Investigation of seemingly pathogen-negative diarrhoea in patients infected with HIV

G M Connolly, A Forbes, B G Gazzard

Abstract
Thirty three consecutive patients infected by human immunodeficiency virus type 1 (HIV1) with persistent diarrhoea which remained undiagnosed after microbiological examination of six stool samples and rectal histology were investigated for malabsorption. All had xylose and Schilling tests, distal duodenal biopsy, comprehensive barium studies, microbiological examination of six further stool samples, and repeat rectal histology. A microbiological or histological diagnosis of infection was made in 12 patients (multiple organisms in three). Cryptosporidia were identified on five occasions, cytomegalovirus on four, Giardia lamblia on two, and herpes simplex, Campylobacter jejuni, Salmonella enteritidis, and Entamoeba histolytica once each. No organism was found when weight loss was less than 5 kg or stool volume less than 400 ml/day (n=9). Pathogens were identified in nine of 13 patients (69%) with weight loss greater than 10 kg and stool volume more than 800 ml/day. Barium studies were normal except for ileal flocculation in two patients with cryptosporidiosis. Evidence for malabsorption existed in 24 patients - impaired xylose absorption (n=19) and abnormal Schilling test (n=21). Of the patients with a severely abnormal Schilling test, a pathogen was identified in 11 (79%) (including all five with cryptosporidiosis, and two of the patients with only moderate diarrhoea and weight loss). A simple scoring system based on degree of weight loss and Schilling test result may help to identify the HIV positive patient with seemingly pathogen-negative diarrhoea in whom further investigations are likely to show a specific cause.

Diarrhoea is a common symptom in AIDS patients and occurs in up to 50% at some time during their illness. In a proportion of patients no pathogenic cause of their diarrhoea is found, and it has been hypothesised that direct human immunodeficiency virus type 1 (HIV1) infection of the gut mucosa leading to malabsorption may be responsible. The frequency of finding a pathogen obviously increases with the number of tests performed but the clinician needs advice to decide when further investigation is unlikely to yield diagnostic information. We prospectively studied a group of 33 patients in whom no pathogen was detected despite examination of six stool samples and rectal biopsy specimen. Further microbiological analysis, histopathology, and tests of malabsorption were performed to see if it was possible to predict which patients would benefit from these tests.

Patients
Thirty three consecutive male patients (median age 28 years; range 18–54) with serum antibodies to HIV1 and with undiagnosed diarrhoea were studied. All had passed more than three loose stools daily for at least one month, with no microbiological diagnosis established from rectal histology and at least six stool examinations performed in a microbiology laboratory accustomed to the particular pathogens associated with this patient population. The CDC criteria for full AIDS were fulfilled by 16 of the 33 patients at the time of presentation (group IV CI in each case); the remaining 17 (by definition) had HIV disease group IV A.

Methods
Daily stool volume (mean of three days) and estimated weight loss from usual premorbid weight (checked against ideal 'weight for height' in cases of premorbid obesity or doubt) were recorded. Upper gastrointestinal endoscopy for distal duodenal biopsy, barium small bowel follow through examination, and double contrast barium enema were performed using standard methods.

Serum concentrations of vitamin B12 and serum and red cell folate concentrations were measured, and double labelled cobalt Schilling tests carried out (normal greater than 11% excreted). Xylose absorption testing (5 g) was performed with assay of a five hour urine collection (normal greater than 1.4 g excreted).

Six further stool samples were examined microbiologically with special attention to opportunistic pathogens; accordingly the modified Ziehl-Nielsen stain and use of both Lowenstein-Jensen and Kirchner media were used routinely as previously described.

Rectal and duodenal histology were stained with haematoxylin and eosin, and with Ziehl-Nielsen stain. Inclusion bodies and an inflammatory response were required for a histological diagnosis of cytomegalovirus infection.

Receiver operating characteristic curves were constructed for each of the precisely quantifiable parameters measured. Sensitivity, specificity, and predictive values were defined and calculated conventionally.

A scoring system based on Bayes' theorem was developed to assist early identification of those in whom an infective aetiology for diarrhoea would eventually be found.

Results
Stool volume varied from 300 ml to more than 3000 ml per 24 hours. Use of arbitrary cut off points at 400 and 800 ml led to division of
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Scattergram to show relation between xylose excretion, Schilling test result, and final evidence for or against infective aetiology for diarrhoea. Xylose results are excretion in g/5/hour urine collection and Schilling results are presented as the percentage of radioisotope excretion; in both cases the horizontal line marks the lower end of the normal reference range.

patients between three approximately equal sized groups: less than 400 ml, n=9; 400–800 ml, n=11; more than 800 ml, n=13. Similarly arbitrary cut off points for weight loss led to distribution of patients between three groups: less than 5 kg loss, n=9; 5–10 kg, n=10; more than 10 kg, n=14. The groupings form natural pairs and in all but one patient the stool volume grouping was ‘equivalent’ to that for weight loss.

Serum B12 and folate and red cell folate concentrations were normal in all patients. Xylose excretion, however, was impaired in 19 patients and in nine of these excretion was less than 50% of the lower limit of normal (severely abnormal) (Figure). Unsurprisingly, the Schilling tests showed no evidence of intrinsic factor deficiency, and both labels were detected at similar radioconcentration: the mean of the two values for each patient was therefore used. In 21 patients abnormal values were detected and in 14 of these, mean values of less than 50% of the lower limit of normal were recorded (severely abnormal).

Infection was finally shown in 12 patients (more than one organism in three): cryptosporidium, n=5; Giardia lamblia, n=2; Entamoeba histolytica, n=1; Campylobacter jejuni, n=1; Salmonella enteritis, n=1; cytomegalovirus, n=4; and herpes simplex virus, n=1. Diagnosis of cytomegalovirus was from biopsy specimen alone. Diagnosis of herpes simplex virus was from biopsy specimen and from swabs taken from rectal vesicles. Cryptosporidium was detected in both biopsy specimen and stool microbiology in two patients. The remaining diagnoses were from stool microbiology alone. All histological diagnoses were obtained from rectal biopsy specimens except for one patient where rectoscopy was normal but typical cytomegalovirus foci were seen in the transverse colon at colonoscopy and a biopsy confirmed cytomegalovirus infection.

CORRELATION OF MALABSORPTION AND INFECTION

An estimated weight loss of more than 10 kg was strongly associated with eventual identification of gastrointestinal pathogens: nine of 14 patients with this extent of weight loss had organisms detected, accounting for nine of the 12 patients in whom an organism was eventually found. When weight loss was less than 5 kg or daily stool volume was under 400 ml, or both, no pathogens were identified. Weight loss of less than 5 kg therefore had a negative predictive value of 100% for an eventual microbiological diagnosis, whereas weight loss of greater than 10 kg had a positive predictive value of 64% and a sensitivity of 75% (Table).

Abnormality of xylose absorption (whether or not severe), however, was less discriminatory – an abnormal result had only a 53% positive predictive value for a final microbiological diagnosis and a normal result only an 86% negative predictive value. Severe abnormality of xylose absorption had a specificity of 80% associated with a sensitivity of only 42% (Table). The receiver operating characteristics curve constructed from the xylose absorption data indicated no better discrimination from any other given level of cut off.

A microbiological diagnosis was made in only one patient with a normal Schilling test but of 14 patients with a severely abnormal result, pathogens were found in 11 (including all five patients with cryptosporidial infecton, and two patients with only moderate weight loss and stool volume). A normal Schilling test therefore had a 92% negative predictive value for a final microbiological diagnosis. A severely abnormal Schilling test, however, had a sensitivity of 92% and a positive predictive value of 79% for subsequent microbiological diagnosis (Table). The receiver operating characteristics curve for Schilling test results indicated that a cut off point at 50% of the lower limit of normal (by chance) corresponded to the best combination of sensitivity and specificity achievable.

A barium follow through study was abnormal in only two patients who had cryptosporidial infection, appreciable weight loss, and severely abnormal Schilling test results. All endoscopies were macroscopically normal.

TREATMENT AND OUTCOME

Patients were treated on the basis of the results obtained, using specific antibiotic drugs for the bacterial infections (ampicillin for S enteritis, erythromycin for C jejuni), metronidazole for giardiasis and amoebiasis, and high dose acyclovir for herpes simplex virus. In all patients relevant treatment eradicated the organism for the duration of microbiological follow up (more than two months) and stool volumes were reduced. The five patients with cryptosporidial diarrhoea were treated with a variety of antimicrobial agents without success, but starting them on zidovudine and opioid anti diarrhoeal drugs was associated with control of symptoms and considerably reduced stool volumes in two

Sensitivity, specificity, and positive predictive value (PPV) for each of the degrees of abnormality of the parameters evaluated, with negative predictive value (NPV) for a normal result (percentages in each case).

<table>
<thead>
<tr>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
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<tr>
<td>Weight loss:</td>
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<tr>
<td>&gt;5 kg</td>
<td>100</td>
<td>43</td>
<td>50</td>
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<tr>
<td>&gt;10 kg</td>
<td>75</td>
<td>90</td>
<td>64</td>
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<tr>
<td>Abnormal xylose absorption</td>
<td>83</td>
<td>57</td>
<td>53</td>
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<tr>
<td>&lt;50% normal lower limit</td>
<td>42</td>
<td>81</td>
<td>56</td>
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<tr>
<td>Abnormal Schilling test</td>
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<td>52</td>
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<td>&lt;50% normal lower limit</td>
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patients. The other three patients failed to respond and died within one month with continued watery diarrhoea. Three of the patients with cytomegalovirus colitis were treated (the fourth was terminally ill by the time of diagnosis). One responded to a single course of phosphonoformate but the last showed no response to antiviral treatment.

SCORING SYSTEM

Stool volume and estimated weight loss provided almost entirely concordant information; they are clearly not independent variables as far as their diagnostic potential is concerned. Since stool volume, despite being a valuable marker of disease activity and response to treatment, is more difficult to measure (particularly in outpatients) and proves relatively expendable diagnostically, we used estimated weight loss as the preferred variable, grouping patients between the three arbitrary bands of severity described above. Of the other variables considered, Schilling test result, with receiver operating characteristics curve-determined cut off at 5-5% or less, showed most promise. Application of Bayes' theorem led to the allocation of the following scores: weight loss less than 5 kg: +0-19; 5-10 kg: +0-04; more than 10 kg: −0-25; Schilling result: more than 5-5%: +0-17; 5-5% or less: −0-48. Fortunately, almost as effective is a simplified version where weight loss is scored with the final result for more than 10 kg loss, 1 point for a loss of 5-10 kg, and 0 points for weight loss of less than 5 kg, and where points are allocated for the Schilling result as follows: more than 5-5%: 0 point; 5-5% or less: 2 points. Retrospective application to the present patients (predictably) shows a 5% frequency of pathogen recognition for patients with scores of 0–2 (n=19), whereas 90% of patients with a score of 4 (n=10) had at least one pathogen identified.

Discussion

Microbiological examination of numerous stool samples is evidently the most useful diagnostic procedure, and this applies equally to the present difficult group as to the generality of HIV-infected patients with diarrhoea. The present results, however, help to indicate the patient in whom it is reasonable to conclude that a specific diagnosis is unlikely to emerge however many samples are studied. The value of a specific diagnosis is not always obvious when no established treatment exists (for example in microsporidiosis), but valuable prognostic information may be obtained (several of the present patients were moved from HIV group IV A to group IV C1 or IV C2) and, increasingly, effective new treatments (for example for cytomegalovirus) are becoming available.

It is acknowledged that the infecting organisms identified may not necessarily have been relevant to the chronic diarrhoea – particularly the bacterial and amoebic infections – as these may have arisen acutely and been superimposed on (still) undiagnosed chronic diarrhoea thus explaining why they were not detected on initial investigation. There was, however, a good and long lasting (more than two months) remission from diarrhoea after antibiotic treatment. Stool electron microscopy in the routine evaluation of these patients may be thought valuable, although in a large series of stools from patients with AIDS and diarrhoea (unpublished observations), no viral particles were observed or other evidence provided for infection missed by other investigations. Electron microscopy of small bowel mucosa, however, is of considerable value in the diagnosis of microsporidial infection, which may be associated with malabsorption and altered small bowel histology but without evidence of the organism in conventional duodenal pinch biopsy specimens. In one recent study of patients similar to those considered here, more than a third had Microsporidia present at jejunal electron microscopy. Good evidence for malabsorption was found frequently in this group of patients, but although there has been speculation that HIV itself causes malabsorption, the present data argue for malabsorption being linked to secondary microsporidal infection in many patients. The normal serum vitamin B₁₂ values is unsurprising given the relatively short total illness period in the patients concerned, but the site of the abnormality in the two positive barium follow through examinations and the normal folate studies suggest that the malabsorption is predominantly of ileal origin. Recent reports of possible relevance here indicate that there may be antibodies to cryptosporidial infection, one where organisms are detected in stool only with difficulty but small bowel involvement and B₁₂ malabsorption are prominent, and the other where oocysts are readily identified in stool and the Schilling test is usually normal.

As we have shown in other contexts, barium radiology and endoscopy are only rarely helpful in patients with AIDS and will not often be indicated, but rectal biopsy is a useful early investigation and remains the only route to diagnosis in some patients.

Based on the results presented here, the diagnostic yield of further microbiological samples in patients who have lost less than 5 kg in weight and have a normal Schilling test is extremely low. A putative scoring system is suggested but this clearly needs prospective evaluation in future patient groups.

We thank our patients and our many colleagues without whose assistance this work would not have been possible, but are particularly indebted to Drs Shanson, Harcourt-Webster, and Gleeson.

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