Reduced carbon monoxide transfer factor (TLCO) in human immunodeficiency virus type I (HIV-I) infection as a predictor for faster progression to AIDS

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Abstract
Background-In addition to the acute fall in carbon monoxide transfer factor (TLCO) associated with Pneumocystis carinii pneumonia (PCP) or other opportunistic lung infections, reduced TLCO occurs in HIV-I seropositive individuals without active pulmonary disease. Abnormal TLCO, in the absence of lung disease, may be a surrogate marker of HIV-I induced immunosuppression and, therefore, a predictor for a more rapid progression to AIDS.

Methods-Eighty four individuals with AIDS, who had regular pulmonary function tests before the diagnosis of AIDS was made, were identified from a cohort of patients with HIV-I infection. None had evidence of active pulmonary disease at the time of initial pulmonary function testing. The relation between the time taken to progress to AIDS and initial pulmonary function tests was examined with life table survival analysis.

Results-Patients with a TLCO value of <80% of predicted normal (n = 46) progressed significantly faster to AIDS, with a median time of 8-0 months compared with 16-5 months for those with a TLCO value of ≥80% (n = 38). When stratified by AIDS defining diagnosis (PCP or non-PCP), median time to PCP was also significantly related to initial TLCO values (TLCO of <80% = 9-0 months, TLCO of ≥80% = 19-0 months). Reductions in other measurements of lung function (FEV1, FVC, Kco) were not temporally associated with the development of AIDS.

Conclusions-HIV-I seropositive individuals with TLCO values of <80% predicted and no evidence of lung disease progress more rapidly to AIDS than those with TLCO values of ≥80%.

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The use of simple pulmonary function tests is a non-invasive method of investigating HIV-I infected individuals with respiratory symptoms and is helpful for assessing the presence and extent of respiratory disease, the need for further investigation, and the response to treatment. An acute reduction in carbon monoxide transfer factor (TLCO) is the most sensitive test determining lung disease in HIV-I infection and AIDS. However, it lacks specificity for the development of Pneumocystis carinii pneumonia (PCP), by far the most important pulmonary complication of HIV-I infection, where a low or falling TLCO has been used as a screening test. The reduction in TLCO that may occur with PCP is a result of alveolar capillary block and improves with treatment of the acute episode. A reduction in TLCO may also occur in HIV-I infection in the absence of overt pulmonary disease. Lower values of TLCO are seen in patients with more advanced HIV-I disease despite the absence of new pulmonary symptoms or radiographic changes. The reduction in TLCO in this instance is poorly understood, but has been previously suggested to relate either to a specific HIV mediated alveolitis or to reflect pulmonary injury caused by non-specific inflammatory events occurring on a background of HIV induced immunosuppression. If, indeed, reduced TLCO in HIV-I infection represents compromised pulmonary defence associated with progressive immunosuppression, abnormal results in pulmonary function tests may equate with a worse prognosis and be valuable as a predictor of advancing disease.

In this study we have examined the role of TLCO in HIV-I infection as a marker of disease progression, rather than its more usual role as an indicator of disease presence and severity. We have correlated the length of time to the development of AIDS with initial pulmonary function data obtained after HIV-I seroconversion but before the onset of AIDS, and have shown that a reduced TLCO value identifies a group of patients who progress more rapidly to AIDS.

Methods
Between 1986 and 1991, all HIV-I seropositive individuals at St Mary's Hospital, London were invited to join a patient cohort to evaluate the long term trends in pulmonary function during HIV-I infection. The vast majority (80–90%) of seropositive individuals seen at St Mary's Hospital are homosexual/bisexual men. HIV-I seropositive individuals were recruited to the cohort from the genitourinary medicine outpatient clinic or on discharge from the HIV ward if they
had been admitted there for investigation or
treatment. HIV-I seropositive individuals at
all Centre for Disease Control (CDC)11 clin-
cial stages up to, and including, AIDS were
eligible for enrolment to the cohort. The
study was approved by the hospital ethical
committee and all patients gave informed
consent before participation. Details of the
long term changes in pulmonary function
that occurred during the study are reported
elsewhere.12

All individuals from the cohort who had
entered at a CDC clinical stage before AIDS,
and who subsequently progressed to AIDS,
were identified for this analysis. The time
interval between initial pulmonary function
tests, performed on entry to the cohort at a
defined CDC clinical stage, and the develop-
ment of AIDS was correlated with the results
of the pulmonary function tests by life table
analysis (Statistical Package for the Social
Sciences [SPSS-X]). Median times to the
development of AIDS were compared (log
rank χ2) between patients whose initial pul-
monary function test results were either
greater than or equal to, or less than 80% of
predicted normal. Predicted values for sex,
age, ethnicity, and height were obtained from
standard references.13 The AIDS defining
diagnosis was subclassified to PCP or “non-
PCP”—that is, any AIDS defining diagnosis
other than PCP (such as Kaposi’s sarcoma,
cerebral toxoplasmosis).

The demographic details, CDC clinical
stage, and smoking habits of the patients
were determined at the time of the initial pul-
nary function tests. Physical examination
and pulmonary radiography were also per-
formed at this time. All patients underwent the
following tests: forced expiratory volume in
one second (FEV1), forced vital capacity
(FVC), alveolar volume (VA), and TLCO. The
diffusion coefficient (Kco) was derived from
the TLCO divided by the VA. Measurements of
FEV1 and FVC were made with a dry bellows
spirometer and the TLCO and VA were deter-
mained by the single breath helium dilution
method (PK Morgan Transfer Test Model C
Machine; PK Morgan, Gillingham, Kent).

Corrections for temperature and haemoglo-
bin13 were included when calculating values
for TLCO, and all measurements were made after
a period of at least 30 minutes rest. All smok-
ers were requested to abstain from smoking on
the day of their tests. To ensure accuracy of
the results over time, the TLCO machine was
calibrated daily with standard gases, and to
confirm that the equipment was capable of
making normal measurements in normal indi-
viduals, TLCO values obtained on the dedicat-
ed machine used in this study were compared
weekly with values of TLCO obtained on a sim-
ilar machine by the pulmonary function tech-
icians acting as biological disease free
controls. The risk of cross infection between
patients was reduced by modifications to the
equipment as described previously.1

All patients who had joined the cohort with
reduced TLCO values (<80% predicted nor-
mal), respiratory symptoms, or radiographic
abnormalities were investigated for an under-
lying cause by induced sputum analysis or
fibroptic bronchoscopy and bronchoalveolar
lavage to obtain specimens for microbiologi-
cal and cytological investigation.14 They
were not included in this study if found to have a
reduced TLCO because of identifiable pul-
nary disease—for example, PCP, pul-
nary Kaposi’s sarcoma, pneumococcal
pneumonia. Patients were similarly excluded
if an AIDS diagnosis was made less than one
month after the initial pulmonary function
tests. By these means we ensured that all
patients studied who had a reduced TLCO
had no evidence of active or acute pulmonary
disease at the time of the initial tests.

Results
Between 1986 and 1991, 516 individuals
with HIV-I infection had pulmonary function
tests performed at a defined clinical stage
of their HIV-I illness and joined the cohort to
evaluate long term changes in pulmonary
function. Those in the cohort had risk factors
for HIV-I seropositivity that mirrored those of
the population seen at St Mary’s Hospital
during this time. In all, 313 seropositive
individuals without an AIDS defining diagnosis
joined the cohort and 203 already had AIDS
at enrolment. Of the 313 HIV-I seropositive
individuals who joined the cohort before the
diagnosis of AIDS, 84 (two women, 82 men)
with a mean age of 36-4 (range 23–59) years
subsequently developed AIDS and addition-
ally fulfilled the criteria for analysis as
described above.

Of these 84 evaluated patients, two were
intravenous drug users, 81 were homosexual
or bisexual men, and one of the female
patients seroconverted after a sexual assault
and had no other risk factors for HIV-I infec-
tion. The baseline pulmonary function data
for the group, expressed as mean (SD) %
predicted normal, was as follows: FVC,
96-9% (18-2%); FEV1, 102-8% (19-8%); TLCO,
78-3% (18-7%); Kco, 87-9% (16-8%).

Of the 84 patients, 21 had asymptomatic
HIV-I infection (CDC group 2), 25 had
persistent generalised lymphadenopathy
(CDC group 3), and 38 had constitutional
symptoms (CDC group 4a). No patients were
in CDC group 1 (acute HIV-I infection).
Although there was a downward trend in
TLCO values for patients in CDC group 2
compared with those in group 4a, the differ-
ce was not significant (table 1).

There was no significant difference in the
number of smokers (n = 43) and non-smok-

<table>
<thead>
<tr>
<th>CDC classification</th>
<th>TLCO</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 2</td>
<td>83-3 (19-4)</td>
<td>21</td>
</tr>
<tr>
<td>Group 3</td>
<td>81-4 (17-8)</td>
<td>25</td>
</tr>
<tr>
<td>Group 4a</td>
<td>73-9 (18-4)</td>
<td>38</td>
</tr>
</tbody>
</table>
Table 2  Median progression times in months to AIDS related to initial values of FEV₁, FVC, KCO, and TLCO

<table>
<thead>
<tr>
<th>Initial result</th>
<th>&lt;80% predicted</th>
<th>≥80% predicted</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>FVC</td>
<td>11.5 (n = 11)</td>
<td>9.7 (n = 73)</td>
<td>NS</td>
</tr>
<tr>
<td>FEV₁</td>
<td>5.5 (n = 9)</td>
<td>10.1 (n = 75)</td>
<td>NS</td>
</tr>
<tr>
<td>KCO</td>
<td>8.8 (n = 27)</td>
<td>10.2 (n = 57)</td>
<td>NS</td>
</tr>
<tr>
<td>TLCO</td>
<td>8.0 (n = 46)</td>
<td>16.5 (n = 38)</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

TLCO—forced vital capacity; FEV₁—forced expiratory volume in one second; KCO—diffusion coefficient for carbon monoxide; TLCO—carbon monoxide transfer factor. p values determined by log rank χ² test.

Figure 1  Life table analysis of progression of patients to AIDS related to initial TLCO.

Table 3  Median progression times to AIDS stratified by initial TLCO and by AIDS diagnosis.

<table>
<thead>
<tr>
<th>TLCO (%)</th>
<th>Diagnosis</th>
<th>Progression time (months)</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;80</td>
<td>Non-PCP</td>
<td>8.0</td>
<td>16</td>
</tr>
<tr>
<td>≥80</td>
<td>Non-PCP</td>
<td>10.0</td>
<td>18</td>
</tr>
<tr>
<td>&lt;80</td>
<td>PCP</td>
<td>9.0</td>
<td>30</td>
</tr>
<tr>
<td>≥80</td>
<td>PCP</td>
<td>19.0</td>
<td>20</td>
</tr>
</tbody>
</table>

Figure 2  Life table analysis of progression of patients to AIDS stratified by initial TLCO and by AIDS diagnosis (Pneumocystis carinii pneumonia (PCP) or non-PCP).

ers (n = 41) in each CDC category or in the mean (SD) TLCO values between the two groups at the time of their initial pulmonary function tests (smokers, mean (SD) TLCO = 75.7% (16.6%) predicted; non-smokers, mean (SD) TLCO = 80% (20.5%) predicted; t test, p > 0.05).

Life table analysis revealed no significant difference in progression time to AIDS when normal and abnormal initial values of FEV₁, FVC, or KCO were compared (table 2). Time to progression to AIDS was, however, significantly related to initial TLCO; for patients with TLCO <80% predicted, median progression time to AIDS was 8.0 months and for those with TLCO ≥80% predicted, median progression time to AIDS was 16.5 months (log rank χ², p < 0.001) (table 2). Life table curves for patients based on initial TLCO measurements are shown in fig 1.

Of the 84 patients, 50 had PCP as their AIDS defining diagnosis (PCP diagnosis), and 34 had either Kaposi's sarcoma or non-pulmonary opportunistic infections (non-PCP diagnosis). The median time to development of AIDS when caused by PCP was significantly related to initial TLCO values (TLCO <80%, median progression time to AIDS = 9.0 months; TLCO ≥80%, median progression time to AIDS = 19.0 months; p < 0.005). Median time to non-PCP AIDS may relate to initial TLCO values but the trend was not significant (TLCO <80% = 8.0 months; TLCO ≥80% = 10.0 months; p > 0.05) (fig 2; table 3).

Discussion
Abnormalities of pulmonary function tests are common in patients with HIV-1 infection. Reduction in TLCO is associated both with PCP and other acute pulmonary diseases, but occurs also in HIV-1 infected individuals without obvious clinical lung disease. The mechanism for reduced TLCO in HIV-1 infection in the absence of overt pulmonary disease is not understood, but does not seem to be due to HIV-1 mediated alveolitis, as previously proposed, nor is it related to the presence of HIV-1 in lung macrophages.

When a reduction in TLCO heralds the development of PCP, this reduction is due predominantly to a fall in the membrane diffusion capacity (DM) component of TLCO. This has been attributed to thickening of the diffusion pathway between alveolar gas and pulmonary capillary blood by the attachment of Pneumocystis trophozoites to the epithelial surface—the so-called "alveolar capillary block". This is in contrast to the chronic reduction in TLCO seen in HIV seropositive individuals who have no evidence of acute or active pulmonary disease. In this case the reduction in TLCO is related to the combination of reduced DM and a reduction in pulmonary capillary blood volume (VC). In addition, there may be an accompanying fall in KCO and FVC. This would indicate a product of both reduced diffusion of carbon monoxide and loss of lung volume.

In this study we have shown that a reduced
TLCO in the presence of HIV-I infection and the absence of lung disease is associated with a more rapid progression to AIDS and, in particular, to the development of PCP. Other pulmonary function measurements which may be abnormal in those who are infected with HIV-I—for example, FEV₁, FVC, and KCO—did not predict the development of AIDS. By excluding all patients who developed AIDS within one month of their initial pulmonary function tests, and by investigating fully those patients with abnormal TLCO and respiratory symptoms or new radiographic changes, we ensured that a reduced TLCO was not related to identifiable pulmonary disease, and, in particular, early "smouldering" PCP. The sensitive technique of the polymerase chain reaction has shown that patients without clinical evidence of PCP do not have a positive amplification signal to P carinii DNA, implying that PCP is a discrete infectious event and is not preceded by an indeterminate prodrome which might lead to a fall in TLCO.

In this study we compared the progression time to AIDS between individuals with pulmonary function of either greater than or equal to, or less than 80% predicted normal. The cut off point of 80% was chosen as the value generally accepted to represent the distinction between "normal" and "abnormal" pulmonary function, and the value most often used in descriptive studies of lung function in HIV-I infected individuals. This is an arbitrary choice and, although this approach has little statistical basis in ascribing normality to pulmonary function, it has the advantage that abnormalities of lung function can readily be classified for individual patients in terms of impairment or disability, and can be appreciated readily by patients and their physicians.

We have recently reported that, in HIV-I seropositive patients without lung disease who have reduced TLCO, pulmonary high resolution computed tomography, technetium-99m labelled diethylenetriamine pentaacetic acid (99mTc-DTPA) lung scanning, exercise testing with transcutaneous oximetry, and measurement of both the alveolar membrane component (DM), and pulmonary vascular component (VC) of the TLCO provides no further information as to the cause of the reduction in TLCO. We have also shown that the detection of HIV-I by the polymerase chain reaction in bronchoalveolar lavage cells has no significant effect on pulmonary function, arguing against a direct HIV-I effect as the cause of reduced TLCO. In the absence, therefore, of pulmonary disease such as infection or neoplasm, a low TLCO during HIV-I infection may relate to inflammatory episodes occurring in the lung on a background of advancing immunosuppression. By reflecting this lung injury, a reduced TLCO may act as a marker of more advanced HIV-I infection. This concept is supported by the previously observed fall in TLCO with advancing immunosuppression, and would explain the power of TLCO measurement in predicting those patients at greater risk of a faster progression to AIDS. Since the KCO did not predict the development of AIDS, it would imply that loss of lung volume—that is, the area available for gas exchange—is the major determinant of the predictive capacity of TLCO.

We have shown, therefore, that a reduced TLCO in HIV-I infection, before an AIDS diagnosis, identifies a group of patients who progress more rapidly to AIDS. This reduction in TLCO during HIV-I infection may represent subtle lung inflammation associated with increasing immunosuppression, pathophysiologically different from the acute fall in TLCO during PCP as a result of alveolar capillary block. Further investigation of the potential of TLCO as a surrogate marker of advancing PCP and, although not predicting specific pulmonary infection is required. Since therapeutic decision making in HIV-I infection (such as when to start zidovudine treatment or chemoprophylaxis for PCP) is commonly made on the basis of deteriorating immunocompetence, the clinical usefulness of TLCO in predicting a more rapid development of AIDS should be compared with established indices of disease progression such as CD4+ cell count and P24 antigenemia.

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