Evaluation of non-HIV-related, drug-sensitive cluster outbreaks of tuberculosis with PCR-based DNA fingerprinting

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OBJECTIVES: To characterize two cluster outbreaks of tuberculosis by DNA fingerprinting and to evaluate the possibility of their association with human immunodeficiency virus (HIV) infection and multiple drug-resistant Mycobacterium tuberculosis.

SETTING: Two clusters of tuberculosis cases in adjoining aboriginal Canadian reserves and a control population.

METHODS: All cases of tuberculosis diagnosed in the two communities and a number of control isolates were fingerprinted using a novel simplified DNA-based technique. Demographic data, purified protein derivative (PPD) skin test response to 5TU PPD, type of disease and HIV serology were also evaluated. Positive cultures were assessed for susceptibility to antituberculous drugs.

RESULTS: Two distinct clusters with five cases in one community and 21 cases in the second were identified. Isolates from both communities differed in their DNA pattern but were identical within communities. Thirteen cases had bacteriologically proven disease and grew susceptible organisms. Serological analysis for HIV infection was carried out in 15 of 18 adults, and all were negative.

CONCLUSIONS: The study shows the potential for the rapid transmission of tuberculosis infection in HIV-negative subjects with susceptible organisms. It further highlights the usefulness of DNA fingerprinting for molecular epidemiology in evaluating particular outbreaks.

Key Words: Cluster outbreaks, PCR-based fingerprinting, Tuberculosis

Évaluation de flambées de cas isolés de tuberculose non liée au VIH et sensible à la médication antituberculeuse, par empreinte génétique de l’ADN basée sur la technique de la PCR

OBJECTIFS : Caractériser deux flambées de cas isolés de tubercu-voir page suivante

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There is concern regarding an increased prevalence of tuberculosis (TB) in both developed and developing countries (1). Much of this increase is thought to be associated with human immunodeficiency virus (HIV) infection (2). In the United States multidrug-resistant TB (MDRTB) has particularly been a problem (3). Recent reports of studies using DNA-based technology have shown rapid transmission of infection and development of MDRTB in the presence of HIV infection (4). In addition, population-based molecular epidemiology studies have shown greater transmission of disease than previously thought (5,6).

Among aboriginal Canadians there is a high prevalence of TB – approximately 10 times the rate among the general population (90/100,000 versus nine/100,000 population, respectively) (7). So far MDRTB and HIV infection have not been major problems in this group (8), although recent trends indicate that this may be changing (9). We recently investigated a number of cluster outbreaks of TB in this population group (10); two were geographically close, occurring approximately 5 km apart.

We were interested in establishing the prevalence of HIV infection in both clusters, the drug susceptibility patterns, and finally whether there was any link between the two outbreaks by using a novel DNA-based technique, which has been recently described (11).

**PATIENTS AND METHODS**

All cases of TB diagnosed in both communities between January and December 1992 were identified. Demographic data, risk factors for TB, response to purified protein derivative of tuberculin (5TU PPD), chest x-ray findings, HIV serology, bacteriological results and drug susceptibility patterns were collected. Sputum and pleural fluid were cultured using the Bactec technique. Specimens that were positive on culture for *Mycobacterium tuberculosis* were processed for strain identification.

A newly described method using a ligation-mediated polymerase chain reaction (PCR) procedure was used to amplify the flanking sequences on both sides of the insertion sequence 6110 from restriction-digested chromosomal DNA. This method has recently been described in detail (11). Briefly, this process involves inactivating the mycobacteria with phenol and breaking open the mycobacteria by vortexing in the presence of glass beads. The debris are then pelleted and the DNA supernatant is purified by adherence to glass powder. After restriction enzyme digestion a linker is ligated to the cut ends. Primers homologous to the insertion sequence 986 and the linker are then used in PCR and the products run on an agarose gel. The results are characteristic patterns of multiple bands of different lengths depending on the distance from the insertion sequences to the external restriction sites.

**RESULTS**

Two distinct clusters with five cases in one community and 21 cases in the second were identified. Isolates from both communities differed in their DNA pattern but were identical within communities. Thirteen cases had bacteriologically proven disease and grew fully sensitive organisms. Serological analysis for HIV infection was carried out for 15 of 18 adults; all were negative. All subjects were interviewed by one investigator, and the contact evaluation records for each outbreak were reviewed in the British Columbia Centre for Disease Control, Tuberculosis Division.

**Cluster A:** Both index and source case in cluster A was a 22-year-old male. Details of other cases diagnosed within the community are given in Table 1. He presented with a self-reported cough of six weeks’ duration in May 1992, but a history of coughing for at least six months was subsequently obtained from family members. He denied a history of intravenous drug use. In this community there were 21 cases (12 male). All but four had symptoms; 13 had documented PPD conversion, having been negative on PPD skin testing at initial screening in the community or having previously been known to be negative from prior tuberculin records. Follow-up PPD skin tests were carried out when subjects presented with symptoms or where follow-up PPD skin testing at least three months after the initial screening was carried out. Recently documented PPD conversion indicated recent infection. Seven others had a positive PPD to 5 TU at presentation. HIV serology was completed in 12 of 15 adults and was negative in all. Disease was confirmed by mycobacteriology in 10 cases and all had drug-susceptible organisms.

**Cluster B:** The index case in cluster B was an infant aged 20
months who presented with recurrent pneumonia. Characteristics of the other cases diagnosed in this community are shown in Table 2. Eventually the infant had gastric washings obtained for mycobacteriology and these grew M tuberculosis. Reverse contact tracing identified the father of the infant’s babysitter as the source case. He had bilateral apical abnormalities noted 18 months previously but at that time he refused to attend for further evaluation. He had a history of intravenous drug use. Three further cases, all asymptomatic, were identified in contact tracing. Two had PPD conversions and one had a positive PPD at evaluation. Three subjects had susceptible, bacteriologically proven disease. The remaining subjects had primary disease. The three adults in this cluster were HIV-negative.

**DNA fingerprinting:** The isolates from both communities were found to differ, showing two unique patterns. Twelve control specimens, drawn at random from cases diagnosed throughout the same period and evaluated at the same time, were all different. The exceptions were isolates from two HIV-negative brothers who had the same PCR pattern, which was different from those in the clusters. The control specimens were obtained to provide internal validity, further blinding the laboratory to the epidemiological data. Gel electrophoresis patterns produced by a sample of control and cluster case isolates from cluster A are shown in Figure 1.

**DISCUSSION**

TB is reemerging as a major health concern in both developed and developing countries (1). Traditionally it has been thought that the majority of newly diagnosed cases represented reactivation of previously acquired infection (12), although exogenous reinfection with a new organism has been described using phage typing (13). New insight into the transmission of TB infection has been gained by evaluating clusters of HIV-associated cases in whom infection with a resistant organism has been acquired while on therapy for drug-susceptible disease (4).
The clustering of cases among aboriginal patients is likely multifactorial and, although our data do not address the issue of housing, it is likely that better housing and community education regarding TB would facilitate a reduction in the overall rates in these communities.

Although much of our attention has focused on HIV-related TB, this report highlights the rapid progression to disease that can occur in the absence of HIV infection and with fully susceptible organisms. The availability of DNA fingerprinting of responsible organisms provides a useful epidemiological tool for evaluating such outbreaks and casts new light upon the pathogenesis of TB.

REFERENCES


In addition, population-based studies using restriction fragment length polymorphism as a fingerprinting technique have indicated that between 30% and 40% (6) of all newly diagnosed cases of disease are related. This indicates that, contrary to the traditional belief that approximately 10% of new cases are due to recent acquisition of infection and progression to disease, much more infection transmission and progression to disease occurs than heretofore thought.

We report two separate clusters of TB cases with a total of 27 cases of active TB. Earlier non-HIV-related studies have tended to concentrate on methodological issues (14-16) and distinguishing among strains, but they have lacked a large number of similar isolates and provided no epidemiological information.

The speed at which infection occurred and cases of active TB disease developed can be inferred from the chronological presentation of the active cases, especially in cluster A, following the diagnosis. Our data highlight the rapidity with which infection can be transmitted and disease develop even in non-HIV-infected subjects. This rapidity is further supported by the high proportion of subjects who had a conversion of their Mantoux PPD skin test responses while being evaluated, indicating that infection had recently been acquired. The two distinct patterns in the clusters support two different sources for the two outbreaks.

Paradoxically the source case in cluster A reported the shorter history of symptoms but generated the larger number of cases. Although the frequent communal events on a North American Indian reserve and the generally poor standard of housing may have facilitated the rapid transmission of infection, the total number of cases identified is significant. It highlights the importance of early diagnosis of TB and prompt initiation of therapy (17). Because of the potential for transmission of infection within the community, it is possible that the homogeneity of isolates evaluated from this cluster outbreak could be due to a tendency of isolates on aboriginal reserves to be similar. This hypothesis is disproved by the heterogeneity of isolates from a number of Northern Alberta communities (unpublished data).

A corollary of these data is that in cluster situations, as described, the risk of progression from infection to disease in certain epidemiological situations is that much greater and reinforces the need to consider prompt chemoprophylaxis (18). It highlights the importance of promptly evaluating contacts of all active cases of TB and promptly initiating chemoprophylaxis. For one patient in cluster A there was a two-month delay between having a normal chest x-ray and PPD conversion, and being offered chemoprophylaxis and initiating it. A short time after chemoprophylaxis was started the subject presented with a pleural effusion. The physician who assessed her did not consider TB as a possibility and continued the patient on isoniazid while the presumptive diagnosis of a parapneumonic effusion was being evaluated. Six weeks later the patient presented with a TB empyema which grew a fully sensitive organism. After decortication and a full course of anti-TB therapy there was complete resolution of the abnormalities.

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Figure 1) Mycobacterial DNA patterns after ligase-mediated polymerase chain reaction. Lane 1 is an unrelated control, lanes 2 to 12 are from patients identified as epidemiologically related to cluster 1 and lane 13 is a molecular weight (MW) marker. Lanes 2 to 10 have identical patterns.


