Bile reflux gastritis and intestinal metaplasia at the cardia

M F Dixon, N P Mapstone, P M Neville, P Moayyedi, A T R Axon

Background and aims: Intestinal metaplasia (IM) at the cardia is likely to be a precursor of cardia cancer. Previous work has shown that it is associated with chronic inflammation attributable to either gastro-oesophageal reflux disease (GORD) or Helicobacter pylori infection. An alternative aetiopathological factor is bile reflux. Duodenogastric reflux brings about histological changes in the gastric mucosa that can be graded and used to calculate a bile reflux index (BRI). We used the BRI to assess whether reflux of bile plays a part in the development of cardia IM.

Methods: Histological changes in simultaneous gastric antrum and cardia biopsies from 267 dyspeptic patients were independently graded by two pathologists. The association between cardia IM and age, sex, clinical group, H pylori status, increased BRI (>14), and inflammation at the cardia were evaluated using logistic regression.

Results: A total of 226 patients had adequate cardia and antral biopsies; 149 had GORD and 77 had non-ulcer dyspepsia. Cardia IM was present in 66 (29%) patients, of whom 28 (42%) had complete IM. Increasing age, male sex, chronic inflammation, and a high BRI emerged as significant independent associations with cardia IM. Clinical group and H pylori status were not independent risk factors.

Conclusions: Histological evidence of bile reflux into the stomach is associated with cardia IM. This could have an important bearing on carcinogenesis at this site.

The cardia is an area of controversy. Conventionally this “small and ill defined” region is described as a zone 0.5–3 cm long “just distal to the lower end of the oesophagus” where it is interposed between oxyntic mucosa and squamous epithelium. However, its existence as a normal anatomical structure has been challenged. Chandrasoma and colleagues claim that the cardia is absent in truly normal individuals so that oxyntic mucosa abuts directly onto squamous epithelium. They argue that cardia-type mucosa is always acquired and is a consequence of gastro-oesophageal reflux disease (GORD). Cardia mucosa closely resembles the gastric antrum and it is suggested that the “acquired” cardia is an example of gastric metaplasia in response to acid reflux.

Intestinal metaplasia (IM) at the cardia has attracted further controversy. Its very frequency (15–22% of dyspeptic subjects with an endoscopically normal lower oesophagus++) argues against it being closely associated with cardia cancer yet by analogy with the stomach, cardia IM is considered to be a premalignant condition. While gastric metaplasia can be readily understood as an adaptive response of squamous epithelium to acid injury, it seems likely that intestinal metaplasia is triggered by other factors. Previous work has implicated gastric Helicobacter pylori infection and the associated “carditis” but other investigators have dismissed this in favour of GORD. The latter conclusion does not necessarily implicate acid as the sole causative agent. There is increasing evidence that GORD is accompanied by duodenogastric reflux so that the oesophageal refluxate can contain bile and other components of pancreaticoduodenal secretion. It would be much more plausible for the change from gastric to intestinal phenotype at the cardia to be related to bile than to acid reflux alone.

To investigate the role of bile reflux in the causation of cardia IM, we used a histological indicator of bile injury to the antrum—the bile reflux index (BRI)—along with the extent of cardia inflammation and H pylori status to search for possible aetiopathological associations. Additionally, as the various subtypes of IM at the cardia are likely to carry different risks for progression to cancer, we sought to relate these to the same factors.

Abbreviations: IM, intestinal metaplasia; GORD, gastro-oesophageal reflux disease; NUD, non-ulcer dyspepsia; BRI, bile reflux index; AB/PAS, alcian blue/periodic acid-Schiff.

Patients and Methods

Over a two year period, the Department of Pathology at Leeds General Infirmary received simultaneous biopsies of the gastro-oesophageal junction and gastric mucosa from 267 patients. The majority (192 cases) consisted of patients referred directly by their general practitioner to an open access dyspepsia clinic. Suitable patients were seen as part of an assessment procedure prior to possible recruitment into a study evaluating the symptomatic response of H pylori negative dyspepsia to pantoprazole. Patients aged 18–80 years were considered. Those taking non-steroidal anti-inflammatory drugs or proton pump inhibitors less than four weeks prior to endoscopy, those who had undergone prior gastric or oesophageal surgery, and patients who were known to have been H pylori positive in the past or had received H pylori eradication therapy were excluded. Prior to endoscopy, patients were assessed by an experienced gastroenterologist (PMN) and categorised on clinical grounds as having either non-ulcer dyspepsia (NUD) or GORD. The diagnosis of NUD was based on a clinical presentation of predominant epigastric pain or abdominal symptomatology (for example, bloating or nausea) with clinically insignificant or absent heartburn. The diagnosis of GORD was based on a clinical presentation of significant heartburn or acid reflux with or without epigastric pain. At endoscopy, four quadrantic biopsies were taken from immediately below the squamocolumnar junction (Z line), two biopsies from the gastric body (one on the lesser curve and one on the greater curve), one from the incisura angulosa, and two from the prepyloric antrum. The local ethics committee gave approval for the biopsies with prior consent.

Classification of patients into NUD and GORD was modified after endoscopy. Patients with active ulcer disease, macroscopic evidence of Barrett’s oesophagus, severe (≥grade 3) oesophagitis, and neoplasia were excluded. Those patients...
who had been classified as NUD on clinical grounds who on endoscopy were found to have Savary grade 2–3 oesophagitis were reclassified as GORD on the grounds that significant endoscopic oesophagitis is a more powerful predictor of GORD than clinical assessment.

The remaining 75 patients had cardia biopsies taken for some clinical indication, for example as an adjunct to the histological diagnosis of GORD or because significant endoscopic oesophagitis was a more powerful predictor of GORD than clinical assessment. Histological assessment
Biopsies were fixed in 10% formalin, processed, sectioned at three levels, and stained by haematoxylin-eosin. Additional sections taken at the second level from all biopsies were stained with alcian blue/periodic acid-Schiff (AB/PAS) to demonstrate IM and by the modified Giemsa stain for *H pylori*. The cardia and gastric biopsies were assessed separately by two pathologists (NPM and MFD, respectively) who were “blind” to each other’s findings and to the clinical details. The “cardia” biopsies were categorised as either squamous, oxyntic, oxyntocardiac transitional zone, or cardia type. We accepted as cardia those biopsies composed of gastric-type foveolar and surface epithelium overlying either antral-like mucous glands devoid of parietal cells or mucous glands with occasional parietal cells. The transitional mucosa was characterised by an admixture of mucous and oxyntic glands. A case was only included when there was at least one cardia biopsy and at least one antral biopsy adequate for assessment.

The presence of acute inflammation (that is, neutrophil polymorph infiltration), chronic inflammation (CI)—lymphocytes and plasma cells—and IM in the cardia and antrum were graded on a 0–3 scale (absent, mild, moderate, and marked), while additionally in the antrum, lamina propria oedema (Oed) and *H pylori* colonisation density (*Hp*) were graded. The grading approach was analogous to that described in the updated Sydney system although small numbers of mononuclear cells at the cardia were considered “normal” (see below). The Giemsa stained corpus biopsies were also examined before a case was declared *H pylori* negative. IM was defined by the finding of fully formed goblet cells in the AB/PAS stain. When detected, a new section was cut and stained with Gomori’s aldehyde fuchsin technique to distinguish between type II and III (sulphomucin positive) IM.

![Figure 1](image_url)  
**Figure 1** (A) Cardia biopsy (squamocolumnar junction) stained with alcian blue/periodic acid-Schiff for mucus showing widespread incomplete-type intestinal metaplasia. (B) Antral biopsy from the same patient (haematoxylin-eosin) showing widespread intestinal metaplasia (complete type) (grade=3), mild lamina propria oedema (grade=1), and mild chronic inflammation (grade=1). Helicobacters were not present in this case. Based on these findings the bile reflux index was 20.

Table 1  
Grades of acute and chronic cardia inflammation according to clinical group, *Helicobacter pylori* (*Hp*) status, and bile reflux index (BRI). Grades 1, 2, and 3 are equivalent to a mild, moderate, and marked increase. For acute inflammation grade 0=absence of polymorphs; for chronic inflammation 0=“normal” numbers of lymphocytes and plasma cells (see text)

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>NUD</td>
<td>77</td>
<td>59</td>
<td>14</td>
<td>4</td>
<td>0</td>
<td>35</td>
<td>26</td>
<td>14</td>
<td>2</td>
</tr>
<tr>
<td>GORD</td>
<td>149</td>
<td>122</td>
<td>20</td>
<td>7</td>
<td>0</td>
<td>83</td>
<td>44</td>
<td>20</td>
<td>0</td>
</tr>
<tr>
<td><em>Hp</em>+</td>
<td>44</td>
<td>10</td>
<td>24</td>
<td>10</td>
<td>0</td>
<td>2</td>
<td>15</td>
<td>25</td>
<td>2</td>
</tr>
<tr>
<td><em>Hp</em>−</td>
<td>182</td>
<td>171</td>
<td>10</td>
<td>1</td>
<td>0</td>
<td>118</td>
<td>55</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>BRI&lt;14</td>
<td>188</td>
<td>151</td>
<td>29</td>
<td>8</td>
<td>0</td>
<td>105</td>
<td>57</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>BRI&gt;14</td>
<td>38</td>
<td>30</td>
<td>5</td>
<td>3</td>
<td>0</td>
<td>16</td>
<td>13</td>
<td>9</td>
<td>1</td>
</tr>
</tbody>
</table>

GORD, gastro-oesophageal reflux disease; NUD, non-ulcer dyspepsia; 
Acute and chronic inflammation of grades 2 and 3 were combined for all analyses. 
NUD versus GORD: acute, Pearson $\chi^2=0.97$, df=2, p=0.62; chronic, Pearson $\chi^2=3.08$, df=2, p=0.01. 
*Hp*+ versus *Hp*− acute, Pearson $\chi^2=114.92$, df=2, p<0.0001; chronic, Pearson $\chi^2=94.95$, df=2, p<0.0001. 
BRI<14 versus BRI>14 acute, Pearson $\chi^2=0.98$, df=2, p=0.61; chronic, Pearson $\chi^2=4.72$, df=2, p=0.09.
Histological findings in the antrum were used to calculate the BRI. BRI was originally derived by stepwise logistic regression analysis of the histological grades found in antral biopsies from 350 subjects in whom gastric juice bile acid levels had been measured. Following analysis, an index comprising $(7 \times \text{Oed}) + (3 \times \text{IM}) + (4 \times \text{CI}) - (6 \times \text{Hp})$ gave the best prediction of a raised gastric juice bile acid concentration. An index above 14 had a sensitivity of 70% and a specificity of 85% for a bile acid level >1.00 mmol/l (the upper limit of “physiological” reflux).

**Statistical analysis**

The effect of clinical group (GORD or NUD), *H pylori* status, and increased bile reflux on acute and chronic inflammation at the cardia was assessed using the Pearson $\chi^2$ test. The association between IM at the cardia and sex, *H pylori* status, increased bile reflux, clinical group, and the presence of acute and chronic inflammation at the cardia were evaluated using Fisher's exact test. The association between these factors and type of IM was assessed using Fisher's exact test while the association between age and IM was assessed using the Student's independent $t$ test. Independent risk factors for IM at the cardia were evaluated using logistic regression with IM at the cardia as the independent variable. The proportion of the variance explained by the logistic regression model was calculated using Nagelkerke $r^2$. A $p$ value <0.05 was considered statistically significant for all analyses, and calculations were performed using SPSS version 9.0 (SPSS Inc. Chicago, Illinois, USA).

**RESULTS**

In 39 of 267 patients biopsied, no cardia-type mucosa was identified, and in a further two cases the antral biopsies were inadequate or missing. Thus there were 226 patients with cardia mucosa available (range 1–4 biopsies, median 2, mean 1.98) and at least one adequate biopsy of antral mucosa (see fig 1). Of these, 149 patients were considered to have GORD and 77 had NUD. The gastric biopsy findings from 47 of these NUD patients who were *H pylori* negative were used in the control group of a previous study on bile reflux gastritis. With regard to *H pylori* status, 182 of 226 patients were negative and 44 (19.5%) were positive. There were no cases where the corpus biopsies were positive and the antral biopsies were negative for helicobacters. Concerning the BRI, 188/226 patients had a BRI $\leq 14$ while 38 had a BRI $>14$. While inflammation at the cardia showed no significant differences between subjects with GORD or NUD, or with an elevated BRI, *H pylori* positive cases were highly significantly different from negative patients for both acute and chronic inflammation (table 1).

IM was found at the cardia in 66 patients (29%); in 28 the cardia IM was predominantly (if not exclusively) type I, in 25 it was type II, and in 13 it was type III.

Unadjusted analyses revealed that *H pylori* infection, BRI $>14$, and acute and chronic inflammation at the cardia were associated with IM (table 2). There was a trend for male sex to be associated with cardia IM but this did not reach statistical significance, while there was no difference between GORD and NUD patients (table 2). Patients with IM at the cardia were older (median age 57 years) than patients without IM (median 47 years) and the difference in mean age was 9.7 years (95% confidence interval (CI) 5.6–13.7). *H pylori* infection and BRI $>14$ were also associated with increasing age and it is possible that these factors are acting as markers for older patients. To evaluate independent risk factors for IM at the cardia, sex, clinical group, BRI $\leq 14$ or $>14$, *H pylori* status, and chronic inflammation at the cardia were entered into a logistic regression model. Acute inflammation was omitted from the model as this exhibited strong colinearity with chronic inflammation. Age, sex, BRI $>14$, and chronic inflammation remained independent risk factors for cardia IM but *H pylori* status became non-significant (table 2). The model explained 25.5% of the variance of the data. *H pylori* status was strongly associated with chronic inflammation and the lack of a statistically significant relationship between *H pylori* infection and cardia IM could be due to colinearity. This is not the case however as *H pylori* was not associated with cardia IM in a logistic model that only included age (odds ratio 1.7; 95% CI 0.8–3.4; $p=0.17$).

The influence of *H pylori* status, BRI $>14$, sex, clinical group, and chronic and acute inflammation on the type of cardia IM

![Table 2](http://gutjnl.com)
DISCUSSION

The most striking finding in this study was that a histological index of bile reflux into the stomach was significantly associated with the presence of IM at the cardia. This is in keeping with a previous study in which we found that patients with Barrett’s oesophagus (that is, IM in the oesophagus) had significantly higher BRI values than those with uncomplicated reflux oesophagitis or simple NUD. Interestingly, we found no statistically independent association between cardia IM and symptoms of GORD or with H pylori infection. While it is evident that a lack of statistical significance does not equate to absence of effect, our finding that increasing age, male sex, and chronic inflammation at the cardia were significantly greater in patients with cardia IM compared with those with NUD, negative status, absence of acute inflammation at the cardia, and female sex. However, these factors did not reach formal statistical significance and the numbers of cases are small, making the data difficult to interpret (table 3). Antral IM was found in 26 of 66 patients with cardia IM (39%) but there were an additional 16 patients with isolated antral IM. While the score for antral IM contributed to BRI, its presence did not predict high values. Equal numbers of patients with metaplasia in the stomach (that is, 21/42) had a BRI above and below 14.

Table 3 Association between incomplete intestinal metaplasia (types II and III IM), sex, bile reflux index (BRI), Helicobacter pylori status, clinical group, and chronic and acute inflammation at the cardia

<table>
<thead>
<tr>
<th>Variable</th>
<th>Total No</th>
<th>% with incomplete IM</th>
<th>Unadjusted odds ratio</th>
<th>95% CI</th>
<th>p Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>32</td>
<td>68.8</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>34</td>
<td>47.1</td>
<td>0.40</td>
<td>0.15–1.11</td>
<td>0.087</td>
</tr>
<tr>
<td>BRI</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤14</td>
<td>45</td>
<td>51.1</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;14</td>
<td>21</td>
<td>71.4</td>
<td>2.39</td>
<td>0.79–7.28</td>
<td>0.18</td>
</tr>
<tr>
<td>H pylori</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>47</td>
<td>63.8</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>19</td>
<td>42.1</td>
<td>0.41</td>
<td>0.14–1.22</td>
<td>0.17</td>
</tr>
<tr>
<td>Group</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GORD</td>
<td>42</td>
<td>57.1</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NUD</td>
<td>24</td>
<td>58.3</td>
<td>1.05</td>
<td>0.38–2.90</td>
<td>1.00</td>
</tr>
<tr>
<td>Cardia CI</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>22</td>
<td>63.6</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>44</td>
<td>54.5</td>
<td>0.69</td>
<td>0.24–1.96</td>
<td>0.60</td>
</tr>
<tr>
<td>Cardia AI</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>44</td>
<td>65.9</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>22</td>
<td>40.9</td>
<td>0.36</td>
<td>0.12–1.03</td>
<td>0.067</td>
</tr>
</tbody>
</table>

*Fisher’s exact test

GORD, gastro-oesophageal reflux disease; NUD, non-ulcer dyspepsia; CI, chronic inflammation; AI, acute inflammation.

With regard to the subtypes of IM found in cardia mucosa, we found a trend towards an association between incomplete IM and a high BRI, H pylori negativity, absence of acute inflammation, and female sex but the groups were too small to yield significant associations. Previous work has examined IM subtypes at the cardia and their disease associations and found significant differences. For instance, Voutilainen and colleagues examined 235 patients with cardia IM and found that complete IM was associated with a pattern of gastritis characteristic of longstanding H pylori infection while incomplete IM was associated with erosive oesophagitis, the lesion which predisposes to Barrett’s oesophagus.

To our knowledge this is the first study that has demonstrated, albeit by indirect means, a link between bile reflux and cardia IM. This has potentially important implications for the development of cancer at the cardia. The connection between exposure to bile acids or their breakdown products and carcinogenesis at other sites is well known. The development of duodenal adenomas and carcinomas in patients with familial adenomatous polyposis and the predilection for adenocarcinoma to arise at the ampulla of Vater are thought to be bile related. The appearance of lymphocytic infiltration, especially close to the squamocolumnar junction, is likely to be “physiological”. As Weinstein puts it “these changes (carditis) could represent the consequence of gastric contents lapping at the shores of the oesophagus in health” (italics added). If it is acknowledged that the cardia has a “normal” complement of mononuclear cells, then the diagnosis of chronic inflammation requires a higher threshold and this has been our approach. Some authors avoid the difficulties in diagnosing chronic inflammation at this site by defining “carditis” as infiltration by polymorphs. Using this criterion, equivalent to active chronic inflammation, we found “carditis” in only 45/226 cases (19.9%). Interestingly this is very close to the 18.4% prevalence of polymorph infiltration found by Der and colleagues yet these authors diagnosed chronic inflammation in all cases where cardia mucosa was present. In our series, acute and chronic inflammation at the cardia were significantly greater in H pylori infected compared with non-infected patients but inflammation bore no significant association to clinical group. These findings run contrary to recent studies that emphasise the role of GORD but are entirely analogous to those reported by others. With regard to the subtypes of IM found in cardia mucosa, we found a trend towards an association between incomplete IM and a high BRI, H pylori negativity, absence of acute inflammation, and female sex but the groups were too small to yield significant associations. Previous work has examined IM subtypes at the cardia and their disease associations and found significant differences. For instance, Voutilainen and colleagues examined 235 patients with cardia IM and found that complete IM was associated with a pattern of gastritis characteristic of longstanding H pylori infection while incomplete IM was associated with erosive oesophagitis, the lesion which predisposes to Barrett’s oesophagus.

dysplasia and adenocarcinoma in gastric remnants following partial gastrectomy, and the development of carcinoma in the stomach of experimental animals subjected to enterogastric reflux, have also been attributed to exposure to bile and/or duodenopancreatic juice. Finally, there is evidence that faecal bile acids in conjunction with dehydrogenating bacteria are implicated in the development of sporadic colorectal cancer. Thus bile acids have the potential not only to bring about IM at the cardia but also to generate the carcinogens which act on this altered mucosa to produce neoplasia. Changes in pH and the composition of the refluxate after regurgitation through the stomach could alter its carcinogenic potential and render it more damaging to the oesophagus. Given the intrinsic instability of junctional sites and their predisposition to neoplasia, it is certainly plausible that exposure of the cardia to duodenogastric reflux could contribute to the development of both IM and adenocarcinoma at this site. Nevertheless, our conclusions must remain speculative until confirmed by direct measurements of intragastric bile content.

ACKNOWLEDGEMENTS

We are grateful to Knoll Ltd for an unrestricted grant in support of Dr Neville during the conduct of this study. Thanks are due to Stephen Ryan and Peter Jackson for technical assistance, Tim Branch for slide retrieval, Paul Moayyedi is funded by a UK MRC Health Services Research Training Fellowship.

Authors' affiliations
M F Dixon, N P Mapstone, Academic Unit of Pathology, University of Leeds, UK
P M Neville, P Moayyedi, A T R Axon, Centre for Digestive Diseases, General Infirmary at Leeds, Leeds, UK

REFERENCES
Bile reflux gastritis and intestinal metaplasia at the cardia

M F Dixon, N P Mapstone, P M Neville, P Moayyedi and A T R Axon

*Gut* 2002 51: 351-355
doi: 10.1136/gut.51.3.351

Updated information and services can be found at:
http://gut.bmj.com/content/51/3/351

These include:

**References**
This article cites 26 articles, 7 of which you can access for free at:
http://gut.bmj.com/content/51/3/351#ref-list-1

**Email alerting service**
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Topic Collections**
Articles on similar topics can be found in the following collections

- Pancreas and biliary tract (1949)
- Stomach and duodenum (1689)
- Campylobacter, Salmonella, Shigella, Escherichia coli (242)
- Dyspepsia (297)
- Gastro-oesophageal reflux (351)
- Helicobacter pylori (218)

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/