Endoscopic and histological pitfalls in the diagnosis of celiac disease: A multicentre study assessing the current practice

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ABSTRACT

Background and aims: the diagnosis of celiac disease requires small bowel biopsies to identify the characteristic mucosal changes. The current biopsy practice among endoscopists for celiac disease is in most part unknown. The aim of this study was to compare the different diagnostic policies in various centers in their current practice.

Method: information from a total of 931 confirmed celiac disease patients was retrospectively obtained retrospectively from nine centers in European and Middle Eastern countries. The number of small-bowel biopsies obtained from the duodenal bulb and the second part of the duodenum was compared among different centers.

Results: the most frequent stage of mucosal changes amongst Iranian subjects was Marsh IIIa whereas in the rest of the study population was Marsh IIIc. Marsh I and Marsh II were more prevalent in adults (P < 0.05) and Marsh IIIc was significantly higher in pediatric ages between 1 and 15 (P < 0.05). The most common number of biopsy specimens obtained from Romanian subjects was 1 (52% of cases), followed by 2 for Iranian (56%), 3 for Lithuanian (66.7%) and 4 for Italian patients (48.3%). For majority of cases, anemia was the most prevalent symptom (18.7%) followed by malabsorption (10.5%), diarrhea (9.3%) and dyspepsia (8.2%), respectively.

Conclusions: despite the evidence-based recommendations, this study revealed a poor compliance with major guidelines on diagnosis of celiac disease. We emphasize that taking adequate number of duodenal biopsies should be implemented for an accurate diagnosis and also for the exclusion of celiac disease.

Key words: Celiac disease. Pitfall. Biopsy specimen. Multicenter.

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INTRODUCTION

Diagnosis of celiac disease (CD) is one of the most challenging aspects of this condition. The immune response to the grains of wheat, rye, and barley leads to progressive damages in the small intestine, affecting differently many individuals and resulting in the production of IgA serum antibodies directed against tissue transglutaminase (tTG) in most but not all (1-4).

Despite the availability of specific and sensitive serological tests, diagnosis of atypical forms of CD could be very challenging (5). Intestinal biopsy is still considered an essential diagnostic tool for detection of CD in different stages (6). Histology also permits to evaluate the response to a gluten-free diet, should there be any doubt about the diagnosis.

The sensitivity of the serological tests available might be correlated with the degree of mucosal abnormalities and according to many studies, a negative serological test may not always exclude the diagnosis of CD (7). The mucosal lesions in CD can be patchy, and if the sampling is insufficient, the risk of missing the diagnosis increases (8,9). Low degrees of mucosal abnormalities like microscopic enteritis in small intestinal specimens are not specific for CD, but a combination of clinical, serological and genetic evaluation may help to confirm the diagnosis (10). If possible, it would be more desirable to avoid endoscopy completely. We might argue that endoscopy may be unnecessary a small subgroup of patients who have classic symptoms and high level positive serology, but we still find that taking a biopsy appears essential in the majority of cases (11). A multicentre endoscopy database study found that most patients undergoing upper GI endoscopy for indications such as anemia, iron deficiency, and weight loss had no duodenal biopsy taken during the procedures (12-14). Moreover, a normal endoscopic appearance lacks sufficient sensitivity to exclude CD (15,16).

As mucosal abnormalities in CD are patchy and the orientation of biopsies are variable, multiple biopsies (4-6 biopsies) from the duodenal bulb and descending duodenum are usually recommended as standard method for CD evaluation (10,17-19). Study results by Pais and co-workers showed that 2 biopsies confirmed CD in 90% of patients, while 4 biopsies established a diagnosis in 100% of cases (10). By using a large, international, pathology database, we assessed the adherence to these proposed guidelines.

In this study, we investigated the procedure-related factors associated with the submission of biopsy specimens and compared the different diagnostic criteria in various centers in Italy, Iran, Lithuania, Romania and the UK respectively. This study elucidates how a guideline is exercised in clinical practice, both in terms of adherence to the recommendation as well as the incremental yield of adherence.

MATERIAL AND METHODS

Patients and settings

In this retrospective multicentre study, data were collected in the period May 2009-May 2011 from Iran (Tehran), Romania (Bucharest), Lithuania (Vilnius), the United Kingdom (Birmingham) and 6 different areas in Italy, including Brescia, Bologna, Desio, Messina, Perugia and Empoli. For each center, at least 100 confirmed CD patients with endoscopic procedures and small bowel biopsies were included.

Diagnosis of celiac disease in different centers

Histopathological findings in all centers were evaluated in accordance with the modified Marsh classification (1). The final diagnosis of celiac disease was based on a combination of clinical, analytical and histological data, and a response to a gluten-free diet, and defined by clinicians in charge of clinical care of patients.

In Iran, IgA-tTG antibody and immunoglobulin A were measured using a commercially available kit (AESKULISA tTG, Germany) and an enzyme linked immunosorbent assay (ELISA) method. In Italy, IgA-anti-endomysial antibodies (EMA) were investigated by indirect immunofluorescence that used human umbilical cord cryostat sections. Sera were tested at the initial dilution of 1:5 and when positive, titrated up to the end point. In Romania, the patients suspected for CD were usually referred directly for the gastroenteroscopy because of lack of funding for human leukocyte antigen (HLA)-typing and serology is performed only in cases with suspected CD at biopsy.

The duodenal specimens in Bucharest were oriented using Endokit and the biopsies in some departments in Italy (Brescia, Desio and Messina) were oriented on acetate cellulose filters.

Duodenal biopsy and histology classification

To determine the number of duodenal biopsy specimens for each biopsy set, we used a free-text search of the pathologist’s description of each sample. When present, specimens from the duodenal bulb were included in the total number of specimens submitted. In each lab, biopsies were interpreted by expert gastrointestinal pathologists (blinded to the clinical data). Centers with the lowest and highest number of biopsy specimens were analyzed separately and also compared in order to investigate causes for those differences.

All centers used the cutoff of > 25 intraepithelial lymphocytes per 100 enterocytes for a Marsh I stage, increased intraepithelial lymphocytes accompanied by crypt hyperplasia (Marsh II), partial (Marsh 3A), subtotal (Marsh 3B)
and total villous atrophy (Marsh 3C) as described in modified classification (1). According to the follow up reports to ordering HLA typing, the genetic data available were incomplete and HLA DQ typing was carried out only in some centers.

**Data collection form**

During the same period, the number of biopsy sample taken from each center and their clinical data were compared. For each specimen, the following information was available: sex and age of the patient, location and sample provider, summary of the clinical history, serology/genetic and histopathological findings.

GI symptoms such as abdominal pain, diarrhea, constipation, nausea and vomiting, weight loss and flatulence as well as additional signs and symptoms such as iron deficiency anemia, osteoporosis, hypertransaminasemia, malabsorption, and other related abnormalities were recorded. Malabsorption was defined by a decreased body mass index, weight loss and low triglyceride, cholesterol, ferritin or serum albumin and also anemia was defined when the level of hemoglobin and hematocrit (HCT) was decreased less than the normal quantity.

**Statistical analysis**

We used the chi-square test to assess the association between adherence to the recommendation of submitting ≥ 4 specimens and the proportion of patients with pathological findings consistent with CD. The results with a p-value < 0.05 were considered statistically significant. Because our data set did not contain information regarding HLA typing for all cases and clinical follow-ups, we defined a priority that results with either blunted villi (Marsh 3A) or flat villi (Marsh 3B/C) were meeting the pathological definition of CD. For assessing the relationship between ordinal categories such as the number of specimens and the pathologic diagnosis of CD, we used the Fisher’s Exact Test for trend.

**RESULTS**

Overall, 658 women and 273 men (mean age at diagnosis 35.4 ± 17.25 years, median age 36 years, age range 1-72 years) with a diagnosis of CD were included. Patients ranged 1-15 years old were considered as children and older as adults. In all participating countries females were predominant (70.7% of patients), and the mean age (± standard deviation [SD]) was 36.6 ± 16.4 years. Lithuanian patients were the oldest, with a mean age of 43.7 ± 18.6 yrs, and Iranian patients the youngest (30 ± 13.6 yrs). There was a statistically significant differences between the genders in studied countries (p = 0.0001). Fifty one patients were excluded based on their incomplete data collection.

Table I showed demographical, clinical, and histopathological data in the study population.

One hundred and four samples (11.17%) were collected from the bulb and the rest was collected from the second duodenal portion. Marsh I and II lesions were noted in 322 individuals (34.5%), Marsh 3A was found in 180 (19.3%), Marsh 3B in 178 (19.1%) and Marsh 3C in 233 study subjects (25%) (Fig. 1). When a pathological diagnosis of CD was defined as that of blunted or flat villi (Marsh 3A/B/C), a total of 591 individuals (63.4%) were categorized as having CD. According to the Marsh classification there were statistically significant differences among studied countries (p = 0.0001). A Marsh 0 stage was recorded in 18 individuals (2%) being most of them were under 15 years old.

According to age groups Marsh IIIc was significantly more prevalent in pediatric patients (aged 1-15) compared to other age groups (p = 0.0001) and Marsh I and II were predominantly present in adult patients (p = 0.0001).

Regarding the number of biopsy specimens taken per CD case, we found that 1 biopsy specimen had led to a CD diagnosis in 63 (6.7%) cases (11 specimens were Marsh I and II and the rest was higher stages), 2 in 212 (22.7%) cases (96 specimens were Marsh II or lower), 3 in 306 (32.9%) cases (153 were Marsh II or lower), 4 in 299 (32.1%) cases (81 specimens were Marsh II or lower) and ≥ 5 in 51 (5.5%) cases (9 specimens were Marsh II or lower). The average number of biopsy specimens per patient was 3 (Table II, Fig. 2). The number of available biopsy samples directly associated with increasing Marsh stages (p = 0.0001).

The most common number of biopsy specimens taken from Romanian subjects was 1 (52%) followed by 2 from Iranian patients (56%), 3 from Lithuanian (66.7%) and British patients (65%) and 4 from Italian patients (48.3%). Villous atrophy (partial, subtotal or total) was the predominant histological lesion in all studied countries (Fig. 1). There was a statistically significant differences regarding number of biopsy specimens taken from studied countries (p = 0.0001).

The most common number of biopsy specimens taken from patients age 1-15 years old (83 cases) and adults age 31-45 (117 cases) was 3. This differences was statistically significant for Italian and Lithuanian patients (p = 0.0001).

The most frequent degree of villous atrophy for Iranian subjects was 3A and for the rest of the study population 3C (Table III). According to the histological variability illustrated above, a statistical correlation was found between type and distribution of the histological lesions and the number of biopsy specimens (p = 0.0001). Four or more duodenal biopsies had been taken for 58.7% of the Italian, 37% of the British, 7% of the Romanian and 5% of the Iranian studied subjects. None of the Lithuanian patients had 4 or more duodenal biopsies.
Routine immune-histochemical evaluation of T lymphocytes with CD3 and CD8 antibodies was used only in Iran and one center in Italy (Brescia).

To confirm the diagnosis of celiac disease, serological assessments were performed in all the participating centers. Anti-EMA resulted positive in 32.3% of Iranian patients; positive anti-tTG IgA was documented on 67.7% on Italians, and positive anti-gliadine IgA antibodies were documented in 6.3% of the UK patients. Regarding to Lithuanian patients 34% of them exhibited positive anti-gliadine IgA, being the remaining positive for anti-tTG IgA antibodies.

HLA typing was not available for Lithuanian subjects, and if performed, there was no access to genetic results of the HLA DQ typing in this country. HLA typing was performed for very few selected cases in Romania. HLA typing for Italian (72/602) and Iranian subjects (52/100) showed that, HLA DQ2 was the predominant genotype for both countries.

The clinical data varied from one region to the other; however, anemia was the main presenting symptom.

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Table I. Demographic, clinical and histopathological information in the study population

<table>
<thead>
<tr>
<th>Countries</th>
<th>Iran</th>
<th>Italy</th>
<th>Lithuania</th>
<th>Romania</th>
<th>UK</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age ± SD</td>
<td>31 ± 13.6</td>
<td>33.3 ± 18</td>
<td>43.7 ± 18.6</td>
<td>38.9 ± 12.3</td>
<td>36.3 ± 15.2</td>
<td>p &gt; 0.05</td>
</tr>
<tr>
<td>Gender</td>
<td>Male (subjects)</td>
<td>35</td>
<td>152</td>
<td>37</td>
<td>21</td>
<td>58</td>
</tr>
<tr>
<td></td>
<td>Female (subjects)</td>
<td>64</td>
<td>347</td>
<td>92</td>
<td>79</td>
<td>45</td>
</tr>
<tr>
<td>Marsh classification</td>
<td>0</td>
<td>0</td>
<td>18</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>I</td>
<td>19</td>
<td>83</td>
<td>83</td>
<td>7</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>II</td>
<td>19</td>
<td>62</td>
<td>22</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>IIIa</td>
<td>27</td>
<td>91</td>
<td>24</td>
<td>19</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>IIIb</td>
<td>13</td>
<td>106</td>
<td>0</td>
<td>25</td>
<td>34</td>
</tr>
<tr>
<td></td>
<td>IIIc</td>
<td>20</td>
<td>141</td>
<td>0</td>
<td>46</td>
<td>26</td>
</tr>
<tr>
<td>No of biopsy</td>
<td>1</td>
<td>0</td>
<td>11</td>
<td>0</td>
<td>52</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>56</td>
<td>49</td>
<td>43</td>
<td>32</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>39</td>
<td>146</td>
<td>86</td>
<td>9</td>
<td>86</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>5</td>
<td>253</td>
<td>0</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>≥5</td>
<td>0</td>
<td>40</td>
<td>0</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Clinical symptoms (subjects)</td>
<td>Abdominal pain</td>
<td>3</td>
<td>18</td>
<td>0</td>
<td>6</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>Diarrhea</td>
<td>32</td>
<td>44</td>
<td>0</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Constipation</td>
<td>8</td>
<td>19</td>
<td>0</td>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>Weight loss</td>
<td>5</td>
<td>7</td>
<td>0</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Anemia</td>
<td>55</td>
<td>72</td>
<td>37</td>
<td>0</td>
<td>6</td>
</tr>
</tbody>
</table>
in 174 (18.7%) patients, malabsorption appeared in 98 (10.5%), diarrhea in 87 (9.3%) and dyspepsia in 77 (8.2%) patients, respectively (Fig. 3). There was a statistically significant correlation between these symptoms and abnormality in histology ($p < 0.05$). as shown in figure 3, anemia and diarrhea associated with marsh 3c stage, malabsorption was predominant in marsh 1, and dyspepsia associated with marsh 3b. Ninety percent of the Lithuanian study subjects were affected by malabsorption. The most frequent symptom in Italian patients was dyspepsia (73 cases [14.6%], probably related with a high-carbohydrate diet), followed by anemia (71 cases [14.2%]) and diarrhea (44 cases [8.8%]). For Iranian subjects the predominant symptoms were anemia (55 cases [55%]), diarrhea (32 cases [32%]) and osteopenia (25 cases [25%]). Romanian subjects predominantly complained of diarrhea (8 cases [8%]), abdominal pain (6 cases [6%]) and weight loss (5 cases [5%]). Statistically significant differences were observed when comparing the reported clinical symptoms in different countries ($p < 0.05$).

Abdominal pain and failure to thrive were significantly reported in pediatrics ($p = 0.01$) and anemia, osteoporosis, osteopenia and dyspepsia were more prevalent in adults ($p < 0.05$) (Table IV).

In spite of the histological variability illustrated in this study, a significant correlation was found between type and distribution of the histological abnormalities and some of the clinical symptoms such as abdominal pain, diarrhea, low vitamin B12/Acid folic serum levels, malabsorption, IgA deficieny, dyspepsia, asthenia and failure to thrive ($p < 0.05$).

Among the 931 study subjects, 419 (45%) displayed typical presentation, 430 (46.2%) atypical presentation,
and 64 (6.9%) had no or only minor GI symptoms. A statistically significant correlation was found between abnormal histology and abdominal pain, dyspepsia and failure to thrive for Italian patients and between abnormal histology and anemia for Lithuanian patients ($p < 0.05$).

**DISCUSSION**

Even though noninvasive antibody testing is an excellent screening tool for CD, endoscopic evaluation via duodenal biopsy in suspected patients remains crucial in the diagnosis of clinical atypical CD.

The typical symptoms associated to CD are abdominal pain, chronic diarrhea, vomiting, weight loss, foul smelling stool, fatigue, failure to thrive or short stature, delayed puberty, osteoporosis and anemia, and the atypical celiac disease symptoms include obesity, dyspepsia, constipation, depression and anxiety (4).

The number of diagnosed patients has increased; this may be due to a greater proportion of outpatients being investigated for relatively minor symptoms, especially elderly people who visit their physicians more often (20).

In this analysis of an international pathology database of duodenal biopsies, only 37.6% of patients had ≥ 4 specimens submitted upon upper endoscopy. Adherence to the proposed standard remained low even for patients with typical CD, with fewer than 40% of such patients having ≥ 4 specimens submitted (1,17,20,21). Italy had the best record of adequate sampling in this study and the rest of participant disappointingly failed to adhere to evidence based recommendations.

Biopsy handling and experience of pathologists in different sites including duodenal bulb (19) was also variable in this study. It is suggested to take multiple biopsies from different duodenal sites in all cases where CD is suspected in order to minimize the risk of misdiagnosis (10). Differences in practice are highlighted and we can see the different approaches and diagnostic workups in countries participated in this evaluation. Biopsy specimens taken from the descending duodenum as well as jejunal specimens produced good results. However, duodenal bulb biopsy specimens despite being very important (17,19), were poorly taken in this study. In addition, a duodenal biopsy should have a reasonable size, be well orientated and adequately stained and should be thoroughly examined by an experienced GI pathologist (22). There was not a reliable tool available to assess all these qualities in the biopsy samples because of the nature of this multicentre study. Although general pathologists might not be aware of the variety of pathological changes seen in CD, in a positive note all the pathologists participated in this study had expertise and interest in histology of CD.

Villanacci et al. suggested that the biopsy taken from anywhere along the duodenum could theoretically lead to the diagnosis of CD if the specimen was correctly obtained and assessed according to the standards (22,23).

In a recent review, Yantiss & Odze (2009) referred to the “optimal method of obtaining biopsies in patients with celiac disease” as “controversial” (24). It may be stressed at this point that the proposed guideline entirely derives from the observation that the histopathologic abnormalities of CD are patchy, and can be missed if an insufficient quantity of specimens is submitted. The recommendation is supported by a single-centre retrospective study of 93 patients with CD, which found that 4 specimens led to a positive diagnosis in 100% of patients, whereas 2 specimens were diagnostic in only 90% of patients (10). The authors concluded that at least 4 duodenal biopsy specimens should be taken to rule out CD. A second study, investigating 56 patients with known CD, found that 3 biopsy specimens were sufficient as long as 1 specimen was obtained from the duodenal bulb; however, 5 biopsy specimens were necessary to recognize the most severe

**Table IV. Different signs and symptoms in pediatric compare to adult celiac patients retrospectively recruited in our multicenter study**

<table>
<thead>
<tr>
<th>Sign and symptoms</th>
<th>Pediatrics (%) n = 127</th>
<th>Adults (%) n = 804</th>
<th>$p$ value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal pain</td>
<td>11 (8.7)</td>
<td>19 (2.3)</td>
<td>0.01</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>9 (7)</td>
<td>75 (9.3)</td>
<td>0.3</td>
</tr>
<tr>
<td>Constipation</td>
<td>7 (5.5)</td>
<td>32 (4)</td>
<td>0.7</td>
</tr>
<tr>
<td>Anemia</td>
<td>24 (18.9)</td>
<td>145 (18)</td>
<td>0.03</td>
</tr>
<tr>
<td>Osteoporosis</td>
<td>0</td>
<td>13 (1.6)</td>
<td>0.003</td>
</tr>
<tr>
<td>Osteopenia</td>
<td>0</td>
<td>30 (3.7)</td>
<td>0.005</td>
</tr>
<tr>
<td>Hypertransaminasemia</td>
<td>0</td>
<td>19 (2.3)</td>
<td>0.01</td>
</tr>
<tr>
<td>Weight loss</td>
<td>4 (3.14)</td>
<td>13 (1.6)</td>
<td>0.83</td>
</tr>
<tr>
<td>Dyspepsia</td>
<td>1 (0.78)</td>
<td>75 (9.3)</td>
<td>0.001</td>
</tr>
<tr>
<td>Bloating</td>
<td>1 (0.78)</td>
<td>12 (1.5)</td>
<td>0.48</td>
</tr>
<tr>
<td>Failure to thrive</td>
<td>15 (11.8)</td>
<td>5 (0.62)</td>
<td>0.0001</td>
</tr>
</tbody>
</table>
extent of villous atrophy. These studies are limited by small sample size and single-center settings (25). Evans et al. recently solve these problems (19). They investigated overall 461 patients including 126 newly diagnosed CD, 85 established CD, and 250 controls. Their study suggests that the optimal strategy for diagnosing CD in both suspected patients with positive serology and those with established CD could only achieve 100% sensitivity by always incorporating a duodenal bulb biopsy (19).

In this study, taking 1-2 biopsy specimens led to the confirmation of diagnosis in 29.5% of cases, while more than 3 biopsy specimens led to the diagnosis in 70.5%. We found a statistical correlation ($p < 0.05$) between the number of biopsies and histological abnormalities. The variation in the number of biopsy samples in this study may be associated with the endoscopists' practice in each region. Endoscopists in Romania, Iran and Lithuania are more likely to perform less and fewer numbers of biopsies compared to endoscopists in Italy.

Because villous atrophy in CD is patchy and the orientation of the specimens is variable, 4 to 6 random biopsy specimens including at least one from the duodenal bulb are recommended in order to maximize the sensitivity and achieve a correct diagnosis of CD (10,17,19,22). This may suggest that if additional biopsy specimens in our study had been taken from subjects CD may have been confirmed in more cases (10).

The incremental yield varied with indication and was greatest when the symptom was anemia (18.7%). However, submitting ≥ 4 specimens also increased the diagnostic yield of CD even when the indication was dyspepsia (8.2%). As H. pylori infection is associated with some mucosal abnormalities in dyspeptic patients, Lecleire et al. suggest that endoscopic markers of villous atrophy are not useful for the detection of CD in patients with dyspeptic symptoms (26).

Retrospective analysis of pathology tissue databases produced high-quality analyses of GI epidemiology and quality measures (27,28). In our study, conducted in a similar manner, we have some limitations. For instance, we did not have access to socioeconomic or racial data to determine whether these individual patient characteristics were associated with the submission of the recommended number of specimens. We also had no access to key variables that influence the likelihood of CD, such as data regarding family history of CD. We looked for including patients with confirmed diagnosis for most of the locations. We did not describe a yield of biopsies because the total number of patients biopsied for suspected CD and control group was not included. Therefore, the false negative rate was not assessed.

Moreover, information regarding the type of sedation used during the procedure and the degree of sedation, which may have impacted the ability to obtain ≥ 4 specimens, was not available. The degrees to which endoscopists adhere to such recommendations in clinical practice and the diagnostic yield of adherence to this standard have not been studied.

Since the study was retrospective and many patients did not have uniform information regarding the degree of adherence to the gluten-free diet in their medical record, we were not able to assess the correlation between seronegativity after treatment and the degree of compliance to the GFD (27). For example in Bucharest, for the majority of the patients that are suspected to have CD, endoscopy is performed prior to serological. After the histological diagnosis compatible with CD, patients tend to have serological tests and, rarely, genetic testing. As methodological approach to the diagnosis of CD in this region may be attributed to high costs involved with genetic testing for DQ2/DQ8 (more than minimum salary/month), HLA typing is rarely performed. A similar situation is present in Iran and Lithuania; because HLA typing is expensive and access to laboratories that can perform genetic testing is limited, HLA typing is rarely carried out, and if performed, then it will be only in specific cases. This may represent the main obstacle for clinicians working without the facility for HLA-typing. However, the situation is not better in UK, as HLA typing is not popular between clinician again due to high cost and National Health Service strict cost saving policies. According to the current literature serology only at high level > 10x above normal value and histology only with Marsh IIIb-c have the highest sensitivity and specificity close to 100% for CD. HLA would be very useful in excluding the diagnosis thanks to its very high negative predictive value.

Another obstacle for correct diagnosis in all investigated geographical locations may be that clinicians generally do not share clinical data or serology results of patients with pathologists. Most pathologists ignore the subtle histological changes like microscopic enteritis as Functional Gut Disorders'.

Regarding to clinical presentation of CD, our results indicate that a classical form of celiac disease is predominant in pediatric patients compared to adult CD, and greater severity and intestinal abnormality are seen in younger patients. In contrast, atypical forms predominate among adults, with lower stages of duodenal damage. Our results are compatible with those by Rodrigo-Sáez et al. (30).

As a conclusion, we should implement the biopsy guidelines with ≥4 biopsy specimens into clinical practice
including or with additional samples from bulb (17). In our multicentre study, only the minority of patients with upper GI endoscopy and duodenal biopsy except Italian cases had ≥ 4 specimens. In the light of the beneficial effect on the histological diagnosis of CD (31), in particular in atypical cases increased adherence to biopsy guidelines is warranted. In typical cases, as suggested by the European Society for Pediatric Gastroenterology and Nutrition (ESPGAN), the endoscopy and biopsy obviously might be avoided. However, the majority of celiac patients present with atypical manifestation where endoscopy and the correct sampling approach are paramount in diagnostic pathway (5).

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REFERENCES