Bile reflux gastritis and Barrett’s oesophagus: further evidence of a role for duodenogastro-oesophageal reflux?

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Abstract

Background—There is increasing evidence that reflux of bile plays a part in the pathogenesis of Barrett’s oesophagus. Bile injury to the gastric mucosa results in a “chemical” gastritis in which oedema and intestinal metaplasia are prominent.

Aim—To determine if patients with Barrett’s oesophagus have more bile related changes in antral mucosa than patients with uncomplicated gastro-oesophageal reflux disease (GORD) or non-ulcer dyspepsia (NUD).

Patients and methods—Patients were identified by a retrospective search of pathology records and those with a clinically confirmed diagnosis of either Barrett’s oesophagus or reflux oesophagitis who had oesophageal and gastric biopsies taken at the same endoscopy and had no evidence of Helicobacter pylori infection entered the study. Control biopsies were taken from H pylori negative NUD patients. Antral biopsies were examined “blind” to clinical group and graded for a series of histological features from which the “reflux gastritis score” (RGS) and “bile reflux index” (BRI) could be calculated. The reproducibility of these histological scores was tested by a second pathologist.

Results—There were 100 patients with Barrett’s, 61 with GORD, and 50 with NUD. The RGSs did not differ between groups. BRI values in the Barrett’s group were significantly higher than those in GORD subjects (p=0.014) which in turn were higher than those in NUD patients (p=0.037). Similarly, the frequency of high BRI values (>14) was significantly greater in the Barrett’s group (29/100; 29%) than in the GORD (9/61; 14.8%) or NUD (4/50; 8%) group. However, agreement on BRI values was “poor”, indicating limited applicability of this approach.

Conclusion—Patients with Barrett’s oesophagus have more evidence of bile related gastritis than subjects with uncomplicated GORD or NUD. The presence of bile in the refluxate could be a factor in both the development of “specialised” intestinal metaplasia and malignancy in the oesophagus.

Barrett’s oesophagus is widely considered to be a consequence of longstanding acid induced injury commencing as an erosive oesophagitis and progressing over years to columnar and intestinal metaplasia of the squamous epithelium. The process of metaplasia represents a non-neoplastic change in cellular phenotype which is thought to be a response to a sustained adverse environment.1 The change may be a consequence of somatic mutation in the epithelial stem cells or an epigenetic event whereby divergent differentiation in progeny cells produces the altered phenotype. Whatever the precise mechanism, the resulting cell lineage has a survival advantage over “native” epithelium so that selection pressures promote the emergence and dominance of the metaplastic population. Examples in the gastrointestinal tract include gastric metaplasia in the duodenum in response to acid injury,1 intestinal metaplasia in the stomach in longstanding helicobacter pylori infection,4,5 and colonic metaplasia in ileal pouches in response to acquisition of a faecal-type flora.6 Following this line of reasoning, the appearance of gastric metaplasia in the lower oesophagus—that is, proximal extension of cardia-type mucosa—is a logical adaptive response to sustained acid reflux. However, acid injury alone would not be expected to induce intestinal metaplasia, the defining feature of Barrett’s oesophagus.

Recent evidence points to a role for bile reflux in Barrett’s oesophagus.7 The presence of bile in the oesophagus obviously necessitates duodenogastro-oesophageal reflux (DGOR) with passage of initially alkaline duodenal contents through the stomach.8 We have previously shown that bile reflux produces consistent histological changes in the gastric mucosa,9 and that an index based on these features was predictive of elevated bile acids in the refluxate.10 More interestingly, we found a stronger association between raised intragastric bile acid levels and incomplete (type III) intestinal metaplasia than with the complete form (type I).10 Incomplete intestinal metaplasia is the phenotype found in Barrett’s oesophagus. These findings suggested to us that bile, as part of DGOR, could have a role in producing intestinal metaplasia in the oesophagus. If this is so, antral histology is more likely to reflect bile related changes in patients with Barrett’s oesophagus.

Abbreviations used in this paper: DGOR, duodenogastro-oesophageal reflux; GORD, gastro-oesophageal reflux disease; NUD, non-ulcer dyspepsia; BRI, bile reflux index; RGS, reflux gastritis score; NSAIDs, non-steroidal anti-inflammatory drugs.
Methods and patients
REFLUX GASTRITIS SCORE AND BILE REFLUX INDEX
In our original study on reflux gastritis, we described a distinctive histological picture comprising foveolar hyperplasia (FH), lamina propria vasodilatation and congestion (VC), oedema (Oed), and a paucity of acute (AI) and chronic (CI) inflammatory cells. Using grades (0–3) for each of these features a “reflux gastritis score” (RGS) was devised which comprised (FH+VC+Oed)−(AI+CI)+6; the addition of “6” is to compensate for the inflammatory cell paucity. Thus cases scoring 3 for both AI and CI will effectively have nothing added whereas biopsies devoid of inflammatory cells have 6 points added. The RGS ranges from 0 to 15 and in the past we have taken scores >10 to be indicative of “reflux gastritis”. A higher frequency of such scores was found in subjects with increased intragastric bile acids. However, we subsequently appreciated that many subjects with a high RGS did not have evidence of bile reflux; a proportion were taking non-steroidal anti-inflammatory drugs (NSAIDs), some had high alcohol consumption, and in many instances no cause was identified. Therefore we sought a more accurate predictor of bile reflux.

The bile reflux index (BRI) was derived by stepwise logistic regression analysis of the histological grades used in the RGS together with intestinal metaplasia (IM) and H pylori colonisation (Hp) found in gastric biopsies from 350 subjects in whom gastric juice bile acid levels had been measured. Following analysis, an index comprising (7×Oed)+(3×IM)+(4×CI)−(6×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). As we wished to investigate only H pylori negative subjects in this study (see below), we used the original data to examine the predictive value of the BRI in negative cases. In the 128 H pylori negative cases, a BRI >14 had a sensitivity of 79.2% and a specificity of 82.7% for a bile acid concentration >1 mmol/l. In so far as H pylori colonisation has a negative effect on BRI, the values obtained in a particular clinical group will be affected by the prevalence of infection in that group. Additionally, the degree of H pylori colonisation and chronic inflammation in the antrum in positive cases will be affected by concurrent treatment with proton pump inhibitors which will differ between groups.

Finally, recognition of lamina propria oedema and other histological features of reflux gastritis are more difficult in H pylori positive cases. For these reasons, we have chosen to compare only H pylori negative subjects.

PATIENT GROUPS
A computer search of pathology records from August 1998 to July 1999 revealed 396 patients whose oesophageal biopsies carried a diagnosis of reflux (erosive) oesophagitis and 474 patients with Barrett’s oesophagus. Of these, a total of 227 had gastric biopsies taken at the same endoscopy. The clinical details of each patient were checked and any not fulfilling the correct clinical and/or endoscopic diagnosis were eliminated, as were subjects with previous gastric surgery. At the same time, on the basis of clinical, endoscopic, and histological findings, patients were allocated to either mild/moderate or severe GORD, and either long or short (<3 cm) segment Barrett’s oesophagus. There were 79 patients with confirmed GORD and 124 patients with Barrett’s oesophagus with adequate gastric biopsies of whom 61 and 100 patients, respectively, were found to be H pylori negative. Gastric biopsies taken from 50 patients presenting with dyspepsia and having (a) negative tests (biopsy urease and histology) for H pylori infection, (b) a clinical history suggestive of non-ulcer dyspepsia (NUD), (c) predominant epigastric pain, and no heartburn or reflux, and (c) no endoscopic evidence of peptic ulceration or Barrett’s oesophagus, were used as controls. The archived slides of the gastric biopsies stained by haematoxylin-eosin and alcin blue/periodic acid Schiff were assessed (by MFD) “blind” to the clinical group. Foveolar hyperplasia, lamina propria oedema, vascular congestion, acute and chronic inflammation, and intestinal metaplasia in antral mucosa were graded on a 0–3 scale (absent, mild, moderate, or marked) and RGS and BRI were calculated for each case (see fig 1). While it was appreciated that NSAIDs and other factors affect RGS, and
There was no significant difference in the distribution of RGSs between groups (\( \chi^2 = 0.05, \text{df} = 2, p=0.98 \)). When the BRIs within the clinical subgroups were compared (table 3), there was no significant difference between GORD patients with either mild or moderate reflux and those with severe reflux, or between Barrett’s patients with short or long segment involvement.

Patients with a BRI >14 were significantly older than patients with a BRI \( \leq 14 \) (mean age difference 8.6; 95% confidence intervals 3.9–13.3 years). The Barrett’s group were also significantly older than the other groups and it is possible that age could be a confounding factor in the association between BRI and Barrett’s oesophagus. To explore this we entered age, sex, foveolar hyperplasia score, vascular congestion score, and clinical group into a linear multiple regression model with BRI as the dependent variable. Clinical group remained independently associated with bile reflux with a 1.4 (95% confidence intervals 0.5–2.3) unit increase in BRI in the GORD group compared with the NUD group and in Barrett’s group compared with the GORD group (p=0.004).

With regard to reproducibility of histological grading, interobserver agreement on the three levels of BRI was poor (38%), with a kappa value of only 0.051. This was largely a consequence of haphazard disagreement\(^1\) on the grade of lamina propria oedema which is a much more subjective feature than chronic inflammation and intestinal metaplasia. Because of the high weighting given to this feature, a one grade difference in oedema translated into a 7 point difference in BRI. When the agreement on two categories of BRI, “low/medium” versus “high” (>14) was examined, overall agreement was improved (78%) but because of high chance agreement the
A combination of acid and bile reflux offers a credible explanation for the histological components of Barrett’s oesophagus. Sustained acid injury to the lower oesophagus would be expected to lead to gastric metaplasia. Gastric metaplasia is apparent as upward extension of the cardia zone and is an acknowledged feature of GORD. Indeed, it may be a general phenomenon that antedates the development of so-called “specialised” Barrett’s metaplasia. The latter consists of goblet cells and intervening mucus rich vacuolated cells which secrete variable amounts of sulphomucin and express a mixture of gastric and intestinal mucin genes. These morphological and histochemical characteristics and the pattern of mucin gene expression are identical to the incomplete type intestinal metaplasia (type III) found in gastric mucosa. Our previous demonstration of a relationship between intragastric bile reflux and type III intestinal metaplasia in gastric mucosa, mainly in the post-surgical stomach, is therefore entirely analogous to a role for acid and bile in the pathogenesis of Barrett’s metaplasia. The absence of type III metaplasia from the antrum of many subjects with Barrett’s oesophagus may be explicable in terms of changes in pH and the character of the refluxate after passage through the intact stomach.

The increased risk of adenocarcinoma in Barrett’s oesophagus is also understandable in terms of progression from a “specialised” metaplasia equivalent to type III intestinal metaplasia under the influence of bile reflux. Type III (incomplete) intestinal metaplasia, but not type I (complete metaplasia), in the stomach carries a much greater risk of gastric cancer development. Furthermore, bile acids, particularly in an alkaline milieu, could be the source of carcinogens which act on the metaplastic mucosa to produce neoplasia. While an association between Barrett’s oesophagus and bile reflux (as indicated by this and other studies) does not necessarily point to causality, it is certainly plausible that the presence of bile derivatives and metaplasia in the oesophagus equivalent to type III intestinal metaplasia are linked to cancer development. This conclusion has potentially important implications for the management of patients with intestinal metaplasia at the cardia. There are clear differences in demographic features and disease associations between subjects with complete and incomplete intestinal metaplasia at the cardia. Extrapolating from our knowledge of metaplasia in the stomach, one can speculate that complete intestinal metaplasia at the cardia has little or no malignant potential while type III intestinal metaplasia is likely to be part of Barrett’s spectrum and will carry an increased risk of adenocarcinoma. If a direct role for refluxed bile derivatives in carcinogenesis becomes accepted, therapy aimed principally at acid reduction cannot be expected to be cancer protective. Barrett’s oesophagus. It is true that proton pump inhibitors diminish bile reflux into the oesophagus, possibly by reducing overall
gastric juice volume, but the concomitant generation of unconjugated bile acids in a relatively alkaline milieu may exacerbate oesophageal mucosal damage. The most effective and physiological way to diminish DGER is by antireflux surgery.

One conclusion from our study could be that those subjects with GORD who have clear evidence of bile related damage to the antral mucosa—that is, those with a high BRI—may be at increased risk of Barrett’s oesophagus. Equally, one could predict that patients with Barrett’s oesophagus and a high antral BRI are at increased risk of dysplasia and adenocarcinoma. However, there are important caveats to such speculation. The association between BRI and Barrett’s oesophagus may not be causal and could be explained by other confounding factors not evaluated in this study. The BRI was derived from previous observations of one observer (MFD) but was poorly reproduced by an experienced second observer. Thus while we believe that the finding of significant differences in bile related changes in antral histology between NUD, GORD, and Barrett’s subjects is valid in the particular context of a bile reflux index based on the same observer’s experience, our findings require confirmation by comparing histological grading to direct measurements of intragastric bile reflux in similar clinical groups.

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