Diagnosis of *Pneumocystis carinii* pneumonia in HIV antibody positive patients by simple outpatient assessments

Don E Smith, Alastair Forbes, Sandra Davies, Simon E Barton, Brian G Gazzard

Abstract

**Background** As increasing numbers of patients with immunosuppression induced by the human immunodeficiency virus (HIV) present with respiratory symptoms it is important to differentiate *Pneumocystis carinii* pneumonia from other chest diseases rapidly and start treatment early. The management of pneumocystis pneumonia could be improved if clinicians could diagnose this condition confidently on the basis of simple clinical assessments.

**Methods** Three hundred and eighteen patients with evidence of immunosuppression due to HIV infection and suspected pneumocystis pneumonia were investigated. A clinical history was taken and arterial blood gas analysis, chest radiography, oximetry during exercise, and sputum induction or bronchoscopy (or both) were performed.

**Results** Pneumocystis pneumonia was confirmed microscopically from induced sputum or bronchoalveolar lavage fluid in 154 patients; 118 had other chest disease. The remaining 46 patients had no definitive diagnosis. The best single independent predictors of a diagnosis of pneumocystis pneumonia were exercise induced oxygen desaturation and obvious interstitial infiltrates on the chest radiograph (odds ratios of 4.88 and 5.44 respectively). The symptom triad of exertional dyspnoea, cough, and fevers; the absence of pneumocystis pneumonia prophylaxis; and resting arterial hypoxaemia were less predictive (odds ratio 2.07, 3.72, and 0.69). An algorithm was developed that gave a positive predictive value for confirmed pneumocystis pneumonia of 95% and also identified those patients with a very small chance of having pneumocystis pneumonia (negative predictive value 85%).

**Conclusions** The diagnosis of an initial episode of pneumocystis pneumonia can be confidently made in a large proportion of immunosuppressed patients with respiratory symptoms on the basis of clinical symptoms, the absence of prophylaxis, chest radiographic appearances, and oxygen desaturation during exercise as shown by oximetry. Using these simple features clinicians can rapidly assign patients to the appropriate type of management at presentation.

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*Pneumocystis carinii* pneumonia remains the most common opportunistic infection seen in patients with the acquired immunodeficiency syndrome (AIDS), occurring mainly in patients with strong laboratory and clinical evidence of immunosuppression. A definitive diagnosis of pneumocystis pneumonia can be made only by identifying this organism in induced sputum samples, which is time consuming, or in specimens obtained at bronchoscopy, which has known risks. Some non-invasive screening tests, including lung function testing, technetium-99m DTPA scanning, and exercise oximetry, have been developed to reduce the need for these investigations.

We have investigated whether simple information and investigations, which could be available at presentation, might obviate the need for definitive tests in a large proportion of patients by defining those in whom the diagnosis is extremely unlikely and those with such a high chance of having pneumocystis pneumonia that further investigation would be superfluous.

**Methods**

**Investigations**

A prospective assessment of all patients presenting to this hospital with suspected pneumocystis pneumonia was conducted. Patients were assessed either at a specialised HIV outpatient centre (Kobler Centre, St Stephen’s Clinic) or during admission through the accident and emergency department. All patients with respiratory symptoms were evaluated if they had clinical markers of immunosuppression (oral candidiasis, oral hairy leucoplaikia, or Kaposi’s sarcoma), were known to have a recent CD4 (T helper lymphocyte cell) count below 0.2 × 10⁹/l, or any previous HIV related illness. We included both those known to be HIV antibody positive and those presenting for the first time and suspected to be positive on the basis of clinical signs and symptoms of immunosuppression (if it was not known whether a patient had HIV antibodies and he subsequently declined an HIV antibody test or had a negative result he was excluded).

Information on exertional dyspnoea, cough,
and fevers over the previous two weeks and on whether a recognised form of pneumocystis pneumonia prophylaxis had been taken over the preceding three months (nebulised pentamidine, oral co-trimoxazole, dapsone, or Fansidar (pyrimethamine-sulfadoxine)) was recorded on a case record form at the time the patient presented by either a doctor (DS) or a research nurse (SD). Patients then had chest radiography, arterial blood gas sampling, and exercise oximetry\(^9\) (performed immediately after chest radiography in patients seen in the outpatient clinic and within 48 hours in those patients admitted through the accident and emergency department).

Sputum induction and bronchoscopy (when sputum induction gave a negative result) were performed over the next five days by standard techniques. All the samples obtained were stained by Giemsa and silver (Grocott) and by immunofluorescent antibody stains\(^9\) (from mid 1988 onwards).

Routine sputum, induced sputum, and bronchoalveolar lavage samples were cultured for bacteria, fungi, and mycobacteria by standard methods.

### Definitions

The following were defined as positive results for the purpose of analysis:

**Predictive clinical criteria**

1. A history of increasing dyspnoea, cough (dry or productive), and fevers within the preceding two weeks.
2. The absence of any pneumocystis pneumonia prophylaxis within the preceding three months.
3. An arterial oxygen tension of less than 11.0 kPa while the patient was breathing room air (normal range 11.7–15.3 kPa).
4. A chest radiographic appearance of obvious diffuse perihilar or interstitial infiltrates. Equivocal changes, focal abnormalities, and normal chest radiographs were defined as negative.
5. A fall in arterial oxygen saturation (\(S\text{ao}_2\)) to 90% or below during standard testing. \(S\text{ao}_2\) was measured by an Ohmeda BIOX 3700 pulse oximeter during 10 minutes of exercise on an exercise cycle set to a standard resistance of 2 kJ (150 watts) and a standard speed of 20 km/h (70 rev/min).\(^8\) Patients unable to complete the full 10 minutes' exercise through coughing or muscular weakness whose \(S\text{ao}_2\) did not fall to 90% were analysed as though they had a negative result.

**Final diagnosis**

Confirmed pneumocystis pneumonia was defined as the presence of more than five typical cysts of \(P\) carinii in an induced sputum or bronchoalveolar lavage sample.

"Clinically suspected pneumocystis pneumonia" was defined on the basis of suggestive symptoms that responded to standard anti-pneumocystis pneumonia treatment when no other diagnosis was uncovered but \(P\) carinii organisms were not found in induced sputum or bronchoalveolar lavage fluid.

Pneumocystis pneumonia was considered to be excluded in those patients with a negative induced sputum or bronchoscopy sample who did not require pneumocystis pneumonia treatment within the following three months, and in those found to have other disease and no evidence of pneumocystis pneumonia. Pulmonary Kaposi's sarcoma was diagnosed on the basis of typical bronchoscopic appearances.

### Statistical Analysis

The \(\chi^2\) test with Yates's correction was used to compare differences in proportions for categorical data. Differences in continuous data were evaluated by Student's \(t\) test (EPI-Info 5.0 software package). Odds ratios were used to estimate the risk of pneumocystis pneumonia associated with the presence of each variable. Multivariate logistic regression analysis was used to identify the independent predictive risk of pneumocystis pneumonia for each variable (Statistical Analysis Software). Sensitivities, specificities, and positive predictive values were calculated for each variable and for combinations of variables.

### Results

Of the 318 evaluated patients, 310 were homosexual or bisexual men (97%), four were heterosexual women, and four were heterosexual male intravenous drug users. Pneumocystis pneumonia was confirmed in 154 patients and "clinically suspected" in a further 46. These latter patients were excluded from further analysis as no definitive diagnosis could be made, but including them as having confirmed pneumocystis pneumonia would not have altered any of the statistical associations presented here. Other diseases were diagnosed in 118 patients (table 1).

There were no significant differences in age or median CD4 counts at the time of assessment between patients with pneumocystis pneumonia (35-8 years, and 50 \(\times\) \(10^4\) cells/l) and those with other chest diseases (35-6 years and 4-6 \(\times\) \(10^4\) cells/l). A history of previous pneumocystis pneumonia was obtained in 16% of patients with pneumocystis pneumonia as their current illness and 22% of patients with

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>No of patients</th>
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<tr>
<td>Confirmed pneumocystis pneumonia</td>
<td>154</td>
</tr>
<tr>
<td>Clinically suspected pneumocystis pneumonia</td>
<td>46</td>
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<tr>
<td>Bacterial chest function</td>
<td>68</td>
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<tr>
<td>Pulmonary Kaposi's sarcoma</td>
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<td>Mycobacterium avium-intracellulare</td>
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<tr>
<td>Myopathy</td>
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Table 1: Final diagnoses in 318 patients presenting with suspected pneumocystis pneumonia
other chest diseases. Fewer patients with than without pneumocystis pneumonia were smokers (45% vs 57%), though the difference was not significant.

Univariate analysis showed that chest radiographic features, desaturation during exercise, absence of prophylaxis (p < 0.001), and classical clinical symptoms (p < 0.001) were strongly predictive of pneumocystis pneumonia. Despite obvious potential associations between these features, multivariate analysis also showed that they remained strongly predictive as independent variables (table 2). Arterial hypoxaemia, however, did not differentiate pneumocystis pneumonia from other chest diseases (p = 0.07) and had a low predictive value (fig 1). The predictive value of these features did not differ between smokers and non-smokers.

Exercise induced oxygen desaturation was the single most useful parameter (fig 1), with a multivariate odds ratio for confirmed pneumocystis pneumonia of 5.43. Although patients were asked to continue exercising for a full 10 minutes, only 38% of patients with pneumocystis pneumonia were able to achieve this compared with 61% of patients without pneumocystis pneumonia. In 80% of patients with pneumocystis pneumonia Sao2 fell to 90% or less within the first three minutes of the test. Only 13 patients with pneumocystis pneumonia completed a full 10 minute test without developing desaturation.

Although diffuse interstitial shadowing was sensitive, it was not specific for pneumocystis pneumonia as similar infiltrations were seen in 60% of patients with pulmonary tuberculosis and 39% of patients with pulmonary Kaposi’s sarcoma. Only 11% of patients with atypical mycobacteriosis, 7% of those with bronchial candidiasis, and 2% of those with bacterial chest infections had similar infiltrates.

None of the features studied was useful in predicting pneumocystis pneumonia in patients who had had at least one previous confirmed episode of pneumocystis pneumonia (table 3).

With the exception of hypoxaemia, the positive predictive value for a correct diagnosis of pneumocystis pneumonia was improved by combining two or more of the defined criteria (table 4). The triad of typical symptoms, arterial hypoxaemia, and diffuse chest infiltrates together gave a high positive predictive value (88%), but were present in only one third of the patients with confirmed pneumocystis pneumonia.

An algorithm of management was developed from the positive predictive values for combinations of features (fig 2). Typical symptoms and knowledge of pneumocystis pneumonia prophylaxis were combined as they will almost always be available from a clinical history. This algorithm was used to place patients into three categories: (1) pneumocystis pneumonia highly likely; (2) chest disease uncertain; (3) pneumocystis pneumonia highly unlikely. For these categories three corresponding management strategies were proposed—namely, start pneumocystis pneumonia treatment, investigate further, treat for bacterial infection, and observe. Had this algorithm been used, empirical treatment would have been correctly
instigated in 73 patients with pneumocystis pneumonia (95% accuracy); four patients with other diseases would have been incorrectly given pneumocystis pneumonia treatment (two with tuberculosis and two with bacterial chest infections). Forty seven patients would have been assigned to the "unlikely" category; of these, seven in fact had pneumocystis pneumonia (85% accuracy). Forty eight per cent of those patients requiring further investigations had an eventual diagnosis of pneumocystis pneumonia.

Discussion

Our intention in this study was to assess the usefulness of readily available data in making a diagnosis of pneumocystis pneumonia in patients infected with HIV before embarking on more extensive or invasive investigations. As pneumocystis pneumonia is most unlikely in the early stages of HIV infection, the sensitivity and specificity of simple features are likely to be of little clinical relevance in this group. Only patients with signs of immunosuppression or a known reduced CD4 count were therefore studied. We hoped that this would provide useful predictive information in a group of patients with advanced immunosuppression in whom the prior probability of pneumocystis pneumonia was relatively high. The five pieces of information were individually of limited predictive value in the accurate diagnosis of pneumocystis pneumonia. Although the absence of pneumocystis pneumonia prophylaxis and the presence of interstitial shadowing on the chest radiograph were sensitive tests, both were relatively non-specific. Arterial hypoxaemia does not provide a sensitive test as it is present in various other respiratory conditions; but arterial blood gas analyses are still needed during the initial evaluation to determine whether adjunctive corticosteroid treatment is necessary.

Exercise desaturation was the most sensitive single index of pneumocystis pneumonia, but was less specific than an indicative chest radiograph, largely because desaturation occurred in two thirds of patients who had had pneumocystis pneumonia previously, whatever the cause of the current respiratory problem. Exercise oximetry contributes most in those patients who have not had previous pneumocystis pneumonia and those without obvious radiographic changes.

None of the features studied was useful in predicting pneumocystis pneumonia in patients who had at least one previously confirmed episode. Although the increased use of prophylaxis in this subgroup is easily explained, the exercise desaturation patterns are more difficult to interpret. The most likely reason for this is that inflammatory scarring and thickening of the alveolar interstitial occurs during resolution of the initial episode of pneumocystis pneumonia, resulting in some degree of respiratory impairment whenever a subsequent pulmonary insult ensues.

The proposed algorithm was devised on the pragmatic basis that the chest radiograph, the arterial blood gas tensions, and the history at presentation will almost always be immediately available to the clinician. This combination provides highly specific features but with a relatively low sensitivity. The specificity is further increased if results of exercise desaturation are also positive, and the criterion of replacing low resting arterial blood gas oxygen tension with exercise induced oxygen desaturation greatly improves the overall diagnostic usefulness of these simple tests.

Further definitive tests for pneumocystis pneumonia may be considered unnecessary in patients assigned by the algorithm to the category "pneumocystis pneumonia highly likely" (positive predictive value over 95%) and many clinicians would wish to treat such patients even if subsequent tests gave negative
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results. Similarly, the negative predictive value of 85% in the “pneumocystis pneumonia unlikely” category identifies a group of patients with such a low likelihood of pneumocystis pneumonia that it is justifiable to manage them by close observation or treating them for a presumed bacterial chest infection (or both). If used in this way the algorithm would have obviated the need for further investigations in 59% of all the patients seen.

Other screening tests, including DTPA scanning, gallium scanning, computed tomography of the chest, and determination of carbon monoxide transfer factor or serum lactate dehydrogenase activity, have been advocated to help in the diagnosis of pneumocystis pneumonia and therefore reduce the need for further investigations. Most such tests, which are either sensitive or specific, are expensive or unavailable outside specialist centres, or yield results only after some delay.

In conclusion, this algorithm has been developed to provide a simple and rapid assessment of the probability of pneumocystis pneumonia in patients with advanced HIV disease. Empirical treatment may be confidently instigated on the basis of risk features ascertained at the time the patient first presents, thereby enabling treatment to start earlier and survival, if it is hoped, to improve. It should also reduce the number of patients requiring more extensive screening and reduce the requirement for invasive definitive tests.

We wish to thank Dr P Easterbrook (AIDS research unit, St Stephen’s Clinic) and Dr K. MacRae (department of professorial medicine, Charing Cross Hospital) for their statistical advice.

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