Comparison of three faecal occult blood tests in the detection of colorectal neoplasia

R L Hope, G Chu, A H Hope, R G Newcombe, P E Gillespie, S J Williams

Abstract

Methods and Aims—For the detection of colorectal neoplasia, 192 consecutive patients had colonoscopy to evaluate the sensitivity and specificity of three faecal occult blood tests (FOBT). Of 160 evaluable patients (96 female, mean age 51.9), 21 patients (13-1%) had adenomas and three patients (1.9%) had colorectal carcinoma.

Results—When comparing all three faecal occult blood tests for the detection of colorectal neoplasia, the sensitivity of Monohaem (43.8%) was superior to both Hemoccult II (25%) and to BM-Test colon albumin (25%). The specificity of Monohaem (94.6%) was greater than both Hemoccult II (88%) and BM-Test colon albumin (89%). Using McNemar’s test, Monohaem was a more accurate FOBT than Hemoccult II and BM-Test albumin (p<0.05). In the 21 patients with adenomatous polyps, FOBT sensitivity seemed to be dependent on polyp size, but not polyp site.

Conclusion—Monohaem, a faecal occult blood test that uses a monoclonal antibody that is specific for human haemoglobin, is a more accurate test in the detection of colorectal neoplasia and should possibly be used in colorectal cancer screening programmes.

Study design

The initial 56 patients were asked to prepare two FOBT kits: (a) Monohaem screen included one card per day over three days and (b) Hemoccult II screen, which included one card per day over three days. The ensuing 136

Methods

Patients

From March 1991 until November 1992, 192 consecutive patients who required colonoscopy for the investigation of gastrointestinal symptoms or for ‘polyp surveillance’ follow up (see Figure), were recruited for study. In no case was colonoscopy being performed for previously positive FOBTs and patients with a history of overt gastrointestinal bleeding were excluded. No patient had undergone previous colonic resectional surgery. Patients were asked to start the testing of stools five days before hospital admission to ensure enough time to complete the three day regimen before bowel preparation. There were no dietary restrictions placed on the patients and regular medications were continued.

Keywords: colorectal neoplasia, faecal occult blood tests, Monohaem, Hemoccult II, adenoma, screening.
patients were asked to prepare three FOBT kits: (a) Monohaem Screen (as above), (b) Hemoccult screen (as above), and (c) BM-Test colon albumin screen, which included two cards each day over three days.

Monohaem uses mouse monoclonal antibodies covalently linked to the Monohaem strip that specifically bind human haemoglobin to detect abnormal amounts of occult blood in stool samples. After application of the sample to the card, the bound haemoglobin is detected by treatment with an aqueous ethanolic solution of gum guaiac followed by hydrogen peroxide. The gum guaiac resin contains α guaiaconic acid, which is oxidised in the presence of peroxide to a blue coloured product by the pseudoperoxidase enzymatic action of haemoglobin. Because the haemoglobin is bound selectively by the localised monoclonal antibody on the card, a positive reaction appears as a blue ring or spot confined to the area of antibody immobilisation.10 The assessment of the colour change is done visually, five to 60 seconds after the application of the hydrogen peroxide solution and there is little ambiguity in the interpretation of a positive or negative test. The antibody specifically selects for human haemoglobin, thus eliminating false positives from other peroxidase active agents.

The Hemoccult II kit consists of a peroxidase substrate impregnated card, which is developed with hydrogen peroxide after application of a faecal smear. Any haemoglobin present in the faecal smear gives a blue coloured reaction by virtue of its pseudoperoxidase activity. The coloured reaction is evident anywhere on the card where haemoglobin is present. As there is no specificity for human haemoglobin, any other peroxidase present can give rise to a positive reaction. HOII test was performed unhydrated.

The BM-Test colon albumin kit consists of a card that is probed with a strip carrying sequentially arranged reagent fields designed to detect any human albumin present in the faecal sample. Elution from the sample proceeds to formation of a complex with galactosidase conjugated antihuman albumin monoclonal antibody and then on to a dye conjugated galactoside releasing the dye to give a colorimetric end point if human albumin is present. A red to violet colouration constitutes a positive result, and yellow to orange-brown colouration constitutes a negative result. The colour change is assessed visually, five to 15 minutes after the reagent is added.

All participants were given verbal and written instructions in the method of specimen collection. Stools were collected on several layers of toilet paper on a folded newspaper, disposable container or child’s potty, ensuring that the stool was not contaminated with water. Samples for the tests were collected immediately after defecation and using a spatula, the windows on the cards were well covered with faecal smear (a drop of water was added to the window on the Monohaem card). A different section of the same stool was sampled to inoculate a second window on the Hemoccult and Monohaem cards. After sampling, the cards were sealed, labelled, and stored in a cool dry place. Patients returned the completed FOBT kits on hospital admission for the colonoscopy.

Kits were analysed at an independent laboratory (Sugerman’s Pathology Pty Ltd, Hurstville, NSW, Australia) to ensure that the colonoscopist was not aware of the FOBT result. Analysis was performed within 48 hours of arrival at the laboratory and, therefore, all kits were processed within 10 days of stool sampling. All kits were processed according to manufacturers’ instructions. A single positive smear was considered a positive test.

Colonoscopy was performed (by SJW, PEG or under their direct supervision) without knowledge of the FOBT result. If colorectal polyps were detected, polyp site was recorded and polypectomy performed. Polyps were examined histologically and size recorded. Colorectal cancer site and histology was recorded when found.

Results

Overall, 160 patients (83.3%) were evaluated as they had completed FOBT kits correctly and had total colonoscopy (to the base of the caecum). Twenty six patients (13.5%) had inoculated the FOBTs incorrectly and were not included in the analysis. Six patients (3.1%) were excluded from analysis as they had either very poor bowel preparation (three patients) or the colonoscopy was incomplete (three patients).

There were 96 women and 64 men, with a mean age of 51.9 (range 24–89). The Figure shows the indication for colonoscopy. Fifteen patients had hyperplastic polyps (9.4%), 21 patients (13.1%) had adenomas (11 proximal to splenic flexure, 10 distal to splenic), and colorectal carcinoma occurred in three patients (1.3% rectosigmoid, 0.6% hepatic flexure).

Test sensitivity for colorectal carcinoma

Overall, three patients had colorectal carcinoma. Both MH and HOII were positive in these three patients, however, BMCA was negative in one patient with a rectosigmoid carcinoma.

Test sensitivity for adenoma

Overall, 21 patients had adenomas. The sensitivity of MH (52.4%) was greater than HOII (28.5%). If a patient had more than one adenoma (12 patients), the most distal polyp site was recorded. The sensitivity for detecting adenoma (MH or HOII) did not seem to be dependent on polyp site (see Table I), although the absolute number of patients with adenomas was small.

The mean adenoma size was 11 mm (range 3–30). The sensitivity of MH was greater in patients with large (≥10 mm) adenomas (70%) than in patients with smaller (<10 mm) adenomas (36.4%). The same trend in FOBT sensitivity was seen with regard to polyp size using HOII
TABLE I Influence of adenoma site and size on FOBT

<table>
<thead>
<tr>
<th>Site</th>
<th>Monohaem sensitivity (%)</th>
<th>HOOI sensitivity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proximal</td>
<td>54.5</td>
<td>27.3</td>
</tr>
<tr>
<td>Distal</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;10 mm</td>
<td>36.4</td>
<td>9.1</td>
</tr>
<tr>
<td>&gt;10 mm</td>
<td>70.0</td>
<td>50.0</td>
</tr>
</tbody>
</table>

Hemoccult II for small adenomas and 9-1% for small adenomas (see Table I).

TABLE II McNemar table for 24 affected patients (with colorectal neoplasia) comparing MH versus HOOI

<table>
<thead>
<tr>
<th></th>
<th>HOOI+</th>
<th>HOOI-</th>
</tr>
</thead>
<tbody>
<tr>
<td>MH+</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>MH-</td>
<td>4</td>
<td>10</td>
</tr>
</tbody>
</table>

TABLE III McNemar table for 136 unaffected patients (no colorectal neoplasia) comparing MH versus HOOI

<table>
<thead>
<tr>
<th></th>
<th>HOOI+</th>
<th>HOOI-</th>
</tr>
</thead>
<tbody>
<tr>
<td>MH+</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>MH-</td>
<td>15</td>
<td>116</td>
</tr>
</tbody>
</table>

TABLE IV McNemar table for 116 affected patients (with colorectal neoplasia) comparing BMCA versus HOOI

<table>
<thead>
<tr>
<th></th>
<th>BMCA+</th>
<th>BMCA-</th>
</tr>
</thead>
<tbody>
<tr>
<td>MH+</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>MH-</td>
<td>1</td>
<td>8</td>
</tr>
</tbody>
</table>

TABLE V McNemar table for 92 unaffected patients (no colorectal neoplasia) comparing BMCA versus HOOI

<table>
<thead>
<tr>
<th></th>
<th>BMCA+</th>
<th>BMCA-</th>
</tr>
</thead>
<tbody>
<tr>
<td>MH+</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>MH-</td>
<td>8</td>
<td>79</td>
</tr>
</tbody>
</table>

Detecting colorectal neoplasia, the sensitivity of MH (43.8%) was greater than BMCA (25%) with 95% confidence intervals for difference in sensitivity -0.096 to +0.433. The specificity of MH (94.6%) was also greater than BMCA (91.1%) with 95% confidence intervals for difference in specificity -0.022 to +0.136.

Discussion

The ideal screening test on a population should combine satisfactory sensitivity with high specificity. Unfortunately, Hemoccult II has quite poor sensitivity for detecting asymptomatic colorectal neoplasms. While dehydration of the faecal sample does increase the sensitivity of HOII, it does so at the expense of decreasing specificity. This loss in test specificity can generate significant burden on a colorectal screening programme and therefore, rehydration is not routinely performed. Our study has shown that the immunochromatographic test, Monohaem, is more sensitive and more specific for detecting colorectal neoplasms than Hemoccult II. It is a more accurate FOBT. This is important, as minor changes in test sensitivity and specificity can have an important impact on efficiency and cost effectiveness of a screening programme.

St John et al suggest that the better performance of HemeSelect (another immunochromatographic test) related to its high chemical sensitivity for haemoglobin (in vitro) and the specificity of the test for human haemoglobin. Immunoreactive haemoglobin is rapidly degraded in the stomach and duodenum, giving these tests a high degree of selectivity for bleeding from the distal gastrointestinal tract. The other obvious advantage of an immunochromatographic test directed against human haemoglobin is that false positive results do not occur when peroxidase containing materials have been ingested. Red meat and fruit and vegetables containing peroxidase need to be avoided when Hemoccult II is used, which may diminish patient compliance. It is not necessary to restrict the diet before Monohaem testing.

Macrae and St John showed that only large adenomas with a diameter of >2 cm cause faecal blood loss of >2 ml/day. The superior sensitivity of both HOII and MH for detecting larger polyps was in keeping with this finding. Given that most colorectal cancers arise from pre-existing adenomas (adenoma-adenocarcinoma sequence), the detection of large adenomas (and subsequent removal) in an asymptomatic population is an important aim.

Our patients were recruited for study after the decision to perform colonoscopy had been made. While our study was not on an asymptomatic population, we were able to obtain valuable information with regard to FOBT specificity. This has been a criticism of previous investigators using symptomatic subjects with known colorectal neoplasms. Specificity, defined as the proportion of screened people without pre-clinical disease who are designated correctly as negative, was...
able to be compared between the three FOBTs. Monohaem specificity (94.6%) was superior to both HOII (88%) and BMCA (89.1%) in 92 patients who were truly free of colorectal neoplasia. These data on FOBT specificity are useful when we consider that the mean age of these patients was 52.3. While recommendations for FOBT screening on an asymptomatic population remain to be established, most would agree that screening will target an ‘average risk’ asymptomatic person between the age of 50–75.

Many factors including accuracy, cost effectiveness, and patient and physician compliance influence the selection of a faecal occult blood test for screening programmes. While cost effectiveness was not a primary aim of our comparative study, it is worth noting that the Monohaem screen is comparatively inexpensive. The initial outlay of approximately US$2 (including card and reagent) per MH test, should be recouped by being a more specific test, and therefore not generating as many costly second phase tests such as colonoscopy.

In summary, the immunochemical test, Monohaem, was the most accurate faecal occult blood test. It combines high test sensitivity for detecting colorectal neoplasia with high test specificity. It should be considered as a more accurate FOBT and possibly used in colorectal cancer screening programmes.


