Campylobacter pylori in a sample of Finnish population: relations to morphology and functions of the gastric mucosa

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SUMMARY The occurrence of Campylobacter pylori (CP) was examined in 179 subjects representing a sample collected from the population of South Finland. In a normal antral and body mucosa CP was present in 5% and 11% and in superficial gastritis (SG) in 71% and 91% of subjects, respectively. In atrophic gastritis (AG) of antrum and body the prevalence of CP decreased significantly with an increasing degree of atrophy, so that CP was not found in severe body AG. Different combinations of antral and body gastritis revealed a characteristic pattern. Campylobacter pylori was lacking when antral and body mucosa were normal, but was present in 41% when normal mucosa was associated with gastritis in the opposite area. In SG affecting diffusely antrum and body, the bacterium was present in every case, but when SG was associated with AG in the opposite area it was lacking in 29% of the subjects. When SG affecting both areas was compared with SG accompanied by different degrees of AG in the body, there was a highly significant decrease of the prevalence of CP in antrum and body along with an increasing degree of AG in the body. This decrease showed a highly significant positive correlation with the acid output. On the whole, acid output correlated well with the occurrence of CP in both antrum and body. Thus the prevalence of CP was 10% in achlorhydria and rose up to 100% in cases with acid output above 30 mmol/h. The prevalence of CP did not correlate with signs of acute inflammation, but correlated significantly with those of chronic inflammation. No correlation was found in the antrum and a significant negative one in the body, between CP infestation and the extension of intestinal metaplasia. It is concluded that increased pH of gastric contents and mucus secreted by intestinalised glands may create unfavourable conditions for survival of the bacteria and might explain the decrease in the prevalence of CP in the more severe degrees of AG. The present results, however, give no definite answer to the question of the pathogenic significance of CP in the development of chronic gastritis.

Campylobacter pylori (CP) has been recently implicated in the aetiology of chronic gastritis. These studies are as a rule based upon data obtained from outpatient series, however, which consist mainly of patients with upper abdominal complaints caused by organic or functional alterations of the foregut. To avoid biases, we have studied the occurrence of CP and its relations to different forms and degrees of chronic gastritis in a sample representing the population of South Finland, paying attention to the occurrence of CP in different phenotypes of antrofundal gastritis, and to the degree of inflammatory reaction and acid secreting capacity of the mucosa.

Methods

SUBJECTS

The series were collected from a South Finnish
population and represent families consisting of first degree relatives of probands computer matched for gastric carcinoma patients. The studies were performed at the Gastroenterological Unit, Second Department of Medicine, Meilahti Hospital, Helsinki.

The original series consisted, probands included, of 454 subjects. Since the collection of the series, however, the specimens have been cut several times for new stainings. For this reason satisfactory specimens for the staining of CP were only available in 179 subjects, which accordingly formed the series proper (Table 1).

Three of the 179 subjects suffered from duodenal ulcer and one from gastric ulcer, 15 had hiatal hernia, two prepyloric erosions, four hyperplastic polyps, and five gall bladder disease. Most of the present subjects (87%) were symptom free with regard to upper abdominal complaints.

The distribution of the present cases with regard to morphology of the gastric mucosa, various gastroduodenal diseases, complaints, age, and sex distributions was roughly similar to that of the original series. The mean age and the male/female ratio in the present and the original series were 53 and 40 years and 1-1 and 1-0, respectively. The occurrence of upper abdominal complaints was similar, 13% and 14%, respectively. The prevalence of all gastritis cases was somewhat, but not strikingly, higher in the present series – for example, in the antrum 74% as compared with 68% in the original series.

All 179 subjects were examined by gastroduodenoscopy and biopsy specimens were obtained from antral (more than four specimens) and from body (more than six specimens) mucosa. On an average, 11-2 specimens were available from each subject. The specimens were fixed overnight in neutral buffered formalin, embedded in paraffin and stained by a modified Giemsa method. Table 1 shows that in 22 cases the presence of CP could be estimated in only one area of the stomach (antrum or body) because of inadequate quality of the specimens. Accordingly, data on CP were available from antrum and body for 157 subjects.

The pentagastrin test was done as described earlier in all cases. Because of the limited number of cases, men and women were not treated separately.

### Table 2  Occurrence of CP in antrum and body in the whole series

<table>
<thead>
<tr>
<th>Status of the mucosa</th>
<th>Antrum</th>
<th>Body</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Number and prevalence of CP-positive cases</td>
<td>Cases (n)</td>
</tr>
<tr>
<td>Normal</td>
<td>2 (5%)</td>
<td>42</td>
</tr>
<tr>
<td>Superficial gastritis</td>
<td>34 (71%)</td>
<td>48</td>
</tr>
<tr>
<td>Light atrophic gastritis</td>
<td>37 (65%)</td>
<td>57</td>
</tr>
<tr>
<td>Moderate atrophic gastritis</td>
<td>9 (41%)</td>
<td>22</td>
</tr>
<tr>
<td>Severe atrophic gastritis</td>
<td>2 (100%)</td>
<td>2</td>
</tr>
<tr>
<td>Cases (n)</td>
<td>84 (49%)</td>
<td>171</td>
</tr>
</tbody>
</table>

### Statistical Analysis

χ² test and Student’s t test were used in statistical calculations. Deviations were expressed as standard deviation (SD). To test simultaneously the relationship of CP infestation to the degree of gastritis and the severity of acute and chronic inflammation we used a log linear model in three way contingency table.

### Definitions

Antral and body gastritis was scored using the loss of normal elements as the main principle of classification as follows: normal mucosa (NM): no loss of glands. This grading closely resembles that of (SG): no loss of glands but round cell infiltration; light, moderate, and severe atrophic gastritis (LAG, MAG, SAG): light, moderate, and severe loss of glands. This grading resembles closely that of Schindler and Rao et al and is largely similar to those of many others (see 17). In the body the grading shows an excellent correlation with different functional parameters.

The amount of round cell infiltration was roughly graded as absent or slight, moderate, and severe. Acute inflammation was not graded because of its spotty occurrence and only distinct changes were taken into account – that is, the more confluent granulocyte infiltrations up to formation of crypt abscesses with or without micro-erosions.

Intestinal metaplasia was graded as slight when only a few metaplastic tubules were present, as moderate when a larger number of tubuli were seen, and as severe when most tubuli were of intestinal type.

### Results

**CAMPYLOBACTER PYLORI IN DIFFERENT CONDITIONS OF ANTRAL AND BODY MUCOSA**

In Table 2 the total number of adequate antral and body specimens approached 171 and 165, respect-
Campylobacter and gastritis

Table 3 Occurrence of CP in different phenotypes of gastric mucosa

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Campylobacteria in antrum and/or body</th>
<th>Cases (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal mucosa in antrum and body</td>
<td>Present in both areas</td>
<td>0</td>
</tr>
<tr>
<td>Normal mucosa in one area, gastritis in the opposite</td>
<td>Present in both areas</td>
<td>5</td>
</tr>
<tr>
<td>Superficial gastritis in antrum and body</td>
<td>Present in both areas</td>
<td>27</td>
</tr>
<tr>
<td>Superficial gastritis in one area, atrophic in the opposite</td>
<td>Present in both areas</td>
<td>25</td>
</tr>
<tr>
<td>Atrophic gastritis in antrum and body</td>
<td>Present in both areas</td>
<td>11</td>
</tr>
<tr>
<td>Cases (n)</td>
<td>Present in both areas</td>
<td>68</td>
</tr>
</tbody>
</table>

17% of the cases. Such dissimilar behaviour of the two areas was particularly evident when one or both areas were affected with AG. Only a completely normal gastric mucosa displayed a uniform behaviour: in all cases CP was lacking in antrum and body. If normal mucosa in one area was associated with gastritis in the opposite one, however, the occurrence of CP was different: about 40% of the cases CP was present in one or both stomach areas. Similarly, when affecting the whole stomach SG showed a distinctly more uniform behaviour than when associated with AG in the opposite area. The differences between diffuse SG and SG affecting only one area were, with regard to complete absence of CP, statistically highly significant (p<0.001). The presence of CP in both areas decreased further, as shown by Table 3, in cases with AG affecting the whole gastric mucosa.

The diverging behaviour of the different phenotypes of gastritis is further illustrated by Table 4, which shows the occurrence of CP in advanced stages of AG representing the more specific phenotypes A (severe AG in the body), B (AG in antrum but no AG in the body), and AB (AG in both antrum and body). In phenotype A CP was lacking in both areas in all cases, but in phenotype B CP was present with one exception in both areas or in either of them. The difference between phenotypes A and B was statistically highly significant (p<0.001).

Because of the diverging behaviour of the antral

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Because of the diverging behaviour of the antral
and body mucosae and the different phenotypes, the behaviour of the various combinations of SG treating separately the antral and body mucosa has been evaluated in Table 5. Table 5 shows that the prevalences of CP in antrum and body decrease with increasing degree of body gastritis. Thus, in addition to the decrease of CP positivity in the more severely affected body with an increasing degree of gastritis, there occurs a similar dramatic and highly significant (p<0.001) decrease in the antrum, which remains at the stage of SG. When on the other hand, the status of the body (Table 5) remains at the stage of SG and the antrum shows different degrees of gastritis, the more affected side (antrum) shows decrease in the prevalences of CP, which, however, is rather slight and significant at a 5% level only. No decrease is seen in the body, which keeps at the stage of SG. Table 5 also shows the excellent correlation between the prevalences of CP and the results of the pentagastrin test.

**Campylobacter pylori in Different Kinds and Degrees of Round Cell Infiltration**

**Signs of chronic inflammation**

In superficial gastritis, round cell infiltration was confined, according to definition, to the more superficial layers of the lamina propria. For this reason we have treated it separately from AG, in which the distribution of the cells was more diffuse. In SG the prevalence of CP-positive cases was rather similar in the different degrees of round cell infiltration. Thus, in slight accumulation of round cells in the body mucosa the prevalence of CP was 91%, in moderate 84%, and in severe 100%.

In AG (Table 6) there was a slight increase of the prevalences of CP along with increasing degrees of round cell infiltration but the differences were not statistically significant. There were no significant differences with regard to occurrence of lymph follicles in the lamina propria between the CP positive and negative cases, either.

**Table 6 Occurrence of CP and degree of chronic inflammation in cases of atrophic gastritis**

| Chronic inflammation | Antrum | | Body | |
|-----------------------|--------|----------------|------|
|                       | Number and prevalence (%) of CP-positive cases | Cases (n) | Number and prevalence (%) of CP-positive cases | Cases (n) |
| Slight or absent      | 6 (38%) | 16 | 1 (17%) | 6 |
| Moderate              | 11 (55%) | 20 | 6 (40%) | 15 |
| Severe                | 29 (62%) | 47 | 16 (42%) | 38 |

**Signs of acute inflammation**

An insignificant correlation was found between signs of acute inflammation and the presence of CP. Acute signs, such as distinct aggregation of granulocytes with or without micro-erosions or crypt abscesses, were found in 54% of the CP positive and 34% of the CP negative cases. The difference was not statistically significant.

The use of a three way contingency table and a logistic model showed a significant positive correlation with signs of chronic inflammation but not with signs of acute inflammation and the degree of chronic gastritis.

**Relationship of Intestinal Metaplasia to the Occurrence of Campylobacter pylori**

Table 7 shows that in cases of AG there is no positive correlation between the presence of CP and the extension of intestinal metaplasia in antrum and a significantly (p<0.01) negative correlation in the body. In fact, in a CP infested mucosa the bacteria were only occasionally seen in the near vicinity of intestinal tubuli.

**Relationship of Functional Parameters to the Occurrence of Campylobacter pylori**

Table 8 shows that the occurrence of CP is significantly associated with acid output as measured by pentagastrin test and expressed in terms of mmol/h. The mean acid output was highly significantly higher (p<0.001) in subjects infested by CP than in those who were free of the bacteria. Likewise, the prevalence of achlorhydria and of acid values below 10 mmol/h was significantly higher (p<0.001) in subjects without CP than in those with bacteria.

From Table 5 it appears that the acid output closely follows the behaviour of the prevalences of CP in the different phenotypes of gastritis. When SG in the antrum was associated with different degrees of gastritis in the body, there was in both antrum and body a highly significant (p<0.001) decrease of the prevalences of CP, along with a highly significant
The occurrence of CP and acid output in cases of atrophic gastritis

<table>
<thead>
<tr>
<th>Acid output mmol/l</th>
<th>Antral AG</th>
<th>Body AG</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Number and prevalence (%) of CP-positive cases</td>
<td>Cases (n)</td>
</tr>
<tr>
<td>Achlorhydria</td>
<td>1 (10%)</td>
<td>10</td>
</tr>
<tr>
<td>≤10</td>
<td>8 (53%)</td>
<td>15</td>
</tr>
<tr>
<td>11–30</td>
<td>32 (65%)</td>
<td>49</td>
</tr>
<tr>
<td>≥30</td>
<td>8 (89%)</td>
<td>9</td>
</tr>
<tr>
<td>Mean acid output in</td>
<td>Antral AG</td>
<td>Body AG</td>
</tr>
<tr>
<td>CP present</td>
<td>20·9 (11·9)</td>
<td>21·4 (11·5)</td>
</tr>
<tr>
<td>CP absent</td>
<td>7·8 (8·6)</td>
<td>4·7 (8·6)</td>
</tr>
</tbody>
</table>

Values of acid output are expressed as mean (SD).

A (p<0·001) decrease of acid output, which followed the increase of AG in the body. On the other hand, when SG in the body was associated with different degrees of gastritis in the antrum, the prevalences of CP remained high in the body but decreased slightly in the antrum concomitantly with a slight but significant (p<0·01) decrease of acid output (from 32 to 22 mmol/h). The acid output, however, still remained within limits that could be considered normal for men and women. The occurrence of CP correlated well, as expected, with fasting serum pepsinogen I concentrations (determined by M I Samloff, LA, Ca). Thus the mean SPG I was higher and the number of cases with low levels was high significantly lower in subjects with CP than in those who were free of the bacteria.

**Discussion**

The present series are selected from a sample considered representative of the population of South Finland, depending upon the availability of specimens for new cutting and staining. It is therefore possible that the present results reflect the behaviour of an over 15 year old Finnish population at large. The occurrence of CP in the present material was largely similar to that reported in the literature.5, 6, 10, 21–27 Thus most cases with superficial gastritis (SG) revealed the presence of the bacterium, while in a normal mucosa it was as a rule lacking. In addition, with an increasing degree of AG there was a significant decrease of prevalences of CP, particularly in the body, where the prevalence of CP dropped from 91% in SG to 0% in severe atrophic gastritis (AG). A decreased occurrence of CP in the more severe degrees of AG has also been noted by other authors.24–28 On the whole the prevalences of CP were very high in the present series and may have been even higher if bacterial cultures had been done but these were not because of the retrospective nature of the study.

When the occurrence of CP was considered simultaneously in antrum and body it appeared that the behaviour of one area of the stomach reflected to some degree the behaviour of the opposite area. Thus SG in antrum associated with SG in the body showed a high prevalence of CP in both areas: about 90%. On the other hand, antral SG associated with various degree of AG in the body revealed a significant decrease of CP positivity both in antrum and body with increasing severity of AG in the body. These findings can hardly be attributed to changing morphological conditions because there was a drop of CP positivity in the antrum where the status remained unchanged at the level of SG. Likewise, in severe AG of the body (phenotype A) CP was lacking, not only in the body, but also in antrum irrespective of the morphology of the antral mucosa. One the other hand, the acid output showed a highly significant decrease and the prevalences of achlorhydria and achlorhydria a highly significant increase concomitantly with a decrease of CP positivity and an increase of the degree of AG in the body.

We believe that the decrease in acid secretion creates an unfavourable milieu for the growth of CP, as also claimed by other authors.28–30 We are well aware, however, of the studies that suggest CP is resistant to achlorhydria induced by short term treatment with H2 blockers.29–30

The increased pH of gastric contents is hardly the sole explanation for the decrease of CP positivity with increasing degree of gastritis. It is possible that intestinalisation of the gastric mucosa which follows the development of AG, may play some part by producing a mucus which changes the growth milieu for the CP. Thus CP has not been found in the small intestinal mucosa and is rarely seen in an intestinalised gastric mucosa.25, 26, 30–32 In the present study also, we did not find the bacteria within the crypts of intestinal type, and there was in the body a significant negative correlation between the extent of metaplasia and the occurrence of CP. Additional factors involved in the occurrence of CP might be the immunological defence mechanism and duodenal regurgitation with or without infestation with other bacteria – that is, coliform bacteria, which might effectively compete with the CP.

The two factors possibly involved in the occurrence of CP in the gastric mucosa – that is, the pH of the gastric contents and intestinal metaplasia, may explain the variability in the CP infestation noted in
different degrees and different phenotypes of chronic gastritis.

An important aspect which still remains unexplained, is the lack of CP in a normal gastric mucosa, in which the acid output is similar to SG and intestinal metaplasia is absent. A simple explanation may be that they have not been exposed to CP infestation. This view is supported by our finding that when normal mucosa in one area is associated with CP infested gastritis in the opposite side, bacteria were also found in the normal side in a considerable proportion of cases. This explanation might not be sufficient, however, because in the present study a significant proportion of cases (about 20%) infested with CP had bacteria only in one part of the stomach. Moreover, Jones et al. found no more than expected frequency of CP in spouses of patients infested with CP.

Another explanation for the lack of CP in a normal gastric mucosa could be the absence of liability to maintain the bacteria despite an exposure to infection. This lack of liability could be in part genetically determined. We have shown earlier that genetic factors play a role in the onset and early progression of gastritis. Indeed, they seem to delay the progression of SG up to middle age, when differentiation into the more specific forms, phenotypes A, B, and AB takes place.

It seems obvious that the present results do not give a definite answer to the most crucial question as to whether CP is the cause of chronic gastritis or simply a parasite associated with a diseased gastric mucosa. Literature data suggest, however, the existence of an aetiologic relationship. A local and a systemic immunoresponse seems to accompany CP infestation: a local accumulation of granulocytes, leukopenia, crypt abscesses and micro erosions have been found in the near vicinity of the bacteria and antibodies against the bacterium have been found in the sera of patients with CP infestation. Moreover, disappearance of the bacteria and signs of acute and chronic inflammation have been reported after antibiotic therapy and treatment with bismuth compounds. We have also found a significant positive correlation between CP and signs of chronic inflammation, but were unable to show a correlation between CP and signs of acute inflammation. We believe that CP could be one of the factors which act as trigger like mechanisms at the onset of gastritis in persons genetically susceptible to the disease. Further progression of gastritis, however, particularly the development of the more specific forms (phenotypes A, B, and AB) seems independent of the presence of CP, as is shown by the significant decrease of the prevalences of CP with increasing degree of AG, and its complete absence in the phenotype A.

We could not find in the present study any evidence to support the existence of a pathogenic relationship between CP and gastric ulcer. Indeed, in the present series active gastric ulcer was found in only one case despite a very high prevalence of CP. Neither did we find any correlation between CP and upper abdominal symptoms: the present series represent a virtually symptomless population and in the few cases with non-ulcer dyspepsia the prevalence of CP did not exceed that found in the symptomless people.

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References

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