Television coverage of AIDS and transmission of the AIDS virus

The introduction of the government framework “The national strategy for sexual health and HIV” is necessary for attempts to prevent sexual causes of premature death and ill health, at a time when transmission of HIV is at a record level and heterosexual intercourse is the most commonly reported mode of transmission. However, policy formation alone will not be effective in the fight against AIDS in the United Kingdom. Commentators have noted television coverage of AIDS has declined as the progression of the AIDS epidemic has fallen from media interest. While television coverage raises AIDS awareness and encourages safer sex, it is unclear if it has an impact on infection rates. I have studied the association between television coverage and HIV transmission.

Television programmes with an AIDS specific content broadcast in the United Kingdom between 1981 and 2000 were identified. Criteria for inclusion were UK-wide terrestrial television programmes promoting AIDS awareness and encouraging the practice of safer sex; documenting the life of a person with AIDS; or a dramatisation featuring an AIDS storyline. Televised public education campaigns were excluded from the search, having been reported elsewhere. Television listings data were collected from a Scottish national newspaper (Daily Record) providing UK-wide television listings. Hand searching was conducted on the 1st, 15th, and 28th days of the months of February, May, August, and November, from 1981–2000: a 3.3% sample of available coverage. The sample was augmented by additional sources. A search for programmes shown as part of a one-off media campaign (as reported in the literature); between 27 February and 6 March 1987 to promote AIDS awareness and encourage safer sex was conducted. Supplementary programmes were obtained ad hoc from reviewing The Guardian CD Rom (from 1989–2000) and The End of Innocence: Britain in the time of AIDS.

Thirty seven UK terrestrial programmes covering HIV/AIDS were identified, the earliest in 1983, most recent in 2000, and a peak of programmes in the late 1980s. Between 1996 and 1999 no programmes covering HIV/AIDS were found. Twelve programmes about AIDS were shown during the media campaign of 1987. The decline in television coverage of AIDS contrasts with the steady increase in transmission of HIV (see fig 1 and table on STI website).

The findings of my study are tentative, but hint that television coverage of AIDS has declined as transmission of HIV continues. This complements Nicoll and colleagues’ argument that AIDS campaigns (often televised) are likely to have reduced HIV transmission in the 1980s. Television is a rich source of information about AIDS, offering a powerful while unevaluated medium for promoting AIDS awareness and safer sex to the general public. Given the current low level of media interest and the ceaseless increase of HIV transmission, it may be beneficial to formally evaluate a national media campaign on this important public health issue using quasi-experimental methods.

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A table of television coverage of AIDS 1981–2000 is on the STI website www.stijournal.com/supplemental

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High frequency of antibodies to syphilis and HIV in hepatitis C virus positive blood donors may reflect its sexual transmission in this region

Hepatitis C virus (HCV) infection is a great cause of concern because of the risk of chronicity and other sequelae. Studies on the prevalence behaviour pattern and sexual transmission of HCV infection among the population are required for formulating strategies to control spread of HCV. The aim of this study was to evaluate the occurrence of HCV in voluntary blood donors as they are known to be a high risk population for transmission of these infectious agents. Comparison was made between the presence of syphilis, HIV, and HCV infection in HCV negative and positive blood donors to confirm these as markers or predictors of HIV infection in a high risk population which may reflect the transmission of HCV by a sexual route.

Voluntary blood donors (n = 3905) from New Delhi, India, were randomly recruited between August 2001 to March 2002 and were unpaid. These donors were screened for antibodies to VDRL, hepatitis B surface antigen (HBsAg), hepatitis C virus (HCV), and HIV using previously validated second generation ELISA kits. Data were analysed using the $\chi^2$ test. Odds ratio (OR) was used to measure strength of an association.

![Figure 1](www.stijournal.com)
The antibodies to HCV were detected in 61 (1.56%), HBsAg in 109 (2.79%), VDRL in 132 (3.38%), and HIV in 40 (1.02%) donors. In HCV negative blood donors, VDRL was detected in 129 (3.35%), HBsAg in 106 (2.75%) and HIV in 31 (0.8%) donors. In HCV positive blood donors HCV positivity increased significantly to nine (14.75%) OR (21.2) and VDRL and HBsAg reactivity increased to three (4.91%) (OR 1.82 and 1.48 respectively). Thus HIV, VDRL, and HBsAg reactivity was found 21.2, 1.48, and 1.82 times more often respectively in HCV positive blood donors (table 1).

In conclusion, the high frequency of antibodies to syphilis and HIV in HCV positive blood donors may confirm this as a marker or predictor of HIV infection in a high risk population and may also reflect the risk of transmission of HCV by the sexual route, which seems to vary with the population studied.

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References

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Table 1 Percentage positivity of HBsAg, VDRL, and HIV in HCV positive and negative blood donors

<table>
<thead>
<tr>
<th>HIV +ve (n=61)</th>
<th>HIV -ve (n=3844)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBsAg (%)</td>
<td>3 (4.91)</td>
<td>106 (2.75)</td>
</tr>
<tr>
<td>VDRL (%)</td>
<td>3 (4.91)</td>
<td>129 (3.35)</td>
</tr>
<tr>
<td>HIV (%)</td>
<td>9 (14.75)</td>
<td>31 (0.80)</td>
</tr>
</tbody>
</table>

\[ \chi^2 = 122.8, p<0.001. \]

\[ OR = \text{odds ratio}, \text{HCV positive versus HCV negative blood donors}. \]


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A survey of STI policies and programmes in Europe

The survey by Dehne et al was carried out in 1998–9 and inevitably does not necessarily represent the current situation. The authors report that the United Kingdom has no national STI programme and management guidelines. The first ever national strategy for sexual health and HIV for England was published in 2001. It contains the two key objectives of ensuring that everyone has better access to information on sexual health and to make services more available and accessible to all those who require them at all ages. The strategy also has specific aims of reducing acquisition, transmission, and the prevalence of undiagnosed HIV and STIs. Specific targets to increase the uptake of HIV testing in hepatitis B vaccination have also been set.

The issue of case management guidelines was not detailed in the strategy since this has been specifically tackled by the specialty of genitourinary medicine. A clinical effectiveness group was set up between the Association of Genitourinary Medicine and the Medical Society for the Study of Venereal Diseases in 1997, with the specific remit of producing evidence based national guidelines and standards for those working in the specialty of genitourinary medicine. These were published in STI in 19997 and updated in 2002.1

I hope these observations will help to update the survey with particular reference to England.

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Polymorphic immune restoration syndrome after effective HAART

HAART is able to suppress HIV replication and restore specific immune responses. We and others8 have demonstrated that previously
untreatable progressive multifocal leukoencephalopathy (PML) may improve and has very rarely been reported following HAART. However, CD4+ increase is also associated with inflammatory reactions to previously silent infectious agents, as well as aberrant immune phenomena.1

Case report
A 27 year old HIV/HCV (hepatitis C virus) patient was observed in 1997 with a CD4+ count of 357 cells ×109/l and HIV-RNA >1,000,000 copies/ml. HAART was started and maintained until 1999 when it was stopped at the patient’s request. On October 2001 he returned asymptomatic, with a CD4+ cell count of 34 ×109/l and HIV-RNA 130,000 copies/ml. Stavudine + lamivudine + nevirapine were then started, together with trimethoprim-sulfamethoxazole. He experienced an optimal response (CD4+ count 159 ×109/l, HIV-RNA <50 copies/ml, WBC 5300 ×109/l, RBC 4550 ×109/l, Hb 16.1 g/dl, PLT: 259 ×109/l, eosinophils 8.3%, AST 369 IU/l, ALT 1446 IU/l, γ-GT 594 IU/l, PT 84%, and raised total plasma IgE 330 IU/ml. Toxicplasma serology, HBsAg, HBsAb, and HAV-IgM were negative; Hbc-tgG and HAV-IgG were positive; and HCV-RNA was 172 ×106 copies/ml (baseline 13 ×109/l copies as at October 2001, before HAART). HAART was stopped and liver biopsy performed, showing chronic hepatitis with cholestasis, marked eosinophilia, parenchymal and periportal infiltrates of predominant lymphocytes, and plasma cells suggestive of allergic disease. Cranial magnetic resonance imaging scans revealed subcortical white matter lesions in both frontal lobes and internal capsule with faint peripheral enhancement after gadolinium contrast. Cerebrospinal fluid analysis was normal, fungal and bacterial cultures, cryptococcal antigen, and polymerase chain reaction for HSV-1/2, varicella zoster virus, Epstein-Barr virus, cytomegalovirus, Mycobacterium sp were negative, while JCV-DNA was positive; a cidofovir standard dose was initiated. Subsequently, progressive improvement was observed (fig 1) and PML stabilised.

Comment
We report a complex polymorphic immune restoration syndrome of PML (demonstrated also by JCV-PCR in contrast with previous reports1) combined with the first case of liver damage related to delayed immune mediated hypersensitivity reaction to nevirapine. Several facts support an immunopathogenic role of aberrant immune recovery in this case: (i) these phenomena occurred together in concomitance with raising CD4+ T cell count under effective HAART; (ii) PML during HAART is exceptional; (iii) both PML during HAART5 and nevirapine hypersensitivity6 occurred during the first weeks of therapy in the case series reported so far; (iv) nevirapine hypersensitivity is frequent in conditions of hypersensitivity is frequent in conditions of restoration syndrome of PML (demonstrated combined with the first case of liver damage related to delayed immune mediated hypersensitivity reaction to nevirapine. This underlines that pathogenic mechanisms and immunological factors associated with immune restoration diseases may be diverse and unexpected. Moreover, nevirapine hypersensitivity is immediate while this case of liver damage occurred after a substantial period on nevirapine and with concomitant elevation of γ-GT enzyme and increase in HCV replication. Therefore, it would have been very difficult to make the correct differential diagnosis with either late onset metabolic idiosyncrasy to nevirapine or relapsing HIV hepatitis without liver histology. Thus, avoiding liver biopsy in such cases may lead to an underestimation of the causative role of nevirapine.

Figure 1  Haematocochelical clinical course of the study patient.
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