Mucin exocytosis: a major target for Helicobacter pylori

I Micots, C Augeron, C L Laboisse, F Muzeau, F Méraud

Abstract

Aims: To determine whether Helicobacter pylori impairs the secretory function of mucous cells.

Methods: The mucus secreting human cell line Cl. 16E, maintained as confluent monolayers on nitrocellulose filters, was infected with H pylori strain CIP 101260. After three hours of incubation with H pylori the monolayers were washed and reincubated with fresh culture medium for various time periods (24, 48, or 72 hours) before evaluating both the morphology and function of mucous cells. For morphological studies, epithelial monolayers were fixed in situ and processed for both standard histochromy on paraffin wax sections, and electron microscopy. To measure mucins secreted from cultured cells, the cells were metabolically labelled with 3H-glucosamine. Undegraded mucins were quantitated as the radioactive glycoproteins blocked at the stacker gel interface after sodium dodecyl sulphate-polyacrylamide gel electrophoresis of the secretory glycoproteins.

Results: Control cultures of Cl. 16E cells grew on filters as homogeneous monolayers of polarised mucous cells secreting a visco-elastic gel of mucins at the apical surface. In infected monolayers H pylori was in close contact with the apical surface of mucous cells. Cell counts and histological evaluation of the monolayers did not reveal any significant deleterious effect of H pylori on the mucous cells. H pylori induced only a modest inhibition of baseline mucin secretion from Cl. 16E cells, this inhibition being significant only at 24 hours. In contrast, the mucus secretory response to agents that raise intracellular cAMP and calcium—forskolin and ionophore A23187—was strongly inhibited. The inhibitory effect of H pylori on the exocytotic response was not paralleled by an inhibition of glycoprotein synthesis.

Conclusion: Considering the fact that the exocytotic response to a variety of secretagogues constitutes the primary line of defence of the gastric mucosa in an emergency, it is suggested that H pylori exerts its deleterious effects by weakening this important physiological defence.

Helicobacter pylori is now recognised as the main cause of antral gastritis and is implicated as an aetiologial agent in duodenal ulcer disease. However, despite fairly intensive studies, the pathogenic mechanisms of H pylori, either direct or indirect, are still poorly understood. Several morphological studies have contributed important information with regard to the interaction of H pylori with gastric epithelial cells. In particular, the finding of a close association of H pylori with the apical surface of mucous cells in the gastric pits strongly suggests that H pylori can interfere with the function of these specialised cells. In view of the fact that mucous glycoproteins provide the first line of defence for gastric mucosa, the hypothesis that H pylori exerts its deleterious effects by impairing the secretory function of mucous cells should be examined.

The ideal cellular model for testing this hypothesis would be a monolayer culture of mucous cells that have retained in vitro the regulatory mechanisms operative in the same cells in vivo. However, it is still impossible to maintain normal human gastric mucous cells in culture.

A human epithelial cell line, Cl. 16E, which is differentiated into monolayers of polarised mucous cells is now available. Several studies have shown that Cl. 16E cells have retained in vitro the regulatory mechanisms of mucus secretion of normal mucous cells in vivo.

Methods

H pylori strain CIP 101260 isolated from an antral biopsy specimen in a patient with chronic antral gastritis was used in this study. The strain was maintained frozen at -70°C before testing. Culture was performed on Wilkins Chalgren agar (Oxoid, Basingstoke, England) supplemented with 10% human blood and antibiotics (vancomycin, cefsulodin, and actidione), and incubated for 24 hours in a microaerobic atmosphere.

The Cl. 16E cell line is a stable, differentiated derivative of the human colonic cancer cell line HT29. Cl. 16E cells are morphologically and functionally differentiated into typical mucous cells. Cl. 16E cells were routinely cultured in 25 cm² plastic flasks (Falcon, Paris, France) in Dulbecco’s modified medium (DME) (Gibco, Paris, France) supplemented with 10% heat inactivated fetal calf serum (FCS) (Gibco). All cultures were free of mycoplasmal contamination, as tested periodically by the method of Chen. Culture on filters was performed as described previously. Experiments began on day 8 after seeding, when the cells were covered with a
visible mucus gel.

The cell monolayers were washed and 100 μl of a suspension of *H pylori* (OD = 0.8 at 600 nm) prepared in sterile brucella broth was added to each dish containing 2.5 ml of culture medium. Control cultures were incubated without *H pylori*. Incubation was at 37°C in the presence of 10% CO₂ (CO₂ gas generating kit, Oxoid). After three hours of incubation, the non-adherent bacteria were removed by washing the cell monolayers with three changes of phosphate buffered saline. The monolayers were reincubated in standard medium (DMEM, 10% FCS). In other experiments D-(6-3H)-glucosamine hydrochloride was added for 24–72 hours to study mucus secretion.

At indicated time periods, filters were fixed in Bouin’s fluid, cut into strips, and embedded in paraffin wax. Cross sections of filters were stained with haematoxylin and eosin, and with alcian blue (pH = 2.5) for mucin staining. Filters were also processed for transmission electron microscopy, as previously described.13

A monolayered cell suspension of the filter grown cells was obtained using a two-step dissociation method1 and cells were counted in a haemacytometer. Cell viability was assessed by trypan blue exclusion. Metabolic labelling of filter grown cells (control cultures and *H pylori*-infected cultures) was performed to measure mucin secretion using the precursor D-(6-3H)-glucosamine hydrochloride (10 μCi/filte, specific activity 20–40 Ci/mmol) (Amersham) in DMEM, 10% FCS. At indicated time periods, the incubation medium was removed using a pipet and the monolayers were rinsed with the spent medium to remove adherent mucin. Then the incubation medium was dialysed for 36–48 hours against several changes of deionised water at 4°C and lyophilised. The secretory glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay”.

The parietal glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay”.

The parietal glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay”. The parietal glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay”. The parietal glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay”. The parietal glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay”. The parietal glycoproteins were then separated by sodium dodecyl sulphate-polyacrylamide gel electrophoresis on 3% gels. After electrophoresis, each lane of the stacking gel containing the 3H-mucin glycoproteins was cut out, then put into a borosilicate phial and incubated overnight at 40°C in the presence of 500 μl of Soluene 350 (Packard). The phials were allowed to cool, and 10 ml of a solution of Pico-fluor 40 (Packard) was added to each phial. The radioactivity was counted in an LKB scintillation counter. This assay was referred to as the “electrophoretic assay".

**Results**

Confluent cultures of Cl. 16E cells on nitrocellulose filters formed homogeneous monolayers of polarised mucous cells. The cell number remained constant throughout the incubation period in control cultures.

Electron microscopic examination of infected monolayers disclosed numerous organisms which were in close contact with the apical membrane of mucous cells (fig 1) as well as in clusters in the mucus layer. Their location was restricted to the apical membrane with apparently no preference for the region close to interactive adherent junctions. They were seen to ionophore A23187 and forskolin was measured using the release of 3H-mucin as an index of triggered exocytosis. First intracellular mucin granules were labelled by incubating cell monolayers with 3H-glucosamine as indicated above. At the end of the incubation period, the filters were washed three times and transferred into the wells of a multiwell culture plate containing Ham-DMEM (600 μl/well) with 0.01% bovine serum albumin (w/v), 4·10⁻⁴ M ionophore A23187 (Sigma Chemicals, St. Louis, Missouri, USA), and 10⁻⁴ M forskolin (Calbiochem Corporation, La Jolla, California, USA). After 45 minutes of incubation, the medium was removed and the monolayers were rinsed with the spent medium to remove adherent mucin. Then the secreted mucins were measured using the “electrophoretic assay”.

Prior to the intracellular incorporation studies, the cell monolayers were incubated with or without (control) *H pylori* for three hours then washed and reincubated for 24 hours in DMEM with 10% FCS, according to the previously described protocol. The monolayers were incubated for three hours in 600 μl of serum-free medium containing 10 μCi/ml of D-(6-3H)-glucosamine hydrochloride. At the end of the incubation period, the filters were washed three times in 0.9% NaCl without glucosamine, and the cells were scrapped from filters with a rubber policeman in 2·5 ml of a 0.9% NaCl solution. Then the cells were sonicated for 30 seconds.

Incorporation of D-(6-3H)-glucosamine into acid precipitable glycoproteins of the cell homogenates was measured by adding cold trichloroacetic acid (TCA) and phosphotungstic acid (PTA) at a final concentration of 10% and 1%, respectively, to an aliquot of the homogenate. After an overnight incubation at 4°C the precipitate was collected by centrifugation, washed with an additional 1 ml of TCA-PTA, then with 1 ml of chloroform-methanol (1:1, vol/vol) and dried. The pellet was moistened by adding 50 μl of water and then dissolved in Soluene 350 (Packard). Radioactivity was determined after addition of 5 ml of Pico-fluor 40 in a liquid scintillation counter.

Results are expressed as the mean (SEM). Comparison between cells incubated with mediators and control cells was assessed with a Student’s t test. Histological evaluation of the monolayers on cross-sections did not disclose major modifications on infection with *H pylori*. Cell monolayers examined 24 hours after inoculation with *H pylori* did not differ from controls. Only minor morphological changes were observed after 48 hours of incubation. They consisted of a slight disorganisation of the monolayers without erosion. Cell counts confirmed the
Mucin exocytosis: a major target for Helicobacter pylori

Figure 1 Transmission electron micrograph of Cl. 16E cells infected with Helicobacter pylori for three days. Numerous organisms are present in close association with the apical membrane of the mucous cells.

Absence of deleterious effects of H. pylori on mucous cells (fig 2), since there was no significant variation in cell number between control and H. pylori-infected cells over a 72 hour incubation period.

As shown in fig 3, baseline secretion from control cultures was linear over a 72 hour period. The amount of radioactive mucus recovered from culture media of H. pylori-infected cells was significantly lower than in control cultures for the first 24 hour incubation period. When incubation was prolonged to 48 or 72 hours, the concentration of mucus in H. pylori-infected cultures did not significantly differ from controls.

To obtain further insight into the effects of H. pylori on mucus secretion, we measured the mucin secretory rate after a 45 minute stimulation of Cl. 16E cells with ionophore 23187/forskolin. As shown in fig 4, a combination of these two agents triggered a strong exocytotic response from control cultures. In contrast, the secretory response triggered by the combination of ionophore 23187/forskolin was inhibited by nearly 50% in infected cells at time points 24, 48, and 72 hours.

In view of our findings showing that the exocytosis of mucus was severely impaired in H. pylori-infected mucous cells, it was important to examine whether H. pylori was also able to inhibit intracellular glycoprotein synthesis.
A23187 forskolin from pylori on difference described in response triggered by ionophore exocytotic secretagogues. Extracellular Ham-DMEM controls (p experiment. 244 (SEM) for incubated uere Cl. 16E cells. and represents measured < "Methods". 45 minutes of 0 3). for inoculation. Ventola.4 "The cell line provides that Cl. 16E cells. and cell loss remains consistent with normal gastric epithelium. These findings are in line with other experimental work showing that human colonic cancer cells may express antigenic determinants which are normally restricted to gastric mucous cells in the human adult.16 In view of these findings it is conceivable that the adherence of the organism to Cl. 16E cells is in relation to the naturally occurring situation. Several important findings regarding both morphology and function of H pylori-infected mucous cells are highlighted by this study: H pylori did not induce major cellular morphological changes nor did it exert a major cytotoxic effect on Cl. 16E cells over a 72 hour incubation period. This is consistent with the results of other ultrastructural studies which led to the conclusion that, apart from some distortion and depletion of microvilli, H pylori do not cause pathological changes in mucosal epithelial cells, even when they are in membrane to membrane contact with each other.4 Given that the principal function of gastric mucous cells is to protect the mucosa by secreting a visco-elastic gel of mucus, it may be assumed that this secretory process is an important target for H pylori. In the normal gastric mucosa the extracellular mucus blanket is contributed by (i) the baseline secretion of the contents of stored intracellular granules and (ii) the accelerated exocytosis of stored granules under acute stimulation. The acute stimulation is triggered by neuroendocrine agents as well as by mediators released by inflammatory cells.17 These various agents act on specific receptors at the cell surface. It is generally assumed that the acute stimulation is the major contributor to a protective layer of mucins in response to an "emergency" situation. It was therefore important to explore these two mechanisms, first by measuring the baseline secretion of mucus from H pylori-infected cells, and second by measuring the exocytotic response of these cells to known secretagogue agents.

For this purpose, the incorporation of "H-glucosamine into intracellular glycoproteins was measured according to the method of Lamont and Ventola."14 The glycoprotein biosynthesis in H pylori-infected cells was not significantly different from control cultures 24 hours after inoculation.

Discussion
This study provides the first demonstration that H pylori infection can directly impair the secretory function of mucous cells. Our evidence is based on a unique experimental approach using a homogeneous human mucous cell line (Cl. 16E). Cl. 16E cells can be maintained in vitro as fully polarised cell monolayers having the kinetic characteristics of an epithelium in steady state—that is, cell production and cell loss are balanced so that the cell population remains constant. These in vitro characteristics are close to the in vivo situation. This contrasts with most of the previously used models which consisted of undifferentiated, exponentially growing cell cultures.

The Cl. 16E cell line may be considered as an appropriate model system for studying functional alterations of H pylori-infected mucous cells for two reasons: (i) several studies have shown that Cl. 16E maintained as floating cultures on nitrocellulose filters have retained in vitro the same regulatory mechanisms of mucus secretion as those operating in vivo;9 and (ii) H pylori was found to adhere to the cell surface of Cl. 16E cells in a way which mimicked the in vivo situation (this study). In particular, the location of H pylori to the apical surface of mucous cells agrees with previous ultrastructural observations in vivo.3 Interestingly, Cl. 16E cells, which are of colonic origin, resemble human gastric mucous cells in their immunocytochemical reactions: support for such a resemblance has come from the demonstration that polyclonal antibodies raised against Cl. 16E secretory mucins were found to react strongly on immunocytochemistry with normal gastric epithelium.15 These findings are in line with other experimental work showing that human colonic cancer cells may express antigenic determinants which are normally restricted to gastric mucous cells in the human adult.16 In view of these findings it is conceivable that the adherence of the organism to Cl. 16E cells is in relation to the naturally occurring situation.

Several important findings regarding both morphology and function of H pylori-infected mucous cells are highlighted by this study: H pylori did not induce major cellular morphological changes nor did it exert a major cytotoxic effect on Cl. 16E cells over a 72 hour incubation period. This is consistent with the results of other ultrastructural studies which led to the conclusion that, apart from some distortion and depletion of microvilli, H pylori do not cause pathological changes in mucosal epithelial cells, even when they are in membrane to membrane contact with each other.4 Given that the principal function of gastric mucous cells is to protect the mucosa by secreting a visco-elastic gel of mucus, it may be assumed that this secretory process is an important target for H pylori. In the normal gastric mucosa the extracellular mucus blanket is contributed by (i) the baseline secretion of the contents of stored intracellular granules and (ii) the accelerated exocytosis of stored granules under acute stimulation. The acute stimulation is triggered by neuroendocrine agents as well as by mediators released by inflammatory cells.17 These various agents act on specific receptors at the cell surface. It is generally assumed that the acute stimulation is the major contributor to a protective layer of mucins in response to an "emergency" situation. It was therefore important to explore these two mechanisms, first by measuring the baseline secretion of mucus from H pylori-infected cells, and second by measuring the exocytotic response of these cells to known secretagogue agents.
From our results, it is clear that *H pylori* induced only a modest inhibition of baseline secretion from Cl. 16E cells, this inhibition being significant only at time point 24 hours. In contrast, *H pylori* had a major inhibitory effect on the stimulated exocytosis of mucins at all time points tested.

Our approach was based on two agents which are known to trigger rapid exocytosis of mucous granules by stimulating two important intracellular signalling pathways of secretory cells—cAMP and Ca**++**. Knowing the intracellular sites of action of these agents, our finding of an inhibitory effect of *H pylori* on the stimulated secretion can be interpreted as resulting from the action of *H pylori* at a step distal to the second messenger system—in the exocytotic mechanism of mucous cells (fig 5).

In the context of these results, it was important to determine whether the exocytotic machinery was the sole target of *H pylori*, or whether this effect was accompanied by a reduction of glycoprotein synthesis. Together our findings point to a specific impairment of glycoprotein exocytosis in *H pylori* infected mucous cells without any significant effect on mucus synthesis.

Interestingly, our results derived from an in vitro approach are in line with a recent preliminary report based on human antral biopsy specimens, suggesting that *H pylori* alters the release of intracellular mucins into the lumen.18

The molecular mechanisms underlying the exocytotic defect are still unknown. The recent observation by Bell and Manning19 that *Campylobacter jejuni* causes a temporary brush border dysfunction in intestinal epithelial cells, has led to the suggestion that these organisms can selectively impair enzymes that are present at the apical surface. In the same way, *H pylori* could affect exocytosis by impairing the fusion of mucous granule membranes to the apical surface. Alternatively, in view of recent findings suggesting that intimate attachment of *H pylori* to cultured human gastric epithelial cells is associated with actin polymerisation,20 it may be hypothesised that these cytoskeletal changes can prevent secretory granules from reaching the exocytotic sites.

Finally, what is the pathological relevance of our findings? The fact that *H pylori* selectively impairs the exocytotic response of mucous cells may have important pathological consequences. In fact, it is important to consider that the exocytotic response to a variety of physiological secretagogues such as inflammatory mediators constitutes the primary line of defense of the gastric mucosa faced with an emergency. In this context, we postulate that *H pylori* may profoundly weaken this defense, thereby leading to cellular damage when the mucosa is exposed to various types of aggressive agents. We suggest that this indirect action may be more important than the direct cytotoxic action of the bacteria on the epithelial cells.

This work was supported in part by the Conseil Scientifique de la Faculté Bichet, by ARC, and by the Conseil Regional d’Aquitaine.