Short Communication

Incidence of malignant neoplasms among HIV-infected persons in Scotland

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Among 2574 persons diagnosed with HIV throughout Scotland and observed over the period 1981–1996, cancer incidence compared to the general population was 11 times higher overall; among homosexual/bisexual males, it was 21 times higher and among injecting drug users, haemophiliacs and heterosexuals it was five times higher, mostly due to AIDS-defining neoplasms. However, liver, lung and skin cancers (all non-AIDS-defining) were also significantly increased.

The expected numbers of malignant neoplasms occurring among the HIV cohort, assuming the same incidence as that for a matched general population in Scotland, was calculated using age-group, sex, year (for the period 1980–1996) and site-specific data held on the Scottish Cancer Registry. The Standardised Incidence Ratio (SIR) was defined as the ratio of the observed to expected number of neoplasms and the confidence interval calculated assuming that the observed number followed a Poisson distribution.

Knowledge of cancer excess in people with HIV has largely been based on (a) case reports or case series (b) matching population-based cancer registries with AIDS registries or (c) following up specific cohorts of HIV-infected individuals. Large relative risks are reported for AIDS-defining cancers, but the possible relationship between HIV and non-AIDS-defining cancers is less conclusive. In Scotland, a high incidence of many cancers and the existence of two high-quality, national, computerised databases on HIV and cancer incidence afforded an opportunity to adopt a computer linkage approach to examine the occurrence of malignant neoplasms (both AIDS and non-AIDS defining) among all HIV–diagnosed persons across Scotland.

MATERIALS AND METHODS

Records on Scotland’s HIV database were linked with those on the Scottish Cancer Registry to determine the incidence of malignant neoplasms among persons diagnosed with HIV in Scotland. The Scottish HIV Database (Goldberg et al, 1992) holds data on all persons who have tested HIV antibody positive and/or been diagnosed with AIDS in Scotland since the early 1980s; data include date of birth, initials, soundex code of surname, AIDS-defining illness and dates of first HIV antibody positive test, AIDS diagnosis and death. The risk category distribution of the database’s diagnoses is relatively even: 34% injecting drug users (IDU), 34% homosexual/bisexual males, 24% heterosexual and 8% other risk groups. The Scottish Cancer Registry holds a range of other risk groups. The linkage exercise was restricted to the period 1980–1996, the era prior to the full introduction of Highly Active Antiretroviral Therapy (HAART) in Scotland. All malignant neoplasms identified through the linkage process were eligible for inclusion in the analysis, provided they were diagnosed after the HIV diagnosis or preceded it by 2 months or less. This latter criterion allowed for the possibility that the discovery of a malignant neoplasm may have instigated an HIV test and subsequent HIV diagnosis. Links between the two databases were identified using probability linkage techniques and utilising the expertise of the Scottish Record Linkage Group (Kendrick and Clarke, 1993).

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RESULTS

The HIV Database comprised 2574 HIV-infected individuals testing HIV antibody positive between 1981 and 1996, of whom 1955 were male and 619 female. Follow-up to death or 31st December 1996 provided 11 677 and 4236 person-years of follow-up for males and females, respectively. The cohort’s median age at HIV diagnosis was 28 years and comprised 58 children (aged 14 years or less at HIV diagnosis). The linkage process identified 162 malignant neoplasms (two occurring in children). The incidence of malignancy among the HIV-infected population was 11 times that of the general population, SIR = 10.8 (95% CI 9.2, 12.6). The incidence was 21 times higher among homosexual/bisexual males and five to six times higher among the IDU, haemophiliac and heterosexual risk groups (Table 1). Of the 138 AIDS-defining
malignant neoplasms, 82 were non-Hodgkin’s lymphomas, 55 Kaposi’s sarcomas and one was an invasive cervical cancer. The SIRs for the 138 AIDS-defining and 24 non-AIDS-defining neoplasms were 100.0 (95% CI 84.1, 118.1) and 1.8 (95% CI 1.1, 2.6), respectively. Most of the non-AIDS-defining SIRs were of the order 1–4, with the exception of those for liver and bone cancers. The SIRs for liver, lung and skin cancers were statistically significant.

**DISCUSSION**

This study confirmed the large expected excess risks of non-Hodgkin’s lymphoma and Kaposi’s sarcoma among HIV-infected persons; the latter finding is consistent with the previous observation of an increased incidence of Kaposi’s sarcoma during the mid 1980s in Scotland (Brewster et al., 1999). Although invasive cervical cancer is an AIDS indicator disease, only one case was diagnosed; its SIR was 1.7 (0.04, 9.26), a considerably lower ratio than the statistically significant ones associated with cancers of the liver and lungs.

An important finding was the SIR of 1.8 (95% CI 1.1, 2.6) for the non-AIDS-defining malignant neoplasms. Other studies identified a 2-3 fold risk of non-AIDS-defining malignant neoplasms among AIDS patients and an excess of liver cancer (Serraino et al, 2000), lung cancer (Grulich et al, 1999; Frisch et al, 2001) and skin cancers (Cooksey et al, 1999) among HIV-infected persons. Unlike several other studies, we did not find a significant excess of Hodgkin’s disease (Cooksey et al, 1999; Grulich et al, 1999; Serraino et al, 2000; Frisch et al, 2001).

The possibility that some of the non-AIDS-defining neoplasms were extranodal lymphomas, or Kaposi’s sarcomas that had been misdiagnosed or miscoded as other types of cancer, cannot be excluded. Most of the cancers in question, however, were recorded as being microscopically verified and the reliability of diagnostic coding for the Scottish Cancer Registry is high (Brewster et al., 1994).

In this study, it was impossible to associate cancers with immunodeficiency, lifestyle, deprivation or coinfection with another virus. Accordingly, the excess of lung cancer may be due to high rates of smoking among HIV-infected gay men and the excess of liver cancer may be related to HIV/hepatitis coinfection, especially among IDUs. In Scotland, the incidence of many cancers varies by deprivation category; however, deprivation scores were not available for the HIV cohort.

The study method is subject to some potential biases. Persons belonging to the HIV cohort might have been subject to closer clinical scrutiny than those in the general population; this could have led to earlier diagnoses in the former. In contrast, emigration of any HIV-infected persons will have tended to underestimate cancer risk due to an overestimation of person-years at risk and the failure to ascertain and register any subsequent cancers arising in emigrants.

Since the aim of the investigation was to determine the natural history of neoplasia incidence among a national HIV cohort, the linkage was not extended beyond 1996, the time when Highly Active Anti-Retroviral Therapy (HAART) became widely administered in Scotland. The investigators plan to repeat the linkage to determine how much the incidence of neoplasia has changed during the era of HAART.

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**Table 1** Observed malignant neoplasms by HIV Risk Group, AIDS-defining status, site and morphology; SIR and 95% CI (SIR)

| HIV risk group | AIDS/non-AIDS defining | Cancer site | Morphology | N    | SIR  | 95% CI SIR |
|---------------|------------------------|-------------|------------|------|------|------------|
| All risks     | AIDS and non-AIDS      | All         | All        | 162  | 10.8 | 9.2, 12.6  |
| Heterosexual  | AIDS and non-AIDS      | All         | All        | 14   | 5.9  | 3.2, 9.9   |
| Homosexual    | AIDS and non-AIDS      | All         | All        | 102  | 21.4 | 17.4, 26.1 |
| IDU           | AIDS and non-AIDS      | All         | All        | 35   | 5.5  | 3.8, 7.6   |
| Haemophilic   | AIDS and non-AIDS      | All         | All        | 5    | 6.1  | 2.0, 14.2  |
| Other/mixed   | AIDS and non-AIDS      | All         | All        | 6    | 9.0  | 3.3, 19.6  |
| All risks     | AIDS                    | All         | All        | 138  | 100  | 84.1, 118.1|
| All risks     | AIDS                    | Skin (45), Connective and soft tissue (5), Head and Neck (4), Lung (1) | Kaposi’s sarcoma | 55  | 2750 | 2067, 3583 |
| All risks     | AIDS                    | Not available | Non-Hodgkin’s lymphoma | 82  | 108  | 85.6, 134.4|
| All risks     | AIDS                    | Cervical    | Squamous cell carcinoma | 1   | 1.7  | 0.04, 9.26 |
| All risks     | Non-AIDS                | All         | All        | 24   | 1.8  | 1.1, 2.6   |
| All risks     | Non-AIDS                | Skin        | Squamous cell carcinoma (3), Basal cell carcinoma (1), Unspecified (2) | 6   | 2.8  | 1.04, 6.2  |
| All risks     | Non-AIDS                | Lungs       | Squamous cell carcinoma (1), Small-cell carcinoma (1), Adenocarcinoma (2), Unspecified (1) | 5   | 4.1  | 1.3, 9.5   |
| All risks     | Non-AIDS                | Liver       | Hepatocellular carcinoma | 2   | 22.0 | 2.7, 80.2  |
| All risks     | Non-AIDS                | Bladder     | Transitional cell carcinoma (1), Papillary cell carcinoma (1) | 2   | 4.2  | 0.5, 15.0  |
| All risks     | Non-AIDS                | Brain and CNS | Astrocytoma | 2   | 3.3  | 0.4, 12.0  |
| All risks     | Non-AIDS                | Not available | Hodgkin’s disease | 2   | 3.6  | 0.4, 13.1  |
| All risks     | Non-AIDS                | Bone        | Ewing’s sarcoma | 1   | 9.1  | 0.23, 50.5 |
| All risks     | Non-AIDS                | Head and neck | Unspecified | 1   | 1.6  | 0.04, 8.8  |
| All risks     | Non-AIDS                | Leukaemias  | Acute myeloid leukaemia | 1   | 2.2  | 0.06, 12.4 |
| All risks     | Non-AIDS                | Stomach     | Unspecified | 1   | 2.9  | 0.07, 15.9 |
| All risks     | Non-AIDS                | Testis      | Seminoma | 1   | 0.7  | 0.02, 3.75 |

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