10 in favor, unanimous

Protocol Review:

Study of Chlamydia Trachomatis in Military Women: Prevalence, Risk Factors and a Cost Benefit Analysis of Early Diagnosis and Treatment
PI: LTC Rose M. Hendrix, MC
DDEAMC 95-17

With regard to Minutes of the Institutional Review Committee Meeting 12 March 1998, the investigator offered the following replies to questions raised by the committee: We propose that not only will the mass therapy option cure chlamydia, but a significant proportion of the additional cases of gonorrhea, and respiratory infections as supported above. The cost of hospitalized pelvic inflammatory disease and ectopic pregnancy are staggering for the military. The prevention of these sequelae diseases and their associated costs is the focus of this proposal. The results of a cost effective decision model we have performed has indicated that the most cost-effective strategy for the Army would be mass therapy, mostly because the diagnostic test is not perfect and the extra infections cured and sequelae prevented offset the extra cost of antibiotic. Anecdotal data from the current study has indicated that in over 1,000 women treated with single-dose azithromycin, there has only been one adverse event that caused a recruit to seek medical attention. It was not classified as serious. Further, a tabulation of 6 studies published from 1992-1998 substantiates a lack of significant differences for serious adverse episodes between individuals treated with azithromycin therapy compared to those receiving doxycycline. Reactions were considered mild to moderate and were noted primarily to be gastrointestinal in nature (diarrhea, nausea, etc). Additionally, use of mass therapy with azithromycin in a large trial (12,000 volunteers; 6,000 each of treatment arm and control arm) in Uganda has not reported any serious adverse events. The 6,000 volunteers have been treated four times. Confusion relating to misclassification of azithromycin as a macrolide (e.g., erythromycin) is frequent. The issue of pregnancy should be of no impact upon this protocol. Upon in-processing at Ft. Jackson, recruits are screened for possible pregnancy and pregnant women are not allowed to continue in basic training (EPTS). Azithromycin is a class B drug and is currently being given in clinical practice to pregnant patients. A modification of the consent form has been appended to clarify that there are no benefits to participating in this study for an individual who has never engaged in sexual activity.
MCHF-CI
SUBJECT: Minutes of the Institutional Review Committee Meeting 9 April 1998

It is well known in behavior research that there exists a high probability for a respondent to misrepresent herself. In this study we have documented 14 chlamydia infections in females who had denied sexual activity. Additionally, in analyzing demographic information on individuals choosing to not participate, it is clear the only significant difference between that group and those participating is in engagement in sexual activity. Any woman engaging in even a single episode of unprotected sexual activity in the past, can be at risk for asymptomatic disease or its sequelae. Although evidence is accumulating that there are no serious drug interactions, the investigators will ask recruits to list medications they are taking if they participate. This information will be carefully studied and tabulated. They think the consent form, which is the routine Army consent form DA Form 5303-R is clear on this point. However, they will instruct the civilian study nurse to reiterate that participation is entirely voluntary. The study nurse reports that when recruits talk to her afterwards they express concern to know whether they are infected since most chlamydial infections are asymptomatic in women. PID, ectopic pregnancy, and infertility are valid concerns of these women. They consider that the high volunteer rate is a reflection of the nurse's excellent teaching skills regarding the potential serious sequelae. (Probability of developing PID after chlamydial infection is 30% Howell et al). The recruits' desire to know about their health status in comparison to the ease of the collection of urine specimen.

The committee reviewed the submitted responses to its questions from the 12 March 98 meeting. Azithromycin is, in fact, an azalide antibiotic which is a subclass of macrolides structurally, but possess a very different milder side effects profile. The principal argument if favor seems to be an economic one. Based on the assumptions of the model, fewer cases of PID or asymptomatic chlamydia would result. The assumptions were challenged that a single dose could have such lasting effects. The training dollars lost could probably not be detailed that precisely. True, EPTS discharges occur but often for undiagnosed abdominal pain and not proven PID. Some are for endometriosis and other menstrual-related dysmenorrhea. Silent chlamydia would show largely during an infertility workup. The need for recycling in training as a result of PID seemed high to members who have been or are currently involved in AIT and basic training. The basis for some of those lost training dollars was questioned.

The principal reason for doing the study would be for a potential policy for future basic trainees. If women (or men) were forced to take an antibiotic and developed a severe side effect, they might have a basis for a future claim against the government despite the current rules against suit by active duty members. The trend in medicine in antibiotic usage is against most prophylactic use except in very limited settings of respiratory or unavoidable threat. Otherwise disease is tested for and then treated, especially when a relatively good test is available. It is only for the false negatives that an issue exists. Repeat testing at a later date might pick up some of these false negatives, but this is not part of the study design. The issue of emerging resistance patterns was also cited. Several committee members expressed the feeling that they would be personally offended at being expected to take a pill to treat a presumed STD. The ethical issues for humans is different than that of a swine herd. The Army does not require mass prophylaxis unless the environmental threat is unavoidable. The clinical standard is still to test and then treat.

The original protocol was an all female study, however, it would be useful to know how the prevalence of chlamydia in the female recruit population compares to that of males in the same population. What they have proposed is that they test approximately 1000 male volunteers following the current protocol. These males would be counseled and tested in single gender groups by the same non-military protocol nurse we are presently using for the females. Those individuals testing positive would be referred for treatment, as are the females currently. Once male volunteers have been tested, they would go back to the original protocol and test only females for the remainder of the study.
9 April 1998

SUBJECT: Minutes of the Institutional Review Committee Meeting 9 April 1998

Recommendation: Approve the male prevalence testing arm and correct minor errors in consent form.

Clinical Investigation Committee Vote: 10 in favor, unanimous.

Human Use Committee Vote: 10 in favor, unanimous.

Recommendation: Disapprove the mass treatment arm.

Clinical Investigation Committee Vote: 10 in favor, unanimous.

Human Use Committee Vote: 10 in favor, unanimous.

Amendments:

A Randomized, Double-Blind, Multicenter comparison for the Efficacy and Safety of Grepafloxacin (Raxar) 400mg or 600mg Once Daily and Clarithromycin (Biaxin) 500mg Twice Daily in the Treatment of Patients with Acute Bacterial Exacerbations of Chronic Bronchitis

PI: LTC Warren L. Whitlock, MC
DDEAMC 98-17

A letter from Pharmaceutical Research Associates, Inc dated 3 Mar 98 was received regarding an amendment to this study. This amendment changes from patients with known moderate or severe hepatic or renal disease to hepatic failure or known moderate to severe renal disease. It also changes the primary efficacy population from patients who are clinically evaluable to clinically and bacteriologically evaluable. And therefore, changing the secondary efficacy population from bacteriologically evaluable to clinically evaluable. The third change in the amendment changes the package inserts for Biaxin and Raxar.

Recommendation: Approve
Human Use Committee Vote: 10 in favor, unanimous.

Periodic Review:

A Randomized, Multicenter, Third Party Blinded Trial Comparing Trovafloxacin with Amoxicillin/Clavulenate (Augmentin) with or without Erythromycin for the Treatment of Community Acquired Pneumonia

PI: LTC Warren L. Whitlock, MC
DDEAMC 97-26

There has been good response to treatments with this drug. Trovafloxacin has now been approved by the FDA to treat Community Acquired Pneumonia and we have been part of that. A total of 15 patients have been enrolled and only one serious adverse event has been reported.

Recommendation: Approve
Human Use Committee Vote: 10 in favor, unanimous.
Use of Ligase Chain Reaction with Urine versus Cervical Culture for Detection of Chlamydia trachomatis in an Asymptomatic Military Population of Pregnant and Nonpregnant Females Attending Papanicolaou Smear Clinics

CHARLOTTE A. GAYDOS, M. RENE HOWELL, THOMAS C. QUINN, JOEL C. GAYDOS,† AND KELLY T. MCKEE, JR.,* Infectious Disease Division, The Johns Hopkins University, Baltimore, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, and U.S. Army Center for Health Promotion and Preventive Medicine, Aberdeen Proving Ground, Maryland, and Preventive Medicine Service, Womack Army Medical Center, Fort Bragg, North Carolina

Received 13 November 1997/Returned for modification 27 January 1998/Accepted 16 February 1998

Ligase chain reaction (LCR) (Abbott Laboratories, Abbott Park, Ill.) with first-catch urine specimens was used to detect Chlamydia trachomatis infections in 456 asymptomatic military women attending clinics for routine Papanicolaou smear tests. Results were compared to results of cervical culture to determine the sensitivity of the urine LCR and the possible presence of inhibitors of amplification in pregnant and nonpregnant women. Discrepant results for LCR and culture were resolved by direct fluorescent antibody staining of culture sediments, two different PCR assays, and LCR for the outer membrane protein 1 gene. The prevalence of Chlamydia in specimens by urine LCR was 7.3% compared to 5% by culture. For 434 women with matching specimens, there were 11 more specimens positive by LCR than by positive by culture, of which all but one were determined to be true positives. There were four culture-positive, LCR-negative specimens, all from nonpregnant women. The sensitivity, specificity, and positive and negative predictive values of urine LCR after discrepant results were resolved were 88.6, 99.7, 96.9, and 99.0%, respectively. The sensitivity of culture was 71.4%. From the 148 pregnant women (prevalence by LCR, 6.8%), there were no patients who were cervical culture positive and urine LCR negative to indicate the presence in pregnant women of inhibitors of LCR. Additionally, a subset of 55 of the LCR-negative frozen urine specimens from pregnant women that had been previously processed in LCR buffer were inoculated with 5 cell culture inclusion forming units of C. trachomatis each and retested by LCR; all tested positive, indicating the absence of inhibitors of LCR in urine from these pregnant women. The use of LCR testing of urine specimens from asymptomatic women, whether pregnant or not, offers a sensitive and easy method to detect C. trachomatis infection in women.

MATERIALS AND METHODS

Populations and specimens. Military women (n = 480) attending clinics for a routine PAP smear test volunteered for a study to compare urine LCR tests to cervical cultures for the detection of C. trachomatis infections. The volunteer rate of the women approached by the civilian research nurse was 71%. The study was approved by the Institutional Review Boards of The Johns Hopkins University, the U.S. Army Medical Research Material Command, Fort Detrick, Frederick, Md., and Womack Army Medical Center, Fort Bragg, N.C. Of 480 women enrolled, 465 provided a urine specimen. All subjects completed a questionnaire for demographic information and behavioral risk factors for sexually transmitted diseases. The data instrument was a one-page, two-sided scanable bubble form (Scantron Corporation, Tustin, Calif.). During the pelvic examination, an endocervical swab was obtained by the attending clinician at the PAP smear clinic, who recorded clinical signs and symptoms on the data form. Culture swabs were placed into 2-macro-phosphate chlamydia transport medium. Commercial transport medium was replaced with in-house transport medium after 1 month of the study due to some toxicity of the former to tissue culture cells. Specimens were stored appropriately (4°C for urine specimens and -70°C for cultures) until shipping of the urine specimens at 4°C and cultures at -70°C. Specimens were made to ensure arrival at the laboratory within 4 days of collection. All specimens, consent forms, and data forms were shipped to Johns Hopkins Chlamydia Research Laboratory.

Approximately 4 million Chlamydia trachomatis urogenital infections occur in the United States annually, and more than 50 million cases occur worldwide (7, 28). Unfortunately, symptoms are often mild or absent among infected men and women, leaving a large reservoir of infected persons to continue transmission to new sex partners (29). Chlamydial infections occur primarily among young sexually active persons. A high prevalence is common to all socioeconomic groups and may range from 5 to 20% in various groups of young adults (32, 33). Because of the high probability of progression of asymptomatic disease to serious sequelae, it has been recommended that individuals at risk for chlamydial infections be screened, especially women who are vulnerable to the serious consequences of genital infections, such as pelvic inflammatory disease, ectopic pregnancy, and tubal infertility (7, 11). Urine can now be used to detect chlamydial infections in women by ligase chain reaction (LCR) (2, 8, 14, 20, 31, 34), which with its easily used to detect chlamydial infections in women by ligase chain reaction (LCR) (2, 8, 14, 20, 31, 34), which with its easily
Laboratory procedures. Urine specimens were processed and tested by LCR (Abbott Laboratories, Abbott Park, Ill.) according to the manufacturer’s instructions. Briefly, 1 ml of urine was centrifuged at 15,000 × g for 15 min. After the supernatant was removed, 1 ml of urine buffer was added to the pellet and the mixture was vortexed. After being heated at 97°C for 15 min, specimens were cooled and 1/3 ul of each specimen was added to an LCR unit “dune.” An appropriate chlamydia-positive control was included for the processing steps for each group of specimens. Additionally, two negative controls and two positive calibrator controls supplied by the manufacturer were used for each LCR assay run. After the amplification step in the automated thermocycler, unit dose tubes containing the specimens and controls were transferred to the automated enzyme immunoassay machine for the detection of amplified products. Tubes containing the amplified products were never opened; the automated enzyme immunoassay process sampled tubes by piercing the tops of the unit dose tubes, which prevented amplicon contamination. In order to prevent other sources of contamination, specimens were processed in a designated room separate from the room used to amplify and detect specimens. Gloves were frequently changed and screw-cap pipette tips and dedicated pipetters were used. Strict quality control measures such as machine maintenance checks, daily cleaning of laboratory areas, and area security tests to monitor contamination were employed.

Culture specimens were stored frozen at −70°C for up to 3 days. Cultures were done in 96-well microwell plates in McCoy cells by standard methods (12). Tissue cultures were stained with genus-specific fluorescein-conjugated antibody (Kallestad, Chaska, Minn.) and species-specific antibody (Boehringer Mannheim/Syva, San Jose, Calif.). Stained cultures were read for the presence of chlamydial inclusion bodies with an epifluorescence microscope. Discrepancy analysis was done for any sample with discordant results between culture and LCR. A sample that was positive by culture and negative by LCR was considered to be a true positive, but the discrepancy was investigated for the presence of inhibitors to amplification by LCR. The urine LCR was repeated from the originally processed specimen and repeated again after diluting the processed specimen 1:10 in urine LCR buffer to check for the presence of inhibitors in the specimen. (Dilution is sometimes used to decrease the concentration of the inhibitor enough to allow a true-positive specimen to be amplified.) Additionally, PCR (Roche Diagnostic Systems, Branchburg, N.J.) was done on an archival aliquot of frozen urine and another LCR was done for a different DNA target, the outer membrane protein 1 (OMP-1) gene. For specimens that were positive by LCR and negative by culture, the culture specimen transport sediment was stained by direct fluorescent antibody (DFA) (Boehringer Mannheim/Syva) for chlamydial elementary bodies. PCR was done on the specimens from the culture transport vials. In addition, PCR was done on the archived urine and an LCR for the OMP-1 gene was done on the previously processed (buffered) urine specimen. Specimens that were positive by one or more of the ancillary tests were considered true positives. An LCR-positive urine specimen which could not be confirmed by another test was considered to be a false positive.

Testing of urine specimens from pregnant women. A subset of all available (n = 465) previously processed (buffered) LCR-negative urine specimens that were from pregnant women were inoculated with 5 inclusion forming units of C. trachomatis and retested by LCR to check for the presence of inhibitors. Additionally, 65 archived LCR-negative unprocessed urine specimens that were available from pregnant women were tested by a research internal control assay to evaluate the presence of inhibitors (9). This assay tested for the ability to amplify an extraneous sequence of DNA which was added as an internal control to the specimen. The assay contained primers for the extraneous DNA internal control as well as the primers for the organism of interest. A positive amplification of the internal control indicated that the specimen contained no inhibitors to the amplification process, while a negative result indicated that the specimen contained something which inhibited the amplification process.

Data analysis. The data from the questionnaire forms were scanned into a data set (D-base III Plus, Ashton Tate, Borland International, Spring Valley, Calif.), and LCR results, demographics, and risk factor information were analyzed by the chi-square test, Fisher’s tests of exaction, and univariate analysis (Intercooled Stata, version 4.0; Stata Corporation, College Station, Tex.).

RESULTS

Patient characteristics. Among the 480 women enrolled, only 1 woman had reported mild symptoms and the remainder were asymptomatic. Approximately half (55.2%) were 25 years or younger; 50.8% were African-American. Over 90% were enlisted personnel, 98.3% reported vaginal sex, 11.3% had a new sex partner in the previous 90 days, 15.2% had more than one sex partner in the previous 90 days, 88.5% reported inconsistent condom use, and 30.8% were pregnant (Table 1). Reasons for clinic visit, clinical presentation, and sexual risk history are presented in Table 1. Of the 465 women who provided an urine specimen, the overall prevalence for chlamydia infection by LCR was 7.3%. The prevalences of infection for other categories based on LCR included 11.0% for women ≤25 years of age, 8.9% for African-American women, and 6.8% for pregnant women. By risk category the prevalences were 15.1% for those with a new sex partner in the previous 90 days, 10.3% for those with more than one sex partner in the previous 90 days, 7.5% for those with inconsistent condom use, 7.4% for those reporting vaginal sex, and 3.6% for those with a prior chlamydial infection.

In univariate analysis only young age (≤25 years) (odds ratio [OR], 4.23; 95% confidence interval [CI], 1.72 to 10.43) and a new sex partner (OR, 2.61; 95% CI, 1.11 to 6.1) were predictors of chlamydial infection (Table 2). However, when we controlled for age, a new sex partner was no longer significant.

Comparison of urine LCR to cervical culture. Of the 465 women, 31 women did not have matched culture specimen results. Ten specimens were toxic for tissue culture and no cervical cultures were collected from 21 women, leaving 434 matched specimens for comparison. After the use of the commercial chlamydia transport buffer was stopped and the in-
TABLE 2. Univariate analysis of results relative to factors associated with positive urine LCRs for military women attending PAP smear clinics

<table>
<thead>
<tr>
<th>Factor†</th>
<th>% with a positive LCR</th>
<th>OR (95% CI)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Factor absent</td>
<td>Factor present</td>
<td></td>
</tr>
<tr>
<td>Age ≤25 yr (254)</td>
<td>2.8</td>
<td>11.0</td>
<td>4.2 (1.72, 10.43)</td>
</tr>
<tr>
<td>African-American (233)</td>
<td>6.8</td>
<td>8.6</td>
<td>1.3 (0.61, 2.7)</td>
</tr>
<tr>
<td>Pregnant (142)</td>
<td>7.4</td>
<td>7.0</td>
<td>0.94 (0.44, 2.03)</td>
</tr>
<tr>
<td>Normal pelvic exam (275)</td>
<td>12.5</td>
<td>6.6</td>
<td>0.49 (0.18, 1.31)</td>
</tr>
<tr>
<td>Prior diagnosis of STD* (127)</td>
<td>8.3</td>
<td>3.9</td>
<td>0.45 (0.17, 1.2)</td>
</tr>
<tr>
<td>Having had more than one sex partner in last 90 days (68)</td>
<td>6.8</td>
<td>10.3</td>
<td>1.6 (1.33, 3.73)</td>
</tr>
<tr>
<td>Having had a new sex partner in last 90 days (53)</td>
<td>6.4</td>
<td>15.1</td>
<td>2.6 (1.11, 6.10)</td>
</tr>
</tbody>
</table>

† Numbers in parentheses represent numbers of individuals with the factor present (n = 465).
* STD, sexually transmitted disease (chlamydia, gonorrhea, syphilis, or trichomoniasis).
† A new sex partner was not significant when we controlled for age.

LCR of urine of pregnant women. There were 148 urine specimens from pregnant women. The prevalence of chlamydia infection by LCR for the pregnant women was 6.8%, and that for the nonpregnant women was 7.8%. There were no culture-positive, LCR-negative results from pregnant women which could have indicated the presence of LCR inhibitors. All four of the culture-positive, LCR-negative specimens were from women who were not pregnant. In addition, a subset of 55 LCR-negative urine specimens, previously processed in LCR buffer and frozen, which were from pregnant women and were inoculated with chlamydia and retested by LCR were all LCR positive, indicating the lack of inhibitors. Of the 65 available archived urine specimens from pregnant women which were LCR negative and tested in the internal control assay, there were 3 (4.6%) that exhibited inhibition based on a negative value for amplification of the internal control.

DISCUSSION

Chlamydia infections were of a higher prevalence than expected from these asymptomatic military women attending a clinic for a routine PAP smear test. An LCR prevalence of 7.3% underscores the necessity for the recommendation to screen all sexually active young women when they are attending a routine health care clinic (7). The high prevalence of 11.0% for those ≤25 years of age confirm the result of studies of others that young age is a significant risk factor for chlamydial infections (13, 17, 22). These results indicate the need for increasing public awareness and education for the need for screening and treatment.

TABLE 4. Resolution of urine-LCR-positive and cervical-culture-negative discrepant results for C. trachomatis in military women attending PAP smear clinics (n = 11)

<table>
<thead>
<tr>
<th>Laboratory no.</th>
<th>Test result</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCR (urine)</td>
<td>DFA*</td>
<td>PCR (urine)</td>
</tr>
<tr>
<td>1264</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>2407</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>3197</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>3659</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>3891</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>5560</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>5570</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>6082</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>6280</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>6966</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>8016</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

* DFA staining of culture transport vial specimen.
for an ongoing chlamydial control program for such female
military personnel as those enrolled in this study. This popula-
tion demonstrated a high degree of sexual behaviors placing
them at risk for sexually transmitted diseases, with 98% being
sexually active, 15% having more than one partner, 11% hav-
ing a new partner in the last 90 days, and 88% using condoms
inconsistently. All of these behaviors have been shown by oth-
eres to be predictive of chlamydial infection (1, 22-24, 36). In
the univariate analysis for this study, both young age (preva-
ence, 11.0%) and having had a new partner (prevalence, 15.1%)
reached statistical significance. However, when we con-
trolled for age, a new sex partner was not significant. Young
age (≥25 years), which is an easily determined risk factor and
which is a nonthreatening question for those women who may
be reticent to answer questions about their sexual behavior,
appears to be an excellent predictor of chlamydia infection and
can be recommended for deciding who should be screened in
clinical or outreach situations (13, 17).

Urinary LCR performed well in this study of asymptomatic
women, with a sensitivity of 88.6%, which is similar to that
demonstrated by others for asymptomatic women (87.5%) (2).
Compared to cervical culture, which had a sensitivity of 71.4%,
LCR detected more infected women. Many reasons can ac-
count for the lower culture sensitivity. Not only can the cold
chain of the transport be interrupted, but the quality of the trans-
port medium is important as well. Initially, a commercially
available transport medium was used in this study, which re-
sulted in many (10) toxic tissue culture results. Quality-control
assays of the remaining lot of uninoculated transport medium
demonstrated that it was toxic to cells in tissue culture. After
switching to the use of our own transport medium, which is
quality controlled in tissue culture, we observed no further
toxicity.

Additionally, the quality of the endocervical specimen, as
measured by the presence of columnar epithelial cells, has
been shown to play a significant role in the numbers of positive
specimens (19, 37). In another study of family-planning clinics
in Baltimore, Md., clinicians obtained adequate specimens
only 72.5% of the time (37). Thus, inadequate cervical swab
specimens could have contributed to the lower sensitivity of
culture in our study. Other studies have demonstrated higher
sensitivities for urine LCR than cervical culture (2, 5, 8, 20, 31,
34). Sensitivities for cervical culture in these studies has ranged
from 45.5 to 46.9% to 55.6 to 65.0% (2, 5, 8, 34). Schachter et
al. have demonstrated that the sensitivity of culture for C.
trachomatis may be increased from 67.1% to 74% by adding a
urethral swab culture, which could be indicative that some
women may be infected only in the urethra and not the cervix
(31). This could help explain the higher number of positives
found by urine LCR, presumably reflecting infections from
both the cervix and the urethra. Because urine is an easy-to-
obtain noninvasive specimen giving accurate results with LCR,
it is ideal for screening asymptomatic individuals who may not
be presenting for a pelvic exam or for outreach screening
programs.

Although our study enrolled only 148 women who were
pregnant, we did not observe any indication of inhibitors in
urine specimens, as evidenced by the lack of urine-LCR-neg-
ative results when the cervical culture was positive. Although
there were four such specimens in this study, they were all from
nonpregnant women. Another study has reported a significant
problem with inhibitors in urine with use of the LCR test; how-
ever, the urine specimens were heated and transported at ambient
temperatures, which may have influenced the LCR results (18,
25, 30). The spiking experiment in our study did not demon-
strate any inhibitors in the 55 LCR-negative, previously frozen
urine specimens from pregnant women. It is possible that
freezing and thawing of these processed urine specimens re-
duced or destroyed some LCR inhibitors. Freezing and thaw-
ing reduced the inhibition from 19 to 16% in one study (35).
Additionally, the experiment which tested the archived urine
of 65 pregnant women demonstrated only three (4.6%) inhib-
ited specimens. This value is of the same order of magnitude as
that reported by others for inhibition in urine specimens (2.6
and 1.8%) for amplification testing (3, 15). Most investigators
now believe that inhibitors to amplification exist for both urine
and cervical specimens (3, 15, 35). A combination of heat treat-
ment (95°C for 10 min) and 10-fold dilution of the processed
specimens reduced inhibition of PCR from 19 to 4% in one
study (35). The pH of the cervical mucosa was partly correlated
with inhibitors (35). Decreased inhibition was found at pH
values of ≥7.5. The degree to which inhibitors to amplification
influence the prevalence detected by LCR and PCR needs to
be further studied. Roche Molecular Systems has addressed
this problem by incorporating an internal DNA control amplifica-
tion and detection assay into their new combination PCR
assay for C. trachomatis and Neisseria gonorrhoeae, which will
prove to be a great advance in the diagnostic capability of
amplification assays. Specimens exhibiting inhibitors can be
diluted or heated and their DNA can be extracted, and tests
can be repeated. The use of the internal control will give a
greater degree of confidence to the validity of a negative am-
plification result. Consideration of the use of an internal con-
roll should be given for amplification tests in the future. The
College of American Pathologists now requires examination of
a control to assess the presence of inhibitors in all amplifica-
tion procedures.

In summary, young sexually active women, including those in
the military, should be frequently screened for chlamydia in-
fecions. Urine LCR offers an easy and sensitive method to
accomplish this, especially for women not presenting for a
pelvic examination. It is cost-effective in preventing the expe-
sive sequelae of pelvic inflammatory disease, ectopic preg-
nancy, and tubal infertility (16).

ACKNOWLEDGMENTS

We thank the study coordinator, Barbara Pare; the research nurses
Eleanor Howard, Kathy Cline, and Bobbi Jones and the staffs of the
Fort Bragg clinical sites for obtaining specimens; the laboratory tech-
nicians Griciaels Jaschek, Laura Welsh, Dinh Pham, Diana Perkins,
Sandy Leister, and Kimberly Crotchfeld for performance of laboratory
tests and data entry; Kathryn Clark for statistical assistance; and Pat
Buist for assistance in manuscript preparation.

Funding for this study was from Department of the Army grant
DAMD 17-95-1-5064.

REFERENCES

1. Addiss, D. G., M. L. Vaughn, R. Gohulnatinikov, J. Pfister, D. F. I. Kartyecz,
and J. P. Davis. 1990. Chlamydia trachomatis infection in women attending
urban midwestern family planning and community health clinics: risk factors,
selective screening, and evaluation of non-culture techniques. Sex. Transm.
Dis. 17:138-146.
Epidemiology Group. 1997. Multiplex AMPLICOR PCR screening for Chla-
mydia trachomatis and Neisseria gonorrhoeae in women attending non-sexu-
for diagnosis of Chlamydia trachomatis genital infections. Mbl. Med. 156:
420-421.
5. Buimer, M., G. J. Van Doornum, S. Ching, P. G. H. Peerbooms, P. K. Pier,
D. Ram, and H. H. Lee. 1996. Detection of Chlamydia trachomatis and
Neisseria gonorrhoeae by ligase chain reaction-based assays with clinical spec-
imens from various sites: implications for diagnostic testing and screening.
Chlamydia trachomatis Infections in Female Military Recruits

C. A. Gaydos and Others
CHLAMYDIA TRACHOMATIS INFECTIONS IN FEMALE MILITARY RECRUITS


ABSTRACT

Background Asymptomatic genital Chlamydia trachomatis infections in women can lead to pelvic inflammatory disease, infertility, and ectopic pregnancy. To design a chlamydia-control program, we conducted a large survey of women in the U.S. military.

Methods From January 1996 through December 1997, urine specimens from 13,204 new female U.S. Army recruits from 50 states were screened by ligase chain reaction for C. trachomatis infection. Information on potential risk factors was obtained by questionnaire. With multivariate analysis, we identified criteria for a screening program.

Results The overall prevalence of chlamydial infection was 9.2 percent, with a peak of 12.2 percent among the 17-year-old recruits. The prevalence was 15 percent or more among the recruits from five southern states. The following risk factors were independently associated with chlamydial infection: having ever had vaginal sex (odds ratio for infection, 5.9), being 25 years of age or less (odds ratio, 3.0), being black (odds ratio, 3.4), having had more than one sex partner in the previous 90 days (odds ratio, 1.4), having had a new partner in the previous 90 days (odds ratio, 1.3), having had a partner in the previous 90 days who did not always use condoms (odds ratio, 1.4), and having ever had a sexually transmitted disease (odds ratio, 1.2). A screening program for subjects 25 years of age or less (87.9 percent of our sample) would have identified 95.3 percent of the infected women.

Conclusions Among female military recruits, the prevalence of chlamydial infection is high. A control program that screens female recruits who are 25 years old or younger with urine DNA-amplification assays has the potential to reduce infection, transmission, and the sequelae of chlamydial infection.

M. S., M.D., M.P.H., CLARK, M.D.; M.P.H., GAYDOS, M.D., M.P.H., AND THOMAS C. QUINN, M.D.

METHODS

Population and Specimens

All female Army recruits who were present on Sundays between January 1996 and December 1997 at the Physical Examination Section, Reception Battalion, Fort Jackson, South Carolina, were invited to participate in this study. The study was approved by the institutional review boards of Johns Hopkins University and Fort Jackson (Eisenhower Army Medical Center, Fort Gordon, Ga.), as well as the Human Subjects Research Review Board of the U.S. Army Surgeon General. Of the 16,593 recruits approached, 13,223 (79.7 percent) volunteered to participate in the study and were given a briefing about the study as well as an educational briefing about chlamydial infections by the civilian research nurse.

All subjects signed an informed-consent form and completed a questionnaire regarding demographic information, home state, and medical care, screening of young, sexually active women has been recommended. In the past, screening for C. trachomatis infections in women has been limited by the need for access to a medical clinic and a pelvic examination. However, C. trachomatis infections can now be detected with high sensitivity (85 to 95 percent) and specificity with DNA-amplification assays performed on urine specimens, allowing cost-effective screening of large numbers of women in nonclinic settings. Few studies of the prevalence of chlamydial infection in U.S. military populations have been published, and there have been no studies using DNA-amplification techniques among women not seeking health care. Because adolescents have the highest prevalence of disease and most military recruits are young, we conducted a large prevalence study and risk-factor analysis of female recruits from throughout the United States who began basic training at Fort Jackson, South Carolina. We performed this study to determine the extent of infection, assess the feasibility of screening urine specimens for C. trachomatis by the ligase chain reaction, and assess which epidemiologic correlates would be useful for implementing an effective chlamydia-control program for female recruits.

and sexual history. The data instrument was a two-sided scannable form (Scantron, Tustin, Calif). To determine the similarity of the study subjects and those who chose not to participate in the study with regard to demographic characteristics and sexual history, 823 of the 3370 women who did not volunteer were invited to fill out an anonymous questionnaire. Nonvolunteers were asked to fill out a questionnaire only during the first week of each month.

Each volunteer was instructed to collect 20 to 30 ml of first-catch urine (the first part of the urine stream). A unique study number was assigned to each volunteer. All urine specimens, consent forms, and questionnaires were shipped to the Johns Hopkins University chlamydia laboratory. Urine specimens were kept at 4°C until processed, within 48 hours.

Laboratory Procedures and Treatment

Urine specimens were processed and tested by the ligase chain reaction (Abbott Laboratories, Abbott Park, Ill.) for chlamydial DNA according to the manufacturer’s directions. Each week a list of infected subjects was sent to the research nurse. The infected subjects were contacted and treated at the Troop Medical Clinic at Fort Jackson by directly observed therapy with a single 1-g dose of azithromycin. The subjects were also tested for coexisting sexually transmitted diseases. The sensitivity and specificity of the ligase chain reaction in urine specimens as compared with cervical culture for chlamydia had been previously determined to be 88.6 percent and 99.7 percent, respectively, in another military population.

Statistical Analysis

Questionnaire forms were scanned into a data base (dBASE III Plus, Borland International, Spring Valley, Calif). The results of the ligase chain reaction, demographic information, and risk-factor information were analyzed as dichotomous variables with the chi-square test. Univariate and multivariate logistic-regression analysis for factors associated with chlamydial infection was performed with Intercooled Stata software (version 4.0, Stata, College Station, Tex.). All independent variables were entered into the model, and a two-sided P value of less than 0.05 was considered to indicate statistical significance. The 95 percent confidence interval for the prevalence value for recruits from each state was calculated with Stata software. A one-way analysis of variance was performed to assess the degree of significance of differences in prevalence between states.

RESULTS

Characteristics of the Subjects

Of 13,223 subjects presenting at the Physical Examination Section on Sundays from January 1996 through December 1997, 19 could not be evaluated because of missing data or insufficient urine. The median age of the 13,204 who could be evaluated was 21 years (range, 17 to 36); 51.3 percent were white, and 31.9 percent were black. The mean age of the remaining nonvolunteers was 21 years (range, 17 to 36); 51.3 percent were white, and 31.9 percent were black. The mean age and the racial distribution of these recruits were not significantly different from those of the volunteers. Only 66.9 percent reported having had vaginal sex, as compared with 93.1 percent of the volunteers (P<0.001). This group differed significantly from the volunteers in four variables, even after adjustment for whether the women reported having had vaginal sex: only 4.0 percent reported prior chlamydial infections (P=0.013), 18.2 percent had had a new sex partner in the previous 90 days (P=0.002), 20.1 percent had partners who consistently used condoms (P<0.001), and 90.7 percent reported no previous diagnosis of a sexually transmitted disease (P=0.001).

Of the nonvolunteers, 17.7 percent had had more than one sex partner in the previous 90 days; the proportion of the volunteers who had had more than

| TABLE 1. CHARACTERISTICS OF 13,204 FEMALE ARMY RECRUITS SCREENED FOR CHLAMYDIA TRACHOMATIS. |
|-------------------------------|------------------|
| CHARACTERISTIC                | VALUE            |
| Age — yr                      | 21               |
| Median                        | 17–39            |
| Race — no. (%)*               |                  |
| White                         | 6,715 (51.0)     |
| Black                         | 4,732 (35.9)     |
| Other                         | 1,726 (24.1)     |
| Ever had vaginal sex — no. (%)†| 12,281 (93.1)    |
| Sexual history in previous 90 days — no. (%)‡| 3,478 (26.7) |
| More than one partner‡        | 4,076 (31.4)     |
| Partner always used condoms†  | 2,115 (16.9)     |
| Previous diagnosis of sexually transmitted disease — no. (%)§| 1,206 (9.1) |
| Chlamydia trachomatis         | 430 (3.3)        |
| Neisseria gonorrhoea          | 74 (0.6)         |
| Syphilis                      | 611 (4.6)        |
| Trichomonas                   | 11,722 (86.1)    |
| None                          | 1,219 (9.2)      |
| Chlamydia-positive — no. (%)  |                  |

*Data were missing for 30 subjects.
†Data were missing for 9 subjects.
‡For 168 subjects, data were missing or subject did not know answer.
§For 225 subjects, data were missing or subject did not know answer.
¶For 684 subjects, data were missing or subject did not know answer.
one sex partner in the previous 90 days was similar after adjustment for vaginal sex ($P=0.189$).

**Prevalence of Infection**

The age-specific prevalence of *C. trachomatis* infection among the 13,204 volunteers is shown in Figure 1. The highest prevalence of chlamydial infection (12.2 percent) was among 17-year-olds. The prevalence declined sharply with increasing age, to below 5 percent for women over 25 years of age. For further analysis, the youngest age groups (17 to 25 years) were combined into a category called “young.” The prevalence in this group was 10.0 percent (1162 of 11,603). In the older-age category (26 to 39 years), the prevalence was 3.6 percent (57 of 1601). The prevalence was 5.5 percent (369 of 6715) for whites, 14.9 percent (707 of 4733) for blacks, and 8.1 percent (143 of 1756) for other races.

**Univariate Analysis**

Univariate analysis identified 10 variables significantly associated with chlamydial infection: young age (17 to 25 years), black race, race other than white or black, ever having had vaginal sex, having had more than one sex partner in the previous 90 days, having had a new sex partner in the previous 90 days, a prior diagnosis of gonorrhea, a prior diagnosis of trichomoniasis, and a history of any sexually transmitted disease (Table 2). A prior diagnosis of chlamydia or syphilis was not significantly associated with being positive for chlamydial infection.

**Multivariate Analysis**

In the complete multivariate model, having had vaginal sex, an age of 25 years or less, black race, having had more than one sex partner in the previous 90 days, having had a new sex partner in the previous 90 days, having had a partner who did not always use condoms in the previous 90 days, and a history of any sexually transmitted disease were independent predictors of chlamydial infection (Table 3).

**Strategies for Selective Screening**

A screening strategy involving all variables identified as independent predictors would require that 100 percent of the population be tested and would detect 100 percent of the positive subjects. In this model, the magnitude of risk associated with having had a new sex partner might vary according to race. For the purpose of a screening program, this would not alter the proportion of the population tested or the percentage of positive subjects detected with this model. Because screening on the basis of race would probably be viewed as inequitable, a strategy excluding race was examined. According to this strategy,

![Figure 1. Mean (±SE) Age-Specific Prevalence of Chlamydial Infection among 13,204 Female Army Recruits, According to Ligase-Chain-Reaction Assays of Urine Specimens.](image-url)
recruits would be tested if they were 25 years of age or less or if they reported on a questionnaire having had more than one sex partner or a new sex partner in the previous 90 days, having had a partner who did not always use condoms, or for whom data were missing. Because most chlamydial infections are asymptomatic in women and because the sequelae of disease present a severe and costly burden, screening women at entry into the Army is an appropriate way to identify infections early and to explore opportunities for a control program.1,6,7,22

Civilian chlamydia-control programs have sought to identify criteria for selective screening,23-27 Most of these control programs have used diagnostic assays that require pelvic examinations and cervical specimens.28 However, it has recently been shown that testing urine specimens by DNA-amplification techniques is cost effective for screening large numbers of persons in different settings.16,28 We used this new technique to determine the prevalence of chlamydia and to identify screening criteria for a program to control chlamydia in the military.10,12,14 Collection of urine specimens in this study was highly acceptable and easily implemented.

Using the ligase chain reaction with urine samples, we found a high prevalence of C. trachomatis infection (9.2 percent). This prevalence was higher than that observed in family-planning clinics28 but not as high as that reported in some adolescent health clinics.29,30 Our data agree with those from previous studies of chlamydial infections in Army women, in which prevalence rates ranged from 8.2 percent to 9.8 percent.17,18 In one large, community-based screening study, the overall prevalence of chlamydia in young women was 8.6 percent, as detected by the urine ligase chain reaction, a prevalence similar to that found in our study.31 Because our population was not clinic-based and was not made up of women seeking health care, the finding of such a high prevalence in these women warrants the institution of a control program for the routine identification and treatment of chlamydial infections in order to prevent sequelae and transmission to sex partners.8

The study population consisted of a young, sexually active group of female recruits with sexual risk factors known to be associated with chlamydial infection.28 Although 91 percent of the subjects reported having had chlamydial infection in the past, this factor was not associated with the risk of current infection. The highest prevalence was observed among 17-year-olds. This prevalence is similar to comparable age-specific rates in other studies, confirming that young age is associated with chlamydial infection.24,31 In our study, young age was associated with being chlamydia-positive in both univariate and multivariate analyses (odds ratio, 3.0). In order to include more positive subjects, we used an age cutoff of 25 years, which allowed the detection of 95.3 per-

### Table 3. Multivariate Analysis of Factors Independently Associated with Chlamydial Infection in Female Army Recruits.

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Odds Ratio (95% CI)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age ≤25 yr</td>
<td>3.0 (2.3–4.0)</td>
</tr>
<tr>
<td>Black race†</td>
<td>3.4 (2.9–3.8)</td>
</tr>
<tr>
<td>Other (nonwhite, nonblack) race†</td>
<td>1.7 (1.4–2.1)</td>
</tr>
<tr>
<td>Having ever had vaginal sex</td>
<td>5.9 (3.2–10.6)</td>
</tr>
<tr>
<td>Having had &gt;1 sex partner in previous 90 days‡</td>
<td>1.7 (1.2–1.7)</td>
</tr>
<tr>
<td>Having had a new sex partner in previous 90 days§</td>
<td>1.3 (1.1–1.6)</td>
</tr>
<tr>
<td>Having had a partner who did not always use condoms in previous 90 days¶</td>
<td>1.4 (1.1–1.6)</td>
</tr>
<tr>
<td>Having ever had a sexually transmitted disease</td>
<td>1.2 (1.0–1.4)</td>
</tr>
</tbody>
</table>

*CI denotes confidence interval.
†The reference group consisted of the white subjects.
‡The reference group consisted of the subjects who did not have >1 sex partner, who answered that they did not know, or for whom data were missing.
§The reference group consisted of the subjects who did not have a new sex partner, who answered that they did not know, or for whom data were missing.
¶The reference group consisted of the subjects who had a partner who always used condoms.

Geographic Variation in Prevalence

There was considerable variation in the prevalence of chlamydial infection according to the state of origin of the recruits (P<0.001 by one-way analysis of variance). The prevalence was more than 15 percent for recruits from South Carolina, Georgia, Alabama, Louisiana, and Mississippi. For New Jersey, North Carolina, Kentucky, Texas, Oklahoma, and Arkansas, the prevalence was 10 to 15 percent, and for 17 other states and Puerto Rico, it was 5 to 10 percent. For five states (Washington, Oregon, Minnesota, Arizona, and Massachusetts), the prevalence was less than 5 percent. Fewer than 100 recruits were tested from each of 17 states, 3 territories, and the District of Columbia, and prevalence figures from these areas were therefore not included in the analysis. The prevalence for the five states with the highest prevalence and the five states with the lowest prevalence differed significantly, since the 95 percent confidence intervals for prevalence did not overlap.

DISCUSSION

Although the diagnosis and treatment of sexually transmitted diseases has always presented a challenge, there has been no routine screening of recruits for chlamydial infections at entry into the U.S. Army.21 Because most chlamydial infections are asymptomatic in women and because the sequelae of disease present a severe and costly burden, screening women at entry into the Army is an appropriate way to identify infections early and to explore opportunities for a control program.1,6,7,22

Civilian chlamydia-control programs have sought to identify criteria for selective screening,23-27 Most of these control programs have used diagnostic assays that require pelvic examinations and cervical specimens.28 However, it has recently been shown that testing urine specimens by DNA-amplification techniques is cost effective for screening large numbers of persons in different settings.16,28 We used this new technique to determine the prevalence of chlamydia and to identify screening criteria for a program to control chlamydia in the military.10,12,14 Collection of urine specimens in this study was highly acceptable and easily implemented.

Using the ligase chain reaction with urine samples, we found a high prevalence of C. trachomatis infection (9.2 percent). This prevalence was higher than that observed in family-planning clinics28 but not as high as that reported in some adolescent health clinics.29,30 Our data agree with those from previous studies of chlamydial infections in Army women, in which prevalence rates ranged from 8.2 percent to 9.8 percent.17,18 In one large, community-based screening study, the overall prevalence of chlamydia in young women was 8.6 percent, as detected by the urine ligase chain reaction, a prevalence similar to that found in our study.31 Because our population was not clinic-based and was not made up of women seeking health care, the finding of such a high prevalence in these women warrants the institution of a control program for the routine identification and treatment of chlamydial infections in order to prevent sequelae and transmission to sex partners.8

The study population consisted of a young, sexually active group of female recruits with sexual risk factors known to be associated with chlamydial infection.28 Although 91 percent of the subjects reported having had chlamydial infection in the past, this factor was not associated with the risk of current infection. The highest prevalence was observed among 17-year-olds. This prevalence is similar to comparable age-specific rates in other studies, confirming that young age is associated with chlamydial infection.24,31 In our study, young age was associated with being chlamydia-positive in both univariate and multivariate analyses (odds ratio, 3.0). In order to include more positive subjects, we used an age cutoff of 25 years, which allowed the detection of 95.3 per-
cent of the chlamydial infections. Other studies have supported age-based screening for chlamydia. 27,28,32

Thus, for this group of female recruits coming from a civilian background, who were tested within three days of starting basic training, young age alone can be recommended as a single indicator of who should be tested for chlamydial infection. Other models considered in this study offered high sensitivity, but the models were more complex and required valid sexual-risk histories. We documented 13 chlamydial infections (prevalence, 1.4 percent) among 914 recruits who denied being sexually active, as well as chlamydial infections in 8.4 percent of those who reported that their partners consistently used condoms. These figures indicate that self-reported sexual-risk histories are not always valid. 23 The lower prevalence of chlamydial infection among recruits for whom the data on condom use were missing, or who indicated on their questionnaires that they did not know whether their partners always used condoms, may be due to lack of sexual activity, because 58.6 percent of the 684 recruits in these categories reported that they had never had vaginal sex. There is a fixed laboratory budget available for population screening in the Army. Young age is the simplest, least expensive, and most easily documented risk factor on which to base a recommendation for a screening program, as well as being highly sensitive. Alternatively, since the use of age as a selective screening criterion would have missed 4.7 percent of the infections, universal screening might be more cost effective from a societal perspective, and future studies of cost effectiveness are warranted. 28

This was one of the largest programs for screening young, sexually active subjects that was not clinic-based and whose results were derived from urine DNA-amplification assays. The geographic variation in prevalence was striking. From more than 15 percent in the five states with the highest prevalence to less than 5 percent in the five states with the lowest, these differences may reflect the levels of disease burden in certain states. These regional variations also appear to reflect regional differences in chlamydial disease, as reported by the Centers for Disease Control and Prevention. 24,35 For example, the prevalence in North Carolina reportedly varied from 10 percent to 17 percent. 35 The prevalence is lower in regions such as Wisconsin and Washington State, where clinic-based chlamydia-control programs are in place and where declining rates of prevalence of chlamydia have been reported. 26,27,32,36 In our study, the prevalence was 11.3 percent for North Carolina and 3.8 percent for Washington State. Our data imply that chlamydial infection remains common in young women across the United States. With a volunteer rate of 80 percent among women who were approached and representation from 50 states and 4 territories, our study had a wide geographic sampling.

One limitation of our study is that it is not known whether the prevalence of risk factors for chlamydial infection differs between young women who decide to join the military and those who do not. However, the demographic and sexual risk-factor characteristics of our subjects appear to be similar to those of other regional and clinic-based populations, 25 as well as those from a large, community-based study. 31 An additional limitation is that the nonvolunteers in our study differed from the volunteers with regard to sexual risk factors for chlamydia. However, the nonvolunteers represented a group who were mostly sexually active, who had had new sex partners in the previous 90 days, and whose partners did not use condoms. Thus, their risk of chlamydial infection may have been as high as that of the subjects in our study.

Although amplified-DNA tests are more expensive than traditional nonculture tests, the savings associated with not having to have a clinician collect specimens from a pelvic examination and the advantages of being able to use urine as a diagnostic specimen may outweigh the extra cost of the test. 15 In addition, it has been demonstrated that amplified-DNA testing of urine specimens is cost effective, and treating chlamydial infections prevents serious complications such as pelvic inflammatory disease, ectopic pregnancy, and infertility. 15,28,37

In conclusion, our study indicates that with the limited funding available at present, young age (25 years or less) would be the best criterion on which to base a screening program using amplified-DNA testing of urine for female Army recruits and perhaps for other young women. Institution of such a control program has the potential to reduce drastically the burden of chlamydial disease in the U.S. Army and to prevent morbidity due to these infections. 37

Supported by a grant (DAMD17-95-1-5064) from the Department of the Army.

We are indebted to Dr. Daniel Scharfstein, Mary Sheppard, Dr. Guillermo Madine, and Dr. Anne Ronapalo for statistical assistance; to Kathryn Kaesna for data entry; to Diane Perkins, Sandra Leisley, Dori Plant, and Graciekon Jaseko for performance of the laboratory tests; to Lt. Col. Cynthia Dodd, Lt. Col. (ret.) Delois Daniels, and Kimberly Riley for treatment of infected women and processing of specimens; and to Patricia Bails for assistance in the preparation of the manuscript.

REFERENCES


©Copyright, 1998, by the Massachusetts Medical Society
Printed in the U.S.A.