Electron microscopic study of association between *Helicobacter pylori* and gastric and duodenal mucosa

L A Noach, T M Rolf, G N J Tytgat

Abstract

**Aim**—To study the ultrastructural appearances of *Helicobacter pylori* in antral and duodenal biopsy specimens and its relation with the epithelial cells.

**Methods**—Endoscopically obtained antral and duodenal biopsy specimens were examined using transmission electron microscopy and freeze fracture analysis.

**Results**—Most bacteria looked curved, but in the duodenal bulb cocccoid bacteria were relatively common. Bacteria were often found around intercellular junctions. Freeze fracture examination indicated abnormalities of the tight junction complexes in patients with *H pylori* infection. In many biopsy specimens bacteria were seen closely attached to the epithelial cell membrane by different forms of adhesion. In addition to what looked like intracytoplasmic penetration by bacteria, several examples of genuine penetration were observed.

**Conclusion**—*H pylori* is commonly found adhering to epithelial cells. Occasionally, *H pylori* may also penetrate cells. These features may contribute to the pathogenic action of the organism.


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*Helicobacter pylori* is ubiquitous and its presence in the stomach and duodenal bulb is strongly associated with gastritis, duodenitis, and peptic ulcers.1–3 The usual appearance of *H pylori* in vivo is that of a curved or S-shaped Gram-negative rod with several sheathed flagella.4,5 Next to spiral-shaped organisms, coccoid forms have occasionally been described.6,7 In a previous study these forms were predominantly found in the duodenal bulb.4 However, other investigators did not find evidence for a specific distribution pattern.6,8 As yet, it is not clear whether these forms reflect degenerative alterations or adaptations to marginal or hostile environments with the capacity for regrowth.4,6–8

*H pylori* is well adapted for residence within or beneath the mucus layer of the stomach. For reasons unknown, the area around intercellular tight junctional complexes seems to be a preferred site.4,6–8 A striking feature of *H pylori* is its exclusive relation to cells derived from gastric-type mucosa. The physicochemical properties of the mucus overlying gastric-type cells may be responsible for this phenomenon.9

Although most *H pylori* are found in the mucus layer, a considerable proportion adhere to epithelial cells.10,11–16 Adhesion is considered to be an important aspect of bacterial pathogenicity.11,17 Evidence shows that adherence affords advantage for toxin-producing organisms and induces degeneration of microvilli, degeneration of the cytoskeleton with actin polymerisation, depletion of mucous granules, and an increase in sialic acid-rich glycoproteins in the apical part of the cytoplasm.12,14,18 Some investigators have also suggested a direct relation between the proportion of adhesion sites and the histological grade of epithelial degeneration.12

Controversy exists over whether *H pylori* is also capable of invasion into epithelial cells. Internalisation of *H pylori* by HEp-2 cells in vitro has been observed.19 In vivo, apart from phagocytosed organisms in neutrophils, intracellular bacteria have been detected only incidentally in intracytoplasmic vacuoles, in lysosomes and closely related to such structures.4,8,9,13,15,20–22 In one study intracytoplasmic *H pylori* were reported to be present in duodenal mucosa in 10% of patients with duodenal ulceration.4,13 In contrast, several other authors did not find any evidence for invasive- ness or internalisation.5,9,10,14,23

Methods

Antral and duodenal mucosal biopsy specimens were obtained during upper gastrointestinal endoscopy from 27 patients (16 men, 11 women; mean age 55 years, range 20–79 years) with dyspeptic complaints and *H pylori* infection. The presence of *H pylori* was assessed by culture and histological examination of antral biopsy specimens.

For transmission electron microscopy, biopsy specimens from the antrum and duodenal bulb were immediately fixed in McDowell’s solution. The specimens were postfixed in 1% osmium tetroxide and dehydrated through an ethyl alcohol sequence until alcohol reached 90%. The tissue was embedded in Epon 812 and 1 μm sections were stained with Richardson’s stain. Areas of interest were selected, trimmed, and cut with a diamond knife into sections of about 600 × 101 μm for transmission electron microscopy. These sections were stained with Reynold’s lead citrate and uranyl magnesium acetate. Ultrastructural examinations were...
performed using a Philips 300 electron microscope at an accelerating voltage of 60 kilovolts.

In some preparations, ruthenium red was added to the fixation fluid to stabilise surface polysaccharides.

The distribution of coccoid forms and curved or spiral forms of *H. pylori* in stomach and duodenum was studied in biopsy specimens from five patients from whom we obtained additional tissue from the duodenal bulb.

Adhesion sites were studied in transmission electron micrographs of antral mucosa from 27 patients. Adhesion was defined as a close attachment between bacteria and epithelial cells, such that virtually no space was visible between them using transmission electron microscopy. On the basis of morphological appearances, the different types of interaction of the organism with the epithelial cells were categorised into five groups: (1) with raised plasma membrane forming an adhesive pedestal; (2) occupying depressions in the plasma membrane; (3) abutting on to the plasma membrane; (4) fusion of bacterial and plasma membrane with dissolution of membranes; (5) internalisation in cytoplasm.

The numbers of adherent organisms were counted and the prevalence and relative frequency of each type of adhesion was calculated.

In biopsy specimens from three patients freeze fracture analysis was performed to investigate the membrane ultrastructure of *H. pylori* and to examine the structure of the intercellular tight junction complexes.

Before freezing the tissue had been fixed in 2.5% glutaraldehyde in 0.1M phosphate buffer (pH 7.2). The tissue was frozen in liquid propane (−196°C), fractured at −100°C (Balzer’s freeze-etch unit), shadowed with platinum carbon, and then digested in filtered commercial bleach. The replicas were washed three times in distilled water, mounted on grids, and examined in a Philips 300 electron microscope.

**Results**

In all biopsy specimens curved as well as coccoid micro-organisms were observed. Of the total number of *H. pylori* organisms counted in biopsy specimens from five patients, from whom specimens from both duodenum as well as stomach were obtained, 13 (17%) of the microbial structures in antral biopsy specimens, and 24 (35%) in duodenal specimens were coccoid in appearance (fig 1).

Investigation of the internal appearance of *H. pylori* by freeze fracture analysis showed that the organism was surrounded by a cytoplastic membrane, which, in turn, was covered by an outer membrane (fig 2). Both the outer and inner face of the two membranes were covered with dot-like structures, presumably corresponding to incorporated proteinaceous structures.

Preliminary freeze fracture analysis in two additional patients with *H. pylori* infection and one *H. pylori* negative patient was suggestive of damage of the junctional strands in the

![Figure 1](image1.png)  *Figure 1* Multiple organisms in the duodenal bulb exhibiting a coccoid appearance.

![Figure 2](image2.png)  *Figure 2* Freeze fracture of membrane surfaces of *H. pylori* showing the cytoplasm (cyt) surrounded by the cytoplasmic membrane (cm). The fracture face of the half of the cytoplasmic membrane associated with the protoplasm is referred to as the protoplasmic face (PF) and the half facing the extracellular matrix as the extracellular face (EF).
Association between *H pylori* and gastric and duodenal mucosa

*H pylori* positive cases (figs 3 and 4). In contrast to the normal delicate network of interconnecting protein strands in the *H pylori* negative patient, the tight junctional structure appeared irregular, thickened, knobby and focally fragmented in the patients with *H pylori* associated gastritis.

At transmission electron microscopy, most *H pylori* organisms in the antrum were seen scattered in the mucus layer or positioned close to the microvillous surface without membrane-to-membrane attachment with the epithelial cells. Sometimes organisms along the surface epithelium were orientated into parallel arrays but, in general, organisms showed no preferential orientation. Moreover, many bacteria were present in the gastric pits. The immediate vicinity of intercellular junctions appeared to be a favoured site.

The presence of *H pylori* was sometimes, but not always, associated with depletion of microvilli and mucin granules. Strands of filamentous material were readily observed projecting from the bacterial surface towards the epithelial luminal membrane (fig 5). This attachment probably involves interaction between surface polysaccharides (glycocalyx) mediated by lectin binding.  

Apart from abundant bacteria in the mucus layer, without intimate contact with the epithelial cell membrane, adherent organisms were detected in 23 (85%) out of 27 infected patients whose antral biopsy specimens were investigated.

The table shows the prevalence of the different types of adhesion and internalisation formed by 231 organisms in these 23 patients.

Plateau-like extrusions of the epithelial cell or adhesive pedestals were seen in seven (30%) patients. Occasionally, pedestal formation was found at more than one site between one bacterium and the cell surface (fig 6). At sites of intimate contact, the plasma membrane was usually devoid of microvilli and of glycocalyx-like material. Other closely adherent organisms were seen occupying depressions in the plasma membrane. The most frequently observed form of adhesion was the abutting type. With this type, the contour of the epithelial membrane at the point of contact with the organism appeared unaltered, apart from loss of microvilli. Usually, adhesion was accompanied by degenerative changes in the adjacent cytoplasm consisting of a more translucent area and disruption of the cytoskeletal elements.

At other sites, a picture suggesting partial membrane fusion between the outer membrane of bacteria and the epithelial plasma

<table>
<thead>
<tr>
<th>Type of adhesion</th>
<th>n (%)</th>
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<tbody>
<tr>
<td>Adhesive pedestals</td>
<td>19 (8)</td>
</tr>
<tr>
<td>Membrane depressions</td>
<td>26 (11)</td>
</tr>
<tr>
<td>Abutting adhesion</td>
<td>164 (71)</td>
</tr>
<tr>
<td>Membrane fusion</td>
<td>16 (7)</td>
</tr>
<tr>
<td>Internalisation</td>
<td>6 (3)</td>
</tr>
</tbody>
</table>
membrane was observed. Moreover, images that suggested intracellular engulfment were occasionally seen (fig 7).

In accordance with previous findings intracellular cystic spaces filled with granular material were occasionally found.\(^1\)\(^6\) Such alterations are perhaps the morphological expression of cytotoxin-induced vacuolisation of the cytoplasmic structures.\(^1\)

Images mimicking intracytoplasmic penetration were seen in several cases. At higher magnifications, filamentous strands were sometimes visible between the bacterial and cytoplasmic membrane. Tangential sectioning may have been responsible for such confusing images. Yet early internalisation cannot be ruled out. Indeed, after an extensive search, six examples were found (in four patients) which were highly suggestive of true internalisation (fig 8).

Images of apparent deep penetration into the intercellular space were readily visualised but appeared instead to be due to excessive ballooning of the apex of the surface mucus cells. These images created a misleading impression of deep penetration whereas the organism remained superficial to the tight junction area (fig 9).

Discussion

There is substantial clinical evidence that \(H\) \(pylori\) has a causal role in the development of active chronic gastritis.\(^1\)\(^3\) \(H\) \(pylori\) has many characteristics which are considered to be prerequisites for pathogenicity: motility in the viscous gastric mucus layer; production and elaboration of urease; production of other harmful enzymes, cytokines, and toxins; and its potency of adhesion to gastric epithelial cells.\(^4\)\(^7\)\(^9\)\(^11\)\(^12\)\(^14\)

Ultrastructurally, \(H\) \(pylori\) looks like a curved or S-shaped organism with a length of about 3 \(\mu\)m and a width of about 0.5 \(\mu\)m. In accordance with previous observations, we also found coccoid-shaped bacteria in duodenum and antrum.\(^4\)\(^4\) It should be emphasised that these observations were based on a small number of biopsy specimens. Moreover, with transmission electron microscopy it is impossible to define three-dimensional shapes and the number of coccoid forms may have been overestimated due to cross-sectioning of curved bacteria. However, our finding that the percentage of organisms with a coccoid profile is seen more often in the duodenum than in the stomach, and that some duodenal biopsy specimens exclusively showed coccoid-shaped \(H\) \(pylori\), may indicate that the duodenal milieu somehow predisposes to coccoid transformation or that coccoid organisms are transported preferentially from the stomach. Why the duodenal bulb may be a preferential site for coccoid transformation is still not clear and a matter of debate. Also the question whether the coccoid form of \(H\) \(pylori\) represents a degenerative or a non-viable form or only a (temporary) dormant state needs further clarification. Jones \textit{et al} suggested that the coccoid structure assists the survival of the
A more stable form of interaction concerns the formation of adherence pedestals. Such pedestals are similar to those formed by enteropathogenic Escherichia coli. At other sites, organisms abut on to the epithelial plasma membrane or occupy depressions in the plasma membrane, or presumably attempt to penetrate cells. Whether the different morphological features of adhesion represent different stages of attachment which will ultimately end in intracytoplasmic penetration or engulfment has still to be elucidated.

Adhesion is generally associated with effacement of microvilli and disruption of cytoskeletal elements. According to some authors, evidence of intraepithelial degeneration is commonly observed. They have described distortion of the plasma membrane, dissolution of mucus granules, increased generation of sialic acid–rich glycoproteins, and the appearance of intracytoplasmic vacuoles and abundant phagolysosomes.

The production of a vacuolating cytotoxin by \( H \) pylori may be responsible for the often observed vacuolation. In one study the presence of epithelial degeneration was associated with a significantly increased proportion of adhering \( H \) pylori. Apart from loss of mucus granules resulting in a translucent empty-looking structure, and occasionally disruption of the plasma membrane, we found no other ultrastructural evidence of intracellular degeneration. It has still to be clarified whether the aforementioned degenerative changes are induced by adhesion of \( H \) pylori per se or whether \( H \) pylori preferentially adheres to damaged cells.

In several studies \( H \) pylori was mentioned as being non-invasive or as penetrating epithelial cells only rarely. The recent findings of Ohkusa et al and Posalaký et al lend further support to this theory. They reported discontinuity and decreased numbers of tight junctional strands in patients with (non-drug induced) gastric ulcers. The changes as observed in a limited number of \( H \) pylori positive patients by freeze fracture analysis in our preliminary study also indicate that damage of tight junctions may be involved in the pathogenesis of \( H \) pylori associated inflammation and ulceration.

Many previous reports have described the ability of \( H \) pylori to adhere to epithelial cells. A variety of putative bacterial adhesins have been reported to be involved in the interaction with the mucosal cell receptor.

Initial adhesion is thought to be represented by the formation of a glycolympath and probably involves a non-specific interaction.

organism by minimising the bacterial surface during unfavourable external conditions. The coccoid form of \( H \) pylori seems to be stable against physical and chemical assaults for about 30 days and has shown the ability to regrow into the curved or spiral shape under certain favourable conditions. A similar phenomenon of transformation to coccoid forms has been observed in Campylobacter jejuni and other campylobacters. The reversion of coccoid to spiral forms has also been reported in Campylobacter fetus.

In agreement with other published data, we found multiple \( H \) pylori organisms lodged between foveolar cells in the neighbourhood of the tight junction complexes. Congregation at these sites may be explained by a higher concentration of nutritive substances. Moreover, local high concentrations may weaken the intercellular junctions. Disruption of this barrier may also facilitate the transport of toxins from the bacteria to the cells and may finally result in epithelial degeneration. The recent findings of Ohkusa et al and Posalaký et al lend further support to this theory. They reported discontinuity and decreased numbers of tight junctional strands in patients with (non-drug induced) gastric ulcers. The changes as observed in a limited number of \( H \) pylori positive patients by freeze fracture analysis in our preliminary study also indicate that damage of tight junctions may be involved in the pathogenesis of \( H \) pylori associated inflammation and ulceration.

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