**RECENT ADVANCES IN ACID PEPTIC DISEASE**

**Hp and pH: Implications for the eradication of Helicobacter pylori**

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**RH HUNT. Hp and pH: Implications for the eradication of Helicobacter pylori.** Can J Gastroenterol 1993;7(5):406-410. Helicobacter pylori infection causes inflammation of the gastric and duodenal mucosae, which results in a disturbance of the regulatory mechanisms for gastrin, gastric acid and pepsin secretion. Acid secretion may be decreased, normal or increased, depending on the stage of infection, although the meal-stimulated gastrin response is invariably elevated. The exact mechanisms involved are not known but may be due to the release of cytokines in response to bacterial toxins. H. pylori colonization is reduced by effective acid suppression with proton pump inhibitors, although it is not eradicated. In combination with amoxicillin, omeprazole, up to 40 mg twice daily, eradicated the organism in up to 82% of cases. This synergistic effect may be due to a direct effect of omeprazole on the organism, the protection of amoxicillin from acid degradation, or enhancement of host defence mechanisms accompanying acid suppression.

**Key Words:** Amoxicillin, Duodenal ulcer, Eradication, Gastric acid, Helicobacter pylori, Omeprazole, pH

**«Hp» et pH: implications pour l’éradication de Helicobacter pylori**

**RÉSUMÉ:** L’infection à Helicobacter pylori provoque une inflammation des muqueuses gastriques et duodénales, qui résulte en une altération des mécanismes régulateurs de la sécrétion de gastrine, d’acide gastrique et de pepsine. La sécrétion acide peut être diminuée, normale ou accrue, selon le stade de l’infection, bien que la réponse de la gastrine stimulée par la prise d’aliments soit invariablement élevée. Les mécanismes précis en jeu sont inconnus, mais pourraient être attribuables à la libération de cytokines en réponse à des toxines bactériennes. La colonisation de H. pylori est réduite par une suppression efficace de l’acide à l’aide d’inhibiteurs de la pompe à protons, bien que l’éradication ne soit pas obtenue ainsi. En association avec l’amoxicilline, l’oméprazole, jusqu’à concurrence de 40 mg deux fois par jour, a entraîné le pathogène chez 82% des cas. Cet effet synergique peut être dû à une action directe de l’oméprazole sur le pathogène, à une protection de l’amoxicilline contre la dégradation par l’acide ou à l’amélioration des défenses de l’hôte qui accompagne la suppression de l’acidité.

Duodenal ulcer has traditionally been considered an acid-related disease, which responds readily to pharmacological or surgical reduction of acid secretion, and a considerable change in thought is required to accept the concept that ulceration in the duodenal cap might be the result of infection with Helicobacter pylori. Colonization with H. pylori leads to a sequence of pathophysiological events that include mucosal inflammation, weakening of the mucus bicarbonate barrier, superficial epithelial cell damage, increased serum gastrin levels with defective feedback control, a possible increase in parietal cell mass, and the development of gastric metaplasia in the duodenal cap.

These observations have led to the concept of a complicated interplay between the organism and the host, whereby the sophisticated and highly integrated normal mucosal defence mechanisms are compromised in the presence of normal or elevated levels of acid and, perhaps, pepsin secretion. Ulceration may then occur as a result of this imbalance.

**RELATIVE ROLES OF ACID AND H PYLORI IN ULCER DISEASE**

H. pylori colonizes the antrum of the stomach with the aid of its urease mechanism and motility generated by spiral structure and multiple flagella. Urease, together with a spectrum of bacterial products, leads to weakening
of the mucus/bicarbonate layer and damage to the surface epithelial cells. The release of cytokines with an inflammatory response results in further damage to mucosal cells and may affect the regulatory mechanisms involving gastric secretion and motility. These changes may lead, over time, to an increase in the parietal cell mass and hypersecretion of gastric acid, with the subsequent development of gastric metaplasia in the duodenal cap. These areas of gastric metaplasia then become colonized with *H pylori* from the gastric antrum, leading to further inflammation in the duodenal cap.

Inflamed gastric metaplasia in the duodenal cap, with altered sialomucin production, and the already compromised functional mucosa, weakens the normally well-orchestrated and buttressed mucosal defences. With ongoing normal or increased acid secretion and peptic activity, mucosal damage and ulceration occur.

**H pylori and Intragastric Acidity**

*H pylori* does not thrive in a hypoacidic environment. The organism colonizes at the interface of the mucus/bicarbonate layer and surface epithelial cells, where the pH is between 6 and 7. In vitro studies show that the organism will replicate in the presence of urea down to a pH of 4.3, and it will survive without replication down to a pH of 2.3. When acid secretion is effectively suppressed, the minimal H⁺ in the lumen of the stomach results in little or no back diffusion of acid across the mucus/bicarbonate layer. The pH of the submucosal environment then rises because of ongoing urease activity and becomes alkaline due to unbuffered ammonia. It seems probable that *H pylori* can tolerate this alkaline pH for only a short period of time.

**H pylori and Gastric Acid Secretion**

**Reduced secretion:** *H pylori* may be associated with hyposecretion, hypersecretion or normal secretion of gastric acid, and this may be determined by the stage of infection. The term 'hypochlorhydria gastritis' was first coined by Oster in 1905 (1), who described an acute dyspeptic illness remarkably similar to that described by two volunteers who ingested *H pylori* and other subjects voluntarily infected (2). In the self-inoculation studies, both volunteers became infected only after predosing with an H₂ receptor antagonist, and in one well-documented case an acute gastric gastritis occurred with prolonged hypochlorhydria, as determined by the pH of basal acid samples (3).

**Epidemic hypochlorhydria:** The natural history of epidemic hypochlorhydria is not well known, but, from a review of the limited studies on this topic, it may last for weeks or months (2). Parietal cell function may be directly affected by *H pylori* or by mediators of the inflammatory response that accompanies the gastritis. In vitro, *H pylori* inhibits acid secretion, as measured by¹⁴C-aminopyrine in isolated guinea pig parietal cells (4) or rabbit gastric glands (5). In the presence of acute inflammation, several cytokines are produced, including interleukin (IL)-1β, and these may also reduce acid secretion (6.7). It is presumed that the reduction in gastric acid secretion that sometimes accompanies acute infection with *H pylori* somehow facilitates its colonization.

As gastritis extends proximally to involve the body and fundus, inflammation, which may or may not be accompanied by an autoimmune component, leads to the gradual atrophy of the parietal cell mass and reduction of acid secretory capacity.

**Hypersecretion:** The results of in vivo studies to determine the link between *H pylori* and gastric acid secretion are unclear, due to the large number of retrospective studies, variability in study design, the mixed populations studied, and the different methods used both to detect the organism and to measure acid secretion (8). In summary, there appears to be no consistent effect on basal acid secretion, although there is a trend towards increased secretion in patients infected with *H pylori*. Two studies of pentagastrin-stimulated acid secretion have suggested that there may be an increase in acid output, but both suffer from the problems alluded to above (9,10).

The data on 24 h pH profiles are also inconsistent with several studies showing no clear changes. However, Wagner et al (11) did show a significantly lower median 24 h pH in patients with duodenal ulcer (*H pylori*-positive, pH 1.34; *H pylori*-negative, pH 1.32), compared with others with gastritis (*H pylori*-positive, pH 1.58; *H pylori*-negative, pH 1.63) and healthy volunteers (*H pylori*-negative, pH 1.66), but the differences were independent of *H pylori* status. Our own study (unpublished data), although limited by small numbers, demonstrated that the median 24 h pH in *H pylori*-positive duodenal ulcer patients (pH 1.6±0.6) was lower than in *H pylori*-positive gastritis patients (pH 1.5±0.68) and *H pylori*-negative normal volunteers (pH 1.36±0.37). Furthermore, the median 24 h pH in duodenal ulcer patients after ulcer healing and eradication of *H pylori* showed a trend towards increased acidity (pH 0.94±0.18). This would be consistent with the hypothesis that the parietal cell mass or response to stimulation is increased in duodenal ulcer patients with *H pylori*, due to the trophic action of gastrin, and does not immediately return to normal.

**H pylori, Acid, Gastrin and Inflammation**

Patients with duodenal ulcer tend to have a pattern of acid secretion that is either normal or increased, and the parietal cell mass may also be increased, contributing to this hypersecretion of acid. An increase in meal-stimulated gastrin release with a defective inhibition of acid was first demonstrated in duodenal ulcer patients by Walsh et al (12). More recently, this exaggerated and prolonged meal-stimulated gastrin release has been reported in patients infected with *H pylori* (13-15). Furthermore, this abnormality is reversed when the infection is eradicated (10,13-16).

The relationship between *H pylori* infection and gastrin release is complex (17). It has been suggested that the exaggerated gastrin response is due to alkalization of the antral mucosa, as a result of the production of ammonia by *H pylori* urease activity interfering with...
the acid feedback on gastrin release by the antral G-cells (18). However, Chitajallu et al (19) have recently shown that gastric alkalization following eradication of H pylori did not cause basal gastrin values to rise to the levels observed when H pylori was present. Furthermore, a recent study with high-dose antibiotics effectively cleared the antrum rapidly of H pylori, but it took several days for the gastrin release to return towards normal, suggesting that this mechanism is altered by inflammatory mediators rather than changes in pH occurring locally in the mucus/bicarbonate layer or at the epithelial cell surface (Graham, personal communication). There is some further evidence in support of this alternative hypothesis. Dogs that had been systemically immunized showed a significant increase in gastrin levels when challenged orally with specific human gammaglobulin, compared with immunized control dogs (20). Also, when isolated antral mucosal cells were incubated with IL-1, a significant dose-dependent increase in gastrin release was observed (21).

Not only are meal-stimulated gastrin levels elevated in H pylori infection, but the area under the curve (AUC) for gastrin over the whole 24 h period is increased, raising the possibility of a continuous trophic effect of gastrin on the parietal cell mass. However, it remains speculative as to whether the increased meal-stimulated gastrin release and raised AUC for gastrin have a trophic effect on the parietal cell mass, leading to increased acid secretion. Studies to date have been indirect and have not confirmed this hypothesis (22). However, experimental designs have been suboptimal, and the studies have not had a sufficiently prolonged follow-up period to determine whether acid secretion falls after eradication.

There has been a suggestion that bacterially derived amines produced by H pylori and other bacteria might be important in stimulating gastrin release (14,23). In support of this hypothesis, gastrin release is reduced dramatically in patients with pernicious anemia and achlorhydria when the stomach is lavaged at pH 7. Furthermore, when gastric juice from these patients is perfused in the stomachs of conscious rats, gastrin secretion is elevated.

In patients given the proton pump inhibitor SKF 96022, plasma was measured before and after eradication of H pylori, and it was shown that levels of gastrin fell significantly (22). This supports the view that the hypergastrinemia found in association with H pylori is not due to an increase in antral mucosal pH caused by bacterial ammonia production, but by other products of bacterial metabolism or by mediators associated with the mucosal inflammation.

H PYLORI, PEP SIN O G EN AND P EPSIN SECR ETION

The pepsinogens are a group of proenzymes that are secreted by the chief cells and are converted to pepsins in the gastric lumen in the presence of acid. The pepsinogens are classified as two immunologically distinct subtypes, pepsinogens A/I and C/II, which are detectable in the blood or serum. Before H pylori was isolated, the serum pepsinogen profile was shown to be elevated in patients with ulcer disease (24), and in patients with chronic gastritis the pepsinogen profile correlates with the histological severity of the gastritis (25). Recently, it has been shown that eradication of H pylori is accompanied by a decrease in both pepsinogen A and C and a rise in the ratio of pepsinogen A/C.

H PYLORI AND THE EFFECTS OF PROTON PUMP INHIBITION

Omeprazole, 20 and 40 mg daily, has suppressed H pylori in some studies but not in others. The reasons for this are difficult to assess, but patient numbers have generally been small, the diagnosis and criteria of patients entered are heterogeneous, and biopsy sites in the stomach variable, thus confounding interpretation.

While treatment with omeprazole alone may suppress H pylori, the same is not found with the H2 receptor antagonists. It is unclear whether the effect results from the greater reduction in gastric acid secretion, or whether it is due to unique characteristics of omeprazole or the proton pump inhibitors as a class.

Antral gastritis improves in patients taking omeprazole, and the bacteria are few at this site; they appear to migrate to the gastric lumen (26) or to the fundus, where the environment may be more acidic.

Omeprazole and lansoprazole have been investigated in vitro for their activity on H pylori (27,28). The minimal inhibitory concentration for omeprazole is between 12.5 and 50 μmol/L (27, Goldie, Van Zanten and Hunt, unpublished data), and it appears to be increased by substitution of position 4 of the pyridine ring by a fluoro-alkoxy group (27). These studies have also suggested that acid-converted rearrangements of the parent compounds might be two to four times more potent than the parent compounds.

RATIONALE FOR EFFECT OF OMEPRA ZOLE COMBINED WITH AN ANTIBIOTIC ON H PYLORI

Omeprazole, 20 mg daily, provides effective inhibition of intragastric acidity, maintaining the pH above 3 for 16 to 18 h (29). The reduction in intraluminal H+ activity will reduce the pH gradient across the mucus/bicarbonate layer, raising the pH at the surface of the epithelial cell - possibly to alkaline levels. This could explain the proximal migration of the organism to the acid-secreting fundus which is seen in patients taking omeprazole (26).

Many antibiotics are rapidly degraded by gastric juice, but amoxicillin monotherapy eradicates H pylori in up to 25% of cases (30-32). Amoxicillin given intravenously does not appear in gastric juice under normal circumstances (33, 34), suggesting that amoxicillin may act topically in the stomach. The prolonged elevation of intragastric pH increases the concentration of acid-labile antibiotics in gastric juice and may also prolong their effectiveness.

It is possible that an elevated intragastric pH enhances the environment for the optimal effectiveness of the host defence mechanisms. The secretion of H pylori-specific immunoglobulin has an indirect correlation with the degree of invasion by the organism (unpub-
lished data). Under normal circumstances, immunoglobulin secreted into the stomach coats any luminal organisms. However, immunoglobulins are rapidly degraded by peptic activity and gastric juice at low pH. Thus, the elevation of intragastric pH above 3 for prolonged periods might extend the half-life and effectiveness of secreted immunoglobulin. Little is known about the pH-dependence of other host responses to H pylori. It is possible that the effective reduction in acidity seen with omeprazole, 40 mg twice daily, optimizes the pH in the mucus bicarbonate layer for neutrophil function.

RESULTS OF OMEPRAZOLE/AMOXICILLIN TRIALS

The results of clinical trials with omeprazole and amoxicillin showed considerable discrepancy in eradication rates. The duration of treatment and dose regimens were very variable, and different criteria for eradication were applied. In most instances, the trials were not randomized or double-blinded, many had a heterogeneous set of disease criteria, and patient numbers were usually small.

One trial by Bell et al (35) showed an eradication rate of 31.3%, but only 16 patients were studied, and the design was not comparable to the study by Unge et al (36), in which the eradication rate was 54%. Patients in the study by Bell et al included those who were metronidazole resistant, and the doses and administration of omeprazole and amoxicillin were different. Bell gave omeprazole, 20 mg once daily in the evening, which provides a shorter duration of pH greater than 3 than when omeprazole is administered in the morning (37), whereas Unge et al administered omeprazole, 40 mg once daily in the morning. Patients in the study by Bell et al also received amoxicillin, 750 mg daily, which is in contrast to the study by Unge et al, who, in addition to omeprazole, gave amoxicillin, 750 mg twice daily.

The study reported by Unge et al is much larger than that by Bell et al, and is probably more representative of the population, with respect to metronidazole resistance. Unge et al used doses that are more appropriate to the known pharmacodynamics and postulated mechanisms of action (see above). However, Unge et al gave omeprazole alone for two weeks before introducing the amoxicillin; a recent study has suggested that this approach might be a disadvantage to the combination of omeprazole and amoxicillin, which might provide better results if given together initially (38).

A recent study by Bayerdörffer et al (39) compared omeprazole, 40 mg twice daily for 10 days, followed by omeprazole, 20 mg daily for the rest of the six-week period, with omeprazole, 40 mg twice daily, plus amoxicillin, 1 g twice daily, for 10 days, followed by omeprazole, 20 mg daily, for the rest of the six-week period. H pylori was eradicated in 82% of the patients in the omeprazole/amoxicillin group. Labenz et al (38) reported that omeprazole, 20 mg twice daily, and amoxicillin, 500 mg four times daily, for two weeks eradicated H pylori in 82.8% of 62 patients. However, such treatment for one week proved much less effective (40).

Logan et al (41) gave omeprazole, 40 mg once daily in the morning, and amoxicillin, 500 mg four times daily, for two weeks to 19 metronidazole-resistant H pylori patients, and an eradication rate of 74% was achieved.

Recent work suggests there is a synergistic effect between omeprazole and triple therapy. In a trial involving 155 duodenal ulcer patients randomly allocated to receive either omeprazole plus bismuth, tetracycline and metronidazole for one week, followed by omeprazole for a further three weeks, or omeprazole alone for four weeks, eradication of H pylori occurred in 95% of patients who received omeprazole plus triple therapy but in only 4% of those taking omeprazole alone (42).

These trials demonstrate a high rate of eradication of H pylori when omeprazole is combined with amoxicillin. Regimens with twice-daily dosing of omeprazole appear to be superior, but the optimal dose, frequency of dosing, and formulation of both drugs remain to be established.

REFERENCES

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