Bile reflux and intestinal metaplasia in gastric mucosa

G M Sobala, H J O’Connor, E P Dewar, R F G King, A T R Axon, M F Dixon

Abstract

Aim: To determine associations between enterogastric bile reflux and gastric mucosal pathology.

Method: A retrospective study using fasting gastric juice bile acid measurements and antral or prestomal biopsy specimens from 350 patients, 66 of whom had previously undergone surgery that either bypassed or disrupted the pyloric sphincter.

Results: Bile reflux was positively associated with reactive gastritis and negatively with Helicobacter pylori density. After stratification for previous surgery, age, and H pylori status, the histological feature most strongly associated with bile reflux was intestinal metaplasia, including all its subtypes. The prevalence of intestinal metaplasia was greatest in patients with both H pylori infection and high bile acid concentrations. Bile reflux was also positively associated with the severity of glandular atrophy, chronic inflammation, lamina propria oedema and foveolar hyperplasia.

Conclusions: Bile reflux is a cause of reactive gastritis. It modifies the features of H pylori associated chronic gastritis. The changes are not confined to patients who have had surgery to their stomachs. The positive associations with atrophy and intestinal metaplasia have implications for models of gastric carcinogenesis.

Methods

Biopsy and gastric juice bile acid results were available from 350 patients who had participated in five published studies. The 168 patients in the first three studies had been selected for their known pathology or previous gastric surgery; 60 of these had also undergone surgery that had destroyed or bypassed the pylorus. The fourth study comprised 135 consecutive patients attending an open access endoscopy clinic, excluding those with neoplastic disease. The 47 patients in the last study were attending diagnostic endoscopy lists and were selected on criteria of convenience of the timing of sample collection and of absence of previous gastric surgery. All patients had at least two biopsy specimens taken from the antrum, or in the 35 with Billroth gastrectomies, from the remnant of stomach within 5 cm of the stoma.

All patients gave informed consent. The studies were individually approved by the local research ethics committee.

Gastric juice was aspirated through a nasogastric tube in the first two studies, and at the time of endoscopy in the latter three. Bile acid concentration was then determined by the steroid dehydrogenase method in the same laboratory throughout.

The biopsy specimens were fixed in 10% formalin. After routine processing sections were taken at three levels and stained with haematoxylin and eosin. Additional sections were stained with alcian blue (pH 2.5) and periodic acid Schiff (AB/PAS) for neutral and acidic mucosubstances and by the modified Giemsa method for H pylori. If intestinal

Enterogastric reflux of bile has long been suspected to affect the gastric mucosa, but most histological studies of this topic have been performed before the discovery of Helicobacter pylori and its major role in the aetiology of chronic gastritis. Few clear conclusions have therefore been drawn, although we previously reported an association between bile reflux and a distinctive reflux (now “reactive”) gastritis, and associations between bile reflux and intestinal metaplasia have also been noted. Furthermore, bile reflux has usually been sought only in the context of the stomach after surgery, and has largely been neglected as a factor in non-surgical patients.

The aim of this study was therefore to determine the role of bile reflux in the causation of gastric mucosal pathology, specifically controlling for the presence of H pylori, in patients both with and without previous gastric surgery. A large study population was required to disentangle the effects of bile reflux from those of H pylori infection. We achieved this by performing a meta analysis on gastric biopsy results and gastric juice bile acid measurements obtained during the course of our studies over the past 10 years. In particular, we sought to examine more closely the association we have previously reported between bile reflux and reflux or reactive gastritis, to report on the relative importance of the individual components of the histological reflux score we have previously proposed, and to confirm and further explore the association between bile reflux and intestinal metaplasia. We also attempted to derive a histological index predicting the presence of abnormal bile reflux.

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metaplasia was detected in the alcian blue/PAS stained section, new sections were cut and stained with the high iron diamine/alcian blue technique to demonstrate sulphomucins.\textsuperscript{10}

The sections were examined by one pathologist (MFD) who was unaware of the bile acid results and clinical condition. The following eight features were noted and graded: chronic inflammatory cell infiltration; polymorphonuclear activity (degree of neutrophil polymorph infiltration); glandular atrophy; intestinal metaplasia; foveolar hyperplasia; oedema of the lamina propria; vasodilatation or congestion; and density of \textit{H pylori} colonization. Each feature was given a numerical score 0–3 equivalent to absent (or normal), mild, moderate and severe. The first four features and \textit{H pylori} density were graded in a manner analogous to that recommended recently in the Sydney system for the classification of gastritis.\textsuperscript{11} The other features were graded so that a “reflux score” could be calculated. The score represents the sum of the grades for foveolar hyperplasia, lamina propria oedema, and vasodilatation or congestion, less the grades for chronic inflammation and activity; plus 6. Thus the minimum score is 0 and the maximum is 15. In the past we have taken scores greater than 10 to be indicative of reflux (reactive) gastritis.

Where intestinal metaplasia was found in the stained sections, the various subtypes\textsuperscript{12} were sought by using the high iron diaminobenzidine/alcian blue sections, and the extent of each subtype (I, II, or III) was also graded on a four point scale.

The bile acid results fell in a skewed distribution and were therefore grouped into six bands before analysis: 0–00–0.05 mmol/l, 0.06–0.20 mmol/l, 0.21–0.50 mmol/l, 0.51–1.00 mmol/l, 1.01–5.00 mmol/l, and >5.00 mmol/l. Associations between bile acid concentrations and the histological scores for chronic inflammation, activity, atrophy, intestinal metaplasia, foveolar hyperplasia, lamina propria oedema and vascular congestion were sought using Cochran-Mantel-Haenszel \textit{X}² tests for non-zero correlation. The analyses were controlled by stratification for the possible confounding effects of previous surgery bypassing or disrupting the pylorus, age, and the presence of \textit{H pylori}.

Odds ratios and their 95% confidence intervals were calculated where appropriate. Differences in bile acids between different diagnostic groups were assessed by the non-parametric two-group Wilcoxon test.

A new histological reflux index was constructed by stepwise logistic regression. The dependent dichotomous variable was the presence or absence of an abnormal gastric juice bile acid concentration of >1.00 mmol/l. The independent variables were the eight graded histological features, including \textit{H pylori} density. The \textit{X}² values for variables to enter and remain in the analysis were set at 0.05. The analysis was terminated when the residual \textit{X}² test for variables not included became non-significant.

Statistical analyses were performed using the SAS package \textit{v}6.04 and EPI-INFO \textit{v}5.0 on an IBM compatible personal microcomputer.

**Results**

Three hundred and fifty patients (212 men) were studied between 1979 and 1990. The median age was 49 years, range 18 to 88.

The fasting gastric bile acid concentrations in each diagnostic group are shown in table 1.

Seven of 158 (4.4%) patients with normal endoscopies had bile acid concentrations greater than 1.00 mmol/l, confirming that this value can be considered the upper limit of the normal range.

Concentrations were substantially higher in patients who had undergone surgery that either bypassed or disrupted the pyloric sphincter. Fifteen of the 43 patients with bile concentrations above 1.00 mmol/l and 25 of the 30 patients with concentrations between 0.50 and 1.00 mmol/l had not undergone such surgery.

**HISTOLOGICAL FEATURES**

Fiftytwo patients had normal gastric mucosa, of whom 3.9% had visible \textit{H pylori} on histological examination, 219 patients had chronic gastritis, (94.0% \textit{H pylori} positive), and 79 patients had reactive gastritis (16.5% \textit{H pylori} positive).

Patients with \textit{H pylori} were less likely to have bile acid concentrations above 1.00 mmol/l (26 of 129 (20.2%) \textit{H pylori} neg-

\begin{table}
\centering
\caption{Bile reflux vs endoscopic diagnosis}
\begin{tabular}{lcccr}
\hline
Diagnosis & \textit{n} & Lower quartile & Median & Upper quartile & \textit{p} Value \\
\hline
Normal & 158 & 0.00 & 0.00 & 0.10 & - \\
Duodenal ulcer & 51 & 0.00 & 0.10 & 0.30 & ns \\
Gastric ulcer & 41 & 0.00 & 0.30 & 0.50 & ns \\
Duodenal and \textit{and} gastric ulcer & 1 & 0.00 & 0.00 & 0.10 & ns \\
Pernicious anemia & 6 & 0.00 & 0.05 & 0.30 & ns \\
Highly selective vagotomy & 23 & 0.00 & 0.00 & 0.10 & ns \\
Vagotomy and pyloroplasty & 12 & 0.00 & 0.20 & 0.95 & 0.07 \\
Vagotomy and gastrojejunostomy & 13 & 0.30 & 0.50 & 3.70 & <0.0001 \\
Billroth I & 19 & 0.00 & 0.40 & 2.70 & <0.0001 \\
Billroth II & 16 & 0.95 & 1.85 & <0.0001 \\
\hline
\end{tabular}
\end{table}

Fasting gastric juice bile acid concentrations (medians and interquartile range; mmol/l) in the 10 different diagnostic groups; \textit{p} values indicate whether results differ significantly from normal group, two tailed Wilcoxon test.
Bile reflux and intestinal metaplasia in gastric mucosa

Table 2  Bile reflux: associations with histological features

<table>
<thead>
<tr>
<th>Histological feature</th>
<th>Mantel-Haenszel $\chi^2$ statistic (df = 1)</th>
<th>p Value</th>
<th>Direction of Association</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic inflammation</td>
<td>4.93</td>
<td>p &lt; 0.05</td>
<td>Positive</td>
</tr>
<tr>
<td>Polyomorphonuclear activity</td>
<td>2.24</td>
<td>NS</td>
<td></td>
</tr>
<tr>
<td>Glomular atrophy</td>
<td>8.24</td>
<td>p &lt; 0.01</td>
<td>Positive</td>
</tr>
<tr>
<td>Intestinal metaplasia</td>
<td>17.29</td>
<td>p &lt; 0.001</td>
<td>Positive</td>
</tr>
<tr>
<td>Foveolar hyperplasia</td>
<td>4.96</td>
<td>p &lt; 0.05</td>
<td>Positive</td>
</tr>
<tr>
<td>Lamina propria oedema</td>
<td>7.78</td>
<td>p &lt; 0.01</td>
<td>Positive</td>
</tr>
<tr>
<td>Vascular congestion</td>
<td>0.73</td>
<td>NS</td>
<td></td>
</tr>
</tbody>
</table>

Cochran-Mantel-Haenszel statistics for non-zero correlations between four-point histological scores and bile acid concentrations, analyses stratified for presence of previous surgery bypassing or disrupting the pyloric sphincter, age and H pylori status.

Table 3  Bile reflux and intestinal metaplasia and reactive gastritis

<table>
<thead>
<tr>
<th>Bile acid concentrations (mmol/l)</th>
<th>n</th>
<th>(a) % Prevalence of intestinal metaplasia</th>
<th>(% Odds ratio (95% CI) for reactive gastritis</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00-0.05</td>
<td>181</td>
<td>13.8</td>
<td>3.3</td>
</tr>
<tr>
<td>0.06-0.20</td>
<td>58</td>
<td>12.1</td>
<td>8.6</td>
</tr>
<tr>
<td>0.21-0.50</td>
<td>58</td>
<td>13.2</td>
<td>2.6</td>
</tr>
<tr>
<td>0.51-1.00</td>
<td>30</td>
<td>23.3</td>
<td>6.7</td>
</tr>
<tr>
<td>1.01-5.00</td>
<td>26</td>
<td>30.8</td>
<td>11.5</td>
</tr>
<tr>
<td>&gt;5.01</td>
<td>17</td>
<td>23.5</td>
<td>11.8</td>
</tr>
</tbody>
</table>

(a) Percentage prevalences of each grade of intestinal metaplasia (focal, moderate, extensive) according to bile acid concentration
(b) Mantel-Haenszel odds ratios and 95% confidence limits for the presence of reactive gastritis according to bile acid concentration, stratified for previous surgery bypassing or disrupting the pylorus. $\chi^2$ for trend = 16.0; p < 0.001.

Table 4  Intestinal metaplasia, H pylori, and bile reflux

<table>
<thead>
<tr>
<th>H pylori present?</th>
<th>Bile reflux &gt; 1 mmol/l</th>
<th>n</th>
<th>% Prevalence intestinal metaplasia</th>
</tr>
</thead>
<tbody>
<tr>
<td>No</td>
<td>No</td>
<td>103</td>
<td>5.8</td>
</tr>
<tr>
<td>Yes</td>
<td>No</td>
<td>204</td>
<td>18.6</td>
</tr>
<tr>
<td>No</td>
<td>Yes</td>
<td>26</td>
<td>30.8</td>
</tr>
<tr>
<td>Yes</td>
<td>Yes</td>
<td>17</td>
<td>23.5</td>
</tr>
</tbody>
</table>

Prevalence of each grade of intestinal metaplasia (focal, moderate, extensive) according to presence of H pylori and of abnormal bile reflux.

Table 5  Logistic regression: predictors of bile reflux

<table>
<thead>
<tr>
<th>Variable</th>
<th>p Value</th>
<th>Odds ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lamina propria oedema</td>
<td>p &lt; 0.0001</td>
<td>4.03 (2.36-6.86)</td>
</tr>
<tr>
<td>H pylori density</td>
<td>p &lt; 0.0001</td>
<td>0.30 (0.17-0.53)</td>
</tr>
<tr>
<td>Chronic inflammation</td>
<td>p &lt; 0.001</td>
<td>2.33 (1.44-3.76)</td>
</tr>
<tr>
<td>Intestinal metaplasia</td>
<td>p &lt; 0.05</td>
<td>1.81 (1.14-2.88)</td>
</tr>
</tbody>
</table>

Results of stepwise logistic regression to predict a fasting gastric juice bile acid concentration of > 1.00 mmol/l. The odds ratios reflect the change in the likelihood of a high bile concentration associated with an single increment in the score of each histological variable.

Discussion

The limitations of this study must be considered before any conclusions are drawn. There were sampling problems. Although analysis of bile acid concentration in fasting gastric juice correlates well with other objective techniques of determining bile reflux, a single measurement does not indicate the duration and may misrepresent the degree of bile reflux experienced by an individual. Histological changes are often patchy and two mucosal biopsy specimens are not always representative. Therefore such a study will underestimate the strength of any true associations between bile reflux and gastric mucosal histology. Any apparent associations may have been due to exposure to substances other than bile in the entero gastric refluxate. Finally, this study is not an independent confirmation of our previously reported findings as there was overlap of subjects. Nevertheless, the results are important as this series is much larger than any published before in this field, with greater statistical power. It is also the

had reactive gastritis. The relation between bile acid concentrations and intestinal metaplasia is shown in table 3.

The prevalence of intestinal metaplasia was greatest in patients with both H pylori infection and a bile concentration of > 1.00 mmol/l (table 4).

Fifty patients had type I intestinal metaplasia, 53 had type II, and five had type III: 24 patients had more than one type. Focal intestinal metaplasia was usually of type II while extensive intestinal metaplasia was largely of type I. Each individual type of intestinal metaplasia occurred more often in patients with bile acid concentrations above 1.00 mmol/l than in those with lower bile acid concentrations (type I, odds ratio = 3.2, 95% confidence interval = 1.4-7.0, p < 0.01; type II, odds ratio = 3.3 (1.5-7.2), p < 0.01; type III, odds ratio = 11.4 (1.3-139), p < 0.05).

Extensive type I intestinal metaplasia in a patient with bile reflux but without previous gastric surgery is shown in fig 1.

Bile reflux and reactive gastritis

Reactive gastritis was diagnosed when the previously defined "reflux score" was greater than 10. Such a diagnosis was more likely when bile acid concentrations were high (table 3). The sensitivity of a diagnosis of reactive gastritis for a gastric juice bile acid concentration of > 1.00 mmol/l was 60%, and the specificity 83%.

New histological bile reflux index

The results of the logistic regression are shown in table 5. A new index predictive of bile reflux was constructed from the regression coefficients: (new index) = (7 × lamina propria oedema) + (3 × intestinal metaplasia) + (4 × chronic inflammation) - 6 × H pylori. A cutoff above 14 gave the highest combined sensitivity (70%) and specificity (85%) for bile reflux of > 1.00 mmol/l.

Discussion

The limitations of this study must be considered before any conclusions are drawn. There were sampling problems. Although analysis of bile acid concentration in fasting gastric juice correlates well with other objective techniques of determining bile reflux, a single measurement does not indicate the duration and may misrepresent the degree of bile reflux experienced by an individual. Histological changes are often patchy and two mucosal biopsy specimens are not always representative. Therefore such a study will underestimate the strength of any true associations between bile reflux and gastric mucosal histology. Any apparent associations may have been due to exposure to substances other than bile in the enterogastric refluxate. Finally, this study is not an independent confirmation of our previously reported findings as there was overlap of subjects. Nevertheless, the results are important as this series is much larger than any published before in this field, with greater statistical power. It is also the
first to grade biopsy specimens in a manner analogous to that of the new Sydney classification of gastritis, although the two corpus biopsy specimens required for a full classification were not available.

Several studies have attempted to correlate bile reflux with chronic inflammatory infiltration in the gastric mucosa, usually unsuccessfully. It is now known that chronic gastritis is aetiologically linked to infection with *H. pylori* and that the intensity of inflammation is related to the histological density of infection. This correlation is strong. Previous studies have failed to take it into account and any effect of bile will have been masked. Controlling for the presence of *H. pylori* infection, we did find a weak correlation between bile reflux and the density of the chronic inflammatory infiltrate. We also observed a negative association between bile and *H. pylori* density and so we conclude that scantiness of *H. pylori* with severe chronic inflammation should alert the observer to the possibility of bile reflux. The logistic regression confirmed that chronic inflammation positively, and density of *H. pylori* infection negatively, predict raised gastric juice bile concentrations. These findings lend some support to our previously stated hypothesis that bile reflux causes a gradual elimination of *H. pylori* and subsequent resolution of the chronic inflammatory response, ultimately giving rise to the distinctive reactive gastritis picture. This phenomenon has so far only been observed in the stomach after surgery. During this process intermediate stages are represented by few or no Helicobacters with persistent chronic inflammatory cell infiltration (Fig. 2).

We also found that many patients with reactive gastritis had little or no demonstrable bile reflux. This highlights the likelihood that other chemical irritants such as non-steroidal anti-inflammatory drugs and possibly alcohol cause the same histological picture. We did not have detailed drug or alcohol history on sufficient patients to control for these factors.

This study casts some doubts on the validity of the construction of the original "reflux score". No association was found between vascular congestion and bile reflux, and the association with chronic inflammation was positive rather than negative. However, the negative emphasis given to activity and chronic inflammation in the original score usefully served to bias it against *H. pylori* associated chronic gastritis: few patients with Helicobacter associated chronic gastritis also have reflux scores greater than 10. Thus reactive gastritis, so defined, is a distinct histological entity ("foveolar hyperplasia, oedema and congestion without cellular inflammation").

The new index derived by logistic regression differs in concept from the original reflux score. This is an attempt to predict the presence of bile reflux, rather than to define a new histological entity. Lamina propria oedema, intestinal metaplasia, and a disparity between *H. pylori* density and the degree of chronic inflammation were found independently to predict bile reflux.

Associations between bile reflux and intestinal metaplasia have been reported before but usually only following gastric surgery. We suggest that bile reflux is also a factor in the production of intestinal metaplasia in the unoperated stomach. The mechanism of this association is debatable. The currently dominant hypothesis is that intestinal metaplasia is a precancerous condition, occurring as a result of exposure to
Duodenogastric bile reflux

Mucosal injury, erosions

Regression

Normal gastric mucosa

Progression

Intestinal metaplasia (chiefly focal, type II)

Intestinal metaplasia (extensive, types I or III)

Figure 3 Diagram summarising hypothesis of the causation of intestinal metaplasia.

It can certainly be argued that mutagens can be contained in or formed from the entero-gastric refluxate. However, we have previously failed to find raised nitrosocompound concentrations in association with intestinal metaplasia or bile reflux. An alternative hypothesis is that intestinal metaplasia arises from divergent differentiation in regenerating epithelium following erosion or ulceration. There is some evidence to support this view. Bile in the presence of acid causes erosions in experimental animals, although this has not been shown in humans. Intestinal metaplasia seems to originate during the regenerative process in the healing of gastric erosions, common in the regenerative epithelium of gastric ulcers, and usually regresses with time. When the injury is repetitive and on a background of *H pylori* associated chronic gastritis, however, we suggest that intestinal metaplasia becomes more extensive and permanent. This hypothesis is summarised in fig 3 and is consistent with the view that intestinal metaplasia is a defence response aimed at protecting the gastric mucosa against repeated bile damage in the same way that gastric metaplasia develops in the duodenum when subjected to a high acid load, and gastric type epithelium appears in the lower oesophagus in response to acid reflux. This hypothesis of the genesis of intestinal metaplasia differs from that proposed by Correa. It merits further study.

23 Ishii T. Experimental study on the atrophic gastritis especially on the cause of the postoperative gastritis and on the histogenesis of experimentally induced atrophic gastritis. Jpn J Gastroenterol 1966;63:1323-37.