Introduction

Diagnostic methods are insensitive only for H2 (all had negative aspirates), 1 responded to antibiotics, 11 had negative aspirates but a 73% antibiotic response. 60 (48%) had a positive breath test—5% H2, 10% CH4 and 35% both gases. 24 (46%) tested positive for both gases. 33% of these had positive D2 aspirates with 3 (50%) responsive to antibiotics. 5 (10%) tested positive for CH4 only, of which 1 had a positive D2 aspirate with a response to antibiotics and 4 (80%) had negative aspirates but 2 responded.

Conclusion

Methane breath testing identifies 10% more patients with SIBO compared to the glucose hydrogen breath test alone. D2 aspirase increases the detection rate by 12%. A trial of antibiotics, other tests negative, benefited 15% of patients. All current diagnostic methods are flawed. Better diagnostic tests for SIBO are required.

Competing interests

None declared.

PTU-156

DIETARY RESPONSE IN COELIAC DISEASE SHOULD BE ASSESSED BY REPEAT DUODENAL BIOPSY

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Introduction

Therapeutic goals in coeliac disease (CD) are poorly defined. Emerging evidence reports worse outcomes with lack of mucosal recovery, suggesting histological normalisation as the aim of treatment. However, few centres routinely re-biopsy and many clinicians depend on serology or symptoms for follow-up. This retrospective study evaluates the role of the follow-up biopsy compared to serology in treated coeliacs and is the largest study of paired biopsies reported.

Methods

Details of patients attending the Addenbrookes Adult CD Clinic are entered onto a database. Patients undergo repeat duodenal biopsy 9–12 months after commencing dietary therapy. Serum Anti-TTG antibodies and total IgA are measured at clinic visits. From the database of around 600 adult patients (50% male, 70% female, average age at diagnosis 48 and 44 yrs respectively) follow-up biopsy results were available for 453 patients and compared using a paired t-test.

Results

Paired initial and follow-up biopsies were available for 368/453 patients (no information about index biopsy in 65). 239 (65%) were re-biopsied within 12 months, 311 (85%) within 24 months. The proportion achieving normalisation of duodenal mucosa (Marsh 0) at 12 months, 12–24 months or after 24 months is 27%, 25% (p=0.61) and 25% (p=0.1) respectively. Those without villous atrophy (VA) (Marsh 0, 1 or 2) is 58%, 58% (p=0.98) and 52% (p=0.24) respectively. Excluding the 2.5% of patients with IgA deficiency, 13% of patients had negative anti-TTG antibody titres at the time of diagnostic biopsy. In the group of patients with positive serology at first biopsy, only 43% with VA on follow-up biopsy were positive for anti-TTG antibodies. Conversely, 13% without VA on follow-up had positive serology. The sensitivity, positive predictive and negative predictive values for serology compared to biopsy at follow-up are 43.4%, 68.3% and 70.5% respectively.

Conclusion

Response to diet in CD might be monitored by symptoms, serology or histology. Many patients diagnosed in adulthood report minimal or no symptoms at diagnosis and this is therefore not a credible marker of response. Here we show that:

- Anti-TTG antibody serology has a sensitivity of 87% at diagnosis
- Anti-TTG serology has a poor sensitivity (43%) on follow-up and is therefore unreliable
- Histological response rates are not increased after 1 year on diet
- Approximately 28% of patients achieve normal duodenal mucosa, 58% show no or minimal signs of coeliac disease on follow-up biopsy.

We suggest that follow-up biopsy on diet is the only reliable form of assessing response to treatment, and that the proportion of patients achieving histological response could be used as a means of auditing quality of care in coeliac clinics.

Competing interests

None declared.

PTU-157

A PROSPECTIVE STUDY OF SERUM VITAMIN A AND D LEVELS IN PRIMARY BILE ACID DIARRHOEA

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Introduction

There is evidence that primary bile acid diarrhoea (BAD) is caused by disruption of the homeostatic mechanism controlling bile acid synthesis, with an overproduction of bile acids leading to chronic watery diarrhoea. Fibroblast Growth Factor 19 (FGF19) has been shown to be central to this homeostatic mechanism, providing negative feedback on CYP7A1, the rate limiting enzyme in bile acid synthesis. Median serum FGF19 levels have been shown to be lower in BAD patients than in diarrhoea controls. In mouse models, it has also been shown that expression of FGF15, the orthologue of FGF19, is induced by the active forms of vitamins A and D. The aim of this study was to investigate a possible causal relationship between low serum levels of vitamins A and D and BAD.

Methods

Patients with chronic diarrhoea were recruited prospectively. All patients underwent routine testing to exclude other causes of diarrhoea and had SeHCAT tests. Patients were classified as having primary BAD, secondary BAD, or unexplained chronic diarrhoea (CD). Serum 25OH-vitamin D was measured by LC-MS using standard procedures. Serum Vitamin A was calculated by HPLC. Results are expressed as medians. Mann–Whitney and Spearman rank correlation tests were used in analyses.

Results

Serum vitamin A levels were available in 145 subjects (primary BAD 46; secondary 29; CD 70). Serum vitamin D levels were available in 150 subjects (primary BAD 50; secondary 29; CD 71). There were no significant differences in the level of vitamin A between the groups with primary or secondary CD (2.10, 4.10, 4.10 ppm). Some patients also had duodenal aspirates collected endoscopically. Positive result: >104 colony forming units/ml. Patients with positive tests were treated with antibiotics. Results were assessed retrospectively.

Results

126 patients, 66 males and 60 females, median group age 61 years (range 35–85), treated for gynaecological (30%), upper GI (25%), lower GI (9%), urological (23%), other cancers (10%)—myeloma, ependymoma, bronchial, breast and lymphoma—were re-biopsied within 12 months, 311 (85%) within 24 months. The proportion achieving normalisation of duodenal mucosa (Marsh 0) at the first biopsy, only 43% with VA on follow-up biopsy were positive for anti-TTG antibodies. Conversely, 13% without VA on follow-up had positive serology. The sensitivity, positive predictive and negative predictive values for serology compared to biopsy at follow-up are 43.4%, 68.3% and 70.5% respectively.
PTU-158 COELIAC DISEASE INVESTIGATION AND FOLLOW-UP

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Introduction Coeliac disease (CD) is an abnormal immune response to gluten affecting 1% of the UK population, resulting in small bowel villous atrophy, malabsorption and GI symptoms. A large number of patients are tested for CD and often repeat requests are made. This retrospective analysis looked at the prevalence of patients with positive CD serology in relation to subsequent gastroenterology referrals and/or small bowel biopsy.

Methods The data were collected from patients who had undergone serological CD testing. All patients with a positive anti-tissue transglutaminase, or a positive endomysial antibody test were included. The data were obtained from the laboratory database at Brighton and Sussex University Hospitals over a period of 1 year.

Results We identified 7569 CD serology requests. The referral source for CD serology was a general practitioner, hospital specialist, gastrointestinal surgeon or gastroenterologist. Overall, 169 (2.5%) individuals had a positive result. From the total number of requests, 6.75% (498) were repeat requests. Of these, 8.23% (41) had a second positive test. 13.86% (69) of the patients who had a repeat serology test were given a clinical diagnosis of gluten sensitive IBS (GS-IBS). Small bowel histology confirmed CD in 52% of the biopsy samples, that is, almost one third of the total number of cases with positive serology.

Conclusion Positive CD serology was recorded in 2.5% of the study group. Although small bowel histology is the gold standard in the diagnosis of CD,1 this was performed on 58% of the patients with positive serology. It is apparent that not all cases with positive serology are being referred to a specialist. We recommend that patients with suspected or incidental diagnosis of CD should be referred to a gastroenterologist for assessment, confirmation of the diagnosis with small bowel biopsy and advice on gluten-free diet.

Competing interests None declared.

REFERENCES

Abstract PTU-158 Table 1 Source of coeliac serology referral

<table>
<thead>
<tr>
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<th>Total serology requests (%)</th>
<th>Positive serology result (%)</th>
<th>Negative serology result (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GP</td>
<td>4502 (61.1%)</td>
<td>82/4502 (1.8%)</td>
<td>4420/4502 (98.2%)</td>
</tr>
<tr>
<td>Other hospital specialist</td>
<td>1716 (23.3%)</td>
<td>41/1716 (2.4%)</td>
<td>1675/1716 (97.6%)</td>
</tr>
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<td>GI surgeon</td>
<td>102 (1.4%)</td>
<td>3/102 (2.9%)</td>
<td>99/102 (97.1%)</td>
</tr>
<tr>
<td>Gastroenterology</td>
<td>1049 (14.2%)</td>
<td>41/1049 (3.9%)</td>
<td>10081049 (96.1%)</td>
</tr>
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</table>

PTU-159 SELF REPORTING OF GLUTEN SENSITIVE GI SYMPTOMS IN PRIMARY CARE: SHOULD WE ACCEPT THE DIAGNOSIS OF COELIAC DISEASE WITHOUT FURTHER INVESTIGATIONS?

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Introduction Many unselected patients presenting to gastrointestinal (GI) clinics self-report that they have symptoms related to the ingestion of gluten. For this reason we undertook a prospective and systematic evaluation of this group of patients to ascertain the subsequent diagnostic yield.

Methods All patients were referred to a dedicated coeliac or gluten sensitivity clinic by GPs through a choose and book system. The referral criteria were “GI symptoms attributed to gluten ingestion.” Investigations included baseline haematology, biochemistry, haematinics, C reactive protein and HLA status for the DQ2/DQ8 haplotypes. In addition, coeliac serology was performed: endomysial antibody (EMA), tissue transglutaminase antibody (tTG), immunoglobulins, as well as duodenal biopsies on a gluten containing diet. A diagnosis of coeliac disease was based on either the presence of villous atrophy or in cases with lesser degrees of the modified Marsh grading, an associated positive coeliac serology and/or HLA typing.

Results 140 patients were investigated over a 5-year period. 80% were women and the median age of presentation was 57 yrs (range 16–88). In patients self-reporting symptoms related to the ingestion of gluten, the diagnosis of coeliac disease was reached in 10% (n 14). 55% (n 119) did not have coeliac disease but fulfilled the ROME criteria for irritable bowel syndrome (IBS). These patients were given a clinical diagnosis of gluten sensitive IBS (GS-IBS). Importantly, organic pathology was found in 5% (n 7) all of whom had additional alarm symptoms—mesenteric ischaemia, bacterial overgrowth, lactose intolerance, bile salt malabsorption, lymphocytic colitis, ulcerative colitis and pyloric stricture. A positive coeliac serology (p<0.0001, exact fisher test) was significantly associated with coeliac disease. All patients with coeliac disease were HLA positive compared to 44% of GS-IBS cases. There was statistically no significant difference in gender, clinical symptoms or baseline bloods (haemoglobin, vitamin B12, folate, ferritin, calcium or albumin) between the groups, (p>0.05).

Conclusion Self-reporting of gluten related GI symptoms only results in a diagnosis of coeliac disease in 10% of cases. The majority of patients do not have overt coeliac disease but may belong to the spectrum of GS-IBS, a relatively new and under researched concept. Moreover, 5% had significant underlying disease. An empirical trial of gluten-free diet prior to referral may be unhelpful and delay the diagnosis in this group of patients. This data suggests that patients who self-report gluten induced GI symptoms should be considered for further investigations.

Competing interests None declared.