Pharyngeal gonorrhoea: the forgotten reservoir

Urethral gonorrhoea (UG) dramatically decreased in Paris between 1986 and 1997 as a consequence of safer sexual behaviour. Thus, only 43 cases of gonorrhoea have been collected in our clinic in 1997, the lowest number since the early 1970s. Since 1998, an increase has been observed, as in other STD clinics in France and in the Renago laboratory network.1" Men who have sex with men (MSMs) represent an increasing number of men with UG. Many of them are HIV seropositive and recognise unprotected oral sex as the only risk factor for gonorrhoea. This finding prompted us to study pharyngeal carriage of Neisseria gonorrhoeae (NG) and Neisseria meningitidis (NM) in this population.

From January 1999 to May 2001, 200 consecutive cases of male UG were observed in our clinic; a pharyngeal smear for culture of NG and NM was suggested as well as a standardised questionnaire aimed at sexual behaviour; 178 gave informed consent. Results are presented in table 1, comparing MSMs and men who also have sex with women (MSWs).

Interestingly, MSMs represent more than 50% of patients with UG (compared to 10% in 1986 and 20% in 1995). One third of them are HIV seropositive (a minimal figure because of a high rate, 9%, of test refusal). Fifty eight per cent admitted unprotected oral sex as the sole risk factor for gonorrhoea. Moreover, 98% of the gonococci cultured in MSMs are serogroup W-2–3 (v 7% in MSWs) and only 1/92 produce penicillinase (v 26% in MSWs), suggesting a homogeneous cluster of strains circulating in the Paris gay community (study ongoing). Finally, pharyngeal carriage of both NG (14%) and NM (20%) is high.

Data concerning MSWs are heterogeneous, UG affects mainly male patients from north (35%) and central (31%) Africa, with oral sex as the only risk factor for gonorrhoea (10%), and pharyngeal carriage of NG and NM (6%) is much lower, but not inconsistent. Pharyngeal gonorrhoea is mostly asymptomatic (all our cases were) and bacteriological diagnosis is uncertain, but we believe that the pharynx acts as an important reservoir accounting for the recent increase in UG, particularly in MSMs using unprotected oral sex as an alleged safer sex act. The high proportion of HIV infected patients is a major cause of concern and information about the hazards of unprotected oral sex is warranted.

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References

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Emerging of dual AIDS associated neoplastic diseases in the era of highly active antiretroviral therapy

Kaposi's sarcoma, non-Hodgkin's lymphoma, and cervical cancer remain the only AIDS associated malignancies, according to the 1993 CDC definition, but other neoplasms were reported throughout AIDS pandemic (Hodgkin's lymphoma, oropharyngeal, oesophageal, gastric, anal, lung, and breast cancer, testicular-ovarian neoplasms, melanoma, skin and thyroid malignancies, multiple myeloma, leiomysosarcomas, angiomyxosarcomas, smooth muscle tumours), with an increasing frequency despite HAART introduction.8

Among 711 AIDS patients notified since 1985, 66 (9.3%) were diagnosed because of an AIDS defining cancer, and 51 more patients (7.2%) developed a malignancy with AIDS, but dual AIDS associated neoplasms were never seen until 2000. A rare combination of lethal Kaposi's sarcoma plus non-Hodgkin's lymphoma was recently observed. Two homo/bisexual men had received multiple antiretroviral lines since 1990, but dual AIDS associated neoplasms was achieved by a CD4+ count of 42–235 cells × 10³ in the first patient, and 68–355 cells × 10³ in the second case. A first AIDS related neoplasm (a cutaneous-mucous Kaposi's sarcoma), was identified 2 and 5 years before death, respectively. Repeated cytotoxic treatment with adriablastine-bleomycin-cyclophosphamide, reduced disease progression, while a number of HIV related opportunistic infections occurred: oesophageal candidiasis and cryptococcosis in the first patient, and pneumonia, zoster, plus wasting syndrome in the second subject. Eleven and 5 months before the lethal outcome, respectively, a Burkitt's B cell lymphoma involving multiple skin sites and complicated by bone marrow, gastroduodenal, gingivobuccal, and pulmonary localisations was detected in the first patient, while the second subject had a high grade non-Hodgkin's lymphoma involving axillary-medistinal lymph nodes, lungs, and pleura. Notwithstanding therapeutic attempts (methotrexate-zidovudine, followed by MNCOP-B), a rapidly fatal course occurred.

The introduction of HAART determined a profound modification of the evolution of HIV disease, but improved patient survival, persisting immune system abnormalities, and co-infection with potentially oncogenic viruses may be responsible for the increased incidence of neoplasms during the HAART era.14 This phenomenon seems to extend beyond typical AIDS defining neoplasms, since other malignancies were reported with an incidence greater than that of the general population, and that of the pre-HAART era, although they may be largely underestimated, owing to the unchanged CDC AIDS classification system. This trend is not uniform for Kaposi's sarcoma,15 probably because of the favourable effects of antiretroviral-antiherpetic medications. The occurrence of dual AIDS associated malignancies remains exceptional: only two patients with a rare and aggressive non-Hodgkin's null cell lymphoma and prior Kaposi's sarcoma were described by Ascoli.16 Although our patients developed “typical” AIDS defining neoplasms, this phenomenon may become of increasing concern, when

### Table 1 Urethral gonorrhoea (UG)

<table>
<thead>
<tr>
<th></th>
<th>MSMs (n=92)</th>
<th>MSWs (n=86)</th>
<th>Total (n=178)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age years (SD)</td>
<td>31.4 (7.3)</td>
<td>33.5 (11.3)</td>
<td>32.7 (9.8)</td>
<td>NS</td>
</tr>
<tr>
<td>White, n (%)</td>
<td>67 (73)</td>
<td>17 (20)</td>
<td>84 (47)</td>
<td>10⁺</td>
</tr>
<tr>
<td>Oral sex as the only risk factor for UG, n (%)</td>
<td>53 (58)</td>
<td>9 (10)</td>
<td>62 (35)</td>
<td>10⁺</td>
</tr>
<tr>
<td>HIV test positive, n (%)</td>
<td>30 (33)</td>
<td>3 (3.5)</td>
<td>33 (18.5)</td>
<td>10⁺</td>
</tr>
<tr>
<td>HIV test negative, n (%)</td>
<td>8 (9)</td>
<td>5 (6)</td>
<td>13 (7)</td>
<td>NS</td>
</tr>
<tr>
<td>PPNG, n (%)</td>
<td>1 (1)</td>
<td>22 (26)</td>
<td>23 (13)</td>
<td>10⁻</td>
</tr>
<tr>
<td>NG pharynx, n (%)</td>
<td>13 (14)</td>
<td>5 (6)</td>
<td>18 (10)</td>
<td>0.06</td>
</tr>
<tr>
<td>NM pharynx, n (%)</td>
<td>18 (20)</td>
<td>5 (6)</td>
<td>23 (13)</td>
<td>10⁻</td>
</tr>
</tbody>
</table>

PPNG = penicillinase producing Neisseria gonorrhoeae; NG = Neisseria gonorrhoeae; NM = Neisseria meningitidis.
involving rare cancers. The increased life expectancy of HAART treated patients, a direct involvement of HIV itself, or abnormalities driven by oncogenic viruses, including EBV, HSV-8, and papillomavirus, might explain the tendency to develop a broader spectrum of long term neoplastic complications. In our experience, a persistent HIV-associated immunodeficiency and an incomplete virological response to HAART, possibly had a pathogenetic role. Clinicians should maintain an elevated clinical suspicion for a broad spectrum of HIV-associated cancer, even after a first diagnosis of AIDS-related neoplasm. Epidemiological studies should give a reliable estimate of the frequency of all HIV-associated tumours, and recognise eventual dual AIDS associated cancers. The pathogenesis underlying AIDS-related malignancies (especially neoplasms immunity and viral onco genesis) deserve careful insight.

Contributors
RM collected and interpreted data and literature evidence during the entire work; LC collected clinical and laboratory data and literature evidences, and revised both data collection and discussion; FC proposed and supervised the report, read and revised both data evaluation and discussion; FC proposed and supervised the report, read and revised both data evaluation and discussion; LC collected and revised the entire work. Correspondence to: Dr Roberto Manfredi, Department of Clinical and Experimental Medicine, Division of Infectious Diseases, University of Bologna “ Alma Mater Studium”, S. Orsola Hospital, Bologna, Italy.

References

Accepted for publication 13 February 2003

Impact of the Sexually Transmitted Infections Foundation course on the knowledge of family planning nurses and doctors

There has been convergence of genitourinary medicine and reproductive healthcare services in the United Kingdom to produce “one stop sexual health clinics,” such as the Sandyford Initiative in Glasgow. As part of service development a number of educational initiatives such as the Sexually Transmitted Infection Foundation (STIF) course have been initiated to ensure that minimum skills and competencies are obtained. Training programmes such as the STIF course coordinated by the Medical Society for the Study of Venereal Diseases (MSSVD) play a vital part in providing staff with the education required to competently extend their roles. The first Scottish STIF course was run in Glasgow in March 2002. The course was developed as a UK-wide initiative to support the implementation of the English national strategy for sexual health and HIV.

In order to evaluate the impact attendance at the STIF course had on the knowledge of family planning staff, a prospective study was performed in Glasgow. Eighteen members of family planning staff (15 doctors and three nurses) were assessed on their knowledge of vaginal and cervical infections before and after attendance at the course, using four clinical case scenarios with accompanying clinical pictures. A maximum score of 12 was awarded for each assessment. The cases comprised candida, trichomomas, bacterial vaginosis, and chlamydia. The participants were asked to provide a provisional diagnosis based on the history and a clinical picture. The vaginal pH was then provided and each participant was given the opportunity to alter their diagnosis in the light of this additional information. They were then asked about the management of each condition. Within 3 months of the STIF course, each doctor and nurse were retested with the initial scenarios. Answers and feedback were provided on completion.

Two sample t tests and confidence intervals for the difference of two means were employed to compare all participants and the doctors and nurses scores. As the numbers in the study were small a subanalysis of the results for different grades of doctors was not performed. Table 1 shows the mean (SD), median precourse and post-course scores, and mean difference in scores. The mean increase in all participants’ and the doctors’ scores were statistically significant (p = 0.001, and p = 0.006, respectively). The mean increase in the nurses’ scores was 5.0% (95% CI 3.6 to 13.6), however the number of nurse participants was small (n=3).

This study suggests that knowledge increased following attendance at the STIF course. Educational initiatives such as the STIF course are important tools for development of staff working in the field of sexual and reproductive health care. A larger study of this type assessing a wider range of subject matter with longer follow up would enable further evaluation of the STIF courses’ impact on knowledge.

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Impact of the Sexually Transmitted Infections Foundation course on the knowledge of family planning nurses and doctors

| Table 1 | The mean (SD), median precourse and post-course scores, and mean difference in scores |
| --- | --- | --- |
| *n* | Precourse score | Post-course score | Mean difference (95% CI) |
| All participants | 18 | | |
| Mean (SD) | 8.2 (2.3) | 10.7 (1.8) | +2.5 (1.1 to 3.9) |
| Median | 9.0 | 12.0 | |
| Doctors | 15 | | |
| Mean (SD) | 8.6 (1.9) | 10.6 (1.8) | +2.0 (0.7 to 3.3) |
| Median | 9.0 | 12.0 | |
| Nurses | 3 | | |
| Mean (SD) | 6.0 (3.0) | 11.0 (1.7) | +5.0 (3.6 to 13.6) |
| Median | 6.0 | 12.0 | |

Improving response rates for self collected urine samples

Chlamydia trachomatis is the commonest bacterial sexually transmitted infection (STI) in Victoria, Australia, with the number of notifications increasing threefold in the past 8 years from 1287 in 1994 to 3977 in 2001.1 As infection with chlamydia is frequently asymptomatic, notification data underestimate population prevalence. Innovative study designs are necessary to investigate chlamydia prevalence and risk factors. We conducted a pilot study among women aged 18–32, to estimate the rate of response to a request to provide a self collected urine specimen for chlamydia testing. Recruitment via mail was compared with recruitment via mail and follow up telephone contact.

Between March and May 2002, the names and addresses of 150 Victorian women aged 18–32 were randomly selected from the electoral roll. These were linked with the Electronic White Pages and telephone numbers obtained where possible, producing two groups: (1) women with telephone numbers identified, and (2) women without telephone numbers identified. All women were mailed a letter of invitation and an information leaflet. Women in group 2 were also mailed a reply paid participation form asking them to indicate whether they wished to participate.

Women in group 1 were telephoned after 1 week and consent sought to mail them a urine kit. Two reminder letters were sent to non respondents in group 2. Women testing positive...
were treated with azithromycin through their nominated doctor.

Participants provided 20 ml first void urine in collection vials provided. Specimens were tested for chlamydia by polymerase chain reaction.

Telephone numbers were found for 70 (47%) women. Among women in group 1, five (7%) were excluded because they were living overseas. Of the remaining 65 women, 35 (54%, 95% CI 41 to 66) agreed to participate and 29 (45%, 95% CI 32 to 57) provided a specimen. One case of chlamydia was diagnosed giving a prevalence of 3.4% (95% CI 0.1 to 17.8) in this group (table 1). Among women in group 2, 11 (14%) were excluded because they were not living at their registered address. Of the remaining 69 women, 16 (23%, 95% CI 14 to 35) agreed to participate and 14 (20%, 95% CI 12 to 32) provided a specimen. No cases of chlamydia were diagnosed.

In this pilot study we showed recruitment via mail and telephone had a significantly higher response than mail alone (45% v 20%, p=0.002). This suggests that telephone communication will increase response in population-based chlamydia research that uses mail contact as the principal recruitment tool. Although the method of recruitment was not randomly allocated, the 25% difference in response is unlikely to be explained by differences between the two groups.

A response rate of 45% for those recruited via mail and telephone compares well with results obtained in similar overseas studies. A response rate of 45% for those recruited via mail and telephone compares well with response is unlikely to be explained by differences in recruitment. Although the method of recruitment was not randomly allocated, the 25% difference in response is unlikely to be explained by differences between the two groups.

**References**

3. J Hocking and this project were supported by the Victorian Health Promotion Foundation (VicHealth).

**Table 1**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Total No (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18-22 years</td>
<td>28 (100)</td>
</tr>
<tr>
<td>23-27 years</td>
<td>23 (100)</td>
</tr>
<tr>
<td>28-32 years</td>
<td>23 (100)</td>
</tr>
<tr>
<td>Total</td>
<td>74 (100)</td>
</tr>
</tbody>
</table>

**Table 2**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Hospital clinic patients No (%)</th>
<th>Outreach clinic patients No (%)</th>
<th>Outreach clinic data (95) 1998</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethnicity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asian</td>
<td>11 (5)</td>
<td>5 (5)</td>
<td>17 (17.4)**</td>
</tr>
<tr>
<td>Black Caribbean</td>
<td>77 (37)</td>
<td>48 (43)</td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>54 (26)</td>
<td>23 (21)</td>
<td></td>
</tr>
<tr>
<td>African</td>
<td>14 (7)</td>
<td>18 (16)</td>
<td></td>
</tr>
<tr>
<td>First time attendees</td>
<td>72 (34)</td>
<td>66 (59)*</td>
<td></td>
</tr>
<tr>
<td>Women</td>
<td>107 (51)</td>
<td>75 (68)*</td>
<td></td>
</tr>
<tr>
<td>Men</td>
<td>102 (49)</td>
<td>36 (32)*</td>
<td></td>
</tr>
<tr>
<td>Median age</td>
<td>28</td>
<td>26</td>
<td>26**</td>
</tr>
<tr>
<td>Age &lt;20 years</td>
<td>22 (11)</td>
<td>21 (19)*</td>
<td>19 (20)**</td>
</tr>
<tr>
<td>Disease</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gonorrhoea</td>
<td>7 (3)</td>
<td>5 (5)</td>
<td></td>
</tr>
<tr>
<td>Chlamydia</td>
<td>12 (6)</td>
<td>9 (8)</td>
<td></td>
</tr>
<tr>
<td>Trichomoniasis (females)</td>
<td>4 (4)</td>
<td>1 (1)</td>
<td></td>
</tr>
<tr>
<td>NGU (male)</td>
<td>15 (15)</td>
<td>3 (8)</td>
<td></td>
</tr>
<tr>
<td>Genital herpes</td>
<td>5 (2)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Genital warts</td>
<td>12 (6)</td>
<td>3 (3)</td>
<td></td>
</tr>
<tr>
<td>HIV test</td>
<td>64 (31)</td>
<td>30 (27)</td>
<td>27 (28)**</td>
</tr>
</tbody>
</table>

* p<0.05, outreach v hospital clinic patients for 2001 data
** p<0.05, 95 outreach v 105 hospital clinic patients, data collected in 1998 for both.
Clearance of HPV infection in middle aged men and women after 9 years’ follow up

The age prevalence of human papillomavirus (HPV) cervical infections is high in young age groups, declining sharply thereafter, reaching a steady state after age 40. Women who remain persistent carriers of HPV DNA are considered at high risk for cervical cancer. To investigate viral persistence over an extended period of time, we re-contacted, in 1997–8, a group of women who participated in case-control studies between 1988–91 in Spain, Colombia, and Brazil. Among women with confirmed normal cervical smears, follow up was scheduled for all women positive for HPV cervical detection (n=91) and for a group of age matched women who were HPV negative (n=254). All but one HPV infection were of high risk types. Husbands of the women in Colombia and in Spain, initially detected to be HPV positive (n=110), were also re-contacted. Follow up data were obtained from personal interview and from HPV DNA tested in cervical and urethral (men) exfoliated cells. The follow up protocol was approved by the institution’s ethics committee and participants signed an informed consent. Finally, 198 women (57.4%) and 42 (38.2%) men were interviewed. Of them, 99 women provided cervical samples and 14 men provided urethral samples. HPV detection was carried out in the same laboratories that tested the initial samples. The Spanish and Colombian samples were tested using the PMY09/11 PCR L1 based method and the Brazilian samples were tested using the GP5+/6+ PCR system.

The average age at entry was 50.8 years for women and 51.9 for men (range 27–79 years). After an average of 9 years of follow up (range 7–11), none of the women examined harboured HPV DNA irrespective of their initial HPV status (table 1). The follow up cervical smear identified three women in Colombia and one in Brazil with a cervical intraepithelial neoplasia grade I. All were HPV negative. Among the HPV positive husbands who were re-examined, two remained positive (14.3%, 95% CI 3.7 to 32.6), one for low risk type HPV 6 and one for high risk type HPV 16. No penile lesions were detected upon clinical examination. An active search in the corresponding cancer registries did not identify any case of invasive cervical or penile cancer in the target population.

The data, albeit limited by small size, suggest that HPV infection in middle age is subject to clearance as is commonly observed in young women. All women with follow up information had no HPV infection after an average follow up period of 9 years. None of the women developed advanced cervical disease in the interval as would be expected in some cases of chronic carriers of HPV infection.

Table 1 HPV detection in women and men at entry and at follow up time

<table>
<thead>
<tr>
<th>HPV status at entry</th>
<th>Spain</th>
<th>Colombia</th>
<th>Brazil</th>
<th>All</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative</td>
<td>13</td>
<td>31</td>
<td>35</td>
<td>79</td>
</tr>
<tr>
<td>Positive</td>
<td>3</td>
<td>3*</td>
<td>12</td>
<td>20</td>
</tr>
<tr>
<td>HPV status at follow up</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>5</td>
<td>36</td>
<td>47</td>
<td>79</td>
</tr>
<tr>
<td>Positive</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Men:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HPV status at entry</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>9</td>
<td>5</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>HPV status at follow up</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>1**</td>
<td>1***</td>
<td>–</td>
<td>2</td>
</tr>
<tr>
<td>Negative</td>
<td>8</td>
<td>4</td>
<td>–</td>
<td>12</td>
</tr>
</tbody>
</table>

*One woman was positive for a low risk HPV DNA, **HPV 6, ***HPV 16.

References


Accepted for publication 20 January 2003
Are all genital Chlamydia trachomatis infections pathogenic? A study in men

Chlamydia trachomatis infection of the genital tract is initially mild and most sufferers do not know they have the infection. However, over a period of time, untreated infections may be associated with considerable pathology.

During a recent prospective survey of 500 men presenting in this department we recorded the Gram stained microscopy results from urethral swabs. These were scored by the pathologist who had no knowledge of the patient. They were scored on a scale of 1–5, corresponding to 0, 1–4, 5–9, 10–14, and >15 polymorphonuclear leucocytes (PMNLs). The results were later correlated with the routine chlamydia ELISA testing. The results are given in table 1.

It can be seen that in the chlamydia positive no<sub>22</sub>% do not have urethritis, defined as >5 PMNLs per high power microscopy field. Similarly, urethritis was found in 22% of men who were non-chlamydia, non-gonococcal (non-GC). This clearly confirms that chlamydia infection does exist in the absence of urethritis. Furthermore, this 34% did not correspond with asymptomatic infection; 55% were symptomatic and 45% asymptomatic. Likewise in those with urethritis, 57% were symptomatic and 43% asymptomatic. The most common symptom was discharge and the peak duration was 21 days. Of the total chlamydia positive group 16% had neither symptoms nor urethritis.

It is therefore possible that not all chlamydia infection leads to pathology and morbidity? Perhaps the non-inflammatory serovars are not harmful and do not produce the pathology that others do. Evidence does exist which suggests that different serovars do produce different pathology. 1, 4

Of the 22% of men who had non-chlamydia non-GC specific urethritis it seems highly likely that these will be due to Mycoplasma genitalium. In future we intend to test for Mycoplasma genitalium and to compare the pathology that these two organisms produce.

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References


Accepted for publication 27 February 2003

Follow up of patients who have been recently sexually assaulted

Follow up rates for victims of sexual assault have traditionally been low, ranging from 10% to 31%. Rates improved if a follow up appointment was arranged at a genitourinary medicine clinic (GUM) clinic—50% of the 70% of patients for whom an appointment was made.

FAMSAC (Forensic and Medical Sexual Assault Care) is a medical sexual assault service that has been integrated into an existing sexual health clinic for the past 19 months. A total of 114 sexual assault patients have used the service since November 2001 (106 females, 8 males). Consent for follow up contact from the nurse coordinator of FAMSAC is sought at the initial consultation; this occurs in the first week after the report of sexual assault.

The following elements of care are addressed at the follow up visit:

- Follow up screening for sexually transmitted infections and hepatitis B vaccination (initiation or continuation)
- Follow up pregnancy testing as necessary (emergency contraception is given at the initial medical examination)
- Management and follow up of injuries as necessary
- Referral to counselling services (patients are offered independent immediate support at the time of medical examination)
- Discussion of legal matters (police action, victim’s compensation, etc)
- Health promotion information and safety awareness strategies.

Patient follow up is the responsibility of the nurse coordinator with medical support as required, other duties include organising the preparation of legal reports, court appearances, and support of the medical officers ensuring continuity of care for the patient and minimal delay in the legal process.

To date we have contacted 97/114 (85%) of our patients. These rates are significantly higher than those reported by Herbert, 1 who reports a loss to follow up of 46% within 24–48 hours. This may be due in part to better access to telephones since her 1988–90 study—53% of our patients own a mobile telephone and 80% of patients gave a home contact telephone number. A total of 17 patients were unable to be contacted.

We offer a further opportunity for contact 3 months after the assault. To date 75 patients have been eligible; of these 59 (80%) have been contacted and 39 have attended (66%). Three patients who received HIV prophylaxis were offered a 6 month follow up appointment; all of those have attended.

The sexual health clinic appears to be an ideal venue for follow up of these patients, who appear to be at higher risk of acquiring a sexually transmitted infection. The nurse coordinator model has enabled the follow up of patients at higher rates than previously reported.

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References


Accepted for publication 10 March 2003

Patients lost to follow up: experience of an HIV clinic

The National Strategy for Sexual Health and HIV aims to reduce the pool of undiagnosed HIV infection in the United Kingdom. Potential benefits of earlier diagnosis include timely initiation of highly active antiretroviral therapy (HAART), prevention of complications of HIV, screening for STIs that are known to enhance HIV infectivity, 3 and psychological support. Patients may not realise these benefits if they are lost to follow up (LFU). Previous studies have found associations between frequent non-attendance (as distinct from LFU) and less severe illness, 4 drug addiction, 5 and patients’ health beliefs. 6

We studied the case notes of all surviving patients who had enrolled in our HIV clinic within a 15 month period but had not received medical care for 12 months. Patients were excluded if they had been transferred to other centres or if the case notes were unavailable. For each case, one control was matched for date of first attendance. Data including demographics, virological and immunological markers, antiretroviral therapy, and psychological and social factors were collected from the notes using a standardised proforma.

Ninety four cases were found. LFU patients were younger than controls (table 1), with a trend towards more patients being born outside the United Kingdom. Cases were about half as likely to be on HAART than controls (RR 0.46, 95% CI 0.32 to 0.66). This

Table 1 Microscopy

<table>
<thead>
<tr>
<th>PMNLs</th>
<th>Chlamydia positive, non-GC</th>
<th>Chlamydia negative, non-GC</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>9</td>
<td>284</td>
</tr>
<tr>
<td>1–4</td>
<td>6</td>
<td>55</td>
</tr>
<tr>
<td>5–9</td>
<td>3</td>
<td>32</td>
</tr>
<tr>
<td>10–14</td>
<td>12</td>
<td>21</td>
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<td>&gt;15</td>
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Total 66% 21 Total 22%
association was true regardless of disease stage. Numbers were too small to analyse any association between LFU and poor adherence to HAART. There were no statistically significant differences in sex, ethnicity, disease stage, or surrogate markers. The number of general practitioners with whom communication was maintained was equally low in both groups.

The number of cases with a history of psychiatric illness, substance abuse, deliberate self harm, or use of counselling or psychiatric services was not significantly different from controls. Further data provided by CDSC showed that at least 29 cases (31%) had attended another clinic for follow up, without correspondence being made between centres. These included only eight of 26 (30%) cases on HAART and eight of 22 (36%) cases with CDC stage C or CD4 count under 200 cells x10^9/l. (Soundex code and date of birth were used for matching records without compromising patient identity, and subsequent treatment locations were not specified.)

This study highlights that patients who are LFU attend those at all stages of disease and are not necessarily those with a lack of clinical need. It is of interest that the association between not being on HAART and being LFU is independent of clinical stage. The patients who discontinued care from our centre were a diverse group in terms of illness, ethnicity, and transmission category, typical of the clinic population as a whole.

In a patient who is symptom free and not on HAART, one might argue that a year without specialist follow up is of no clinical importance. Such patients might be better managed in primary care or in a setting which focuses less on the patient’s disease stage than on their wellbeing. An appropriately designed study might further elucidate reasons that lead patients to default from follow up. Interventions need to be in place to prevent loss of follow up of patients who are at high risk of disease progression or who are on HAART.

Acknowledgements
Nina Fudge helped to retrieve the data. Andrew Copas provided statistical advice.

Contributors
LH developed the study, retrieved and analyzed the data, and co-wrote the text; SE co-wrote the text; DM conceived the study and provided comments on the text; KS provided further data from CDSC.

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References

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Interactive continuing medical education (CME) and its influence on the working practices of genitourinary clinicians

Didactic lectures are the traditional vehicle used by the MSSVD for updating clinicians on developments in the specialty, but there is mounting evidence that this sort of educational format is unlikely to change clinical practice, whereas a format which more actively involves participants can produce measurable changes. The MSSVD decided to formally assess the impact of combining the lecture format with an interactive approach at one of its national update meetings. The subject under review at this meeting was human papillomavirus (HPV) infection. Specialist attendees were asked to vote electronically on a combination of knowledge base and treatment strategy questions. They were then presented with information on the correct answers to the questions, and on currently preferred treatments. Feedback questionnaires invited comparison with the usual didactic approach. Participants were also asked whether their clinical practice would change as a result of the meeting. Seventy MSSVD members signed for CME, and 43 returned feedback questionnaires at the end of the event. A small majority of 51% preferred the new format to the usual didactic format, while a minority of 21% preferred the traditional approach. Despite only a small majority preferring the interactive over the customary didactic lecture format, a clear majority of respondents, 70%, felt that the interactive format was better able to maintain their concentration and interest, and 60% felt the new format was more likely to induce reflection and stimulate change. To our surprise, 74% of respondents planned to make some change in clinical practice as a result of attending the event. Three months later, the majority of respondents were still planning to do so as soon as circumstances permitted. A large proportion of respondents stated explicitly what changes had been made. The major influence was on a cessation in the use of podophyllin, and an increased use of the topical wart treatments imiquimod and podophyllotoxin.

The incorporation of hand held electronic response units to facilitate audience participation in educational events in the future could considerably enhance the value of such events solidifying research results and reducing the need for national meetings. Continuing medical education courses could be considerably extended to offer on-going update and consultation. Educational meetings could be considerably augmented by the incorporation of interactive technology which was employed here powerful: firstly, each participant communicates directly not only with the lecturer, but also anonymously with all his peers; secondly, the event has to be formatted in such a way as to directly engage participants by requiring them answer clinically related questions; and thirdly, the organiser has to focus to a higher degree than normal on how everything that is said will be perceived.

We have been encouraged by the outcome of this event, and we believe that clinicians would benefit from increased utilisation of this interactive educational method.

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Reference

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Lower motor neuron syndrome and HIV infection

A 33 year old right handed male injection drug user presented with a 4 week history of progressive shoulder and upper arm weakness and difficulty speaking. Together with a 2 week history of fever and a productive cough. He had been HIV-1 antibody positive for 16 years, had no AIDS defining illness, and was on no antiretroviral therapy. The CD4 count was 110 cells $\times 10^9/l$ and viral load was 56 000 copies/ml. There was no past history of, nor had the patient recently been vaccinated against, poliomyelitis; the patient was HB antigen and hepatitis C antibody positive. There was no family history of neurological disease.

On examination there were signs of a right basal pneumonia; general examination was otherwise normal. Neurological examination showed he was alert and oriented, dysarthric, and had bilateral facial weakness worse on the left; the palate deviated to the right and there were fasciculations of the tongue, which was not wasted. Neck flexion was weak. The other cranial nerves were normal. In the limbs tone and sensation was normal. In the arms power was 4−4 but in the legs power was 4−3. Reflexes were absent in the left biceps, triceps, and supinator but otherwise they were intact; plantar reflexes were the plantar test.

Blood cultures grew Streptococcus pneumoniae; with broad spectrum antibiotics the patient improved with antiretroviral therapy.

The patient declined further investigation and therapy.

PostScript

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References


Accepted for publication 24 March 2003

Syphillis specific antibodies in newborn infants in Lower Saxony, Germany 1993–2001

In 1979, a test to detect syphillis specific antibodies was added to the neonatal screening programme for metabolic diseases in Lower Saxony.1 We report test results for the period 1993 to the end of 2001 (684 156 samples). Analysis of data allowed calculation of annual incidence rates, dependency of maternal age, and birth weight of infants on the incidence of positive results.

Neonatal screening for syphillis specific antibodies does not aim (and is not able) to diagnose congenital syphilis. The goal of our test is to remind physicians responsible for the infants of the positive history of the mother, so they can check whether sufficient treatment of the maternal infection can be proved or whether further measures are necessary.

Material was eluted from dried blood samples collected on filter paper. Syphillis-specific antibodies were identified using Treponema pallidum haemagglutination (TPHA) test (Fujiokjiki, Tokyo, Japan) or, since 1999, Treponema pallidum particle agglutination (TPPA) test (Fujiiebie Inc, Tokyo, Japan). Extract and suspension of sensitised erythrocytes or particles were mixed at a dilution of 1:80. All samples showing a reaction at this dilution were assayed as being positive and retested for quantification of antibodies. Both test versions are based on indirect particle agglutination caused by 7S-IgG and 19-IgM antibodies against Treponema pallidum.

During the observation period the incidence of infants with a positive test result increased significantly from 11.05 cases per 10 000 infants in 1993 to 19.73 cases per 10 000 children in 2001 (R$^2$ for the linear regression: 0.75, p = 0.003) (fig 1). The level of significance would be even higher if it took into account that the formerly used TPPA test produced a small but not exactly defined number of false positive results.

In former years, the incidence of syphillis antibodies in newborns increased with maternal age. Recently, age distribution of mothers of antibody positive babies changed: a large number of young mothers had babies with positive antibodies. Data for the years 1993−7 were compared with those of 1998−2001 by using the χ$^2$ test (two tailed p values). The most obvious change occurred in the group of 20–24 year old mothers and 25−29 year old mothers (p 0.001).

A significant correlation between birth weight and the probability of a positive test result was found. Data on birth weight were available for 405 786 newborns (81.2%). The incidence (cases per 10 000 tested infants) of syphillis antibodies in very low birthweight infants (<1500 g) and in low birthweight infants (<2500 g) was significantly higher (4.54 × 2.53, p<0.001 v 0.002) than in the total group (19.2).

Although the syphillis test was not introduced for epidemiological reasons, some conclusions can be drawn with respect to epidemiology. Firstly, there has been a significant rise in the incidence of syphillis antibody positive infants in the past 9 years, indicating an increasing number of young women who had or have a syphillis infection. Therefore, physicians in charge of newborn infants must be aware of the increasing probability of finding a congenital infection. Secondly, the probability of a positive test result is higher in low birthweight infants and in infants of younger mothers.
package and not be an optional extra. This is particularly important for women who, at a vulnerable time, may not be aware of the wider health benefits of screening.

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References

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Recent pilot studies of chlamydia screening

The recent pilot studies of chlamydia screening in Portsmouth and the Wirral show that there is a substantial burden of chlamydial infection in young women and that high uptake of screening and good coverage of the target population can be achieved. This is important. However, the pilot studies do not demonstrate the effectiveness of chlamydia screening in reducing either morbidity or the prevalence of infection (nor were they designed to do this). In fact, further screening (in the recall study) of the same target group in the same settings, approximately 16 months after the pilot screening had ended, shows no change in chlamydia prevalence: 11.2% (pilot) v 11.9% (recall) in the Wirral and 9.8% v 11.4% in Portsmouth. Opportunistic screening continued after the pilot in family planning clinics in the Wirral, but there has been no reduction in chlamydia prevalence (11.4% during March-August 2000 compared with 12.4% during March-August 2002).

It would be wrong to conclude that opportunistic screening does not work. The incidence of chlamydia in the United Kingdom appears to be rising and it may be that the prevalence found in the recall study would have been higher still in the absence of earlier screening and treatment. Thus, controlled studies are needed to determine effectiveness empirically. Economic modelling is important for assessing the long term effects of different screening sce-

References
4 Catchpole M, Robinson A, Temple A. Chlamydia screening in the United Kingdom. Sex Transm Infect 2003;79:3–4

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7th European Society of Contraception Seminar

An ESC seminar entitled: “Contraception practice in Europe: differences in availability and accessibility” will be held in Budapest, Hungary, on 12–13 September 2003. Further details: ESC Central Office, Essentrastraat 77, B-1740 Ixelles, Belgium (tel: 32 2 582 08 52; fax: 32 2 582 55 15; email: esccentraloffice@contraception-esc.com and website: http://www.contraception-esc.com/).

8th European Society of Contraception Congress

The 8th European Society of Contraception Congress will be held from 23–26 June 2004 in Edinburgh, Scotland, United Kingdom. Further details: ESC Central Office, c/o Organised Congress Office, Essenstraat 77, B-1740 Ixelles, Belgium (tel: 32 2 582 08 52; fax: 32 2 582 55 15; email: orgamed.ann@pandora.be and website: http://www.contraception-esc.com/edinburgh.htm).
Improving response rates for self collected urine samples

J Hocking, S Tabrizi, D Jolley, S M Garland and C K Fairley

Sex Transm Infect 2003 79: 346-347
doi: 10.1136/sti.79.4.346-a

Updated information and services can be found at:
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