Incidence of *Helicobacter pylori* strains activating neutrophils in patients with peptic ulcer disease

H Rautelin, B Blomberg, H Fredlund, G Järnerot, D Danielsson

Abstract

A total of 61 human gastric isolates of *Helicobacter pylori* were studied for their ability to induce an oxidative burst in human neutrophils measured by luminol enhanced chemiluminescence. About one third of the strains induced strong and rapid chemiluminescence in neutrophils even without serum opsonins and agglutinated these cells on glass slides within two minutes. For other strains complement was required, although even then the reactions remained at a lower level. The activating and agglutinating property was bound to the cells, heat labile, and sensitive to several enzymes but resistant to acid. Strains possessing such activity were more common in patients with peptic ulcer disease than in patients with active chronic gastritis only (p=0.0261, Fisher’s exact test, two tailed). The activity shown might be a new indicator for ulcerogenic strains and could also partly explain the accumulation of neutrophils in the gastric mucosa during *H pylori* infection.

(The association of *Helicobacter pylori* (H pylori) with gastroduodenal disease was first shown by Warren and Marshall. The association of *Helicobacter pylori* (H pylori) with peptic ulcer disease has been established. A large number of *H pylori* strains, capable of activating and agglutinating neutrophils, were found significantly more often in patients with peptic ulcer disease than in those with active chronic gastritis only.

Patients and methods

**HELCOBACTER PYLORI STRAINS AND PATIENTS**

A total of 61 isolates of *H pylori* from human gastric biopsies were included in the study. All the strains were oxidase, catalase, and urease positive, and showed typical morphology with Gram stain. The individual strains were stored at −70°C before use. For the in vitro experiments the strains were grown for 48–72 hours on GC agar plates (GC II agar base, BBL, Cockeysville, MD, USA, supplemented with 1% bovine haemoglobin (BBL), 10% horse serum, and 1% IsoVitaleX) without antibiotics. All the strains were tested for the induction of chemiluminescence in neutrophils and for agglutination of these cells. Six strains were selected and used in extended experiments and will be referred to as reference strains; one strain designated NCTC 11637 from the National Collection of Type Cultures, one Canadian strain designated C 7050, two Finnish strains, F 6 and F 88, and two Swedish strains, S 206 and S 210. The rationale for selecting these six strains was three-fold: in preliminary experiments they behaved differently in the chemiluminescence, they originated from patients in four different countries, and were shown in a previous study to express different serological patterns with homologous and heterologous hyperimmune rabbit *H pylori* antisera. The remaining 55 strains were isolated from gastric biopsies from an unselected group of 210 consecutive patients referred for upper gastrointestinal endoscopy. The endoscopic findings and the presence or absence of peptic ulcer disease were recorded. The gastric biopsies were evaluated histologically for the type of gastritis as described. The medical records of *Helicobacter pylori* positive patients in whom the endoscopy did not show peptic ulcer were examined to find those in whom peptic ulcer disease had been diagnosed earlier.

The six *H pylori* reference strains had undergone multiple subcultures whereas the 55 recent isolates were subcultured four or five times before being studied. The reference strains were tested for hydrophobicity with the salt agglutination test, and for mannose resistant and manganese sensitive haemagglutination of erythrocytes from humans, sheep, and rabbits as described elsewhere. Organisms of these strains were subjected to heat treatment at temperatures of 45°C to 90°C or boiling for 30 minutes or to different treatments for 30 minutes at 37°C - namely, in NaH₂O₃ (0.5% w/v in 0.05 M Tris, pH 7.0), trypsin (Sigma, St Louis, MO, USA, 0.5% w/v in 0.03 M Tris, pH 7.0), pepsin (Sigma, 0.5% w/v in 0.03 M Tris, pH 2.3), glycine hydrochloride (0.2 M, pH 2.2), or formalin (3% in phosphate buffer saline (PBS)). After treatment the organisms were washed and suspended in PBS and then used in tests to induce chemiluminescence in neutrophils and to agglutinate these cells in parallel with live organisms of the same strains. Also, organisms of the six reference strains were disrupted by ultrasonic treatment with an MSE 20 Kc ultrasonic oscillator operated at maximum efficiency. Outer membrane preparations were made from three of the six reference strains, NCTC 11637, C 7050, and F 88 according to the method of Tam et al.
suspensions of disrupted cells, the supernatants and the sediments obtained after centrifugation at 10 000 g for 30 minutes as well as the outer membrane preparations adjusted to correspond to an original concentration of 10^8 organisms/ml were used in parallel with live organisms of the same strains to induce chemiluminescence in neutrophils and to agglutinate these cells.

**HELCOBACTER PYLORI SERUM ANTIBODIES**

Serum was taken from six patients with known *H pylori* state, from two children, and from six members of our laboratory staff who were also used as donors of PMNLs. The serum samples were divided into small portions and kept frozen at −70°C before use. All serum samples showed normal complement mediated haemolytic activity. The concentrations of IgG, IgA, IgM, and subclasses of IgG were within normal limits.

*Helicobacter pylori* serum antibodies of IgG, IgM, and IgA classes were measured with an enzyme immunoassay (EIA) with the use of an acid glycine extract from *H pylori* strain NCTC 11637 as the antigen. For the absorption of *H pylori* antibodies 0.5 ml of a serum sample was mixed with an equal volume of packed bacteria (obtained by centrifugation of strains NCTC 11637 or C 7050 for 20 minutes at 2000 g, incubated at 0°C for 30 minutes, and centrifuged twice to remove bacteria. No specific antibodies were found after the absorption whereas the complement mediated haemolytic activity of the serum was retained.

**SERUM BACTERICIDAL TESTS**

For serum bactericidal tests 0.2 ml of serum diluted 1:10 (untreated or heated at 56°C for 30 minutes, or absorbed with *H pylori*) was mixed with 0.2 ml of freshly prepared *H pylori* organisms (10^9/ml, in a total volume of 0.8 ml of the buffer used (Hank’s balanced salt solution with 0.1% gelatin, HBSS-gel)). The mixtures were incubated at 37°C and viable colony forming units (CFU) were counted at time 0, and after 15, 30, and 90 minutes by serial dilution in sterile PBS and plating on GC agar plates. Controls consisted of serum replaced by the buffer used. The time 0 value was used to calculate the log_{10} values of the decreases or increases of CFU/ml.

**PREPARATION OF POLYMORPHONUCLEAR LEUCOCYTES**

Heparinised blood was taken from *H pylori* antibody negative or antibody positive members of our staff. For each series of experiments on a particular day, blood from a single donor was used. To obtain PMNLs the method of Böyum was used with minor modifications. The purity and viability of the PMNL suspensions was more than 95%.

**PHAGOCYTOSIS ASSAYS**

The phagocytosis and killing of *H pylori* by PMNLs were studied by mixing 0.2 ml of bacteria (10^9/ml) with 0.2 ml of PMNL (10^9/ml) and 0.2 ml of serum (diluted 1:10), untreated or heated at 56°C, in a final volume of 0.8 ml of HBSS-gel. Controls consisted of serum or PMNL or both replaced by HBSS-gel. The mixtures were slowly tumbled at 37°C and aliquots were removed at 0, 15, 30, and 90 minutes to measure the CFU/ml. The time 0 value of the control mixture with bacteria alone in HBSS-gel was used to calculate the log_{10} values of the decreases or increases of CFU/ml.

The uptake of bacteria by PMNLs in the presence or absence of *H pylori* positive or negative serum was also examined by microscopy. In these experiments one volume of 2×10^7 bacteria/ml was mixed and incubated with one volume of 10^5 PMNLs/ml and serum. Slides were prepared at times 0, 5, 10, 15, 30, and 60 minutes by placing 0.05 ml of the mixture on a glass slide treated with egg white and glycera. The slides were dried in air, gently fixed by heat, and then stained with acridine orange. The slides were examined with a fluorescence microscope with oil immersion objectives (Zeiss epiplan, 50 w halogen lamp).

**CHEMILUMINESCENCE**

The induction of chemiluminescence in PMNLs by *H pylori* in the presence or absence of serum was studied by luminol enhanced chemiluminescence. For each experiment 200 µl of PBS, 100 µl of serum, 100 µl of the PMNL suspension (10^9/ml), 50 µl of 10⁻¹ M luminol (Sigma), and 50 µl of the bacterial suspension (10^9/ml) were added to each test tube (LKB, Bromma, Sweden). The measurements with a luminometer (LKB Wallac 1251, Turku, Finland) were always started within one minute after the bacterial suspension had been added, and all tests were run within three hours of collection of the blood samples for the preparation of PMNLs. The assays were performed at 37°C with continuous mixing in the measurement chamber. Chemiluminescence from each sample was measured at 70–90 second intervals for at least 30 minutes. A *Staphylococcus epidermidis* strain (strain L–9 of our strain collection), inducing stable and reproducible chemiluminescence responses of PMNLs, was run in each test as a control.

**SLIDE AGGLUTINATION TEST OF PMNLs AND HELICOBACTER PYLORI**

One drop of a suspension of PMNLs (10^9/ml) was mixed on a glass slide with one drop of a bacterial suspension (10^9/ml). The slide was then tilted about once a second for two minutes, after which the reactions were recorded as positive or negative.

**Results**

**TWO PATTERNS OF INTERACTION WITH NEUTROPHILS**

After opsonisation with anti *H pylori* positive or negative serum the six *H pylori* reference strains regularly induced chemiluminescence responses in PMNLs (Fig 1). Under corresponding conditions they were phagocytosed (judged by micro-
Incidence of Helicobacter pylori strains activating neutrophils in patients with peptic ulcer disease

scopy) and killed in the phagocytic killing experiments (Fig 2). Serum bactericidal tests showed that the strains were sensitive both to anti *H pylori* positive and negative serum (Fig 2), even after serum was absorbed for specific antibodies. This serum sensitivity remained after Clq was removed by treatment with 10 mM ethylene glycol tetra-acetate (EGTA) but was abolished after the serum was heated at 56°C for 30 minutes, which also resulted in no phagocytic killing. Organisms opsonised with EGTA treated serum gave comparable chemiluminescence results with untreated serum whereas heat treated serum was a poor opsonin.

Organisms of the strains NCTC 11637, F 6, and S 206 but not those of C 7050, F 88, and S 210 induced even stronger chemiluminescence responses unopsonised (Fig 1), whether the neutrophils were from donors with or without *H pylori* antibodies. Although there were some day to day variations in the peaks of chemiluminescence responses with PMNLs from different neutrophil donors and also within the same donor, the patterns were consistent. The strains NCTC 11637, F 6, and S 206 thus induced strong and rapid chemiluminescence responses with peaks, measured in mV, within five to 15 minutes. The strains C 7050, F 88, and S 206 induced weak (<30% of mean mVs) of strains NCTC 11637, F 6, and S 206) and slow responses with peaks (mV) after 15 minutes. These two patterns were also found in the unselected clinical isolates.

Studies by microscopy showed that unopsonised *H pylori* organisms of the strains NCTC 11637, F 6, and S 206 attached rapidly to the neutrophils, and were phagocytosed (Fig 3). These findings were confirmed by electron microscopy (data not shown). No phagocytic killing was noted, however, unless complement was added (Fig 2). Agglutination of neutrophils could also be seen on a glass slide within two minutes of mixing and tilting one drop each of bacteria and neutrophils. No such agglutinations were found with neutrophils and *H pylori* of C 7050, F 88, and S 210 strains. Slide agglutination and chemiluminescence tests of the 55 unselected *H pylori* isolates also agreed.

**Figure 1:** Chemiluminescence induced in polymorphonuclear leucocytes (PMNLs) by Helicobacter pylori strains NCTC 11637 and C 7050, either without opsonising serum, or in the presence of serum with (SeAb) or without (SeAb) specific antibodies. Results shown from a representative test run.

**Figure 2:** Serum bactericidal activity against Helicobacter pylori and phagocytic killing. For bactericidal tests *H pylori* NCTC 11637 organisms were added to serum with (SeAb) or without (SeAb) *H pylori* specific antibodies. For phagocytic killing polymorphonuclear leucocytes (PMNLs) were added to mixtures of bacteria and serum with or without *H pylori* antibodies. Control mixtures included bacteria alone in buffer, or bacteria with PMNLs. Aliquots were removed from each mixture after 0, 15, 30, and 90 minutes of incubation, serially diluted, and plated to determine the decrease of colony forming units (CFU/ml) at each time point. Results are shown from a representative experiment.

**Figure 3:** Interaction of Helicobacter pylori NCTC 11637 and C 7050 with neutrophils stained with acridine orange after incubation for 15 minutes. Unopsonised organisms of NCTC 11637 have agglutinated neutrophils that also show phagocytosis (A). Neutrophils and unopsonised organisms of C 7050 are randomly distributed, with neither agglutination nor phagocytosis (B).

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trypsin (pH 7.0), pepsin (pH 2.3), formalin, or heat ≥50°C. Outer membrane preparations of *H. pylori* cells could not activate or agglutinate PMNLs, nor could the sediment or the supernatant of organisms disrupted by ultrasonication. The activation and agglutination properties of *H. pylori* were, however, preserved after exposure to glycine hydrochloride and to NaCN.

The PMNL agglutination showed no correlation with haemagglutination patterns of the six *H. pylori* reference strains, five of which showed mannosae resistant haemagglutination of human, rabbit, and sheep erythrocytes, and one (S 210) which did not agglutinate sheep red cells but showed weak mannosose resistant agglutination of human and rabbit red cells. All strains showed low relative hydrophobicity (salt aggregation test ≥5-12) regardless of the chemiluminescence patterns.

**Neutrophil activating and agglutinating property of isolates of *H. pylori***

<table>
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<th>H. pylori positive patients</th>
<th>No with peptic ulcer disease</th>
<th>No with chronic gastritis only</th>
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<tr>
<td>Present</td>
<td>14</td>
<td>6</td>
<td>20</td>
</tr>
<tr>
<td>Absent</td>
<td>13</td>
<td>22</td>
<td>35</td>
</tr>
<tr>
<td>Total</td>
<td>27</td>
<td>28</td>
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**Discussion**

Our study showed that *H. pylori* strains could be divided into two main groups on the basis of their interaction with PMNLs. In the absence of opsonins, about one third of the *H. pylori* strains isolated from gastric biopsies of an unselected and consecutive group of patients with gastro-duodenal disease induced a rapid and strong oxidative burst in the neutrophils, which were also rapidly agglutinated by the organisms of these particular strains. The unopsonised bacteria were obviously phagocytosed, as confirmed by electron microscopy, but no phagocytic killing occurred. The oxidative burst in phagocytosis might, however, liberate free radicals and so cause cell damage. It is therefore of particular clinical interest that 14 out of the 20 (70%) strains that activated and agglutinated neutrophils, occurred in patients with peptic ulcer disease. This finding may indicate that some *H. pylori* strains are more ulcerogenic than others. The factors responsible for the described activity were found to be bound to the cells, as well as sensitive to heat (≥50°C) and several peptidic enzymes. These properties indicate that they might be proteins or peptides. Formalin treatment also abolished the activity that was preserved after treatment of the organisms with acid glycine.

Other research groups have also studied chemiluminescence induced in PMNLs by *H. pylori* organisms and found results corresponding to ours with opsonised organisms. They reported only low responses, however, with unopsonised organisms. The small number of isolates tested by Das et al might have belonged to the group of *H. pylori* strains that does not have the properties described in the present study. Other researchers used heat killed organisms. This might explain why they did not detect this heat labile activity in their isolates.

Both types of *H. pylori* strains were phagocytosed and the number of organisms significantly reduced (≥99-9%) in phagocytic killing tests in the presence of serum with or without *H. pylori* antibodies. These findings are in agreement with those described by others. We also found that the killing was mainly due to the sensitivity of *H. pylori* organisms to normal serum and that this sensitivity was in fact dependent on the alternative complement pathway as reported by Das et al. These observations do not, however, preclude the fact that specific *H. pylori* antibodies enhance the complement dependent killing of *H. pylori* organisms.

Although *H. pylori* infection is strongly associated with active chronic gastritis and peptic ulcer disease, the mechanisms of pathogenesis are far from understood. The sensitivity of *H. pylori* to normal serum might explain why these organisms do not cause septicemia and remain localized. *H. pylori* has been shown to produce chemotactic factors that might account to some extent for the accumulation of neutrophils in the gastric mucosa. These factors have been shown to be soluble or present in sonicates whereas the activity described in our study was found to be bound to the cells and not present after sonication. It is tempting to speculate that the neutrophil activating and agglutinating properties in one third of our strains might be a relevant virulence factor in the ability of such strains to challenge neutrophils both in the absence and in the presence of serum with or without specific antibodies.

Leunk et al described an extracellular cytotoxin produced by some *H. pylori* strains that was shown to be associated with duodenal ulcer
Incidence of Helicobacter pylori strains activating neutrophils in patients with peptic ulcer disease

603


