How should *Helicobacter pylori* positive dyspeptic patients be managed?

N J Talley

**Introduction**

Dyspepsia, according to the internationally accepted Rome criteria, refers to pain or discomfort centred in the upper abdomen; patients with predominant heartburn are excluded from this group, although minor or infrequent heartburn is commonly associated with dyspepsia. It is an important condition not only because it is common and costly, but because it may indicate the presence of serious disease such as peptic ulcer or gastric cancer. However, the most frequent causes of dyspepsia are functional dyspepsia and gastro-oesophageal reflux disease. The discovery of *Helicobacter pylori* has resulted in important advances in the management of dyspepsia. The clinician faced with a patient who has persistent or recurrent dyspepsia needs to differentiate clearly those patients who have not been previously investigated from patients documented to have functional dyspepsia after investigation (fig 1). Here, the management of *H pylori* positive dyspeptic patients who have and have not been fully investigated will be reviewed.

**Uninvestigated dyspepsia**

In new patients with dyspepsia who have alarm features (such as older age at first onset (>50 years), regular ingestion of non-steroidal anti-inflammatory drugs, or symptoms such as weight loss, dysphagia or bleeding) or a fear of serious disease, prompt upper endoscopy is always indicated if available. In younger patients without alarm features, management options include a trial of empirical therapy, prompt endoscopy for all, *H pylori* testing followed by endoscopy in positive cases, and testing for *H pylori* and treatment of those patients found to be infected with the bacterium.

Early endoscopy does have some advantages. It is more accurate than a barium meal and biopsy specimens can be taken to detect *H pylori* as well as from any suspicious areas to rule out malignancy. Peptic ulcer can be identified and then adequately managed. It is also more reassuring to patients than empirical therapy. However, endoscopy is also expensive and cancer is rarely found in otherwise healthy dyspeptic patients. Moreover, endoscopy is often delayed because of waiting lists or carried out when patients are on ulcer medications; this may lead to an erroneous diagnosis of functional dyspepsia in patients with ulcer disease or oesophagitis.

The practice of testing for *H pylori* in younger patients with new onset dyspepsia who do not have alarm features is becoming more widespread. A number of studies from Europe as well as New Zealand have reported that among dyspeptic patients who are infected with *H pylori*, between 20 and 60% have an underlying peptic ulcer at endoscopy (fig 2). In view of the overwhelming evidence that eradication of *H pylori* eliminates the ulcer diathesis, treatment of *H pylori* in these cases is very likely to be of value. Moreover, some without ulcer disease are likely to develop the condition over a lifetime so treating these cases now may be useful. However, it is not firmly established that those who are dyspeptic and have the infection are significantly more likely to develop an ulcer than those with the infection who are symptom-free. In Western countries patients with dyspepsia who do not have alarm features also have a very low probability of serious structural disease, including gastric cancer.

This knowledge has led to the concept that testing for *H pylori* and treating all infected cases (test and treat) should be efficacious because it will largely eliminate the pool of ulcer patients. The preliminary results of controlled trials have in general supported the safety and efficacy of a test and treat approach. Heaney et al randomised 104 *H pylori* positive patients based on breath testing to empirical eradication therapy without endoscopy or endoscopy with treatment based on the results. Overall, 25% in the empirical eradication therapy group proceeded to endoscopy because of no improvement in dyspepsia over six months. Symptom scores improved in both arms by six months but were significantly better in the empirical therapy group. Jones et al randomised 233 patients to *H pylori* test and treat or endoscopy. Only 12% of patients in...
A number of prestigious working parties have recommended, based on present evidence, that test and treat represents optimal clinical practice. However, there are limitations that clinicians need to keep in mind. The predictive value of diagnostic testing for *H pylori* peptic ulcers (arguably the key condition that test and treat addresses) depends on the sensitivity and specificity of the test as well as the prevalence of the disease in the population being assessed. The sensitivity and specificity of serology varies notably, and only a locally validated test with at least 90% sensitivity and specificity for the infection should be relied upon. Urea breath testing has produced consistently superior results compared with serology in the published literature. However, even if the test is excellent, the predictive value for relevant disease may be poor. For example, if the prevalence of *H pylori* infection in dyspepsia is 40% but only one in 10 *H pylori* positive cases has an ulcer, then the positive predictive value of the test for ulcer disease (with 95% sensitivity and specificity for *H pylori*) is only 45%, compared with 90% if four in 10 *H pylori* infected dyspeptic patients have ulcer disease. With a rise in *H pylori* negative ulcers in some parts of the world and the declining incidence of *H pylori* overall in Western countries, the value of a test and treat strategy will need careful scrutiny and likely revision in the future.

**Investigated dyspepsia**

The first question to address is should all patients with dyspepsia undergoing endoscopy, the “gold standard” investigation to exclude clinically relevant pathology, be tested for *H pylori* if no clear explanation such as a peptic ulcer is discovered? The next question to consider is should all infected patients with functional dyspepsia should be offered eradication therapy? The answers to these questions are controversial because of the limitations of the available evidence.

The key issue is whether identification and cure of *H pylori* cures the symptoms of functional dyspepsia. Older short term studies provided disparate results and have been roundly criticised. However, new well designed trials have not yet provided a definitive answer. In the past year, large trials have reported new results (fig 3). McColl et al from Scotland reported the findings from a single centre randomised controlled trial of 318 patients. They compared triple therapy (omeprazole, metronidazole and amoxicillin) with omeprazole for one week, with 12 months’ follow up. Overall, 87% in the active treatment group were cured of the infection compared with 4% on placebo. At one year, 21% had symptom relief in the *H pylori* treatment arm compared with 7% in the comparison group, a significant difference. The investigators concluded that one in five patients with functional dyspepsia will benefit from eradication therapy. However, the results from equally well conducted multicentre trials have reached different conclusions (fig 3). Talley et al conducted a trial in Europe and Australasia;
275 patients were randomised to triple therapy (omeprazole, clarithromycin and amoxicillin) or placebo for one week and followed for one year. An eradication rate of 84% and 4% was achieved in the triple therapy and placebo arms, respectively. At one year, 24% on active treatment and 21% on placebo had relief of dyspepsia, a non-significant difference.34 Similarly, Blum et al randomised 348 patients to triple therapy or omeprazole for one week, and then followed patients for one year. Overall, 27% on active treatment and 21% on placebo had symptom relief, a difference that failed to reach significance.35 Koelsz et al randomised 181 patients with H pylori who had failed to respond to treatment with proton pump inhibitors; they received either omeprazole and amoxicillin or omeprazole for two weeks, and were followed for six months. In this selected population, the proportion of responders was almost identical in each arm although data on symptom relief were not provided.36 The new trial results generally suggest that H pylori eradication is no more successful than placebo in relieving functional dyspepsia. Hence, a test and treat strategy for H pylori cannot rely on providing benefits in cases of functional dyspepsia, who usually comprise the majority of patients suffering from dyspepsia.

Working parties have generally recommended that H pylori infected patients with functional dyspepsia be offered treatment on a case-by-case basis, after careful explanation of the risks and benefits.5 9–12 This translates into begrudging acceptance that most clinicians will accept treatment to these people once infection is documented, despite some lingering concerns about rising antibiotic resistance rates and side effects. Cost benefit on relief of dyspepsia in recent trials.

Conclusions
The management of H pylori positive dyspeptic patients has now changed; in general, there is widespread acceptance that treatment of the infection should represent first line therapy in those who do not require further investigation, including those also suffering from functional dyspepsia. The increasing trend to offer eradication therapy to H pylori infected patients with documented functional dyspepsia, despite the lack of a clear benefit on relief of dyspepsia in recent trials. However, cost effective strategies need to be put in place when symptoms fail to resolve or there is only temporary relief. Trials specifically targeted at those who have failed to obtain symptom relief after H pylori eradication have still to be undertaken.

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