Perceived transmissibility of STIs: lack of differentiation between HIV and chlamydia

Sexually transmitted infections (STIs), such as HIV and chlamydia, differ widely in their transmissibility. The estimated probability of HIV transmission from an infected heterosexual man to a woman in one act of unprotected vaginal intercourse is 0.1%, whereas the same probability for chlamydia is 35%.

This research examines college students’ knowledge about the per act transmission probabilities for HIV and chlamydia.

Previous studies reported median perceived transmission probabilities of 50% and 33.4% for HIV for one act of unprotected receptive vaginal intercourse with an infected man. These findings were interpreted as demonstrating “badly overestimated per act transmission probabilities” (Pinkerton et al. 1999). However, the distributions of the estimates were not provided. If estimates are widely dispersed across the entire probability range from 0% to 100%, interpretations of averages are meaningless and interpreting the data as indicating a systematic overestimation of the transmission probability would be unfounded. We studied this possibility in a sample of college students.

In all, 234 undergraduate university students (145 women, 85 men, mean age 21.14 years, SD 2.82, four did not report their age and sex) enrolled in a variety of academic programmes were randomly selected and individually approached after classes. Aside from their age and sex, participants were asked in two separate questions: “What do you think is the probability, in percentages, of a woman becoming infected with HIV (chlamydia) from one unprotected act of vaginal intercourse with an infected man?” The order in which people were asked the two questions was counterbalanced.

Figure 1 presents the distribution of the estimates, showing that they are widely and quite equally dispersed across the entire range from 0% to 100%, and that distributions do not differ between the two infections (Kolmogorov–Smirnov Z = −0.73, p = 0.46). No age or sex differences were found. Only 3.9% and 5.6% of the estimates for HIV and chlamydia, respectively, come close to the correct probabilities if “correct” is defined as smaller than 0.5% for HIV and between 30% and 40% for chlamydia. In all, 34.8% of the participants falsely estimate that chlamydia has a lower transmission probability than HIV, 39.5% correctly estimate that chlamydia has a higher transmission probability than HIV, and 25.8% provide exactly the same percentage estimate for both STIs.

The data show that a large majority of college students clearly lacks knowledge of the transmission probabilities of HIV and chlamydia and does not know that chlamydia is more infectious than HIV. Previous reports of statistical averages of the perceived transmissibility and their interpretation as indicating a systematic overestimation bias may be unfounded. The results highlight the importance of inspecting response distributions and restraining from reporting statistical averages when distributions are widely dispersed. Furthermore, they highlight that information about transmission probabilities should be incorporated into sexual health programmes in order to make people more aware of STIs that are considerably easier to contract than HIV.

Acknowledgements

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Contributors

The study was jointly conceptualised and designed by BK and RK; the data were collected by BK; RK should be credited for assistance. BK analysed the data and led the writing; both authors jointly interpreted the findings, reviewed drafts of the manuscript, and approved the final version.

B Knäuper, R Kornik
McGill University, Montreal, Canada

References


A new method for extended trichomonad storage

With the introduction of the InPouch test for Trichomonas vaginalis, T. gallinae, and T. foetus, it was desirable to have a procedure available for maintaining extended culture viability. The three trichomonads are viable after 8 days by subculture in the InPouch at 33°C. Extended viable storage of these three trichomonads is the subject of this letter.

We have evaluated various procedures involving freezing 24 hour InPouch cultures at −70°C. We now report a procedure that has demonstrated storage of viable trichomonad cultures for more than 2 years.

The freshly subcultured trichomonads are incubated at 35°C for 24 hours, which should produce a viable count of approximately 1.0 × 10^7/ml. It is important to note that subsequent subculture will require an adequate nutrient available for growth in the pouch. Then 0.1 ml of pure sterile glycerol is added to the medium in the pouch and thoroughly mixed employing the “shoe-shine” technique. It is important to immediately place the pouch in a −70°C freezer.

After freezing most of the trichomonads in the pouch are non-viable, but successful subculture is routinely achieved upon thawing. When the pouch is removed from the freezer, it should be immediately placed in an incubator at 35–37°C. After 3 days a few viable trichomonads will be observed, and after 4 days it may be subcultured.

This procedure has been effective for T. vaginalis, T. gallinae, and T. foetus.

K A Borchardt
CBLS, San Francisco State University, San Francisco, CA, USA

J H Hall
BioMed Diagnostics, Inc, San Jose, CA, USA

Figure 1 Distribution of estimates for HIV and chlamydia for one act of unprotected vaginal intercourse with an infected man.
Intriguing condition in a patient.

Penile involvement secondary to urethral tuberculosis is rare and its presentation with perirectal fistulas leading to “water can” penis is unknown. We report this rather unique condition in a patient.

A 40 year old male agricultural labourer presented with a 1 year history of purulent discharge per urethra with multiple discharging sinuses on the tip of the penis. The patient was asymptomatic about a year ago, when he developed multiple nodules on the penis. He had a purulent discharge per urethra with multiple discharging sinuses on the tip of the penis. The patient was otherwise healthy and had no systemic complaints. He was married with two children and had no history of extramarital contact or genital ulcers.

On physical examination, the penis shape was like a saxophone. The prepuce and glans penis were oedematous and indurated. The penis was like a saxophone. The prepuce and glans penis had multiple sinuses around the urethral meatus (fig 1). On squeezing the meatus, pus was expressed from the meatus and the sinuses. The glans penis also showed areas of depigmentation (vitiligo). The distal part of the shaft of the penis showed induration involving corpora cavernosa whereas the proximal part was devoid of any lesion.

On ultrasonography of abdomen and prostate, the prostate was normal on rectal palpation. The prostate was normal on rectal examination. The routine haemogram revealed an elevated erythrocyte sedimentation rate of 100 mm in the first hour. Liver and renal function tests were normal. The discharge smear stained with Gram stain and Ziehl-Neelsen stain. The Gram stained smear revealed numerous pus cells and acid fast stain showed abundant acid fast bacilli. Culture for Mycobacterium tuberculosis grew contaminants. A roentgenogram of the chest and penis was unremarkable. An intravenous pyelogram was normal. Voiding cystourethrography revealed glandular urethral stricture with urethrococintaneous fistulas. Ultrasonography of abdomen and prostate was normal. Mantoux skin test was strongly positive (30 mm) (fig 2). His veneral disease research laboratory test (VDRL) and HIV serology was non-reactive.

Based on these clinical features, positive Mantoux test and acid fast bacilli in the urethral smear, the diagnosis of urethral tuberculosis with urethrococintaneous fistula was made. The patient was started on anti-tuberculous treatment comprising isoniazid 300 mg, rifampicin 600 mg, pyrazinamide 1500 mg, and ethambutol 800 mg per day. The patient showed marked improvement after 4 weeks of treatment. The sinuses closed and discharge ceased. Patient was referred to urology for management of stricture, which was planned after the anti-tuberculous treatment. The patient tolerated anti-tuberculous treatment and completed 9 months of treatment with remarkable recovery in the swelling of the penis.

Genital involvement occurs in 50% of male patients with urogenital tuberculosis. Penile tuberculosis is rare with less than 1% of patients having penile involvement. Tuberculosis of the penis usually presents as ulcers, tubercular cavernosis, or nodules. In most cases, the lesion appears as a superficial, solitary, painless ulcer on the glans penis. It can be clinically indistinguishable from malignant disease. Rarely, lesions may persist as solid nodule or cavernousitis with ulceration. Papuloncrotic tuberculide may also present as an ulcer on the penis. Penile involvement may occur secondary to co-existing urinary tract tuberculosis. The transmission occurs secondary to bacilluria in these patients. Infection of the penis may occur by direct contact at the time of intercourse with a partner having urogenital tuberculosis.

Tuberculosis of male urethra is an uncommon condition and presents as urethral strictures, periurethral abscesses, or fistula formation. Fistulas can occur in the perineum leading on to “water can” perineum. Similar occurrence of fistulas in penis can be aptly be designated as “water can” penis. In our case, penile involvement occurred secondary to urethral tuberculosis. Such involvement of the penis by tuberculosis is unique and not reported in the literature. “Water can penis” is also known to occur with gonorrhoea but our patient had a negative urethral smear for Gram negative diplococci and had features suggestive of urethral tuberculosis. Further, the strictures, fistulas, and lymphoedema had led to “saxophone” deformity of the penis. Such deformity is well known with lymphogranuloma venereum, but is unknown in tuberculosis.

K Karthikeyan, D M Thappa, K N Shivswamy
Dermatology and STD Department, JIPMER, Pondicherry - 605006, India
Correspondence to: Professor D M Thappa, Dermatology and STD Department, JIPMER, Pondicherry - 605006, India; dmtthappa@jipmer.edu

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1 Vipulkumar M, Thappa DM, Kaviyarasan PK. Papuloncrotic tuberculide of the glans penis (correspondence). Sex Transm Infect 2001;77:147.

South Asians with HIV in London: is it time to rethink sexual health service delivery to meet the needs of heterosexual ethnic minorities?

Recent conservative estimates suggest that at the end of 2002, 4.8 million people were living with HIV/AIDS in South Asia including 4.8 million in India. In the United Kingdom there are estimated to be 1.5 million people of south Asian ethnicity. While the National Strategy for Sexual Health aims to improve health care in those who have HIV through earlier diagnosis, studies have shown that that other ethnic minority groups present with advanced disease and not through routine genitourinary medicine (GUM) screening. We studied the case notes of all adults self defining as of Indian, Pakistani, Bangladeshi, or Sri Lankan ethnicity diagnosed HIV positive from
Table 1  Characteristics of presentation of study population at time of HIV diagnosis (n = 117)

<table>
<thead>
<tr>
<th></th>
<th>Heterosexual men (n = 45)</th>
<th>Homosexual men (n = 36)</th>
<th>Heterosexual women (n = 27)</th>
<th>Other risk groups (n = 9)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AIDS illness at presentation</td>
<td>16 (36%)</td>
<td>6 (16%)</td>
<td>2 (7%)</td>
<td>1 (11%)</td>
</tr>
<tr>
<td>Median CD4 cell count ×10^3/μl (range)</td>
<td>178 (3–1,023)</td>
<td>381 (4–810)</td>
<td>377 (10–1,104)</td>
<td>151 (50–795)</td>
</tr>
<tr>
<td>Median HIV viral load copies/ml (range)</td>
<td>24 500 (50–1,000 000)</td>
<td>24 636 (425–3 000 000)</td>
<td>7822 (173–489 184)</td>
<td>12 870 (6676–57 530)</td>
</tr>
<tr>
<td>Reasons for HIV test AIDS/symptomatic</td>
<td>27 (60%)</td>
<td>11 (31%)</td>
<td>7 (26%)</td>
<td>2 (22%)</td>
</tr>
<tr>
<td>Known HIV+ sexual partner</td>
<td>3 (7%)</td>
<td>4 (11%)</td>
<td>12 (44%)</td>
<td>0</td>
</tr>
<tr>
<td>Routine screen for sexually transmitted infections</td>
<td>1 (2%)</td>
<td>16 (44%)</td>
<td>1 (4%)</td>
<td>0</td>
</tr>
<tr>
<td>Patient request</td>
<td>7 (15%)</td>
<td>3 (8%)</td>
<td>2 (8%)</td>
<td>2 (22%)</td>
</tr>
<tr>
<td>Child positive</td>
<td>3 (7%)</td>
<td>0</td>
<td>3 (11%)</td>
<td>0</td>
</tr>
<tr>
<td>Insurance/visa purposes</td>
<td>3 (7%)</td>
<td>1 (3%)</td>
<td>0</td>
<td>2 (22%)</td>
</tr>
<tr>
<td>Antenatal screening</td>
<td>0</td>
<td>0</td>
<td>2 (7%)</td>
<td>0</td>
</tr>
<tr>
<td>Other</td>
<td>1 (2%)</td>
<td>1 (3%)</td>
<td>0</td>
<td>3 (34%)</td>
</tr>
</tbody>
</table>

The median age at diagnosis was 38 years (range 19–64 years) for men and 28 years (range 20–55 years) for women. Forty-five patients (38%) had originated from Africa, 28 (24%) from India, and 18 (15%) from the United Kingdom. The majority were of Indian ethnicity (95/117; 81%) with the next largest ethnic group being Sri Lankan (12/117; 10%).

The primary mode of transmission was heterosexual sex (72/117; 62%) with transmission through sex between men accounting for a further 31% (36/117) of cases. Four infections were acquired through blood transfusion, two through injecting drug use, one from a needle stick injury, and in two cases risk behaviour could not be identified.

There were substantial differences in the reasons for testing between individuals in the main risk groups. In particular, heterosexual men and women were both significantly less likely than homosexual men to be diagnosed via routine attendance at a GUM clinic (2% and 4%, compared to 44%, respectively, p < 0.001, Fisher’s exact test). Among heterosexuals, the main reason for testing in men was symptomatic HIV infection/AIDS (60% of men but only 26% of women), whereas women were more likely to be tested through partner notification of a known HIV+ sexual contact (44% v 7% in males) (table 1).

The median CD4 count at presentation overall was 300 (range 3–1104) cells ×10^3/μl. However, male heterosexuals presented with significantly lower CD4 counts (median 178, range 3–1023 cells ×10^3/μl ) than either homosexual men (median 381, range 4–810 cells ×10^3/μl; p = 0.01) or heterosexual women (median 377, range 10–1104; p = 0.02).
Table 2 Details of STIs diagnosed in men and women

<table>
<thead>
<tr>
<th>STI</th>
<th>Male</th>
<th>p Value using $\chi^2$ test</th>
<th>Female</th>
<th>p Value using $\chi^2$ test</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>37 (0.5)</td>
<td>39 (0.8)</td>
<td>0.061</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>B1</td>
<td>262 (3.8)</td>
<td>190 (2.9)</td>
<td>0.002</td>
<td>41 (0.9)</td>
</tr>
<tr>
<td>C4a/C4c</td>
<td>234 (3.5)</td>
<td>179 (2.7)</td>
<td>0.005</td>
<td>177 (3.9)</td>
</tr>
<tr>
<td>C4h</td>
<td>683 (9.9)</td>
<td>479 (7.2)</td>
<td>&lt;0.001</td>
<td>–</td>
</tr>
<tr>
<td>C10a</td>
<td>89 (1.3)</td>
<td>55 (0.8)</td>
<td>0.009</td>
<td>111 (2.3)</td>
</tr>
<tr>
<td>C11a</td>
<td>264 (3.8)</td>
<td>254 (3.8)</td>
<td>0.998</td>
<td>147 (3.1)</td>
</tr>
<tr>
<td>Total diagnosed with an STI at this episode</td>
<td>1579 (22.8)</td>
<td>1210 (18.2)</td>
<td>&lt;0.001</td>
<td>486 (10.1)</td>
</tr>
</tbody>
</table>

*Value using $\chi^2$ test with Yates’s correction.

(A1) Primary diagnosis of syphilis; (B1) gonorrhoea; (C4a, C4c) uncomplicated chlamydia; (C4h) non-gonococcal urethritis; (C10a) first attack of genital herpes; (C11a) anogenital warts.

References


Prevalence of HSV-1/HSV-2 antibodies in HIV seropositive patients in Coventry, United Kingdom

The seroprevalence of herpes simplex virus (HSV) antibody among HIV patients within the United Kingdom is unknown. We therefore conducted a HSV seroprevalence study in HIV patients attending our genitourinary medicine clinic from January 2000 to December 2001. Our previous study revealed an overall prevalence of HSV-1 (60%), HSV-2 (20%), and both HSV-1 and HSV-2 (12%) among male and female genitourinary medicine clinic attendees who were either HIV negative or whose HIV status was unknown.

Serum samples from 96 consecutive ethnically diverse HIV patients were collected during routine investigations, and tested for HSV type specific antibodies by monoclonal antibody blocking enzyme linked immunosays. Out of 96 patients, two HSV-1 and three HSV-2 antibody test results were equivocal in four individuals. These were excluded from the analysis and results are presented here for 92 patients.

There were 56 men and 36 women in the study. 46 (50%) were white, 43 (47%) black African, and three were from other ethnic groups. All the black Africans were heterosexuals and 71% of men were homosexuals. The median age was 35 years (range 21–80).

HSV-1 seroprevalence was 86% among men and 97% among women ($p = 0.14$). HSV-2 seroprevalence was 50% among men whereas it was 94% among women ($p = 0.0001$). There was no statistically significant difference between the seroprevalence of HSV-1 between white and black people. However, seroprevalence of HSV-2 and both serotypes was significantly higher among black than among white people.

This study shows very high seroprevalence of HSV-1 (90%), HSV-2 (67%), and both HSV-1 and HSV-2 (64%) among our HIV positive cohort in Coventry. The high prevalence of HSV-2 in women is possibly because most of them were black African and acquired HIV through sex. These findings may have important public health implications as the high rate of HSV-2 is therefore likely to act as a cofactor in HIV transmission.

P S Allan, S Das
Department of GU Medicine, Coventry and Warwickshire Hospital, Coventry CV1 4FH, UK;
risa.allen@yahoo.co.uk

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Effective Sexual Health Interventions: Issues In Experimental Evaluation


HIV spreads more every day and there are epidemics of other STIs in both the developed and developing world at least in part because the fear of HIV appears to be receding in the population. Our current strategies to contain these problems are meeting with limited success and treatment of people who are already infected, important though that is in controlling bacterial infections, is much less effective with continuing viral infections. There is an urgent need to develop and to test better
methods of helping people to reduce their risky sexual behaviour.

This book is excellent, brief, fairly comprehensive, and very readable. Its focus is designing studies on the effectiveness of sexual health interventions. If we are to get anywhere in improving behavioural interventions it is essential that what is done is carefully evaluated.

The first three chapters of the book are concerned with methodology, particularly whether randomised controlled (RCTs) trials are an appropriate method for evaluating interventions in this area. While this section of the book is well argued on all sides it doesn’t really break any new ground. The strengths and weaknesses of RCTs in behaviour change are pretty much what they are in any other area of medicine. Methodologies don’t exist as stand alone phenomena, whether an RCT or some other methodology is appropriate depends simply on what question one is seeking to answer.

The second section of the book covers models of behaviour change and the choice of design and outcome measures. It is clear that one of the main problems in intervening in sexual health is the poor quality of the available psychological models and our real lack of understanding about why people behave as they do. Without understanding why people behave as they do it is difficult to help them to change. It is interesting that models of health behaviour never seem to get discarded, even the ones that are known to be weak. There are particularly strong chapters on cluster randomisation, an approach which probably gives rise to more inappropriate statistics than any other and on complex behavioural measures. The latter should be required reading for anyone measuring any behaviour change programme would be of no use in developing countries if it was labour intensive and dependent on highly skilled staff for its delivery.

I would recommend this book to anyone planning a trial or simply seeking to understand the existing literature. I would however caution that to make sense of it you will have to look at some of the available reviews of the behaviour change literature since the book assumes some knowledge, or willingness to acquire knowledge, of these.

The book ends by looking at generalisability in its broadest sense. Generalisability is an area that tends to get overlooked. Even a highly successful behaviour change programme would be of no use in developing countries if it was labour intensive and dependent on highly skilled staff for its delivery.

I would recommend this book to anyone planning a trial or simply seeking to understand the existing literature. I would however caution that to make sense of it you will have to look at some of the available reviews of the behaviour change literature since the book assumes some knowledge, or willingness to acquire knowledge, of these.

J Green
St Mary’s and Imperial College Hospital,
London, UK; mail@john-green.com

Australasian Sexual Health Conference 2004: Behind the Mask

This conference will be held at the Adelaide Convention Centre, South Australia, on 31 March to 3 April 2004. For further details please contact Dart Associates (tel +61 2 9418 9396/97; email dartconv@mpx.com.au; and website http://www.acshp.org.au).

8th European Society of Contraception Congress

The 8th European Society of Contraception Congress will be held from 23–26 June 2004 in Edinburgh, Scotland, UK. For further details please contact ESC Central Office, c/o Orga-Med Congress Office, Essenestraat 77, B-1740 Ternat, Belgium (tel: +32 2 582 08 52; fax: +32 2 582 55 15; email: orgamed.am@pandora.be; and website: http://www.contraception-esc.com/edinburgh.htm).

CORRECTIONS


The authors of a letter in the December issue of STI (Dave SS, Johnson AM, Fenton KA, Mercer CH, Erens B, Wellings K. Male circumcision in Britain: findings from a national probability sample survey. Sex Transm Infect 2003;79;499–500) were listed in the wrong order. The correct author list should be as follows: Dave SS, Fenton KA, Mercer CH, Erens B, Wellings K, Johnson AM.

In the corresponding author’s address of a letter published in the December issue (Bhatia R, Prabhakar S, Shedde D, et al. Coexistent cranial tuberculomas and tuberculosis of the cervix in a postmenopausal woman. Sex Transm Infect 2003;79;496–7) All India Institute of Medical Sciences was incorrectly printed as AU India Institute of Medical Sciences.
"Water can" penis caused by tuberculosis

K Karthikeyan, D M Thappa and K N Shivaswamy

Sex Transm Infect 2004 80: 75
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