Audit of endoscopic surveillance biopsy specimens in HIV positive patients with gastrointestinal symptoms

S G Lim, M Cl Lipman, S Squire, D Pillay, S Gillespie, E A Sankey, A P Dhillon, M A Johnson, C A Lee, R E Pounder

Abstract
An audit of upper gastrointestinal endoscopy in HIV infected patients with gastrointestinal symptoms assessed the frequency of disease detected by endoscopy and routine laboratory analysis of surveillance biopsy specimens. Sixty nine consecutive endoscopies were performed in 59 HIV infected patients. Endoscopic biopsy specimens were taken from the lower oesophagus, gastric antrum, and third part of the duodenum for virology, histopathology, parasitology, bacteriology, and mycobacterial culture. Endoscopic appearances detected disease in 25/59 (42.4%) patients (oesophageal candida, 14; oesophageal ulcer, 3; Kaposi’s sarcoma, 4; others, 4), but only 4/43 (9.3%) specimens showed evidence of disease in the absence of endoscopic abnormality. Virology for cytomegalovirus (detection of early antigenic fluorescent foci and culture) was positive in 6/59 (10.2%) patients, but parasitology and mycobacterial culture were negative in all cases. Histopathology was abnormal in 11/52 (21%) oesophageal biopsy specimens, 13/47 (28%) gastric biopsy specimens, and 4/65 (6%) duodenal biopsy specimens. Abnormal findings were found predominately in those with advanced HIV disease (CDC Stage IV) (21/33 patients (64%)) compared with those with early HIV disease (CDC Stage II) (5/26 (19%)). In conclusion, upper gastrointestinal endoscopy detects macroscopic disease in AIDS patients and those with low CD4 counts, but routine surveillance biopsy specimens of apparently normal bowel in early HIV disease (or where CD4 counts are greater than 0.2 X 10^9/l) are of little value.

Methods
PATIENTS
An upper gastrointestinal endoscopy was performed in 59 consecutive HIV infected patients for a variety of clinical indications (Table I). All but four of the patients were men; the group had a median age of 35 years (range 17 to 61). When staged using the revised Center for Disease Control (CDC) criteria, 26 of the patients were CDC stage II, three were CDC IV A (AIDS related complex), and 30 were CDC IV C (AIDS). Diarrhoea was defined as having more than two loose bowel actions a day for two weeks, and weight loss was defined as a greater than 10% loss of body weight.

ENDOSCOPY AND BIOPSY
Upper gastrointestinal endoscopy was performed with an IT 20, QX 10, or QX 20 endoscope (Olympus, Keymed, Southend-on-Sea, Essex, UK). The agreed protocol was that all abnormalities seen during the endoscopic procedure were to be biopsied for analysis by histopathology and virology, and also for mycobacterial culture. In addition, in all patients regardless of endoscopic findings, two specimens were taken from the lower oesophagus, three from the gastric antrum, and four from the distal duodenum. One biopsy sample from all three sites was sent for histopathological and virological analysis. One specimen from the stomach and duodenum was sent for microbiological analysis, and one duodenal specimen was sent for mycobacterial culture.

HISTOPATHOLOGICAL METHODS
Biopsy specimens sent to histopathology were fixed in 10% buffered formalin, embedded in paraffin wax blocks, and 6 μm sections were cut and stained with haematoxylin and eosin and special stains (Giems, Ziehl Neilsen, Gram, Schiff-periodic acid). In addition, immunohistochemistry was performed for cytomegalovirus.

VIROLOGICAL METHODS
Biopsy specimens were sent to virology in viral transport medium. The samples were homogenised and cultured on human embryonic lung fibroblasts. Detection of early antigenic fluorescent foci (DEAFF) was performed after 12 to 16
TABLE I Results of endoscopy and biopsy in 59 HIV infected subjects (*inflammatory changes without a definite diagnosis have not been included)

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>Endoscopic diagnosis (no)</th>
<th>Laboratory analysis of biopsy specimen* (no)</th>
<th>Outcome (no)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dysphagia heartburn or retrosternal pain (18)</td>
<td>Ulcers (3) Candida (7) HH (1) Normal (7)</td>
<td>CMV (3) Candida (7) CMV (1) HLO (2) No diagnosis (4)</td>
<td>Ganciclovir-improved (3) Flucanazole-improved (7) Improved on ranitidine (1) Acyclovir – improved (1) Tonsil ulcer (1) Gradual improvement (4) Lost to follow up (1)</td>
</tr>
<tr>
<td>Abdominal pain (14)</td>
<td>GU (1) GU and KS (1) Candida (3) Normal (9)</td>
<td>CMV (1) Candida (3) CMV and villous atrophy (1) HLO (5) No diagnosis (3)</td>
<td>Ganciclovir – improved (1) Healed on ranitidine (1) Candida improved (3) No treatment (1) CMV biliary sclerosis (1) Triple therapy – improved (2) HLO epon. improved (1) Presumed gut vasculitis (1) No diagnosis (2) Gradual improvement (1)</td>
</tr>
<tr>
<td>Dyspepsia (5)</td>
<td>Normal (5)</td>
<td>Giardia (1) HLO (1) No diagnosis (3)</td>
<td>Metronidazole – improved (1) Giardia in stool (1) ? sphenomegaly (1) Gradual improvement (1) Lost to follow up (1)</td>
</tr>
<tr>
<td>Diarrhoea (9)</td>
<td>KS (1) Candida (1) Cryptosporidia and Candida (1) Normal (6)</td>
<td>Candida (1) No diagnosis (5)</td>
<td>Interferon for KS, but diarrhoea persisted (1) Cryptosporidia previously diagnosed in stool Candida improved (1) Orange juice related (1) Gradual improvement (2) Persisting (1) Lost to follow up (1)</td>
</tr>
<tr>
<td>Vomiting (4)</td>
<td>Candida (1) Erosions (1) Normal (2)</td>
<td>CMV/Candida/papilloma virus HLO (1) No diagnosis (1)</td>
<td>Treated for CMV/Candida Indomethacin ceased Alcohol related (1) Drug related (1)</td>
</tr>
<tr>
<td>Multiple symptoms (5)</td>
<td>Candida (2) DU (1) Normal (2)</td>
<td>CMV (1) Candida (2) HLO (1) No diagnosis (2)</td>
<td>Fluconazole – improved (1) Caecal TB (1) Died MAI within 1 month (1) Died in intensive care (1) HIV encephalopathy (1)</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>KS staging (2) Anemia (2)</td>
<td>CMV (1) (incidental) Normal (2)</td>
<td>KS treated (2) Anemia resolved (2)</td>
</tr>
</tbody>
</table>

KS=Kaposi's sarcoma, Candida=candida oesophagitis, CMV=cytomegalovirus, HLO=Helicobacter pylori, TB=tuberculosis, DU=douodenal ulcer, GU=gastric ulcer, PUO=pyrexia of unknown origin, MAI=mycobacterium avium intracellulare, HH=hiatus hernia.

hours, to identify infection with cytomegalovirus. In addition, cultures were maintained for 21 days or until a positive result, whichever occurred first.

MICROBIOLOGICAL METHODS
Impression smears of specimens sent to microbiology were assessed microscopically by Auramine (cryptosporidium) and Giemsa (giardia) stains for ova, cysts, and parasites. They were then placed on Sabouraud's medium for yeasts and Robertson's cooked meat medium for bacterial pathogens; these cultures were read after 24 to 48 hours. Biopsy specimens sent for mycobacterial culture were placed on two Lowenstein-Jensen slopes, one supplemented with pyruvate and one with glycerol. Biopsy specimens were, in addition, cultured in Kirchner's broth.

CYTOLOGICAL METHODS
Brush cytology of the oesophagus was performed when candida was suspected endoscopically using a disposable sheathed endoscopic brush (Olympus, Keymed, Southend-on-Sea, Essex, UK). The brush was smeared onto four glass slides and fixed for five minutes in 74% industrial methylated spirits. The fixative was drained and the slides dried before Papanicolaou staining for candida.

Results
The audit showed that a total of 69 endoscopies were performed in 59 patients over 24 months. Table I lists the endoscopic findings, laboratory results, and clinical outcome. Table II lists results of the laboratory analysis of the biopsy samples. The diagnostic yield of the biopsy procedures was much higher in patients with endoscopic disease (25/69 biopsies in 26 patients) than in those without endoscopic abnormality (4/43 in 33 patients excluding the presence of Helicobacter pylori). Of the four positive biopsy specimens in patients with a normal endoscopy, two had cytomegalovirus isolated, one had candida, and the last had giardia. Additionally two patients had unsuspected dual (Kaposi's sarcoma and CMV) and triple disease (cytomegalovirus, candida, and Papilloma virus). In only five of 26 patients with CDC stage II disease was a positive endoscopic appearance or laboratory result found (three of these had oesophageal candida seen at endoscopy, which proved to be their AIDS defining illness), while gastrointestinal abnormality (either on endoscopy or analysis of
Audit of endoscopic surveillance biopsy specimens in HIV positive patients with gastrointestinal symptoms

<table>
<thead>
<tr>
<th>Table II</th>
<th>Results of laboratory analysis of upper gastrointestinal mucosal biopsy specimens from 59 patients with HIV infection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Virology culture</strong></td>
<td><strong>Bacteriology culture</strong></td>
</tr>
<tr>
<td><strong>Results (no)</strong></td>
<td><strong>Total no positive</strong></td>
</tr>
<tr>
<td>CMV positive</td>
<td>6 (10-5%)</td>
</tr>
<tr>
<td>CMV (3)+HSV (1)</td>
<td>4 (6-7%)</td>
</tr>
<tr>
<td>Staphylococci (5)</td>
<td>16 (13-8%)</td>
</tr>
<tr>
<td>Streptococci (5)</td>
<td></td>
</tr>
<tr>
<td>Miscellaneous (6)</td>
<td></td>
</tr>
<tr>
<td>Candida, enterococcus</td>
<td></td>
</tr>
<tr>
<td><strong>CMV=cytomegalovirus, HLO=Helicobacter pylori, KS=Kaposi's sarcoma, HSV=herpes simplex virus, DEAFF=detection of early antigenic fluorescent foci.</strong></td>
<td></td>
</tr>
</tbody>
</table>

Discussion

The prevalence of H pylori in our study was somewhat higher at 24-3% compared with previous reports of 4%14 and 14%15 – which were considerably lower than in the normal ‘healthy’ population. Interestingly

![Correlation of pathological findings, clinical stage of HIV disease, and peripheral blood CD4 count. ARC=AIDS related complex.](http://gut.bmj.com/)

Correlation of pathological findings, clinical stage of HIV disease, and peripheral blood CD4 count. ARC=AIDS related complex.
the Figure shows that almost all cases of *H pylori* occurred in patients with CD4 counts $<0.3\times10^7$, but only 6/11 *H pylori* positive patients had AIDS. This seems to be at variance with Francis’
but, while our data suggest that *H pylori* may increase with immunosuppression, this may be the result of the small population sample. The diagnostic success of endoscopic duodenal biopsy in patients with diarrhoea was low in this study; the most sensitive test remains stool analysis for pathogens and ova, cysts, and parasites, but if these are negative then additional sampling methods should be undertaken — Crosby capsule biopsy, jejunal fluid sampling, brush cytology of jejunum to improve diagnostic yield.

There was no diagnosis of atypical mycobacteria either by histology or by biopsy culture despite the duodenum being the most common affected site. One of these patients died from disseminated atypical mycobacteria infection one month after his endoscopy, having had negative histopathology and duodenal culture for the bacterium.

Multiple biopsy specimens were taken to exclude occult cytomegalovirus infection as a potential cause of gastrointestinal symptoms. The findings of our study show that cytomegalovirus is unlikely to be discovered in the absence of visible pathology. Without mucosal pathology, a positive culture may be of dubious clinical significance, as shown by two patients who did not develop clinical cytomegalovirus disease despite having positive culture results. While attention has been focused on the potential for cytomegalovirus to cause abdominal pain, the diagnosis of cytomegalovirus disease remains difficult. Morphology remains the standard method of diagnosis, but there have been attempts to improve diagnosis by immunocytochemistry, electron microscopy, and viral culture. Unfortunately the detection of cytomegalovirus antigens (whether late or early) or virus in culture, shows the presence of the virus and not disease. Pathological infection with cytomegalovirus must remain a clinical or morphological diagnosis. As a result of this audit, the clinical practice in management of HIV infected patients at upper gastrointestinal endoscopy has been changed at the Royal Free Hospital. Routine surveillance biopsies are no longer performed in those patients with CDC II disease or CD4 counts more than $0.2\times10^7$. Patients with diarrhoea remain an exception to this rule: they have more intensive investigations with culture, histology, electron microscopy, and brush cytology of the jejunal mucosa.

The authors would like to thank Mr O Billington for processing specimens for mycobacterial culture, Mrs A Smith and Mr A Ramsey for processing paraffinology specimens, and Dr Allan Deary for reporting the cytology specimens.

17 Francis ND, Boylston AW, Roberts AH, Parkin JM, Pinching AJ. Cytomegalovirus infection in gastrointestinal tracts of patients infected with HIV-1 or AIDS. *J Clin Pathol* 1989; 42: 1055-64.