Towards at least one more
Canadian consensus

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Achieving a Canadian consensus represents a challenge of the minds, fruit for the lion-hearted, not for the frivolous or the faint. The future rests on a 55/45 verdict. What a remarkable accomplishment that a 40/0 accord could be accomplished on a practical approach to the management of patients with gastroesophageal reflux disease (GERD – page 277). From the discussion, the debate, came social pressures, unchallenged principles, decision points of opinion not substantiated by data but rather by belief. And yet, in any process for change whether it be for a better tomorrow or a just society, we need a beginning from which we may develop a model for examination and review. This was what was accomplished, a sensible beginning for practice, safe, reasonable practice guidelines for all practitioners. Not self-serving either, for the very endoscopists whose livelihood might be affected by a proposal to reduce the indications for a favourite procedure, recognized the need for constraint, a focusing of resources in the hope that these very resources would still be there when clearly and urgently indicated.

Remarkable that the Europeans also have a similar pragmatic approach to Managing Life with Acid Reflux Disease (First United European Gastroenterology Week, Athens, Greece, September 28, 1992). The prevalence of reflux symptoms among the adult population ranges from 26% for a six-month prevalence of heartburn and regurgitation (1), to 37 to 40% for a 12-month period (2,3). If we are to base a diagnosis of GERD on symptoms, and compare this with a ‘gold’ or at least ‘silver standard’ of 24 h pH monitoring, we need to know sensitivity, specificity and positive predictive value of heartburn and regurgitation: 24 to 38%, 63 to 89% and 66 to 81%, respectively (4,5). The assumption of the Canadian Consensus decision tree (Figure 1) is based on these figures, allowing the comfort of using empirical treatment without initially subjecting all persons to costly, and apparently often unnecessary, investigations.

"But", you may say, "my office is booked full with patients with the irritable bowel syndrome, not with GERD. If the prevalence data is correct, something doesn’t fit." To make a good fit, we need to recognize a further important factor – not acid, pepsin or sphincter pressures – but rather the determinants of health-seeking behaviour. Those persons with GERD-symptoms who consult a physician are more likely than nonconsulters to be worried that they have cancer, heart disease or some other possibly serious or fatal condition (6). So, part of our management of persons/patients with GERD must be to address quality of life issues, to ascertain what is the source of the anxiety.

The correlation between symptoms and pH-metry is reasonable (7), but not so for endoscopy (8,9). So why are we so often gazing at a video screen to grade the severity of esophagitis? Where is the evidence for endoscopic healing being associated with fewer recurrences or fewer complications? Certainly, recurrence rates for symptoms and for esophagitis are high so there may well be a good reason to identify the presence of esophagitis. Further investigation is appropriate for those persons with (a) alarm symptoms (dysphagia, weight loss, anemia, bleeding – worry: cancer or stricture); (b) persisting symptoms after eight to 14 weeks of treatment; or (c) recurring symptoms after recently successful therapy (worry: erosive esophagitis requiring more potent acid inhibition and/or maintenance therapy). Dr John R Bennett has provided a flow chart for empirical symptomatic
treatment of GERD (Figure 2) which is remarkably similar to that obtained through our own independent Canadian consensus (10). H2-receptor antagonists 'provide similar effective symptom relief when given in equivalent dosage regimens'. While superior symptom improvement can be achieved with a proton pump inhibitor rather than an H2-receptor antagonist (11), this study was not designed to control rigorously for the assessment of follow-up endoscopies, nor is it generally accepted that more potent acid suppression be undertaken initially in all persons. Why not? Because, why? Well, you decide - the adverse effect profile for short term use of proton pump inhibitors and H2-receptor antagonists is equally impressive. One could argue that an agent of superior efficacy and equivalent cost and safety profile be used for an endoscopy, but that was not the consensus. Perhaps to identify better those persons requiring an endoscopy to identify erosive disease, which is often associated with recurrences and may require appropriate discussion of the use of maintenance therapy. But patients with erosive esophagitis treated with either an H2-receptor antagonist or a proton pump inhibitor for eight to 12 weeks and who then come off therapy will quickly recur - some even suggest that those coming off acute treatment with a proton pump inhibitor might recur more quickly, so that an endoscopic study would be done at that point anyway. Some further reflection is needed on this contentious point. And for the patient who has a rapid return of frequent symptoms after coming off acute therapy (again with no alarm symptoms), what do we do if endoscopy is normal? Probably more of the same, another course of an H2-receptor antagonist (or a proton pump inhibitor) for eight to 12 weeks. And then? Off treatment again. What if the symptoms recur yet again? Some suggest that it is unnecessary to perform a second procedure. Yet do we know that the underlying assumption is correct, that once a normal endoscopy for GERD, always a normal endoscopy? A Wayne's World 'NOT'! So why not simply use intermittent therapy, cycles of H2-receptor antagonists or proton pump inhibitors for cycles of recurrent heartburn. Then, if symptoms frequently recur, consider maintenance therapy with a proton pump inhibitor or an H2-receptor antagonist or a prokinetic such as cisapride (12,13). The dose of acid-inhibitory therapy may need to be adjusted to normalize the percentage of time the esophageal pH is greater than 4 (14), but where is the evidence that this stepwise increase in dose of H2-receptor antagonist or proton pump inhibitor cannot be done based just on patient response, without further endoscopies?

Figure 1) Canadian GERD Consensus decision tree
Are there lessons from the management of patients with duodenal or gastric ulcer disease which can be applied to patients with GERD? Like GERD, peptic ulcer disease is prone to a long duration of symptoms, remaining active for 25 years or more in about 25% of individuals (15), with the ever-present risk of hemorrhage or perforation. Many different factors predispose to recurrence (Table 1), and the likelihood of duodenal ulcer disease recurrence depends on the number of risk factors. We do not yet know if the same factors, or principles, present in GERD, but this is certainly a reasonable working hypothesis, subject to future testing – even the possibility that Helicobacter pylori infection may be more common in persons with GERD. Of course, in GERD (let alone duodenal ulcer disease) the time has not yet come for indiscriminate attempts to stamp out H pylori (16). While symptomatic recurrences of GERD and duodenal ulcer disease can both be prevented with continuous therapy, it is only in the latter that factors have been identified which have an adverse influence on recurrence rates during continuous H₂-receptor antagonist treatment: these include smoking, stress, previous history of frequent ulcer relapses, and duration of disease for more than 10 years (17). While there is clear evidence for all the H₂-receptor antagonists being superior to placebo for maintenance therapy in duodenal ulcer disease (18), and evidence for lower duodenal ulcer recurrence rates with ranitidine than with cimetidine (19,20), we do not yet have comparative data for different H₂-receptor antagonists in GERD. We know that the risk of duodenal ulcer disease recurrence is kept low for as long as nine years with ranitidine maintenance therapy (21). We do not yet know how long to continue maintenance therapy in GERD; certainly good one-year remission rates may be achieved using 150 or 300 mg ranitidine bid. One advantage of using ranitidine 300 mg versus 150 mg for maintenance therapy in duodenal ulcer disease is the lowering of recurrence rates in smokers to the level seen in nonsmokers. There is the preliminary suggestion that prolonged treatment with H₂-receptor antagonists reduces the risk for ulcer recurrence once therapy has been discontinued (22,23). However, others argue that maintenance therapy must be continued for many years, perhaps at least 15 years, when the natural history of the disease has allowed it to become ‘burned out’ (15,24).

The maintenance dose for duodenal ulcer disease is one-half the standard ulcer-healing dose, but patients with GERD may need to be maintained on full doses of H₂-receptor antagonists. Does this have adverse effects? Clinically important hypergastrinemia has not been described in patients with duodenal ulcer disease after five years of treatment with ranitidine (25), and we await with great interest the outcome of a major duodenal ulcer disease maintenance study using omeprazole.

A new field is under cultivation – pharmacoeconomic considerations. There are important direct as well as indirect costs, and it is inefficient to consider just the cost of medication without also establishing the cost of investigations, institutional care and lost productivity, let alone the value of a person remaining pain-free. About one-third of direct costs relates to drugs (26), and direct costs may be less than half of the total cost (27). In a French multicentre study, maintenance therapy for duodenal ulcer disease with ranitidine 150 mg/day for 12 months resulted in a 29% decreased total cost of duodenal ulcer disease compared with no maintenance therapy (28). Continuous treatment with ranitidine was twice as cost-effective as intermittent treatment. In the United States, continuous versus intermittent therapy with ranitidine reduces the cost of one successfully treated patient by 12%, or about $100 (Canadian) per year (29). Another American study has shown that for at least 15 years continuous and intermittent treatment are of similar cost, and cheaper than surgery (30).

Many unknowns, but we have begun to sketch out the map for the future, to frame the problem in a context which may be tested, and improved upon. Please read the Consensus Document in this issue (page 277) with interest, and send us your comments.
REFERENCES


11. Cleator IG, Williams CN, Archambault AP, Taylor DW, Thomson ABR, Hunt RH, Sutherland LR, with the Canadian Collaborative Study Group. A comparison of the clinical efficacy and safety of Losec® (omeprazole) and Zantac® (ranitidine) in the clinical management of uncomplicated duodenal ulcer and gastroesophageal reflux disease. (In press)


