Management of coeliac disease: a changing diagnostic approach but what value in follow up?

Monica Acalovschi, V Jayanthi, C S J Probert, J F Mayberry

Abstract

Objective — To assess the management of patients with coeliac disease in relation to a change in diagnostic method from jejunal suction biopsy to endoscopic biopsy.

Design — 16 item questionnaire survey of consultant members of the British Society of Gastroenterology.

Subjects — 359 consultant physician and gastroenterologist members of the society.

Main measures — Type of routine biopsy; repeat biopsy after gluten withdrawal; gluten rechallenge; follow up measurements; screening for malignancy; and methods of follow up, including special clinics.

Results — 270(70%) members replied; 216(80%) diagnosed coeliac disease routinely by endoscopic duodenal biopsy, 30(11%) by jejunal capsule biopsy, and the remainder by either method. Only 156(58%) repeated the biopsy after gluten withdrawal, though more did so for duodenal than jejunal biopsies (134/216, 62% v 13/30, 43%; p < 0.02). Follow up biopsies featured more duodenal than jejunal biopsies (133/156, 82% v 23/156, 15%; p < 0.02). Regular follow up included assessments of weight (259, 96%) and full blood count (238, 88%) but limited assessment of serum B-12 and folate (120, 44%) and calcium (105, 39%) concentrations. Routine screening for malignancy is not performed, and there are few specialist clinics. 171(63%) respondents thought that patients should be followed up by a hospital specialist and 58(21%) by family doctors.

Conclusions — The practice of diagnosing coeliac disease varies appreciably from that in many standard texts. Many patients could be effectively cared for by their family doctor.

Implications — The British Society of Gastroenterology should support such management by family doctors by providing clear guidelines for them.

Introduction

Coeliac disease is a relatively common condition in western Europe, its prevalence being 50–74/100 000 in the United Kingdom1 and between 50 and 250/100 000 in other Western countries.2 A changing pattern of coeliac disease has been noticed both in adults and in children. Many patients have only mild symptoms and no signs of malabsorption. Presently, only 12% of adults with coeliac disease have overt malabsorption, compared with 63% when the technique of jejunal biopsy was first introduced 30 years ago. This changing pattern is obviously due to an improved diagnostic approach. As a consequence of the increased use and easy performance of upper gastrointestinal endoscopy with duodenal biopsy coeliac disease is now diagnosed in asymptomatic people or those with only minor symptoms. Increased awareness of the disease might be responsible for an apparent increase in its prevalence but also leads to earlier diagnosis and hence a decline in severity at diagnosis.1

Ten years ago the prevalence of members of the Coeliac Society in England and Wales correlated well with regional sales of jejunal biopsy capsules.3 Jejunal suction biopsies have been progressively replaced by duodenal endoscopic biopsies. It is now accepted that the diagnostic adequacy of multiple specimens taken through an endoscope is comparable to or even better than that of specimens taken with suction capsules.4 5 The advantages of endoscopic biopsies are their easy and rapid performance as well as visualisation of the loss of duodenal folds, a finding suggestive of coeliac disease which has 88% sensitivity and 83% specificity.3

As the clinical picture is only rarely classic and histological abnormalities are characteristic but not always pathognomonic a definite diagnosis should be based on histological examination before and after a gluten free diet.6 7 8 A gluten free diet can improve the histological appearance within a few weeks, but normalisation of the mucosa may take from a few months to up to one to two years; it is now accepted that gluten rechallenge to confirm coeliac disease is not mandatory in adults8 whereas it is regarded as standard practice in children.9 The recently introduced rectal gluten challenge is a single, safe, and reliable test which could become a welcome alternative to oral gluten challenge.9

Up to 14% of patients with coeliac disease may develop malignancy,8 10 11 but there is no pattern of haematological or biochemical abnormality which enables lymphoma or other associated tumours to be diagnosed easily and clearly. The role of routine annual faecal
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occult blood testing which has been used to
screen healthy populations for early tumours
has yet to be evaluated in coeliac disease.

Methods
To assess the current management in Britain
of adults with coeliac disease a 16 item
questionnaire (box) was sent to the 359
members of the British Society of
Gastroenterology, who were either consultant
physicians or gastroenterologists. More junior
doctors, of senior registrar grade or lower,
were excluded. Subjects were asked to indicate
which alternative corresponded most closely to
their clinical practice; no response was a third
option.

Results
Two hundred and seventy completed
questionnaires were received (response rate
75%). The specialty of the respondents was
medicine 103(38%) and gastroenterology
46(17%); no comment was given in 121 cases
(45%). Most doctors (151, 56%) cared for
between 10 and 49 patients with coeliac
disease, although 76(28%) managed more
than 50.

Two hundred and sixteen respondents
(80%) diagnosed coeliac disease by
endoscopic duodenal biopsy; only 30(11%)
continued to use jejunal capsule biopsy as
their technique of choice. The remaining
24(9%) used either method depending on the
patient and the facilities available. One
hundred and fifty six respondents (58%)
repeated the biopsy after gluten withdrawal:
98 doctors did so at six months and 33 after
a year; the other 25 doctors had no fixed time
for repeating the biopsy. Doctors who initially
chose endoscopic biopsy to reach the diagnosis
repeated the biopsy after gluten withdrawal
significantly more often than those performing
jejunal suction biopsy (134/216, 62% v 13/30,
43%; $\chi^2 = 8.6; p < 0.02$). For follow up
endoscopic biopsy was more commonly used
than jejunal capsule biopsy (133/156, 82% v
23/156, 15%; $\chi^2 = 7.3; p < 0.02$). Only seven
doctors would consider rechallenging their
patients with gluten once they were in
remission, mainly in the case of patients who
were children or for research. Assessment of
the small bowel by radiological examination
was performed by 67(25%) consultant
members of the society.

Patients with coeliac disease are regularly
followed up with body weight assessment
(259, 96%) and haematological and
biochemical reviews. Two hundred and thirty
eight doctors (88%) asked for routine full
blood counts and 120(44%) for measurement
of serum B-12 and folate concentrations. Liver
function tests and calcium estimations were
regularly performed by 105(39%) of
respondents. There was no routine screening
for malignancy, except by annual clinical
review in 13 clinics (5%) or by carcino-
embryonic antigen in three centres (1%).

The last eight items of the questionnaire
referred to methods of patient follow up. As a
rule there was no separate clinic for patients
with coeliac disease (13(5%) exceptions) and
only 75(28%) of doctors offered an open
access service. Only one respondent had a
specialist nurse for patients with coeliac
disease but almost all (96%) encouraged
membership of self help groups and used
information booklets for these patients (248,
92%). Only a few doctors (18, 7%) made use of
information videos for their patients, but
almost a third (79) referred patients to fellow
sufferers to discuss mutual problems.

The last question dealt with the future role
of general practitioners in long term care. One
hundred and seventy one consultants (63%)
considered that patients with coeliac disease
should be followed up by a hospital specialist
and 58(21%) by general practitioners; others
(13, 5%) thought that this decision depended
on the general practitioner and that it should
be possible to develop guidelines for referral
to hospital in that situation.

Discussion
During the past five years there has been a
dramatic change in the standard practice for
diagnosing coeliac disease. Whereas the
traditional approach had been to use jejunal
biopsies, which were often unpleasant for the
patient, time consuming, and required
radiological screening, widespread recognition

<table>
<thead>
<tr>
<th>Coeliac disease (CD): its diagnosis and management</th>
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<tbody>
<tr>
<td><strong>Specialty: medicine/gastroenterology (please specify)</strong></td>
</tr>
<tr>
<td><strong>Approximate number of patients with coeliac disease (CD) under your care?</strong></td>
</tr>
<tr>
<td>0–9</td>
</tr>
<tr>
<td><strong>Diagnosis and management:</strong></td>
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<tr>
<td>1. What type of biopsies do you routinely take? Endoscopic duodenal/jejunal capsule biopsy</td>
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<tr>
<td>2. Do you routinely perform a small bowel radiological examination?</td>
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<tr>
<td>3. Do you routinely repeat small bowel/duodenal biopsy after gluten withdrawal? If yes when?</td>
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<tr>
<td>4. Do you routinely perform a gluten rechallenge test?</td>
</tr>
<tr>
<td>5. Do you follow up patients with CD by tests?</td>
</tr>
</tbody>
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| 6. Do you have computerised follow up for checking:
| (1) calcium | (2) folic acid | (3) anaemia |
| 7. Which of the following do you perform each time you review a patient (please circle)? Weight/full blood count/B-12 and folic acid concentrations/ liver function tests including calcium/faecal occult blood test |
| 8. Do you have a screening system to detect malignant change? |
| 9. Do you have a separate clinic for patients with CD? |
| 10. Do you have a specialist nurse for patients with CD? |
| 11. Do you have an open access clinic for CD? |
| 12. Do you encourage membership of self help groups? |
| 13. Do you make use of information booklets for your patients with CD? |
| 14. Do you make use of information videos for your patients with CD? |
| 15. Do you refer patients with CD to fellow (consenting) patients to discuss problems? |
| 16. Do you think, under the new contract system, patients should be followed up by their general practitioners rather than by a hospital specialist? |
of the value of endoscopic duodenal biopsies has led to a much more rapid procedure which has greater patient acceptability. In experienced hands endoscopic biopsy can take less than 10 minutes compared with 30 minutes or more commonly required for jejunal suction biopsy. Such a significant diagnostic change will probably influence the incidence of diagnosis in adults; endoscopy in children requires a general anaesthetic and carries less advantage. Despite these comments the incidence of coeliac disease in adults has remained fairly constant during the past decade. With the introduction of open access endoscopy to which family doctors can directly refer patients, the diagnosis of coeliac disease has been opened up to those other than specialist gastroenterologists. With this in mind, how important is the long term care of patients with this condition by such specialists? One aspect of this survey was an attempt to define the role of gastroenterologists in long term care. If they have a role it must be in the early detection of biochemical and haematological abnormalities or of malignant change. These approaches include measurement of serum folate and calcium concentrations and are well established. Coeliac disease predisposes the sufferer to both gastrointestinal lymphomas and oesophageal cancer. All intestinal neoplasms tend to bleed microscopically, and faecal occult blood testing can detect abnormalities of this variety, although not specifying their nature. The role of faecal occult blood testing in detecting neoplasia in coeliac disease has not been investigated. Its role in the early detection of colorectal and other gastrointestinal cancers in healthy populations, however, is clear and well established. Compliance is often a problem in these groups, although less so in groups at increased risk of cancer, such as siblings or patients with a history of cancer. We wished to see if such an approach had been adopted by any gastroenterologists working with patients with coeliac disease. In practice, less than half the consultants in the United Kingdom routinely monitored these blood tests and less than 10% had any cancer or lymphoma detection programme.

Although diagnostic practice in coeliac disease has changed significantly, there has not been a comparable change in management. The two main aspects of this study were who should provide continuing care and the possible role of early detection of neoplastic change. Body weight assessment seemed to be the main objective assessment by consultants during routine follow up. Family doctors could probably provide as effective a service at less cost. Indeed, with their experience of immunisation programmes and health screening they might achieve more complete follow up and patient compliance. In the present climate of care profiles and resource management the justification for such a hospital based approach needs to be continually re-evaluated. Family doctors could perform these tests and be given guidelines to suitable indicators of a need for referral. Before such an approach is adopted, however, patients themselves should be approached to ascertain their wishes. Until now there has been only limited acknowledgement by hospital practitioners of the social needs of such patients. This is well exemplified by the fact that only one clinic provided a nurse based counselling service.

The time seems to have come for the British Society of Gastroenterology to produce guidelines for the management of coeliac disease, which could be made available to non-specialists for use in general practice. These guidelines should identify which variables should be regularly reviewed and indicate when patients should again be referred to hospital.

A responsible family doctor could probably manage a patient with coeliac disease, with a significant cost saving without any deterioration in care. This, of course, contrasts with the current practice of most gastroenterologists in Britain.

Dr Monica Acalovschi was a visiting fellow of the Scotland Romania Medical at Leicester General Hospital.

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