Acid suppression and gastric atrophy: sifting fact from fiction

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Summary
Prolonged pharmacological acid suppression is associated with various histological changes in the gastric mucosa, particularly in Helicobacter pylori infected patients. In a number of subjects these changes include a shift in the gastric inflammation from the antrum to the corpus. This finding has been interpreted as gastric atrophy, and the possibility that acid suppression accelerates the progress of lesions that may lead to gastric cancer has been considered. Two recent studies on the relation between treatment with proton pump inhibitors and atrophic gastritis have yielded apparently contradictory results. These studies are reviewed in detail here and some of the possible reasons for the discrepant conclusions are explored. In particular, the way the terms “gastric atrophy” and “atrophic gastritis” are used is examined critically.

What is gastric atrophy?
As seen in histological preparations, the mucosa of the normal gastric corpus consists of a layer of mucous cells invaginating into shallow pits, which merge with the oxyntic glands. The latter are separated by little or no extracellular matrix in which few mononuclear cells may be interspersed (fig 1A). When the mucosa is infiltrated by inflammatory cells (mostly plasma cells and lymphocytes, in the case of H pylori infection) the glands become separated and pushed aside, and may become invisible, or “lost” (fig 1B). In some circumstances glands may be destroyed and substituted with either intestinal metaplastic epithelium or fibroblasts and extracellular matrix, or a combination of the two. In such cases the glands are invisible for the obvious reason that they are not there—that is, they are truly “lost”.

Most definitions of gastric atrophy, including Whitehead’s, Correa’s; the original Sydney System, and the updated Sydney System have emphasised gland loss without specifying what replaces the missing glands. Therefore, mucosa showing the histopathological features depicted in figure 1B would be considered atrophic by many observers, but not by those who would want to know whether the missing glands are really lost or just displaced by the inflammation. Functional studies do not help clarify the issue: inflammation in the oxyntic mucosa interferes with the somatostatin–gastrin axis and depresses acid production.

Of course, the destruction of oxyntic glands results in hypochlorhydria. Consequently, these two types of gland loss (one apparent and one real, but both with similar pathophysiological consequences) have not been considered as separate entities in studies that have looked at the relation between structure and function of the gastric mucosa. In the absence of stringent histopathological criteria, it is hardly surprising that pathologists have generally failed to agree on whether the missing glands are really lost or just displaced by inflammation.7

The importance of being atrophic
Soon after H pylori was recognised as the cause of chronic gastritis, it became apparent that the epidemiological paradigms that linked atrophic gastritis and gastric cancer could be applied to H pylori infection.8–10 Populations with a high prevalence of H pylori infection and atrophic gastritis (such as Colombia11 and Japan12) have a high incidence of gastric cancer; in contrast, populations with low prevalence of H pylori13 or low prevalence of atrophic gastritis13,14 have low a incidence of gastric cancer. Furthermore, several studies have shown quite convincingly that long-standing H pylori infection results in atrophy and intestinal metaplasia in a portion of the infected subjects.15

The association between atrophic gastritis and gastric cancer detected in pre-helicobacter studies could just as well be applied to H pylori infection: no longer gastritis, but its principal cause—H pylori—could be named as the carcinogen, and both epidemiological and biological evidence became available to support this role.16 The former consist of a large body of data showing that infection with H pylori increases the cancer risk between three- and 10-fold, depending on the population studied.17,18
latter include a broad variety of observations linking particular histopathological features of the gastric mucosa with the development of intestinal-type adenocarcinoma. These histopathological features include alterations of the mucosal architecture, loss of glands, fibrosis, and replacement of epithelial elements by an intestinal-type metaplastic epithelium (“intestinal metaplasia”). The constellation of these features has been variously called “atrophic gastritis” and “gastric atrophy,” but these terms remain poorly defined and, consequently, loosely used. The same lack of agreement that exists on the definition of atrophy also exists about the concept of atrophic gastritis. What is atrophic gastritis? What is the atrophic gastritis considered a precursor of gastric cancer? The classic studies that defined atrophic gastritis (also called “multifocal atrophic gastritis”) were performed on gastrectomy specimens. With the entire stomach available for examination it was possible to see large patches of atrophic and metaplastic epithelium extending proximally and distally along the lesser curvature. The extension of these patches was so obvious that there appeared to be no need for a strict topographical definition. In the past two decades, however, effective antacid therapies and treatment for *H pylori* have greatly reduced the need for gastrectomy and, consequently, the availability of resected stomachs for histopathological examination. The diagnosis of atrophic gastritis must now be made based on the examination of a limited number of biopsy specimens that, even in the most audacious sampling protocols, still represent only a minimal portion of the gastric mucosa. When only two or three samples from the stomach are available for examination it may be imprudent to make a diagnostic statement which has much broader topographical implications, such as “multifocal atrophic gastritis.” Although it is very difficult to produce an accurate topographical definition of atrophic gastritis (the multifocal type associated with chronic *H pylori* infection, unknown environmental factors, and gastric cancer), the best approximation can only result from studies involving the observation of multiple mapped biopsy samples. If such a definition is generated, tested, and found to be satisfactory, it should then be disseminated and pathologists could be encouraged to use it.

**The Dutch Study**

Based on the observation that profound suppression of gastric acid is associated in some subjects with increased severity of gastritis caused by *H pylori*, the authors hypothesised that acid suppression might also increase the risk of atrophic gastritis.

To test their hypothesis, the authors studied patients from two separate cohorts who were being treated for reflux oesophagitis. One cohort consisted of 72 Swedish patients (mean age 53 years) treated with fundoplication who did not receive acid suppressive therapy after their operation. The second cohort consisted of 105 Dutch patients (mean age 62 years) treated with omeprazole (20 to 40 mg once daily). The presence of *H pylori* was assessed at the first visit by histological evaluation in the fundoplication group and by histological and serological evaluation in the omeprazole group; however, none of the patients was treated for *H pylori* infection. In both cohorts, the patients were followed for three to eight years; all patients underwent a gastroscopy at the beginning of the study as well as repeated gastroscopies during the follow up period. At each gastroscopy biopsy specimens were obtained from the gastric corpus for blinded histological evaluation. The histological examination included the semi-quantitative evaluation of the features of *H pylori* gastritis according to the Sydney System. The criteria used by the authors for the diagnosis of atrophic gastritis were not stated. However, they did state that among the patients treated with fundoplication, atrophic gastritis did not develop in any of the 31 who were infected with *H pylori* at baseline or the 41 who were not infected. Among the patients treated with omeprazole, none of whom had atrophic gastritis at baseline, atrophic gastritis developed in 18 of the 59 infected with *H pylori* and two of the 46 who were not infected. None of the patients in either group developed intestinal metaplasia.

**The Swedish study**

In this study 155 patients were randomised to the omeprazole treatment: 116 were men and 39 women; their median age was 54 years; 66 of these patients had *H pylori* infection at the initial examination; 151 patients were randomised to the antireflux surgery: 116 were men and 35 women; their median age was 51 years; 73 of these patients had *H pylori* infection.

None of the patients without *H pylori* infection in either randomisation arm developed atrophy during the follow up period. In the group on omeprazole, of the 62 *H pylori* infected patients who had no atrophy at the...
beginning of the study, five developed mild/moderate atrophy and three developed severe atrophy. In this group, two patients had mild/moderate atrophy at the beginning of the study; at the end of the study, one of them had no atrophy, whereas the other one had severe atrophy. Two subjects who had mild/moderate atrophy remained unchanged, and of the two with severe atrophy initially, one remained unchanged whereas the other became normal. Similarly to the Dutch study, no patient in either group developed any intestinal metaplasia.

The authors concluded that long term omeprazole therapy was no different from fundoplication in its effects on the development of gastric atrophy. Furthermore, they stated that, as no intestinal metaplasia developed in any of these patients and only atrophic gastritis associated with intestinal metaplasia is considered a precursor of gastric cancer, there is no evidence to support the hypothesis that treatment with proton pump inhibitors in subjects with H pylori infection may increase the risk of gastric cancer.

Reactions
The authors of the Dutch study had previously investigated the same issue and had found that long term proton pump inhibitors in patients with H pylori infection significantly decreased the inflammation and bacterial colonisation in the antrum, leading to negative antral cultures in 61% (20 of 33) patients. In contrast, the inflammation of the corpus mucosa significantly increased despite stable bacterial counts. The phenomenon, depicted schematically in figure 2, is commonly observed in clinical practice. Although its significance remains unclear, the authors’ suggestion that H pylori infected patients in need of long term acid suppressive therapy should receive bacterial eradication therapy seems sensible and has been espoused by a recent European Consensus Conference held in Maastricht. However, the unexplained leap from the term “corpus gastritis” (used in the first article) to “atrophic gastritis” (used, but not defined, in the second paper) and an editorial that followed a few months later caused a great degree of consternation among regulatory agencies in the United States. The question of whether proton pump inhibitors may accelerate the progression of preneoplastic lesions in the stomach of H pylori infected subjects was submitted to an advisory panel. The panel concluded that the evidence was insufficient and that no suggestions for change in the usage of proton pump inhibitor therapy should be made.

What proton pump inhibitors really do to the gastric mucosa
In addition to the shift from a predominantly antral to a predominantly corpus inflammation that occurs roughly in half of the H pylori infected patients on long term proton pump inhibitors (fig 3A), in a number of subjects on long term proton pump inhibitors the oxyntic mucosa (irrespective of the H pylori status) acquires a characteristic appearance, variously described as “hypertrophic,” “pseudo-hypertrophic,” or “hyperplastic” (fig 3B). Recently, however, this change has been described also in patients with gastric ulcer apparently not receiving acid suppressing therapy. In a small percentage of H pylori infected individuals (perhaps less than 10%), bacteria become visible in the deeper third of the oxyntic glands, and sometimes they appear to be lodged inside the canaliculi of the parietal cells (fig 3B). No increase in the deep mucosal inflammatory response is seen in these cases. The significance of this curious finding is unclear.

Another effect of prolonged treatment with proton pump inhibitors which is believed to occur in less than 10% of patients without H pylori infection and in up to 30% of those with the infection is the development of significant argyrophil cell hyperplasia. This is not, however, accompanied by an increased risk of developing malignant carcinoids.

Conclusions
Although the issues discussed in this article are apparently disparate, they all revolve around one poorly understood concept, the concept of atrophy. If a clear definition existed and a set of criteria were established for the diagnosis of atrophic gastritis, many of the confusing and sometimes acrimonious debates that originated from the study of the H pylori infected mucosa in subjects treated with proton pump inhibitors would have been avoided.

With this aim in mind, a group of pathologists, gastroenterologists, and gastrointestinal physiologists with a commitment to the study of gastric atrophy gathered in Houston, Texas, in February 1998. Their aim was to agree on...
one usable definition of atrophy and one of atrophic gastritis and to disseminate these definitions to a wide audience of gastroenterologists and pathologists. The deliberations and conclusions of the meeting will be reported in a major pathology journal.


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