Assessment of *Helicobacter pylori* status by examination of gastric mucosal patterns: diagnostic accuracy of white-light endoscopy and narrow-band imaging

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**ABSTRACT**

**Objectives** *Helicobacter pylori* infection is a common cause of chronic gastritis worldwide and an established risk factor for developing gastric malignancy. The endoscopic appearances predicting *H. pylori* status are an ongoing area of research, as are their diagnostic accuracies. This study aimed to establish the diagnostic accuracy of several mucosal features predictive of *H. pylori* negative status and formulate a simple prediction model for use at the time of endoscopy.

**Design** Patients undergoing high-definition upper gastrointestinal (GI) endoscopy without magnification were recruited prospectively. During the endoscopy, the presence or absence of specific endoscopic findings was noted. Sydney protocol biopsies were used as the diagnostic reference standard, and urease test if taken. The results informed a logistic regression model used to produce a simple diagnostic approach. This model was subsequently validated using a further cohort of 30 patients.

**Results** 153 patients were recruited and completed the study protocol. The prevalence of active *H. pylori* infection was 18.3% (28/153). The overall diagnostic accuracy of the simple prediction model was 90.0%, and 100% of patients with active *H. pylori* infection were correctly classified. The presence of regular arrangement of collecting venules (RAC) within the stomach includes the presence of regular arrangement of collecting venules (RAC) showed a positive predictive value for *H. pylori* naive status of 90.7%, rising to 93.6% for patients under the age of 60.

**Conclusion** A simple endoscopic model may be accurate for predicting *H. pylori* status of a patient, and the need for biopsy-based tests. The presence of RAC in the stomach is an accurate predictor of *H. pylori* negative status, particularly in patients under the age of 60.

**Trial registration number** The study was registered with ClinicalTrials.gov, No. NCT02385045.

**INTRODUCTION**

*Helicobacter pylori* infection is a major cause of gastritis worldwide and is well established as a causative agent in gastric atrophy, intestinal metaplasia and dysplasia, culminating in the development of gastric adenocarcinoma.\(^1\)

Detection of *H. pylori* infection and subsequent eradication therapy has been shown to reduce symptoms of dyspepsia, and to stabilise or reduce the risk of developing gastric cancer.\(^2\,3\)

Several techniques exist for diagnosing *H. pylori* infection, and include non-invasive methods such as urea breath testing, faecal antigen testing and serological tests. Tissue samples may also be obtained during endoscopy and can be evaluated by microscopy, culture or rapid urease test.\(^4\) In recent years,
the widespread adoption of high-resolution endoscopy platforms has dramatically improved the level of mucosal detail visible to the endoscopist, and advanced imaging modalities such as narrow-band imaging (NBI) can improve visual contrast to better examine fine mucosal features.5

It is recognised that H. pylori naïve, infected or eradicated states, may be associated with distinct mucosal appearances, and the clinical utility of identifying these patterns is an area of ongoing study.6 With adequate accuracy, it could be appropriate to either initiate eradication therapy for suspected H. pylori, or identify patients at higher risk of long-term complications, that is, those with current or previous H. pylori infection.

A significant area of interest is the presence of a 'regular arrangement of collecting venules' (RAC), first described by Yagi et al, under magnifying and standard endoscopy.7 This is seen as numerous dots, or starfish-like points evenly distributed in the gastric body, as shown in figure 1. Further study of the RAC has suggested that when present, the RAC has a negative predictive value (NPV) of >90% for H. pylori negative status.8-10 This may approach 100% when the stomach is examined with image enhancement, such as NBI.11-13 Different types of abnormal RAC pattern have been described14; this study has attempted to simplify the diagnostic process by specifying only the presence or absence of the normal RAC, when examined without magnification using both white light and NBI.

However, the applicability of the RAC for diagnosis of an H. pylori naïve state may not be universal. Alaboudy et al15 demonstrated that the presence of RAC decreases with age, particularly for patients over the age of 60. The prevalence of H. pylori infection also varies widely worldwide, from as low as 24.4% in Oceania, to 70.1% in Africa. Prevalence in Northern Europe varies from 22.1% in Denmark, to 82.5% in Estonia.16 In addition, a large proportion of the studies examining RAC are performed in Asian populations by expert endoscopists, and relatively few in Western populations.17 18 The use of image-enhanced endoscopy such as NBI may also be of benefit in improving the contrast and delineation of gastric mucosal features and visualising the RAC, although requires further validation.19 20

Positive predictive findings for active H. pylori infection include diffuse redness, (figure 2) mucosal oedema, (figure 3) antral nodularity, enlarged gastric folds and sticky mucus, all of which may suggest a diagnosis of H. pylori infection.10 13 21-23 The significance of gastric erosions and haem flecks is less clear, with some studies reporting that gastric erosions are a predictor of H. pylori naïve status; other reports suggest that presence of gastric erosions may predict active H. pylori infection.24-26

It is also important to recognise patients with previously eradicated H. pylori infection, both to avoid unnecessary treatment and to guide decisions regarding biopsy for assessment of gastric atrophy or intestinal metaplasia.27

Figure 1  The endoscopic appearance of the RAC in the distal stomach, in a Helicobacter pylori naïve patient. Examined at medium distance without magnification under white-light endoscopy (WLE) (A) and narrow-band imaging (B), and on closer inspection without magnification under WLE (C) and NBI (D). NBI, narrow-band imaging.

Figure 2  The endoscopic appearances of diffuse redness, in a Helicobacter pylori infected patient.32

Figure 3  The endoscopic appearances of mucosal oedema, in a Helicobacter pylori infected patient. (A) The gastric body34 and (B) under closer inspection near the antrum.
Exclusion criteria included patients with previous gastric surgery or liver disease, contraindications to biopsy, acute GI bleeding and with known malignancy of the GI tract. Patients were also excluded if they were unable or unwilling to give informed consent, or if consent was withdrawn after the procedure.

Gastroscopy protocol
Patients fasted overnight for procedures scheduled in the morning, and for 6 hours for afternoon procedures. Procedures were completed using either pharyngeal anaesthesia (Xylocaine spray) or sedation (midazolam+fentanyl) or both, according to patient and clinician preference.

Gastroscopies were performed using HD endoscopes (GIF-H290, Olympus) without magnification. The procedures were performed by six experienced endoscopists (>1000 procedures) following familiarisation with the gastroscopy protocol, and data were recorded by members of the research team present during the gastroscopy. Following a standard examination of the duodenum and stomach by the endoscopist, the antrum, incisura and fundus of the stomach were carefully examined under white light with good insufflation of the stomach, and adequate washing of the mucosa if required. The presence or absence of RAC in the lower part of the stomach, diffuse redness, mucosal oedema, map-like redness, erosions, haem flecks, oesophagitis, antral nodularity and gastric or duodenal ulceration, were recorded in real time. These features are associated with H. pylori infection, as well as showing regression after H. pylori eradication. The NBI mode of the gastroscope was then activated, and careful examination of the antrum, incisura and fundus was repeated, for assessment of the same characteristics. Biopsies were then taken according to the updated Sydney Protocol (lesser and greater curvatures at the antrum and body, and at the incisura angularis). The gastroscopy was then completed including any other biopsies (including urease test if clinically indicated) at the discretion of the endoscopist.

Data recording
Patient demographics and baseline characteristics were recorded including age, sex, ethnicity and the experience level of the endoscopist performing the procedure. All assessments of mucosal features were recorded immediately following the end of the procedure and blinded to the results of a urease test (if taken). The mucosal surface patterns were classified as normal or abnormal based on the appearances described by Yoshii et al and Inui et al. H. pylori status was determined using the mucosal biopsies obtained according to the updated Sydney protocol; these were labelled according to their anatomical location and analysed by an expert GI histopathologist. Giemsa staining was performed for all patients. In addition, if a urease test was performed during the gastroscopy (Biohit H pylori UFT300), the results of this were recorded, and

METHODS
Recruitment
We performed a retrospective and prospective validation of the diagnostic performance of endoscopic findings reported in real-time in the upper GI tract, using conventional white-light endoscopy (WLE) and NBI. After recruitment of an initial cohort of patients we carried out multinomial logistic regression modelling to identify the predictive features associated with H. pylori status in a European population, and to provide early prospective validation of a simple endoscopic classification system.

Previous studies have identified a mucosal pattern of ‘map-like redness’ (figure 4) as being predictive of H. pylori eradicated status or gastric metaplasia. It has also been shown that some changes can regress following H. pylori eradication, including diffuse redness and antral nodularity.

This study attempts to evaluate the diagnostic accuracy characteristics of defined mucosal features as predictors of H. pylori status in a European population, and to provide early prospective validation of a simple endoscopic classification system.

Figure 4 The endoscopic appearances of map-like redness, in a patient with previous Helicobacter pylori eradication.

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a patient was classified as *H. pylori* positive if either one or both tests demonstrated the presence of *H. pylori*.

### Statistical analysis

Sample size calculations were performed according to Buderers method, with assumptions that the RAC would have sensitivity 90% and specificity 90% for an *H. pylori* naïve status. A prevalence of 35% was estimated, based on recent demographic meta-analysis for Northern Europe. A sample size of 120 patients was calculated to be required.

We present the sensitivity, specificity, positive and NPVs and accuracy of RAC and other mucosal features, with 95% CIs. All statistical analyses and modelling were performed in IBM SPSS V.25.

The primary outcomes were the accuracy of the RAC and of the other identified mucosal features, for prediction of the *H. pylori* status. The secondary outcome was the accuracy of a simple endoscopic system for classifying patients *H. pylori* status.

### RESULTS

#### Study participants and baseline characteristics

Approximately 350 patients were screened for inclusion in the study, and 157 patients were recruited; the remainder did not meet inclusion criteria (due to, eg, the procedural indication such as anaemia or dysphagia, previous gastric surgery or liver disease. A further four patients underwent gastroscopy but were subsequently withdrawn from the study due to food residue within the stomach or otherwise incomplete procedure. One patient did not have the biopsy protocol performed; they did, however, have a positive urease test and were therefore included in analysis.

A total of 155 patients were included for analysis; the age range was from 19 to 90 years with a mean age of 54.2 years (±16.8). The participant flow diagram is presented in figure 5.

### *H. pylori* prevalence

Using the patient history, histological presence of *Helicobacter*, or positive urease test, 111 of the 153 patients were *H. pylori* naïve (72.5%), 28 had active *H. pylori* infection (18.3%) and 14 were *H. pylori* eradicated. 32% of patients underwent urease test, of which 7 were positive; all of these patients also had evidence of *H. pylori* on histopathological analysis. There were no significant differences in prevalence by age or gender, although the highest prevalence of infection (11/29, 37.9%) was seen in patients from a Middle-Eastern background. Patient characteristics and *H. pylori* prevalence are reported in table 1.

#### Diagnostic performance of the RAC

To date, the most extensively studied feature of the stomach for prediction of *H. pylori* status is the RAC (figure 1). Using WLE, the RAC was identified in 108/153 (70.6%) of this cohort of patients. All patients with RAC identified under WLE, also showed RAC under NBI.

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**Table 1** Patient characteristics and *Helicobacter pylori* prevalence

| Patient characteristics | H. pylori naïve | H. pylori active infection | H. pylori eradicated |
|-------------------------|-----------------|---------------------------|----------------------|
| Age                     |                 |                           |                      |
| All patients            | 54.2±16.80 (19-90) | 111 (72.5%) | 28 (18.3%) | 14 (9.2%) |
| <50                     | n=60 (39.2%)    | 44 (73.3%) | 13 (21.7%) | 3 (5.0%)  |
| ≥50                     | n=93 (60.8%)    | 67 (72.0%) | 15 (16.1%) | 11 (11.8%) |
| Gender                  |                 |                           |                      |
| Male                    | n=66 (43.1%)    | 46 (69.7%) | 13 (19.7%) | 7 (10.6%)  |
| Female                  | n=87 (56.9%)    | 65 (74.7%) | 15 (17.2%) | 7 (8.0%)   |
| Ethnicity               |                 |                           |                      |
| European                | 90 (58.8%)      | 71 (46.4%) | 11 (7.2%)  | 8 (5.2%)   |
| South Asian             | 17 (11.1%)      | 13 (8.5%)  | 4 (2.6%)   | 0 (0.0%)   |
| East Asian              | 3 (2.0%)        | 3 (2.0%)   | 0 (0.0%)   | 0 (0.0%)   |
| Afro-Caribbean          | 14 (9.2%)       | 9 (5.9%)   | 2 (1.3%)   | 3 (2.0%)   |
| Middle Eastern          | 29 (19.0%)      | 15 (9.8%)  | 11 (7.2%)  | 3 (2.0%)   |

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**Figure 5** Participant flow diagram.
NBI examination allowed identification of the RAC in a further four patients, all of whom were *H. pylori* negative.

The presence of the RAC in the lower part of the lesser curvature of the stomach has been shown in many previous studies to be a strong predictor of an *H. pylori* negative status, and our results agreed with this, showing a diagnostic OR of 6.53 (95% CI 2.70 to 15.80, p<0.001) for *H. pylori* negative status (either naïve or eradicated) in the presence of RAC.

In 78.4% of *H. pylori* negative individuals, the RAC was present (under either WLE, NBI or both). By contrast, the RAC was only present in 35.7% of *H. pylori* positive patients.

Diagnostic performance measures are shown in table 2 for the presence of RAC under combined WLE and IEE examination, as well as for different cut-off values of age, as it has previously been demonstrated that the RAC becomes less visible with increasing age. For patients below the age of 55, the presence of RAC carries an NPV of 93.2% (95% CI 88.2 to 92.4).

Regression modelling of significant endoscopic findings

A multinomial logistic regression model was employed to evaluate the predictive capabilities of each endoscopic sign being studied (RAC, diffuse redness, mucosal oedema, map-like redness, erosions, ulceration, haem flecks and oesophagitis). For the first cohort of 123 patients, the presence or absence of each feature was recorded during endoscopy, as was any history of *H. pylori* infection or treatment, and the final histological diagnosis. Patients were stratified into ‘naïve’, ‘active infection’ or ‘eradicated’ groups based on their history and the results of biopsy-based tests taken at endoscopy. Results of the endoscopic and histological findings from the patient cohort are presented in table 3.

A predictive model including all the endoscopic features demonstrated a reasonable goodness-of-fit (multinomial logistic regression modelling, SPSS V.25) with a Pearson χ² statistic of 16.48 (Sig. 1.00). The processing summary of the final model is presented in online supplemental data. The overall predictive accuracy was 81.3%. Adding the patient age as a covariate did not increase the predictive accuracy of the model, but when the model was applied only to patients under the age of 55, the expected accuracy rose to 86.4%.

We attempted to produce a predictive model that would preserve accuracy without requiring the endoscopist to record a large number of endoscopic findings; a two-stage predictive model taking previous history into account was therefore developed, after the approach of Gupta et al. but requiring fewer endoscopic signs to be assessed. The selected features were presence of RAC, diffuse redness, mucosal oedema and map-like redness (as a marker of atrophy), as these features showed the greatest positive or NPVs for *H. pylori* status (figures 3–5). Regression modelling suggested a preserved goodness-of-fit, with Pearson χ² of 21.8 (Sig. 0.005), when including only these findings. The algorithmic approach to the diagnostic prediction is presented in figure 6. Patients in whom none of the included mucosal features could be identified, were classified as ‘naïve’ or ‘eradicated’ based on any previous history of *H. pylori* infection.

Prospective validation of the diagnostic prediction model

A further 30 patients were recruited following the development of the diagnostic predictive model. These patients underwent gastroscopy as per the standard protocol. As previously, endoscopists were informed of any history of *H. pylori* infection, but were blinded to results of recent pre-test stool or serological tests for *H. pylori*, and to the results of urease tests taken during endoscopy.

Endoscopists were required to state whether each sign in the predictive model was present, and to predict the *H. pylori* status of each patient based on the algorithmic approach detailed previously.

An overall diagnostic accuracy of 80.0% was demonstrated, the results are presented in table 4. Six patients had active *H. pylori* infection; all of these were classified correctly. Seventeen patients were *H. pylori* naïve, of which 12 were classified correctly, and the remainder (5 patients) were classified incorrectly as having active infection. Three patients were classified as ‘uncertain’ based on the absence of all studied endoscopic findings but were classified correctly when taking previous history of *H. pylori* into account. No patients with active infection were classified as ‘naïve’ or ‘eradicated’.

| Table 2 | Diagnostic characteristics for RAC presence as a predictor of *Helicobacter pylori* negative status (either naïve or eradicated) |
|---------|--------------------------------------------------------------------------------------------------------------------------------|
| N       | Sensitivity % (95% CI) | Specificity % (95% CI) | Positive predictive value % (95% CI) | Negative Predictive Value % (95% CI) | Accuracy % (95% CI) |
| RAC (all patients) | 153 | 78.4 (70.2 to 85.3) | 64.3 (44.1 to 81.4) | 90.7 (85.5 to 94.2) | 40.0 (30.2 to 50.7) | 75.8 (68.2 to 82.4) |
| RAC (patients aged <50 years) | 60 | 85.1 (71.7 to 93.8) | 76.9 (62.1 to 90.0) | 93.0 (83.1 to 97.3) | 58.8 (40.4 to 75.1) | 83.3 (71.5 to 91.7) |
| RAC (patients aged <55 years) | 74 | 84.8 (73.0 to 92.8) | 73.3 (44.9 to 92.2) | 92.6 (84.4 to 96.7) | 55.0 (38.4 to 70.6) | 82.4 (71.8 to 90.3) |
| RAC (patients aged <60 years) | 86 | 82.9 (72.0 to 90.8) | 75.0 (47.6 to 92.7) | 93.6 (86.0 to 97.2) | 50.0 (35.7 to 64.3) | 81.4 (71.6 to 89.0) |

Examined using combined WLE and NBI. Results are presented for different age cohorts.

NBI, negative predictive value; RAC, regular arrangement of collecting venules; WLE, white-light endoscopy.
DISCUSSION

Changes in the endoscopic appearance of the stomach in relation to *H. pylori* infection are an ongoing area of research, and several characteristic features have been identified to predict *H. pylori* naïve, positive and eradicated status.

A significant feature of the *H. pylori* naïve stomach is the presence of RAC. Our study supports the previous observations that the presence of this pattern is the strongest single predictor for *H. pylori* naïve status and therefore careful examination using WLE could be considered as part of a diagnostic approach to ‘rule out’ *H. pylori* infection.

### Table 3

|                  | *H. pylori* naïve (n=111) | *H. pylori* infected (n=28) | *H. pylori* eradicated (n=14) |
|------------------|---------------------------|-----------------------------|-------------------------------|
| **Prevalence of endoscopic findings:** |                           |                             |                               |
| RAC              | 90 (81.1%)                | 10 (35.7%)                  | 8 (57.1%)                     |
| Diffuse redness  | 27 (28.8%)                | 13 (46.4%)                  | 1 (7.1%)                      |
| Mucosal oedema   | 9 (8.1%)                  | 7 (25.0%)                   | 0 (0%)                        |
| Map-like redness | 8 (7.2%)                  | 3 (10.7%)                   | 6 (42.9%)                     |
| Erosions         | 15 (13.5%)                | 4 (14.3%)                   | 1 (7.1%)                      |
| Gastro-duodenal ulceration | 3 (2.7%) | 4 (14.3%)                   | 1 (7.1%)                      |
| Haem flecks      | 10 (9.0%)                 | 0 (0%)                      | 0 (0%)                        |
| Oesophagitis     | 1 (0.9%)                  | 1 (3.6%)                    | 0 (0%)                        |
| Antral nodularity | 0 (0%)                    | 1 (3.6%)                    | 0 (0%)                        |
| **Prevalence of histological findings:** |                           |                             |                               |
| Histologically normal | 65 (58.6%)             | 0 (0%)                      | 5 (35.7%)                     |
| Acute gastritis  | 7 (6.3%)                  | 20 (71.4%)                  | 0 (0%)                        |
| Chronic gastritis| 37 (33.3%)                | 17 (60.7%)                  | 7 (50.0%)                     |
| Gastric atrophy  | 2 (1.8%)                  | 0 (0%)                      | 4 (7.1%)                      |
| Gastric metaplasia | 4 (3.6%)                  | 2 (7.1%)                    | 4 (28.6%)                     |
| Helicobacter-like organisms | 0 (0%) | 28 (100%)                   | 0 (0%)                        |

NB: Figures do not sum to 100% as findings may coexist in patients. RAC, regular arrangement of collecting venules.

Figure 6  The diagnostic algorithm used for the second cohort of patients to predict classification of *Helicobacter pylori* status. RAC, regular arrangement of collecting venules.
infection. Training in the recognition of the RAC would be of benefit in Western endoscopic practice. It is also important to consider that histopathological assessment, as with all biopsy-based techniques, may not diagnose all cases of *H. pylori* if organisms are not present within the biopsy sample area.

Previous similar studies have reported test sensitivity as high as 100% for *H. pylori* when RAC is present; that is, no patients with RAC present were *H. pylori* positive. Interestingly, our study has found that a significant proportion of *H. pylori* positive patients (10/23) were found to have RAC present within the stomach. The reasons for this are unclear; possibilities include that *H. pylori* infection does not always lead to RAC loss, that RAC loss is not immediate or complete, or that RAC was misidentified by the endoscopists in those patients.

We have attempted to produce a diagnostic model to aid in the endoscopic prediction or classification of *H. pylori* status. This model has been validated in a small initial cohort of patients and demonstrated accuracy of 80%. Importantly, no cases of active *H. pylori* infection were classified as naïve or eradicated by this method.

When designing this model, it was important to avoid overburdening an endoscopist with a large number of signs and features to identify. Other groups have demonstrated high diagnostic accuracy (88.6%–93.4%) for an algorithmic approach using history and endoscopic findings, though this was with a machine-learnt model using up to 16 different findings per patient. The endoscopic features in this approach differ slightly from those in the Kyoto classification, in part due to removing the feature of antral nodularity, which was encountered uncommonly in this population, and also by removing the differentiation between intestinal metaplasia and map-like redness, which is likely to show similar histological findings.

We have proposed a simpler approach which we believe will be more applicable in routine practice. This may aid endoscopists as a screening tool for *H. pylori* naïve patients, and as a decision-making aid for patients who may require further investigation for *H. pylori* or its sequelae, such as atrophic gastritis and metaplasia.

This study has some limitations. The prevalence of *H. pylori* infection was substantially lower (18.3%) than the 35% anticipated based on population studies. This is lower than most globally reported populations and may influence the generalisability of our results. The selection criteria may also limit the wider applicability; this study included only patients with symptoms of dyspepsia or abdominal pain and therefore will not include patients with asymptomatic *H. pylori* infection who have attended for endoscopy for an unrelated reason (such as anaemia or dysphagia).

Similarly, the rates of histologically proven gastric atrophy in *H. pylori* negative patients were low, with only two patients in the *H. pylori* naïve group (2/111) and four patients in the *H. pylori* eradicated group (4/14/) showing histological findings. The reason for this is not clear, although could have important implications for external validity if the rates of gastric atrophy in our cohort differ significantly from those in the wider population.

We also do not know the rates of spontaneous eradication of *H. pylori*; five patients in the initial cohort had no history of infection but did show signs of map-like redness, although no evidence of *H. pylori* was present on biopsy. Additionally, this was a small single-centre study including endoscopists who were motivated and interested to take part. We attempted to assess intraoperator variability but this proved impossible due to the lower image quality of saved files, compared with real-time high-resolution endoscopic assessment. This is an important factor, as features such as mucosal oedema are highly subjective and interoperator and intraoperator agreement should be quantified for these.

If extrapolated to a larger study, it may be appropriate to examine inter-operator variability and the learning curve for applying the proposed algorithm. Furthermore, the prediction algorithm does not consider confounding factors, as each endoscopic feature included is assessed individually, and treated with equal weight. A further limitation is that few participants underwent *H. pylori* stool antigen or urea breath test, and it is understood that biopsy-based diagnosis may carry a higher false-negative rate than these methods. These, however, were not available as patients were recruited as part of their normal routine healthcare.

Another important question is of the effect of PPI treatment on the gastric mucosal appearances. We have included PPI-naïve and PPI-exposed patients within the analysis, which may have implications for the interpretability of the mucosal findings.

It is becoming clear that endoscopic predictors of *H. pylori* status exist, and when used appropriately can contribute to accurate diagnosis by the general

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**Table 4** Results of prospective validation of the diagnostic algorithm

| Predicted *Helicobacter pylori* status according to the diagnostic algorithm | Naïve | Active infection | Eradicated | Total |
|---|---|---|---|---|
| Naïve | 12 | 0 | 0 | 12 |
| Active Infection | 5 | 6 | 0 | 11 |
| Eradicated | 0 | 0 | 7 | 7 |
| Total | 17 | 6 | 7 | 30 |
endoscopist. We suggest that a close white-light examination of the stomach should be attempted in patients with symptoms of dyspepsia and no known previous history of *H. pylori*, to attempt to identify the RAC. Our study also suggests that further examination under NBI may identify patients with RAC present, which was not appreciated under WLE. Presence of the RAC should be a reliable sign that *H. pylori* is not present and may avoid the need for gastric biopsies. If the RAC cannot be identified, the presence of diffuse gastric redness or mucosal oedema suggest the possibility of *H. pylori* infection and may warrant biopsy, stool antigen or breath testing. In the absence of any of these features, if map-like redness is present then an *H. pylori* eradicated status can be suspected, and consideration of future surveillance may be justified based on the patient risk of gastric cancer. Other aids to assessment of these features could include mucolytic agents to ensure good mucosal views, and we suggest their presence or absence could be included in the gastroscopy report for all patients.

**CONCLUSIONS**

This study introduces a simple algorithm for classification of *H. pylori* status that may be applied during endoscopy and acts as a decision aid for further diagnostic tests or actions. This algorithm has been prospectively validated in a small sample of patients, in a single centre.

We propose that further research in this area includes a larger patient cohort for prospective validation of this approach. This validation should ideally be conducted in a range of centres with endoscopes from varied manufacturers and undertaken by endoscopists with a range of experience. It will also be beneficial to evaluate the learning curve and performance curve for adoption of this procedure, and to assess interoperator variability.

**Contributors**

NP planned and supervised the study. BG and NP conducted data collection and analysis. JT, BG and NP interpreted and analysed the data. BG prepared the report for submission; NP and JT reviewed and approved the submission. BG submitted the report and is responsible for the overall content as guarantor.

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**Competing interests**

None declared.

**Patient consent for publication**

Not required.

**Ethics approval**

The study was registered with the local research ethics committee. No. 14/LO/1525.

**Provenance and peer review**

Not commissioned; externally peer reviewed.

**Data availability statement**

Data are available on reasonable request. The data shall be available on reasonable request from contacting the corresponding author on bglover@imperial.ac.uk.

**Supplemental material**

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Supplementary data: Processing summary of final logistic regression model. NB: Mucosal Oedema was incorporated with Diffuse Redness to be considered as a single variable (i.e. either DR or MO or both are present) as results of initial analysis indicated these should be merged.

### Case Processing Summary

|                             | N   | Marginal Percentage |
|-----------------------------|-----|---------------------|
| True status based on history and results |     |                     |
| naive                       | 94  | 76.4%               |
| active infection            | 22  | 17.9%               |
| eradicated                  | 7   | 5.7%                |
| Is the RAC present?         |     |                     |
| no                          | 30  | 24.4%               |
| yes                         | 93  | 75.6%               |
| Is there diffuse redness?   |     |                     |
| no                          | 87  | 70.7%               |
| yes                         | 36  | 29.3%               |
| Is there map-like or patch redness? | |       |
| no                          | 116 | 94.3%               |
| yes                         | 7   | 5.7%                |
| Valid                       | 123 | 100.0%              |
| Missing                     | 0   |                     |
| Total                       | 123 |                     |
| Subpopulation               | 91a |                     |

a. The dependent variable has only one value observed in 83 (91.2%) subpopulations.

### Model Fitting Information

| Model         | Model Fitting Criteria -2 Log Likelihood | Likelihood Ratio Tests |
|---------------|------------------------------------------|------------------------|
|               | Likelihood Ratio Tests                   |                        |
|               | Chi-Square | df | Sig.   |
| Intercept Only| 152.075  |    |        |
| Final         | 130.189  | 21.886 | 8 | .005   |
| Goodness-of-Fit | Chi-Square | df | Sig. |
|-----------------|------------|----|------|
| Pearson         | 186.224    | 172| .217 |
| Deviance        | 118.156    | 172| .999 |

| Pseudo R-Square | . |
|-----------------|---|
| Cox and Snell   | .163 |
| Nagelkerke      | .220 |
| McFadden        | .132 |
| True status based on history and results* | B   | Std. Error | Wald | df | Sig. | Exp(B) | 95% Confidence Interval for Exp(B) |
|-----------------------------------------|-----|------------|------|----|------|--------|----------------------------------|
| naive                                   |     |            |      |    |      |        |                                  |
| Intercept                               | 3.278 | 2.471     | 1.760 | 1  | .185 |         |                                  |
| Patient age                             | -.017 | .024      | .501  | 1  | .479 | .983   | .937 1.031                       |
| [Is the RAC present?=0]                 | .018  | 1.200     | .000  | 1  | .988 | 1.018  | .097 10.708                     |
| [Is the RAC present?=1]                 | 0^b  | .        | .     | 0  | .    | .      |                                  |
| [Is there diffuse redness?=0]           | -.886 | 1.156    | .587  | 1  | .444 | .412   | .043 3.977                      |
| [Is there diffuse redness?=1]           | 0^b  | .        | .     | 0  | .    | .      |                                  |
| [Is there map-like or patch redness?=0] | 1.114 | 1.274    | .764  | 1  | .382 | 3.046  | .251 36.982                    |
| [Is there map-like or patch redness?=1] | 0^b  | .        | .     | 0  | .    | .      |                                  |
| active infection                        |     |            |      |    |      |        |                                  |
| Intercept                               | .947  | 2.706     | .122  | 1  | .726 |         |                                  |
| Patient age                             | -.020 | .028      | .533  | 1  | .465 | .980   | .927 1.035                     |
| [Is the RAC present?=0]                 | 2.339 | 1.260    | 3.448 | 1  | .063 | 10.371 | .878 122.476                  |
| [Is the RAC present?=1]                 | 0^b  | .        | .     | 0  | .    | .      |                                  |
| [Is there diffuse redness?=0]           | -.635 | 1.246    | .260  | 1  | .610 | .530   | .046 6.087                    |
| [Is there diffuse redness?=1]           | 0^b  | .        | .     | 0  | .    | .      |                                  |
| [Is there map-like or patch redness?=0] | 1.125 | 1.478    | .579  | 1  | .447 | 3.079  | .170 55.793                   |
| [Is there map-like or patch redness?=1] | 0^b  | .        | .     | 0  | .    | .      |                                  |

a. The reference category is: eradicated.
## Likelihood Ratio Tests

| Effect                               | Model Fitting Criteria | Likelihood Ratio Tests |
|--------------------------------------|------------------------|------------------------|
|                                      | -2 Log Likelihood of Reduced Model | Chi-Square | df | Sig. |
| Intercept                            | 130.189*a              | .000                   | 0  | .   |
| Patient age                          | 130.778                | .590                   | 2  | .745|
| Is the RAC present?                  | 147.547                | 17.358                 | 2  | .000|
| Is there diffuse redness?            | 130.985                | .796                   | 2  | .672|
| Is there map-like or patch redness?  | 130.867                | .678                   | 2  | .712|

## Classification

| Observed  | Predicted     | Percent Correct |
|-----------|---------------|-----------------|
|           | active infection | eradicated | naive | 6 | 0 | 93.6% |
| naive     | 88            | 6              | 0     | 90.2% |
| active infection | 17          | 5              | 0     | 22.7% |
| eradicated | 6            | 1              | 0     | 0.0% |
| Overall Percentage | 9.8%        | 0.0%           | 75.6% |

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