Value of DNA image cytometry in the prediction of malignant change in Barrett’s oesophagus

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SUMMARY DNA image cytometry was performed on Feulgen stained sections from 91 biopsies obtained during prospective endoscopic surveillance of 55 patients with Barrett’s oesophagus. Aneuploid cells were detected in specimens from six of these patients. Four subsequently developed dysplasia and adenocarcinoma but, in the other two, biopsies had been reported as showing specialised epithelium only, with no apparent dysplasia and no evidence of malignancy on clinical follow up to date. In two of the four patients who subsequently developed carcinoma, aneuploid cells were only found in biopsies showing overt dysplasia or carcinoma but in the two other patients aneuploid cells were present in biopsies taken early in the clinical course before any dysplasia had been identified on the original reports. The presence of aneuploid cells on cytometry of these ‘benign’ biopsies allowed us, on histological review, to identify areas of atypia which were interpreted as mild dysplasia. In this series aneuploidy was always associated with some morphological abnormality varying from mild dysplasia to frank carcinoma. Aneuploid cells were not shown in material from one patient who had an oesophagectomy for dysplasia or in biopsy material from four patients showing ‘indefinite dysplasia’. DNA cytometry combines an objective assessment of epithelial atypia with the advantage of detecting rare cellular aneuploidy and the ability to correlate these events with morphology. It should assist in the more accurate diagnosis of dysplasia and prove useful in identifying those patients with Barrett’s oesophagus who are at greater risk of subsequently developing malignancy.

The columnar epithelial lined oesophagus (CELO), described by Barrett in 1950,1 is the result of severe, prolonged gastro-oesophageal reflex2 and it carries a risk of developing adenocarcinoma.3 The magnitude of this risk differs widely in reported series4 and indeed some have been criticised for exaggerating the malignant potential.5 As yet there is no reliable marker to indicate which patient will develop adenocarcinoma and mucin histochemistry of Barrett’s mucosa is not sufficiently discriminating to predict subsequent malignancy.6 7 At present dysplasia appears to be the only reliable indicator8 9 but the demonstration of aneuploidy by flow cytometry may be helpful in some cases.10

The present study attempts to assess microdensitometric DNA quantification as an indicator for detecting developing malignancy in patients with CELO.

This technique uses a computerised image analysis system which converts the size and staining intensity of nuclei in Feulgen stained sections into a DNA value for the cells measured. In contrast with flow cytometry, the measurements are made under direct visualisation which allows a correlation between the areas showing cellular aneuploidy and the morphology in adjacent conventionally stained sections. Flow cytometry will only identify a significant population of abnormal cells whereas cytometry will identify a single abnormal cell. The presence of aneuploid cells was taken to indicate malignancy or a malignant potential and was related to the known outcome in all cases.

Methods

PATIENTS From 1977 all patients with a diagnosis of Barrett’s oesophagus were entered into a programme of
endoscopic surveillance with random biopsy. Those in whom carcinoma was present at first examination were excluded from the study. The diagnosis was made when the columnar mucosa extended circumferentially up the oesophagus for 5 cm or more from the oesophagogastric junction which was defined endoscopically. At each examination the level of the squamocolumnar junction was measured from the incisor teeth and biopsies were taken randomly from the full extent of the columnar lined oesophagus. Latterly four quadrantic biopsies were taken at longitudinal intervals of 2–3 cm. The clinical features, histology, and mucin histochemistry of the first 56 patients have been published.4

Adequate biopsy material was obtained from 55 patients and to date 194 biopsies were available for study. Biopsies were fixed in neutral buffered formalin, routinely processed to paraffin and stained with haematoxylin and eosin. Mucosubstances were demonstrated using alcian blue pH 2.5/periodic acid schiff and high iron diamine-alcian blue pH 2.5 stains. Each biopsy was examined for the histological type of epithelium present,11 the presence and type of intestinal metaplasia,12 and for dysplasia or carcinoma.

A system for grading dysplasia in inflammatory bowel disease has been proposed13 which has been applied to Barrett’s oesophagus.5,14,15 The nomenclature used in this classification is shown in Table 1. Dysplasia is defined as an unequivocal neoplastic alteration in the columnar mucosa which is distinguishable from reactive or regenerative abnormalities.13 When a definite distinction is not possible, the changes are categorised as ‘indefinite for dysplasia’.

DNA analysis was performed on at least one biopsy from each patient, the most recent biopsy containing adequate amounts of columnar epithelium was selected. In some unselected cases DNA analysis was performed on two or more biopsies. All biopsies showing any degree of cellular atypia and all biopsies from patients who subsequently developed malignant disease were studied. Five micron sections were cut, hydrolysed (5 N hydrochloric acid at 23°C for 50 minutes) and then stained with Schiff’s reagent according to the Feulgen method.16

<table>
<thead>
<tr>
<th>Table 1 Classification of dysplasia in Barrett’s oesophagus</th>
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<tbody>
<tr>
<td>Negative for dysplasia</td>
</tr>
<tr>
<td>Indefinite for dysplasia</td>
</tr>
<tr>
<td>Possibly negative</td>
</tr>
<tr>
<td>Unknown significance</td>
</tr>
<tr>
<td>Probably positive</td>
</tr>
<tr>
<td>Positive for dysplasia</td>
</tr>
<tr>
<td>Low grade</td>
</tr>
<tr>
<td>High grade</td>
</tr>
</tbody>
</table>

**DNA IMAGE CYTOMETRY**

Determinations of nuclear DNA were performed using the Leitz Miamed DNA system which has been developed from the Leitz Texture Analysis system. Slides were visualised by a Leitz Ergolux Orthoplan microscope which had fully automated functions controlled by a microcomputer with television image analysis system. At least two levelled sections from each biopsy were examined.

The diploid standard (2c) of each biopsy was produced by counting at least 20 lymphocytes or as many as required to produce a standard error of less than 3% (coefficient of variation <12 in all cases), a correction factor of 1.19 was used between the DNA values of lymphocytes and epithelial cells.18 The DNA content of at least 50 epithelial cells per biopsy was determined, nuclei were selected preferentially if they appeared morphologically large or atypical. Nuclei could be measured individually using grey level thresholding and touching or tightly packed nuclei could be separated and measured by outlining the nuclei with the aid of a rollerball ‘mouse’.

Aneuploid cells were identified where the DNA content exceeded 5c and abnormal measurements

**Fig. 1 Aneuploid mitosis (arrow) with DNA value >5c in a mucosal biopsy originally reported as ‘histologically’ benign. Feulgen stain.**
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was repeated at least three times to ensure their accuracy. A biopsy was deemed abnormal only when three separate aneuploid cells were identified and where this observation was confirmed on a further section from the same biopsy. Overlapping nuclei, a common feature of dysplastic epithelium, precluded measurements in some areas. Mitotic figures, which

were often present, could be measured as these were situated towards the luminal aspect of the columnar epithelium away from the epithelial nuclei (Fig. 1). A DNA histogram was produced for each biopsy.

Results

Sixty one biopsies of histologically benign columnar mucosa examined from 48 patients (one biopsy 38 patients, two biopsies seven patients, three biopsies three patients) showed normal DNA profiles (Figs 2 a and b). These results included two biopsies each from four patients in which 'indefinite dysplasia probably negative' had been reported. In two patients with previously reported benign mucosa, the most recent of two biopsies examined showed aneuploid cells (Figs 2 c, 3). Review of these biopsies showed foci of mild dysplasia corresponding to the areas in which aneuploid cells were detected (Fig. 4).

Four patients developed adenocarcinoma during the study period and aneuploid cells were present in all the biopsies which were originally reported as...
showing dysplasia or carcinoma (Figs 2 d, e). In two of these patients aneuploid cells were present in the biopsies taken at presentation and no dysplasia had been identified in the original biopsy reports. Review of the biopsy material in the light of the cytometry findings, however, showed areas of mild dysplasia which had been previously overlooked or dismissed. In both these patients the presence of aneuploid cells persisted in subsequent biopsies which had been reported as showing either ‘indefinite dysplasia probably negative’, ‘indefinite dysplasia probably positive’, overt dysplasia or adenocarcinoma. In the two other patients with adenocarcinoma no aneuploid cells were detected in the dysplastic biopsies. Table 2 shows details of biopsy interval, histology, original and review grade of dysplasia and DNA profile in the four cases of malignant CELO.

A further patient had an oesophagectomy for severe dysplasia but on review of the biopsy material the changes were interpreted as ‘indefinite dysplasia probably negative’. DNA analysis showed no aneuploid cells in three biopsies examined or in the oesophagectomy specimen.

**Discussion**

The columnar epithelial lined oesophagus (CELO) is clearly a precancerous condition but reports of the magnitude of this risk differ widely and prevalence studies, which include patients presenting with adenocarcinoma, tend to exaggerate the risk of malignancy. The incidence among the 56 patients from this unit was one per 54.4 patient-years follow up which contrasts with two studies from New Zealand in which no malignant changes occurred. Cameron, however, reported two carcinomas in 102 patients giving an incidence of one case per 441 patient-years and Spechler reported two cases of carcinoma in a series of 105 patients, a rate of one per 175 patient-years. A further series reported one

Table 2  Biopsy interval, histology, original, and review grade of dysplasia and DNA profile of four cases of malignant CELO

<table>
<thead>
<tr>
<th>Time</th>
<th>Histology</th>
<th>Original</th>
<th>Review</th>
<th>DNA</th>
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<tr>
<td>Case 1</td>
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<td>J+S</td>
<td>Neg</td>
<td>LGD</td>
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<td>LGD</td>
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<td>44</td>
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<td>60</td>
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<td>HGD</td>
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</tr>
<tr>
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<td>HGD</td>
<td>HGD</td>
<td>Aneuploid</td>
</tr>
<tr>
<td>69</td>
<td>–</td>
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<td>IM Ca</td>
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</tr>
<tr>
<td>72</td>
<td>–</td>
<td>IM Ca</td>
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<td>78</td>
<td>Oesophagectomy</td>
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<td>57</td>
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<td>IM Ca</td>
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<td>–</td>
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<td>IV Ca</td>
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<td>Aneuploid</td>
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</table>

*Biopsy interval in months from presentation; †Sq=squamous; J=junctional; J+S=junctional and specialised; ‡Neg=negative for dysplasia; IPN=indefinite for dysplasia, probably negative; IPP=indefinite for dysplasia, probably positive; LGD=low grade dysplasia; HGD=high grade dysplasia; IM Ca=intramucosal carcinoma; IV Ca= invasive carcinoma; §Patient unfit for surgical resection.

Fig. 4  Adjacent section to that shown in Fig. 3. Large, pleomorphic epithelial cells with multiple nuclei are present. Nuclei in adjacent pyloric type glands (arrows) for comparison. Haematoxylin and eosin.
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903

case per 166 patient-years. Endoscopic surveillance is
advisable in all patients with CELO but changes in
symptomatology or gross endoscopic appearance
are often absent even after adenocarcinoma has
developed.4 Biopsy is therefore essential for the early
diagnosis of malignancy.

The mucin histochemistry of Barrett’s mucosa has
been studied6-44-28 and these reports show that
incomplete intestinal metaplasia and sulphomucin
secretion are common features. A recent study
showed intestinal metaplasia and sulphomucin
secretion in 64% and 58% of biopsies respectively, both
being present in at least one biopsy from 78% of the
patients studied. Sulphomucin secretion in specialised
mucosa is not a sufficiently sensitive marker to
predict those at risk of developing adenocarcinoma.44 All our patients, however, who sub-
sequently developed malignant change showed
specialised epithelium and sulphomucin secretion in
earlier biopsies.

At present dysplasia appears to be the only reliable
marker for adenocarcinoma9,10 and a long latent
period, varying from many months to several years,
may occur between its onset and the development of
adenocarcinoma.44 There are, however, wide inter-
observer variations in the assessment of dysplasia27
therefore, at present, it lacks objectivity and the
malignant potential of early changes may be missed.

The advent of DNA analysis using flow cytometry
has provided an objectivity which had hitherto been
lacking in the assessment of morphological abnor-
malities in the gastrointestinal tract. Reid10 found
aneuploid cells in 10 patients with Barrett’s oeso-
phagus; nine of these had dysplasia, carcinoma or
both but one patient had only specialised metaplastic
epithelium. McKinley15 found aneuploidy in 79% of
oesophageal malignancy and in seven patients with
CELO, two of whom had dysplasia. Aneuploid
however, is not present in all gastrointestinal
tumours subjected to flow cytometry10,28 and
aneuploid cells were reported to be present in only
two of four cases of dysplastic Barrett’s.9 This may be
because these cells form a minority population, too
small to be detected by flow cytometry and this has
raised doubts about the value of DNA analysis in
Barrett’s oesophagus.39

The present study utilises DNA image cytometry19
which has been shown to correlate well with the
results obtained by flow cytometry.30 This technique
has been shown to be of value in dysplasia in
longstanding ulcerative colitis31 and gastric
adenomas.32 The advantages of image cytometry over
other methods are the ability to detect a single ‘rare
event’ aneuploid cell30 and the direct visualisation of
abnormal cells which can be related to morphological
changes on the Feulgen or adjacent haematoxylin
and eosin stained sections. This latter facility can be
used to train morphologists to detect and improve
evaluation of small or subtle areas of atypia which
may otherwise be overlooked or dismissed as
insignificant. This is shown in the present study where
the presence of aneuploid cells in apparently normal
biopsies caused us to review the histological material
and reassess the degree of dysplasia present (see
Table 2). When more material has been studied, it
may be necessary to reevaluate the criteria for
diagnosing dysplasia in the light of the small areas of
cellular atypia identified by cytometry.

This technique is time consuming and labour
intensive, one to two hours were needed to ade-
quately examine some of the larger biopsies and there
were advantages in having a trained morphologist
perform the analyses. The development of small ‘in
microscope’ systems for DNA quantitation, how-
ever, could make this a routine evaluation. Some
degree of caution was necessary in interpretation as
abnormal measurements due to overlapping nuclei
could not be excluded on every isolated measure-
ment. Therefore a biopsy was deemed abnormal only
when three separate aneuploid cells were identified
and where this observation was confirmed on a
further section from the same biopsy. In some
biopsies overlapping nuclei within dysplastic foci
precluded adequate cell estimations although
mitoses, when present, could be measured. These
constraints would tend to underestimate the number
of aneuploid cells present.

A normal DNA profile was found in all our cases
with a benign clinical course, these included eight
biopsies from four patients showing ‘indefinite
dysplasia probably negative’. In all four patients who
developed carcinoma, aneuploid cells were present
in the biopsies which showed dysplasia or carcinoma.
On the basis of this small series DNA analysis would
seem to provide an excellent discriminator between
benign and malignant biopsies as well as assisting in
the diagnosis of atypia. One patient had an oeso-
phagectomy for severe dysplasia which, on review
was downgraded to ‘indefinite dysplasia probably
negative’ and no aneuploid cells were detected in the
three biopsies examined, nor in samples taken from
the oesophagectomy specimen. In two patients with
biopsies which were originally reported as showing
apparently benign columnar epithelium, aneuploid
cells were detected in the most recent biopsies but
there is no further follow up to date. Review of the
biopsy material in these two cases showed small foci
of mild dysplasia which had been previously over-
looked. Review of the two cases of malignant
Barrett’s, in which aneuploid cells had been found in
biopsies originally reported as ‘non dysplastic’, also
showed foci of mild dysplasia. This technique,
because of the detailed observation required, provides sensitivity and objectivity to simple morphology.

At present it is not possible to make an adequate evaluation of DNA analysis as an adjunct to biopsy in the endoscopic surveillance of Barrett’s oesophagus. Our study, which is the first to report the use of DNA image cytometry in Barrett’s oesophagus, suggests that this technique is useful in the positive identification of dysplasia and the absence of aneuploid cells is reassuring evidence that areas of atypia are ‘negative for dysplasia’. The finding of aneuploid cells in apparently normal biopsies should alert the morphologist to the possibility of small areas of dysplasia which have been overlooked. In two of the four patients in our series who developed adenocarcinoma no aneuploid cells were seen before the onset of dysplasia. This may in part be because of biopsy sampling but further work is required to evaluate DNA analysis as a screening procedure.

We wish to thank Ms Mandy Orchard for typing the manuscript, Mr Bill Brackenbury for photomicrography, Mr Trevor Grey for technical assistance, Dr Jim Lowe and Professor David Turner for reviewing the manuscript.

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*Gut* 1989 30: 899-905
doi: 10.1136/gut.30.7.899

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