ORIGINAL ARTICLE

Gastric Histopathology of Chronic Gastritis in Obese Patients Undergoing Laparoscopic Sleeve Gastrectomy: A Local Experience

Murad A. Alturkustani, FRCPC, MBBS
Department of Pathology, Faculty of Medicine
King Abdulaziz University, Jeddah, Saudi Arabia

Correspondence
Dr. Murad A. Alturkustani
P.O. Box 80205, Jeddah 21589, Saudi Arabia
E-mail: maltarkustani@kau.edu.sa

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Abstract
A retrospective review was conducted using the database at King Abdulaziz University Hospital, to determine the most common cause of chronic gastritis among obese individuals. One hundred and thirty-one specimens were examined, obtained from patients who had laparoscopic sleeve gastrectomy and diagnosed with chronic gastritis. Helicobacter pylori was detected in 15 cases only. The most common diagnosis was “Chronic inactive gastritis without Helicobacter pylori infection” in 57 (43.50%) cases, then “Chronic gastritis with lymphoid follicular hyperplasia” in 54 (41.20%) cases, and lastly “Chronic active gastritis” in the remaining 20 (15.20%) Cases. The low incidence of Helicobacter pylori infection even in cases of gastritis with lymphoid follicular hyperplasia support the existence of the new proposed entity (i.e., obesity-related gastritis). Biopsy specimens for cases diagnosed as chronic gastritis were used as control group to confirm the appropriate rate of detecting Helicobacter pylori organisms by morphological examination in the same laboratory. In these biopsies, Helicobacter pylori was present in (50.70%), and reactive follicular hyperplasia was detected in (6%) of total controls.

Keywords
Gastritis; Obesity; Laparoscopic Sleeve Gastrectomy; Helicobacter pylori

Introduction
The recent increase of bariatric surgeries offered an excellent source for obtaining gastric specimens of obese patients. Histological examination of these specimens resulted in many incidental findings, the most common was features of chronic gastritis[1,2]. However, the most common cause of this gastritis (i.e., no specific etiology) was different from the most common cause of chronic gastritis in gastric biopsies (i.e., Helicobacter pylori (H. pylori) infections)[3]. The latter observation proposed a new form of chronic gastritis, the obesity-related gastritis[4].

Furthermore, chronic gastritis with follicular hyperplasia is consistently associated with H. pylori infection. The organisms are often detected on biopsy examination of patients’ specimens. Accordingly, the presence of follicular hyperplasia was thought to be exclusive to H. pylori infection[5]. In addition, these reactive lymphoid follicles were also considered as the
precursor for gastric Mucosa Associated with Lymphoid Tissue (MALT) lymphoma. However, this association was not this strong in gastric specimens obtained from bariatric surgeries.

This study aims to determine the most common cause of chronic gastritis among obese patients who had laparoscopic sleeve gastrectomy (LSG), attending a tertiary care hospital in western Saudi Arabia, and establish the association between chronic gastritis with follicular hyperplasia, and H. pylori infection in the specimens of obese individuals.

Methods

Approval from the Research Ethics Board of King Abdulaziz University was obtained prior to the study. A retrospective review was conducted using the electronic database of the pathology department at King Abdulaziz University Hospital. As the study focused on obese patients, data was collected from all specimens of LSGs, performed in the period between 2011 and 2015. Histopathological reports upon examining gastric tissue specimens of these patients were reviewed. Within the pathological report, attention was given for gross examination, microscopic findings, performance of special stains to detect H. pylori and the final diagnosis. This was followed by targeted selection of cases to re-examine the specimens, and confirm the diagnosis, after which we clarified any obscure points in the pathological report. I included only cases with the pathological diagnosis of chronic gastritis, with or without lymphoid hyperplasia and excluded cases with specific etiology of the chronic gastritis other than H. pylori infection (e.g., autoimmune gastritis). Another cohort of the 150 routine gastric biopsies, diagnosed in 2015 as chronic gastritis, was included in the study as controls only to confirm the appropriate rate of detecting H. pylori organisms by morphological examination in the same laboratory. The diagnosis of controls was confirmed using the same laboratory testing of the cases, and we also estimated the frequency of H. pylori infection and follicular gastritis in the control specimens.

The following definitions were used in this study for different types of gastritis. Chronic gastritis; “Gastric mucosa infiltrated by a significant number of lymphocytes and plasma cells in the lamina propria (i.e., more than only scattered cells)”. This can be classified into A. Chronic inactive gastritis: when the chronic inflammation is not associated with acute inflammatory cells in the gastric pits and foveolae; B. Chronic active gastritis: where morphological features of chronic gastritis are detected, in addition to the presence of acute inflammatory cells (i.e., neutrophils) in the gastric pits and foveolae; and C. Chronic gastritis with follicular hyperplasia (also known as follicular gastritis): when this inflammation is accompanied by the presence of lymphoid follicles with germinal centers. Helicobacter pylori detection in this study was mainly performed using hematoxylin and eosin (H&E) special stain, and by cresyl fast violet special stain if the organisms were suspected but not primarily detected by H&E stain.

Results

During this period 131 specimens, obtained from LSGs met the inclusion criteria, and under the broad diagnosis of chronic gastritis. These included 98 females and 33 males. The age ranged between 18 - 66, with a mean age of (34.4) year. Helicobacter pylori was detected in 15 cases only. The most common histopathological diagnosis reported was “Chronic inactive gastritis without H. pylori infection” in 57 (43.5%) cases. (Fig. 1A, B) The second most common diagnosis was “Chronic gastritis with lymphoid follicular hyperplasia” (Fig. 1C, D) assigned to 54 (41.2%) cases. For the latter cases, H. pylori organisms were present in only four out of the total 54 cases. “Chronic active gastritis” was the diagnosis reported for the remaining 20 (15.2%) cases, and H. pylori organisms was present in 11 out of these 20 cases. For the control group H. pylori was present in 76 (50.7%) out of 150 cases, and reactive follicular hyperplasia was detected in only nine (6%) cases of the total controls.

In summary, the most significant findings from testing specimens from LSG were: the lower incidence of H. pylori as a cause of chronic gastritis, the high frequency of chronic gastritis with lymphoid follicular hyperplasia, and low association of the latter finding with H. pylori infection.

Discussion

Chronic gastritis refers to the chronic inflammation of gastric mucosa, whether due to infectious or a non-infectious etiology. Among these, H. pylori infection is considered the most common cause. Histopathological examination of the gastric mucosa provides important clues about possible causes, and can present conclusive features such as: corpus-restricted inflammation with atrophy in autoimmune
gastritis, prominent intraepithelial lymphocytic infiltrate without significant neutrophils in lymphocytic gastritis, granulomas in different types of granulomatous gastritis, and reactive mucosal changes that lack significant inflammation in different forms of gastropathy. Despite the above, a small proportion of chronic gastritis remains with no identifiable etiology, even when obtaining detailed clinical information and performing meticulous histological examination\[7\]. This proportion is more prominent in gastric specimens from morbidly obese patients who underwent bariatric surgery.

Chronic gastritis with follicular hyperplasia is defined as: “The presence of reactive lymphoid follicles (i.e., follicles with germinal center), which is easily recognized in a biopsy screening, and considered a strong indicator of \textit{H. pylori} presence. The detection of this organism requires careful and high-power examination of the slides. The frequency of detection of \textit{H. pylori} in cases of chronic gastritis with follicular hyperplasia ranges from very high, to almost all cases after thorough slides’ examination\[8\]. In fact, presence of follicular hyperplasia was thought to be exclusive to \textit{H. pylori} infection\[3\]. Lymphoid follicles are associated with specific response to chronic antigenic stimulation elicited by the organisms, and thus support the specificity of this type of reaction to \textit{H. pylori} infection\[8\]. Furthermore, lymphoid follicle formation is also proved to be the precursor for MALT lymphoma, and thus \textit{H. pylori} is strongly associated with this type of lymphoma\[6\].

\textbf{Figure 1.} Histopathology of chronic gastritis. (A, B) Chronic inactive gastritis with lymphoplasmacytic infiltrate in the lamina propria with no acute inflammatory cells (H&E; original magnification A: 200x, B: 400x). (C, D) Chronic gastritis with lymphoid follicular hyperplasia showing multiple lymphoid follicles with reactive germinal centers and chronic inflammatory cells in the lamina propria (H&E; original magnification A: 40x, B: 100x).
In LSG specimens, chronic gastritis with follicular hyperplasia has a weak association with *H. pylori* infection, as found in the current study, with only 4 (7.4%) of 54 cases demonstrating identifiable organisms. This finding is consistent with the lower rates of *H. pylori* infection in this type of specimens collected from different populations, and as previously demonstrated in two more studies. The first study found only 26 cases of *H. pylori*, in a total of 95 (27%) chronic gastritis cases with follicular hyperplasia[2], while the second study reported only 13 cases of *H. pylori* in 78 (16.6%) cases of chronic gastritis with follicular hyperplasia[1][9]. On the other hand, a more recent study found a high percentage of *H. pylori* infection (84%), among individuals diagnosed with follicular gastritis, and who underwent LSG[2]. *Helicobacter pylori* was a very common finding in this study (40.9%), despite treating all patients with a positive Campylobacter-like organism test, with the standard triple therapy prior to the surgery, which is expected to lower the prevalence of *H. pylori* in these specimens. Furthermore, the frequency of *H. pylori* was also high, and namely (74%), among specimens with lymphoid aggregates only[2]. This may explain the high percentage of *H. pylori* infections in their cohort, as opposed to ours and the other two studies.

Chronic gastritis with follicular hyperplasia is strongly associated with lymphoid tissue lymphoma (MALT) of the gastric mucosa, and chronic gastritis with follicular hyperplasia is considered exclusive to the *H. pylori* infection[2]. The fact that chronic gastritis with follicular hyperplasia occurs in obese individuals, and in the absence of *H. pylori* infection, raises the possibility to consider morbid obesity as a risk factor for MALT lymphoma. Supporting this possibility is a recently published case report documenting the presence of gastric MALT lymphoma in morbidly obese patients with no *H. pylori* infection, and after complete remission and successful weight reduction by gastric bypass and resection[1][10]. Further research is needed to assess this association, and draw better conclusions.

Although *H. pylori* is reported as the most common cause of gastritis[3], it is not thoroughly documented among chronic gastritis patients. Of the limited instances where it is examined, *H. pylori* infections were of a high percentage among patients with chronic gastritis. Furthermore, it was shown to be the prime cause of chronic gastritis in 292 (65.9%) patients in one study[11], and in 78% of gastritis patients in another study[12]. This comes in agreement with our study, where the control cohort reported a prevalence of 50.7% of *H. pylori* in the biopsies of chronic gastritis. This is in contrast with the LSG specimens, where *H. pylori* was only present in 15 out of 131 cases, i.e., (11.5%). Congruent to this are results from a recent review by Verma et al.[13]. They found that the frequency of *H. pylori*, although generally low in these specimens, had a variable prevalence across subgroups ranging from 11.5% - 66.7%. Other studies which came after this review again confirmed this lower rate (Table 1), except for the study by Safaan et al.[2] as discussed above. The latter observation along with higher incidence of chronic gastritis in specimens from bariatric surgeries, led to a proposal by Yamamoto et al.[14] for a new entity of gastritis, i.e., the obesity-related gastritis. The latter could explain the discrepancy between *H. pylori* incidence between all biopsies and those from patients who underwent LSGs.

Obesity is considered a state of chronic low-grade inflammation[15][16]. Recent studies linked the inflammatory state of obesity, to the many co-morbidities associated with it. This includes hypertension, diabetes and several neoplasms[15]. Another observation confirming this inflammatory characteristic of obesity, is the fact that weight loss is often associated with reductions of

<table>
<thead>
<tr>
<th>Study</th>
<th>Total No. of Cases</th>
<th>Chronic Inactive Gastritis</th>
<th>Chronic Active Gastritis</th>
<th>Follicular Lymphoid Hyperplasia</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Total</td>
<td><em>H. pylori</em></td>
<td>Total</td>
</tr>
<tr>
<td>Present study</td>
<td>131</td>
<td>57</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>Safaan et al.[2]</td>
<td>1555</td>
<td>512</td>
<td>NA</td>
<td>105</td>
</tr>
<tr>
<td>Almazeedi et al.[2]</td>
<td>656</td>
<td>488</td>
<td>6</td>
<td>49</td>
</tr>
<tr>
<td>Raes et al.[3]</td>
<td>248</td>
<td>30</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Vrohie et al.[3]</td>
<td>87</td>
<td>NA</td>
<td>NA</td>
<td>20</td>
</tr>
</tbody>
</table>

Abbrev.: *H. pylori* = Helicobacter pylori; NA: Not Available; *Only studies that specified the number of follicular lymphoid hyperplasia under chronic gastritis were included in this table.

*Active & inactive*
the above comorbidities\cite{13}. This type of inflammation is not linked to infections or autoimmune stimulus. This inflammatory status was attributed to several elevated inflammatory mediators such as; Interleukin six (IL-6) and Tumor Necrosis Factor α (TNFα). Accordingly, it is likely that these chronic inflammatory cells which are flared up in obesity, can explain the higher incidence of chronic gastritis in bariatric surgery specimens, namely those not likened to \textit{H. pylori} infections.

A major limitation to acknowledge is related to the detection of \textit{H. pylori} organisms in the sampled specimens. The strong association of \textit{H. pylori} and chronic gastritis with follicular hyperplasia was demonstrated, and this emphasizes the thoroughness needed when examining specimens. However, the large size of specimens derived from LSG may have affected the findings due to the following reasons: (1) The large surface area of the LSG specimens available for examination. This may give a better chance to identify follicular hyperplasia, especially when it is easily detectable by low power examination of the slides. (2) Microscopically, these specimens are not optimal for detection of \textit{H. pylori}, and the larger the surface area is, the more difficult to examine it thoroughly by high magnification. (3) The nature of these specimens such as the antrum, where it is the most likely area to harbor \textit{H. pylori} organisms, were not included in these specimens. Despite the latter, the histopathological detection of \textit{H. pylori} organisms in these specimens was not likely affected by these factors, and this was confirmed when detection by other sensitive means such as serology, also revealed similar \textit{H pylori} rates among these patients. The latter was emphasized by Verma \textit{et al.}\cite{15} in a review of previous studies focusing on \textit{H. pylori} infections in bariatric patients. Another limitation was the lack of data on status of patients who were diagnosed and treated for \textit{H. pylori} prior to the LSG. However, none of the included patients had a gastric biopsy in our institution before the surgery for this purpose, and the high percentage of chronic gastritis in LSG specimens favor that they did not receive a treatment for \textit{H. pylori} prior to the procedure.

In conclusion, \textit{H. pylori} infection was found to be a much less common cause of chronic gastritis among patients who underwent LSG, and most of the chronic gastritis cases were of no identifiable cause, and thus supporting the proposal of an obesity-related gastritis. Chronic gastritis with follicle formation has only weak association with \textit{H. pylori} infection in LSG specimens, proposing that this type of inflammation is part of a morphological spectrum in the obesity-related gastritis, and that chronic gastritis is not restricted to \textit{H. pylori} infection alone. The latter may also mark obesity as a possible risk factor for MALT lymphoma. However, more researches in this field is required to prove or deny this possibility.

\section*{Conflict of Interest}
The author has no conflict of interest.

\section*{Disclosure}
The author did not receive any type of commercial support either in forms of compensation or financial for this study. The author has no financial interest in any of the products or devices, or drugs mentioned in this article.

\section*{Ethical Approval}
Obtained.

\section*{References}


التهاب المعدة المزمن مع التضخم الجريبي في عينات المرضى الذين يعانون من السمنة المفرطة والناجحة عن استئصال جزء من المعدة بالمنظار

مراد عبدالكريم التركستاني
قسم علم الأمراض، كلية الطب، جامعة الملك عبدالعزيز
جدة - المملكة العربية السعودية

المستخلص. تم إجراء مراجعة بتأثر رجعي عينات المرضى الذين يعانون من السمنة المفرطة والناجحة عن استئصال جزء من المعدة بالمنظار باستخدام قاعدة البيانات في مستشفى جامعة الملك عبد العزيز لتحديد السبب الأكثر شيوعاً لإتهام المعدة المزمن. تم فحص مائة وثلاثين عينة وتم العثور على جرثوم المعدة المزمن في (15) حالة فقط. كان التشخيص الأكثر شيوعاً هو إتهام المعدة غير النشط المزمن من غير جرثوم المعدة (43.5٪) حالة. ثم إتهام المعدة المزمن مع تضخم الجربير اللا filmy (54٪) حالة. وأخيراً إتهام المعدة المزمن النشط في (20٪) حالة. (52٪) العدد المنخفض لجرثوم المعدة مع حالات إتهام المعدة المزمن مع تضخم اللمفاوي الجريبي تدعم وجود النوع الجديد المقترح لإتهام المعدة المزمن (التهاب المعدة ذات الصلة بالسمنة). تم استخدام عينات خزعة المعدة للحالات التي تم تشخيصها على أنها التهاب المعدة المزمن كمجموعة مراقبة لتأكيد المعدل المناسب للكشف عن جرثوم المعدة عن طريق الفحص التنسيجي في نفس المعمل وكانت جرثوم المعدة موجودة في (7.5٪)، وتم الكشف عن تضخم جريبي في (2٪) من هذه المجموعة.