The Significance of CD3 Marker in the Diagnosis of Celiac Disease

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ABSTRACT
Background: Celiac disease is a common, permanent and reversible health problem of small intestine occurring all over the world in genetically predisposed individuals and in combination with other environmental factors. It causes chronic inflammation of small intestine which is of autoimmune origin. The histopathological features of Celiac disease in duodenal biopsy was stated according to the modified Marsh classification. The immunohistochemical application of CD3 marker in duodenal biopsy could facilitate the count and the distribution of intraepithelial lymphocytes along the villi, which is regarded as a key for the correct diagnosis of Celiac disease in early stages (Marsh 1).

Objectives: This study was conveyed to correlate the significance of CD3 immunohistochemical expression of intraepithelial lymphocytic population to histopathological changes in Celiac disease, to identify the distribution of CD3 marker along the villi (crescento or decrescendo) and (diffuse or patchy) in duodenal biopsy and to delineate the age and sex of Celiac disease in our locality.

Materials and methods: This prospective and retrospective case series study was carried out on 100 cases of endoscopically obtained duodenal biopsies. Data obtained from archives of the pathology department, at AL-Jamhory, AL-Khansaa and AL-Salam Teaching Hospitals/ Mosul city, and collected in a period spanning from January 2019 to May 2020. The information included age, sex and duodenal biopsy location. Modified Marsh classification was assessed histologically and immunohistochemically.

Results: In a total of 100 duodenal samples, patients age was ranged 1 to 69 years (mean age of 20.74 years), with a female to male ratio (2.2:1).

By applying modified Marsh classification: Marsh 0 was detected in 8 % of the cases, Marsh 1 in 30 % of the cases, Marsh 2 in 10 % of the cases, while Marsh 3 a in 20 % of the cases, Marsh 3 b in 17 % of the cases, Marsh 3 c in 15 % of the cases and Marsh 4 in 0 %. Immunohistochemical expression of CD3 in the sampled cases i.e. CD3 + ≥30 /100 epithelial cells was detected in 79 % of the cases. There was a statistically significant difference between CD3+ immunohistochemical study and modified Marsh classification by Hematoxylin & Eosin (P Value<0.001) for detection of intraepithelial lymphocytosis.

Conclusion: There is a significant relationship between the count of CD3+ T-lymphocytes per 100 epithelial cells and the histopathological changes in the duodenal biopsy according to modified Marsh classification. So, the immunohistochemical expression of CD3 in intraepithelial T-lymphocyte could lead to a definite assessment in 43.3 % of the sampled cases with Marsh type 1. All the positive cases are of crescento pattern of distribution of CD3+ T-lymphocytes as the distribution is more important than the actual count and they distributed diffusely except that associated with Helicobacter pylori infection observed with patchy distribution. In addition to that, the IHC expression of CD3+ marker provides a hint about the distribution of CD3+ marker within the lymphocyte whether global surface or clonal surface and intracytoplasmic to diagnose Refractory Celiac disease.

On the other hand, females were more affected than males with CeD and there is a significant relationship between the gender of the sampled cases and the histopathological changes in the duodenal biopsy. The disease can be diagnosed at any age and there is no significant relationship between the age distribution of the sampled cases and the histopathological changes in the duodenal biopsy.

Keywords: Celiac disease, intraepithelial lymphocytes, modified Marsh classification, CD3, Immunohistochemistry.
The Significance of CD3 Marker :

INTRODUCTION
Celiac disease (CeD) is a common, permanent and reversible health problem of small intestine occurring all over the world in genetically predisposed individuals and in combination with other environmental factors. It causes chronic inflammation of small intestine which is of autoimmune origin. Celiac disease has a wide spectrum of clinical features gastrointestinal and non-gastrointestinal manifestations. It affects both children and adult age groups 

There are many methods to diagnose Celiac disease like serological tests, upper gastrointestinal endoscopy, histopathological features of the biopsies and others 

الخلاصة
الخليقي: اداء الزلاقي هو مشكلة صحية شائعة دائمة وقابلة للانكسار في الامام المقترح يحد في كافة أنحاء العالم عند الأشخاص المتغذين عيانا. لا يوجد تشابه بين الأسباب الرئيسية. كما أنها تؤدي إلى التهاب متصرف في الأعصاب المنكوبة ويعود ذو مصدر مناعي ذاتي. التشريح المرضي لداء الزلاقي في الجراحة الأثنتي عشر يصف حسب تصنيف مارش المعدل.

استخدام الفحص المناعي السيحي الكيميائي لواحة عقود التمايز الثالث للكزيزات الإثنتي عشر من الممكن أن تشمل عملية عدد ومعرفة توزيع الخلايا الملفافية داخل الطبقة الطلائية على طول الزغبات. كما يعتبر مفتاح للاطفال لتشخيص الصحيف للداء الزلاقي في مراحل الالزقاء (مارش (1).

الهدف من الدراسة: أظهر العلاقة اللاحقة لاهية التعبير عن الفحص المناعي السيحي الكيميائي لعقد التمايز الثالث للللمفاوي والتعابير السيحي في الداء الزلاقي. لتحديد توزيع عقود التمايز الثالث على طول الزغبات (تصاصي تعني ) (ومنشأ أو مركع في) في الجراحة الأثنتي عشر والتنقيط للمعجم المعروف في موقع الجرع.

المواد وطرق العمل: هي دراسة مبكرة ومقيمة أجريت على متى حالة من حالات أثنتي عشر المتواجدة تشخيصها بالداء الزلاقي. المعلمات التي تم الحصول عليها من كيف فروع الأعراض. في مختبرات مستشفيات مدينة الموصلي (الجامعى) (الخصم والسلام) (التعليمية) تم جمعها في الفترة الممتدة من كانون الثاني 2019 إلى يار 2020 (المعلومات شملت: العمر، الجنس، وموقع خزيحة أثنتي عشر عن تصنيف مارش المعجم تم تقييمه نسبياً والفحص المناعي السيحي الكيميائي.

النتائج: من متى خزيحة أثنتي عشر، كانت أعمار المرضى تتراوح بين 1 إلى 6 سنة (متوسط العمر 2.2 سنة) (نسبةً من الحالات إلى الذكور 80% والنساء 20%) . حسب تقبيل تصنيف مارش المعجم كان مارش 3: 0% من الحالات، مارش 1: 50% من الحالات و مارش 2: كان 10% مارش 3: 20% مارش 3: 17% من الحالات، كان 5% مارش 3: 20% من الحالات.

الفحص المناعي السيحي الكيميائي لعقد التمايز الثالث على عودات، كان عقود التمايز الثالث موجبة أو < 50 لكل 100 خليلة طليانية في 97% من الحالات. يوجد علاقة أخصائية ذات أهمية بين الفحص المناعي السيحي الكيميائي لعقد التمايز الثالث وبيرو الحفظ المحمل بصبغة الهيماتوكسيلين والأدينين (قيمة احتمالية أقل من 0.001) لتحديد الخلايا الملفافية طليانية، بين الطبقية الطلاانية.

الاستنتاجات: يوجد علاقة إحصائية ذات أهمية بين عدد الخلايا الملفافية بيرو الموجبة لعقد التمايز الثالث لكل متى خزيحة طليانية، بين التغييرات السيحي في المرضية في خزيحة الأثنتي عشر حسب تصنيف مارش المعجم لهذا فالتعبير عن الفحص المناعي السيحي الكيميائي لعقد التمايز الثالث في الخلايا الملفافية في من الممكن أن يؤدي إلى تقلير جسم لم تصل. 43% من العينات وهي تقع بين المريض التمايز ليغدو في جميع الحالات الإجمالية بعدما كانت ذات توزيع مركع وهي متمائلة بعد كزريا العصبية المطلوبة البالية بالإضافة إلى ذلك تكون توزيع صبغة السيحي الكيميائي بشكل مكلا على سطح الخلايا الملفافية الإجمالية لعقد التمايز خلال أو توزيع السيبي للعصب (على سطح الخلية، وهي في حالة السيرتيلين للخلايا الإجمالية لعقد التمايز الثالث يمكن أن يشير إلى الداء الزلاقي المقاوم. من جهة أخرى، لوحظ بأن الألم قد تكون أكثر تأثيرا من الذكور إلى الداء الزلاقي، يمكن لهذا المرض أن يثير في أي عمر.

المفاتيح المفيدة: الداء الزلاقي، الخلايا الملفافية خلال الطبقة الطلائية، تصنيف مارش المعجم، عقود التمايز الثالث، الفحص المناعي السيحي الكيميائي.
Histopathological features depend on the number and distribution of intraepithelial lymphocytes (IELs), presence or absence of crypt hyperplasia and villous atrophy. Then a conclusion stated according to the modified Marsh classification 5-7.

Sometimes there is increase in intraepithelial lymphocytes infiltration without crypt hyperplasia and villous atrophy in duodenal biopsies (Marsh 1) which is not specific for Celiac disease and other causes should be excluded like infections by Helicobacter pylori bacteria, Giardia lamblia, Tropheryma whippeli, Cryptosporidium and many others 8-10.

When immunohistochemistry (IHC) is used for CD3 in Celiac disease, it facilitates counting of IELs and noticing the distribution along the villi (crescendo or decrescendo) and (diffuse or patchy), also it provides a hint about the distribution of the staining (surface or intracytoplasmic) to diagnose Refractory Celiac disease and differentiate between its two types as type II may precede to enteropathy associated T-cell lymphoma (EATL) 3,11-14.

MATERIALS AND METHODS

Patient Selection
This study is a retrospective and prospective case series study which was performed on 100 consecutive cases suspected to have Celiac disease over a period of 16 months started from January 2019 to May 2020. A tissue block of each case was chosen for histopathology and IHC.

This work was maintained a routine protocol approval to access the clinicopathological data from archives of the department of pathology at AL-Jamhory, AL-Khanssa and AL-Salam Teaching Hospitals in Mosul city, and these included: age, sex and duodenal biopsy location.

Histopathological and IHC Staining

Use of formalin fixed, paraffin embedded blocks for each case. Once the slides prepared by two methods: H&E and IHC stains, prepared for target retrieval prior to IHC procedures, using Dako Target Retrieval Solution (10x) that prepared by diluting the concentrate 1:10 in distilled or deionized water together with water bath based Dako PT Link, for deparaffinization, re-hydration and epitope retrieval to increase staining intensity with many primary antibodies by unmasking of the antigen. Then staining by automated staining instrument which is Dako autostainer link 48 slide stainer that used for efficient IHC staining and thermoscientific CD3 early T-cell marker for qualitative IHC that is of clone designation SP7. Positive controls as well as negative controls run simultaneously using the same protocol as the patient’s specimens.

The CD3 marker was optimized using tonsils (T-lymphocytes in the interfollicular areas and dispersed T-cells in the mantle zones and within the germinal centers) as a positive control which displays at least a moderate to strong and distinct membranous staining reaction, with no staining must be seen in the germinal center B-cells which was considered as negative control.

Scoring and Analysis:
All cases are microscopically examined for interpretation of histopathological and IHC stains. So each case stained by H&E designated according to modified Marsh Oberhuber classification of histologic findings in CeD 5-7.

Stage 0 normal (preinfiltrative mucosa).
Stage 1 lymphocytic enteritis (infiltrative).
Stage 2 lymphocytic enteritis + crypt hyperplasia (hyperplastic).
Stage 3 lymphocytic enteritis + crypt hyperplasia + villous atrophy (destructive).

3a mild atrophy.
3b marked atrophy.
3c complete atrophy.

Stage 4 flat mucosa + crypt hypoplasia + mild inflammation (atrophic).

Then by antiCD3 IHC study.

If 30 IELs and more positive for CD3, the case was considered positive.
If 26-29 it is borderline.
If 25 and less it is negative 8,9,15, and their distribution whether in crescendo pattern i.e. increased in number of IELs from the base to the tip of the villous or decrescendo pattern i.e. decrease in number of IELs from the base to the tip of the villous.

The distribution of CD3 positive IELs was noted whether diffused or patchy. Differentiation between surface CD3 and intracytoplasmic CD3 immune stain or both according to its distribution i.e. if it is global (the entire membrane of the CD3+ lymphocytes are of surface CD3 immune stain ) or clonal distribution (both partial membranous and intracytoplasmic CD3 immune stain ) 3.
Statistical analysis

This descriptive study for the collected data was analyzed using Statistical Package for Society Study (SPSS) statistical software version 19.0. Qualitative data were evaluated using the frequency and related percentage by using Chi-square test and a P value equal or less than 0.05 was considered statistically significant with confidence interval of 95%.

RESULTS

Duodenal biopsies have been taken from 100 cases suspected to have CeD where their ages ranges between (1-69) years with a mean age of 20.74 years and median (17) years. Children represented 41 (41%) of the sampled cases (27 female and 14 male) with a female to male ratio 1.9:1, while adult 59 (59%) (42 female and 17 male) with a female to male ratio 2.4:1. Among the 100 cases, there were 69 (69 %) females and 31 (31%) males, female : male ratio is (2.2:1).

By applying modified Marsh’s classification on the mucosal changes by H&E stain on 100 (100%) cases: Marsh 0 was detected in 8 (8 %) cases, Marsh 1 in 30 (30 %) cases, Marsh 2 in 10 (10%) cases, while Marsh 3 a in 20 (20%) cases, Marsh 3 b in 17 (17%) cases. Marsh 3 c in 15 (15%) cases and Marsh 4 : 0 (0%), all stages of modified Marsh classification were shown in figure (1).

Lamina propria in cases with features of CeD shows mild-moderate or heavy infiltration of chronic inflammatory cells (lymphocytes and plasma cells), and some of them associated with neutrophils and eosinophils.

The sex and age distribution of cases suspected CeD according to modified Marsh classification were shown in tables (1 and 2).

Immunohistochemical expression of CD3 in the sampled cases i.e. CD3 + ≥30 IEL/100 epithelial cells were detected in 79 (79 %) cases. Seventeen out of 30 cases (56.7 %) that diagnosed Marsh 1 were CD3 + while 2 out of 30 cases were border line. The last 11 out of 30 cases were negative CD3 immune stain, as shown in table (3).

All the CD3 + cases that have been detected in 79 (79 %) cases are of cresendo pattern of distribution from the base to the tips of the villi while the border line and negative immune stain cases are of decrescendo pattern of distribution from the base to the tips of the villi, as shown in table (4).

The diffuse distribution of CD3+ intraepithelial lymphocytes among the villi was noted in 78 out of 79 cases of CD3 + cases, while only one case was observed with patchy distribution and associated with Helicobacter pylori infection, all stages of modified Marsh classification with CD3 IHC stain were shown in figure (2).

The distribution of staining of CD3 + intraepithelial lymphocytes was global surface (sCD3+) in 98 (98%) cases and clonal surface (sCD3+) & intracytoplasmic (iCD3 +) in < 20% of IEL in 2 (%) cases, that need flowcytometric analysis to confirm the diagnosis of Refractory Celiac disease, was shown in table (5) and figure (3).

Age and sex distribution of CD3 + intraepithelial lymphocytes in cases suspected CeD were shown in tables (6 and 7).

Table 1: The sex distribution in cases suspected CeD according to modified Marsh classification (N=100) (P Value 0.005).

<table>
<thead>
<tr>
<th>Stage</th>
<th>Male</th>
<th>Female</th>
<th>Total</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>3</td>
<td>5</td>
<td>8</td>
<td>8 %</td>
</tr>
<tr>
<td>1</td>
<td>8</td>
<td>22</td>
<td>30</td>
<td>30 %</td>
</tr>
<tr>
<td>2</td>
<td>8</td>
<td>2</td>
<td>10</td>
<td>10 %</td>
</tr>
<tr>
<td>3a</td>
<td>7</td>
<td>13</td>
<td>20</td>
<td>20 %</td>
</tr>
<tr>
<td>3b</td>
<td>4</td>
<td>13</td>
<td>17</td>
<td>17 %</td>
</tr>
<tr>
<td>3c</td>
<td>1</td>
<td>14</td>
<td>15</td>
<td>15 %</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0 %</td>
</tr>
</tbody>
</table>

Table 2: The age distribution in cases suspected CeD according to modified Marsh classification (N=100) (P Value 0.081).

<table>
<thead>
<tr>
<th>Stage</th>
<th>Children</th>
<th>Adult</th>
<th>Total</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>5</td>
<td>3</td>
<td>8</td>
<td>8 %</td>
</tr>
<tr>
<td>1</td>
<td>11</td>
<td>19</td>
<td>30</td>
<td>30 %</td>
</tr>
<tr>
<td>2</td>
<td>7</td>
<td>3</td>
<td>10</td>
<td>10 %</td>
</tr>
<tr>
<td>3a</td>
<td>10</td>
<td>10</td>
<td>20</td>
<td>20 %</td>
</tr>
<tr>
<td>3b</td>
<td>5</td>
<td>12</td>
<td>17</td>
<td>17 %</td>
</tr>
<tr>
<td>3c</td>
<td>3</td>
<td>12</td>
<td>15</td>
<td>15 %</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0 %</td>
</tr>
</tbody>
</table>

Table 3: The count of CD3 + intraepithelial lymphocytes per 100 epithelial cells according to modified Marsh classification (N=100) (P Value < 0.001).

<table>
<thead>
<tr>
<th>Stage</th>
<th>≥30/100</th>
<th>26-29/100</th>
<th>≤25/100</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>1</td>
<td>17</td>
<td>2</td>
<td>11</td>
<td>30</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>0</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>3a</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>3b</td>
<td>17</td>
<td>0</td>
<td>0</td>
<td>17</td>
</tr>
<tr>
<td>3c</td>
<td>15</td>
<td>0</td>
<td>0</td>
<td>15</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>79</td>
<td>2</td>
<td>19</td>
<td>100</td>
</tr>
</tbody>
</table>
The Significance of CD3 Marker

Table 4: The distribution of CD3 + intraepithelial lymphocytes per 100 epithelial cells crescendo vs. decrescendo from the base to the tips of the villi (N=100).

<table>
<thead>
<tr>
<th>Distribution</th>
<th>≥30/100</th>
<th>26-29/100</th>
<th>≤25/100</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decrescendo</td>
<td>0</td>
<td>2</td>
<td>19</td>
<td>21</td>
</tr>
<tr>
<td>Crescendo</td>
<td>79</td>
<td>0</td>
<td>0</td>
<td>79</td>
</tr>
<tr>
<td>Total</td>
<td>79</td>
<td>2</td>
<td>19</td>
<td>100</td>
</tr>
</tbody>
</table>

Table 5: The distribution of staining of CD3 + intraepithelial lymphocytes according to modified Marsh classification (N=100).

<table>
<thead>
<tr>
<th>Stage</th>
<th>Global sCD3 +</th>
<th>Clonal sCD3 + &amp; iCD3 + &lt;20%</th>
<th>Clonal sCD3 + &amp; iCD3 + &gt;20%</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>8</td>
<td>0</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>1</td>
<td>30</td>
<td>0</td>
<td>0</td>
<td>30</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>0</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>3a</td>
<td>20</td>
<td>0</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>3b</td>
<td>17</td>
<td>0</td>
<td>0</td>
<td>17</td>
</tr>
<tr>
<td>3c</td>
<td>13</td>
<td>2</td>
<td>0</td>
<td>15</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>98</td>
<td>2</td>
<td>0</td>
<td>100</td>
</tr>
</tbody>
</table>

Table 6: The age distribution of CD3 + intraepithelial lymphocytes in cases suspected CeD (N=100).

<table>
<thead>
<tr>
<th>Age</th>
<th>CD3 + ≥30/100</th>
<th>CD3 + 26-29/100</th>
<th>CD3 + ≤25/100</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Children &lt;15 years</td>
<td>33</td>
<td>0</td>
<td>8</td>
<td>41</td>
</tr>
<tr>
<td>Adult ≥15 years</td>
<td>46</td>
<td>2</td>
<td>11</td>
<td>59</td>
</tr>
<tr>
<td>Total</td>
<td>79</td>
<td>2</td>
<td>19</td>
<td>100</td>
</tr>
</tbody>
</table>

Table 7: The sex distribution of CD3 + intraepithelial lymphocytes in cases suspected CeD (N=100).

<table>
<thead>
<tr>
<th>Sex</th>
<th>CD3 + ≥30/100</th>
<th>CD3 + 26-29/100</th>
<th>CD3 + ≤25/100</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>27</td>
<td>0</td>
<td>4</td>
<td>31</td>
</tr>
<tr>
<td>Female</td>
<td>52</td>
<td>2</td>
<td>15</td>
<td>69</td>
</tr>
<tr>
<td>Total</td>
<td>79</td>
<td>2</td>
<td>19</td>
<td>100</td>
</tr>
</tbody>
</table>

Figure 1: Duodenal biopsies stained by H&E stain X 40 show all stages of modified Marsh classification.

Figure 2: Duodenal biopsies CD3 IHC stain X40 in relation to modified Marsh classification.
DISCUSSION

Until now the duodenal biopsy is still necessary and remains the gold standard for the diagnosis and follow up of patients with CeD, which is usually supplemented by serologic tests. Accordingly, apart from grading villous atrophy and crypt hyperplasia, the assessment of IELs is essential and regarded as a key for the correct diagnosis of CeD in early stages ( Marsh 1) because sometimes the contrast between the blue color of lymphocyte nucleus and epithelial cell nucleus is inconclusive by H&E stain.

Moreover, the use of sophisticated method such as IHC to differentiate intraepithelial lymphocytes from epithelial cells is essential. As IELs are CD3 positive cells, performing IHC staining against CD3 can aid in estimating the exact number and the distribution of IELs more precisely than using conventional histological stain (H&E).

Age and Gender

In the current study among 100 cases suspected to have CeD and underwent endoscopic biopsy. Children < 15 years old represented 41% of the sampled cases, while adults ≥ 15 years old were 59%, which reflects that adults are affected by CeD more than children. Previously, CeD was considered a disease of childhood because the majority of the cases were less than 2 years of age. However, the disease is common in adults and can be diagnosed at any age. Radiovic, 2013 in Serbia mentioned that the classical form of the disease is mainly seen in infants and children and atypical in later ages and in adults. Some studies were found that its frequency is twice in children than in adults while others showed adult prevalence become more frequent, sometimes reaching similar. This attributed to various environmental factors in addition to age of introduction of gluten and the amount of it in the food, which have been implicated in the pathogenesis of CeD. In the present study the discrepancy in the age of the sampled cases may due to the recording center that focused on an adult more than pediatric population during sample collection.

Gender distribution is variable in different parts of the world, some geographical areas show equal division in both sexes, while others are more to one of them. In the present study, there were 69% females and 31% males, with female to male ratio is (2.2:1). This finding was observed in many studies. While other study done by Ifitkhar et al., 2016 in Pakistan show that 84% of the cases were males and 16% of the cases were females.

The differential rates of diagnosis among gender is thought to reflect several factors, such as hormonal factor, a higher rate of autoimmune disease among women in general, more regular health care interaction in female than male subjects and a higher likelihood of symptomatic disease among women than men.

Histological analysis of duodenal biopsy

Modified Marsh classification was valuable for diagnostic purposes and for follow up of patients with CeD by assessing the histologic recovery in response to gluten withdrawal. This can be accomplished by reviewing the previous histological report and comparing it with the present report, this will give an idea about the progress of CeD and extent of response to gluten challenge.

In this study and according to modified Marsh classification which is used by many studies on the mucosal changes by H&E stain on 100 cases, Marsh 0 was detected in 8% of the cases and considered as chronic nonspecific duodenitis, while another study done in Turkey by Sezgin et al., 2016 showed that half of the cases were Marsh 0, on the other hand, Paul et al., 2019 was detected Marsh 0 in 70% of the cases. While Shihab & Enaya, 2020 in their study in south of Iraq, Marsh 0 was not detected, this difference depends on the awareness of the population to the disease and the availability of the diagnostic facilities.

Marsh 1 was observed in 30% of the sampled cases. This finding is within the range of that observed by Brown et al., 2006 and Lauwers et al., 2015 studies in which Marsh 1 were noted in...
The exact prevalence of Marsh 1 is difficult to determine because there are many causes of increased IELs with normal villous architecture. So, an increase in IELs infiltration alone in duodenal biopsies (Marsh 1) is not specific for CeD and other causes should be excluded like Helicobacter pylori, Giardia lamblia, cryptosporidium and others. Thus, negative serology does not exclude a diagnosis of CeD with a Marsh 1 pattern, a morphologic evaluation remains essential.

In the present study, crypt hyperplasia observed in 62% (Marsh 2 and Marsh 3) while in Iftikhar et al., 2016 study noted in 78%. Villous atrophy (Marsh 3) was observed in 52% of the current study, while in Iftikhar et al., 2016 study was 74% of the cases.

In Turkey Sezgin et al., 2016 found that half of the cases in their study was Marsh 2 and 3, while Paul found in 11% of the cases Marsh 2 and 3. These differences may be due to the fact that in Iraq, the patients generally do not seek medical advice early and this may be due to poverty, illiteracy and parents negligence and therefore, there is a delay of patients’ attendance to clinical observation. Moreover, patients with CeD have a long duration of symptoms and may see many physicians before diagnosis by duodenal biopsy. As a result, there is a high rate of dissatisfaction about this diagnostic delay that is physician based. Failure of physician to recognize the diverse clinical presentation of CeD, less awareness of disease among clinicians as well as for quality of diagnostic tools and their availability contributes to the diagnostic delay. Also CeD regarded as difficult to be diagnosed because of the alternative diagnosis (often irritable bowel syndrome), in addition to the diversity of symptoms and even sometimes associated with a period of latency. Some physicians are unaware of the condition and there are several “myths,” such as CeD is rare, occurs in Caucasians, it is mostly in Europe and the United States, it is disease of childhood, it occurs only with chronic diarrhea and it can be cured after a period of treatment.

Marsh 4 was not detected in the present study. This finding is in agreement with that observed by Dickson and his colleagues, 2006 who consider Marsh stage 4 is a rare stage of the disease.

### CD3 immunohistochemical stain

The use of CD3 IHC expression in the sampled cases of the present study represents a sensitive and specific tool to distinguish IELs from epithelial cells especially in Marsh 1 cases because the occurrence of IELs by itself is not specific for CeD and can be observed in forms of intestinal inflammations or other causes. Therefore, the presence of IELs should be seriously taken into consideration.

In the current study, CD3 positive cases i.e. CD3 + ≥30 IEL /100 epithelial cells were detected in 79% of the cases and this result within the results of other studies, as Shihab & Enaya, 2020 in the South of Iraq were 100%, Iftikhar et al., 2016 in Pakistan were 100% and Mubarak et al., 2015 in Netherland were 68.55%.

This variation in the percentage of CD3 positive cases of these studies may be due to variation in the IHC staining technical operation with different manufacturers, variation of kits and types of antibodies which were used, genetic and environmental factors, in addition to the variation of patient’s age group.

Mubarak et al., 2015 study showed that from 40.88 % of the cases with Marsh 0 by H&E, only 10.69 % of the cases show positive results by CD3 immune stain, and out of 3.77 % of the cases diagnosed as Marsh 1 by H&E, 0.62 % of the cases was negative CD3 immune stain and 55.34 % of the cases Marsh 2 & 3 only 0.62 % was negative CD3 immune stain.

Iftikhar et al., 2016 study on histomorphological and immuno-histochemical analysis of small intestinal biopsies in adults suspected of CeD, as they count IELs both by H&E stain and immune stain CD3 and CD20; found that CD3 immune marker was positive for IELs in all cases establishing the fact that IELs were of T-cell origin, while CD20 immune marker (B-lymphocytes) have shown the focal positivity in areas with lymphoid follicle formation with germinal center.

Shihab & Enaya, 2020 study on 60 cases, CeD was confirmed by both the routine histopathological examination and immunohistochemical analysis in all cases.

All of CD3 positive cases in the present study were crescendo pattern of distribution from the base to the tip of the villi. The same result was obtained by Iftikhar et al., 2016 in Pakistan and by Shihab & Enaya 2020 in the south of Iraq.

In this study 30 % out of 100 cases diagnosed as Marsh 1 by H&E, 17 out of 30 cases (56.7%) were CD3+ positive and they were in crescendo pattern while 11 cases CD3+ less than or equal to 25/100 epithelial cells and of decrescendo pattern. The last two cases were border line and...
The Significance of CD3 Marker

of decrescendo pattern i.e. regarded negative for CD3 immune stain as the distribution pattern of IELs within the epithelium is more valuable than the actual counts \[8,10,11,15\]. So 13 out of 30 cases of Marsh 1 regarded negative for CeD after CD3 immune stain. While Goldstein N.S.& Underhill J.,2001 \[24\] in their study on cases with potential CeD noticed that 9 out of 12 cases with CeD were of decrescendo pattern of distribution from the base to the tip of the villi. On the other hand, in Iftikhar et al.,2016 \[12\] study showed all the CD3 positive cases were in crescendo pattern also.

In the present study, the CD3 + IELs in duodenal biopsy was distributed diffusely and in crescendo pattern from the base of the villi to the tip in all cases except one case that is of patchy distribution and associated with H.pylori. This finding was also observed by Shihab’s study \[19\].

The IHC expression of CD3 marker plays an important role in the diagnosis of early stage of CeD (Marsh 1) which is if misdiagnosed as chronic inflammatory changes by H&E and not treated as CeD, it can therefore develop into overt CeD with the passage of time \[8,10,11\]. So, this study showed that in compared to H&E stains alone, CD3 immune stains could lead to a definite assessment in 13% of cases with Marsh 1. This observation was also noted by a study done by Mubarak and colleagues who showed that compared to H&E stain, CD3 immune stains could lead to definite assessments in 12.6% of cases \[6\].

The IHC expression of CD3+ marker provides a hint about the distribution of CD3+ marker within the lymphocyte whether global surface or clonal surface and intracytoplasmic to diagnose Refractory CeD and differentiate between its two types because type II may precede EATL \[1,4\].

In the current study, it is difficult to differentiate surface CD3 from intracytoplasmic CD3 exactly by CD3 immune stain only. Flowcytometric analysis is better for this situation \[40\], as it is regarded the gold standard for the detection of aberrant phenotype of IELs \[13,14,41\].

**CONCLUSION**

There is a significant relationship between the count of CD3+ T-lymphocytes per 100 epithelial cells and the histopathological changes in the duodenal biopsy according to modified Marsh classification. So, the immunohistochemical expression of CD3 in intraepithelial T-lymphocyte could lead to a definite assessment in 43.3% of the sampled cases with Marsh type 1. All the positive cases are of crescendo pattern of distribution of CD3+ T-lymphocytes as the distribution is more important than the actual count and they distributed diffusely except that associated with H. pylori infection observed with patchy distribution. In addition to that, The IHC expression of CD3+ marker provides a hint about the distribution of CD3+ marker within the lymphocyte whether global surface or clonal surface and intracytoplasmic to diagnose Refractory Celiac disease.

On the other hand, females were more affected than males with CeD and there is a significant relationship between the gender of the sampled cases and the histopathological changes in the duodenal biopsy.

The disease can be diagnosed at any age and there is no significant relationship between the age distribution of the sampled cases and the histopathological changes in the duodenal biopsy.

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