Reproducibility of wireless capsule endoscopy in the investigation of chronic obscure gastrointestinal bleeding

Dimitrios Christodoulou MD1,2, Gregory Haber MD FRCPC3, Umar Beejay MD4, Shou-jiang Tang MD1, Simon Zanani MD1, Rima Petroniene MD3, Maria Cirocco MSc1, Paul Kortan MD FRCPC1, Gabor Kandol MD FRCPC1, Athina Tatsioni MD3, Epameinondas Tsianos MD2, Norman Marcon MD FRCPC1

BACKGROUND: Capsule endoscopy (CE) is a valuable tool in the diagnostic evaluation of obscure gastrointestinal bleeding, but limited information is available on the reproducibility of CE findings.

OBJECTIVE: To compare two successive CE studies with push enteroscopy (PE) in patients presenting with chronic obscure gastrointestinal bleeding.

METHODS: A prospective study was conducted. Ten patients (seven men and three women) with chronic obscure gastrointestinal bleeding and no contraindications for CE were administered and completed the trial. For each patient, the first capsule was administered on day 1, the second capsule was administered on day 2 and PE was performed on day 3. Endoscopists were blinded to the capsule findings. Capsule findings were assessed independently by two investigators blinded to PE findings.

RESULTS: A potential small intestinal bleeding source was found in 60% of the patients when all the studies were combined. A bleeding source was found in four patients in both CE studies. The second CE also identified a bleeding source in a fifth patient. Interobserver agreement by kappa analysis was 0.642 to 1.000 (P ≤ 0.05) for the CE studies. PE identified a potential small bowel bleeding site in four patients, including one patient who had negative CE studies.

CONCLUSIONS: This study confirmed the reproducibility of CE findings on successive studies. Some patients did not have a source of bleeding in the small intestine, and all studies found this.

Key Words: Capsule endoscopy; Obscure gastrointestinal bleeding; Push enteroscopy; Reproducibility

La reproductibilité de l’endoscopie capsulaire sans fil dans l’exploration de saignements gastro-intestinaux occultes chroniques

HISTORIQUE : L’endoscopie capsulaire (ECP) est un outil précieux pour l’évaluation diagnostique des saignements gastro-intestinaux occultes, mais on possède peu d’information sur la reproductibilité des résultats de l’ECP.

OBJECTIF :Comparer deux études d’ECP successives avec entéroscopie poussée (EP) chez des patients qui consultent en raison de saignements gastro-intestinaux occultes chroniques.

MÉTHODOLOGIE : Les auteurs ont procédé à une étude prospective. Dix patients (sept hommes et trois femmes) atteints de saignements gastro-intestinaux chroniques ont été admis à l’étude. Les deux premières études d’ECP ont été exécutées le jour 1, et le jour 2, puis l’EP a été exécutée le jour 3. Les endoscopistes ne avaient pas au courant des résultats des capsules, qui ont été évalués de manière autonome par deux chercheurs non informés des résultats de l’EP.

RÉSULTATS : On a découvert une source potentielle des saignements du petit intestin chez 60% des patients une fois toutes les études combinées. On a découvert une source de saignement chez quatre patients dans les deux études d’ECP. La deuxième ECP a également permis de repérer une source de saignement chez un cinquième patient. La concordance entre observateurs par analyse kappa variant entre 0.642 et 1.000 (P ≤ 0.05) pour les études d’ECP. L’EP a permis de repérer un foyer de saignement dans l’intestin grêle de quatre patients, y compris un patient dont les résultats des études d’ECP étaient négatifs.

that incorporates an ingestible wireless camera (3-8). There is increasingly more evidence reporting the utility of capsule endoscopy (CE) in evaluating patients with COGB and other small bowel pathologies (9-11). However, there are no published data regarding the reproducibility of CE given consecutively to the same patient. The purpose of the present prospective study was to assess the reproducibility of CE given consecutively to the same COGB patient and to compare the findings with those of conventional PE in COGB patients.

**METHODS**

**Ethics**

The study protocol was approved by the research ethics board at St Michael's Hospital (Toronto, Ontario).

**Patients**

Patients 18 years of age or older with a history of COGB, who could give written informed consent, were eligible for the study. Exclusion criteria included known or suspected gastrointestinal (GI) obstruction, strictures or fistulas, the presence of cardiac pacemakers (although CE currently appears to be safe for patients with pacemakers, the test has not been approved for such patients) or other implanted electromedical devices, and a positive pregnancy test for women of child-bearing age. Demographic parameters included sex, age, race, weight and height. Related current and past medical history included prior surgery, comorbid conditions and medications. A standard physical examination was performed. Each patient was given two M2A capsules on two consecutive days and PE was performed on the third day.

**CE**

The patients fasted for 12 h before each capsule study. For improved bowel preparation, 2 L of GoLYTELY (Braintree Laboratories Inc, USA) were administered the previous day. Ease of swallowing the capsule was assessed using a visual analogue scale. CE was performed on day 1 (CE1) with the Given M2A video capsule. On the following day, the patient returned for a second CE (CE2). The second M2A capsule was administered similarly.

The findings of CE were arbitrarily classified into definite, indeterminate and incidental. Definite findings included angiodysplasias (AVMs), tumours, fresh blood and melena (ie, changed blood). Indeterminate findings included non-bleeding red lesions and tiny red spots. Finally, incidental findings included phlebectasias, lymphangiectasias, small polyps and lymphoid nodules. AVMs were classified as red lesions larger than 1 mm in size, with a distinct border or spider-like projections and a bright red colour. Indeterminate red lesions included red lesions 1 mm to 3 mm in size without the asteroid configuration or bright red colour that is characteristic of AVMs. Finally, tiny red lesions were classified as pinpoint red lesions smaller than 1 mm in size (Figure 1).

The findings of CE1 and CE2, as recorded by the first and second investigators, were divided into proximal, middle and distal according to the location of the lesion in the small bowel. The subjective estimation was based on the transit times, as defined by passage through the pyloric sphincter and the ileocecal valve. The localization software helped to identify the timeline of the capsule motion through the small bowel.

**PE**

On the third day of the present study, highly experienced endoscopists (GH, PK, GK, NM) performed PE on patients who were under conscious sedation, using a standard Olympus enteroscope PE1 (Olympus America Inc, USA) or an Olympus pediatric colonoscope PCF-160L (Olympus America Inc, USA). The enteroscope or the pediatric colonoscope was advanced as far as possible into the small bowel until the shaft of the instrument was fully inserted. Any lesion observed during insertion or withdrawal of the enteroscope or colonoscope was carefully documented in terms of its nature, location and size. For treatable bleeding lesions found during PE, appropriate endoscopic intervention was performed with argon plasma coagulation (APC).

Two investigators (UB and ST), both with previous experience in CE, reviewed all capsule images independently while blinded to PE findings and to each other’s findings. Finally, two reviewers (GH and DC) coordinated the study, collected CE findings from both investigators, decided on a final diagnosis when there was a discrepancy in capsule interpretation by the two investigators and analyzed the data.

On day 4, the patients were asked to complete a questionnaire by using a visual analogue scale to assess pain or discomfort during the procedures. The patients were also asked...
TABLE 1
Results of the subjective assessment of capsule endoscopy (CE) and push enteroscopy (PE)

<table>
<thead>
<tr>
<th>Question</th>
<th>CE rating, mean</th>
<th>PE rating, mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>How would you rate the swallowing/insertion of the instrument?</td>
<td>3.4 2.3</td>
<td></td>
</tr>
<tr>
<td>Did you experience pain during the procedure?</td>
<td>4.0 2.9</td>
<td></td>
</tr>
<tr>
<td>Did you experience discomfort during the procedure?</td>
<td>3.8 2.4</td>
<td></td>
</tr>
<tr>
<td>Did you experience pain after the procedure?</td>
<td>4.0 2.9</td>
<td></td>
</tr>
<tr>
<td>Did you experience discomfort after the procedure?</td>
<td>4.0 2.9</td>
<td></td>
</tr>
<tr>
<td>How would you rate the procedure?</td>
<td>3.7 2.0</td>
<td></td>
</tr>
<tr>
<td>Rate the overall convenience of the test</td>
<td>2.6 1.3</td>
<td></td>
</tr>
<tr>
<td>If you were given the possibility to select an examination for diagnosing your problem, would you choose this procedure?</td>
<td>3.7 2.3</td>
<td></td>
</tr>
</tbody>
</table>

Ratings are based on the visual analogue scale, with a range of 0 to 4, where 0=worst and 4=best. *Possible answers included 0=no, 1=possibly, 2=probably, 3=very probably and 4=yes

RESULTS

Ten patients (seven men and three women) were enrolled in the study. The mean age of the study group was 74.2 years (range 64 to 86 years). The mean (± SD) height of the patients was 167.5±10.6 cm (range 152.0 cm to 185.4 cm) and the mean weight was 77.8±18.94 kg (range 54.6 kg to 113.0 kg). The duration of COGB was 29.7±19.32 months. The manifestations of COGB included intermittent melena or obvious blood loss in six patients, and iron deficiency anemia with positive fecal occult blood in four patients. The mean hemoglobin level was 96.6±18.4 g/L (normal values: 125 g/L to 185 g/L) and the mean ferritin level was 35.3±51.3 pmol/L (normal values: 30 pmol/L to 230 pmol/L). A mean of 41.6±42.8 units of blood was transfused, with a median of 17 units of blood (range, zero to 110 units of blood). The patients had been hospitalized a mean of 8.4±6.9 times for the diagnostic investigation and treatment of their COGB. The mean number of previous procedures per patient before undergoing CE are shown in Figure 2.

Upper endoscopy showed that three of the 10 patients were treated for AVMs in the past. Colonoscopy showed diverticulosis in two patients and a cecal AVM that was treated in one patient. Previous PE showed jejunal AVMs in three patients. The lesions were treated by bipolar or APC but the chronic blood loss was continued. Small bowel follow-through showed a duodenal diverticulum in one patient. A nuclear scan showed positive findings in one patient without identifying the bleeding site, and angiography showed the suspected bleeding site in the small bowel in the patient.

Two patients had also undergone intra-operative enteroscopy and in both of those cases, jejunal AVMs were seen and were treated with bipolar coagulation, but chronic bleeding continued.

All 10 patients completed the study. The M2A capsule was swallowed without any difficulty in all 20 CE procedures. There were no complications and no patient complained of any symptoms during or after CE (Table 1). The capsule reached the cecum during the recording period in 17 of the 20 CE procedures. Of the three cases in which the capsule did not reach the cecum during the recording period, in one case (patient 5, CE2), the capsule remained in the stomach for the whole recording time. In the other two cases, there was prolonged gastric retention of the capsule before crossing the pylorus. The mean small bowel transit time for CE1 was 3 h 49 min ±1 h 10 min, and the mean small bowel transit time for CE2 was 3 h 28 min ±1 h 7 min. Likewise, PE was carried out successfully.

The findings of CE1, CE2 and PE are presented in Tables 2 and 3. The measurement of agreement between the two investigators and between the two CE procedures was calculated by using the kappa analysis, as described above (Table 4).

The comparison of the CE findings showed a good degree of agreement between the two investigators. Especially for the significant findings (Table 4), the range of kappa values was 0.642 to 1.000 (P=0.05), showing a substantial to good degree of agreement between the two investigators. For the nonsignificant findings, the range of kappa values was 0.374 to 1.000 for the statistically significant comparisons that showed a moderate to good agreement between the investigators. Despite the overall good rate of agreement between the investigators,
one major significant finding (a tumour in patient 6) was missed by one investigator on CE 2, while it was identified by both investigators on CE1 (Table 2).

The comparison of findings of the investigators showed a variable degree of agreement between CE 1 and CE2. For the A VMs and tumour lesions, the degree of agreement between CE1 and CE 2 was substantial to good (k=0.769 to 1.000, P<0.05). For the presence of fresh blood, the agreement between CE1 and CE2 was fair (k=0.357), likely because it may have been obvious that there was bleeding on one day but not obvious on another day. For the insignificant findings, the degree of agreement between CE1 and CE2 was moderate to good (k=0.500 to 1.000).

A potential SI bleeding source with the combination of all studies was found in 60% of patients (n=6) (Table 3). CE 1 found a bleeding source in four of the 10 patients. The lesions included A VMs, ranging in number from one to five in three of those four patients (Figures 3 to 5). Three of those four patients with positive findings had bleeding manifested by fresh blood (n=2) or melena (n=1). On CE1, one patient had an irregular area with fresh blood in the distal ileum, which

### Table 2

<table>
<thead>
<tr>
<th>Patient</th>
<th>Age/sex</th>
<th>Presenting symptom</th>
<th>Hb/Ferritin, (g/L)/(pmol/L)</th>
<th>Transfused units of blood</th>
<th>CE1 First investigator</th>
<th>CE1 Second investigator</th>
<th>CE2 First investigator</th>
<th>CE2 Second investigator</th>
<th>Push enteroscopy</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>68/M</td>
<td>Melena</td>
<td>76/10</td>
<td>110</td>
<td>3 AVM, fresh blood</td>
<td>3 AVM, fresh blood</td>
<td>2 AVM</td>
<td>3 AVM</td>
<td>Negative</td>
<td>–</td>
</tr>
<tr>
<td>2</td>
<td>69/M</td>
<td>Hematochezia</td>
<td>95/165</td>
<td>2</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Bleeding from stoma site</td>
</tr>
<tr>
<td>3</td>
<td>75/F</td>
<td>Melena and hematochezia</td>
<td>100/85</td>
<td>14</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Diverticulosis</td>
</tr>
<tr>
<td>4</td>
<td>64/M</td>
<td>Iron deficiency anemia</td>
<td>84/9</td>
<td>0</td>
<td>5 AVM, melena</td>
<td>5 AVM, melena</td>
<td>6 AVM, fresh blood</td>
<td>6 AVM, fresh blood</td>
<td>3 AVM</td>
<td>–</td>
</tr>
<tr>
<td>5</td>
<td>82/M</td>
<td>Melena</td>
<td>90/12</td>
<td>12</td>
<td>Capsule stayed in stomach</td>
<td>Capsule stayed in stomach</td>
<td>Negative</td>
<td>Negative</td>
<td>1 AVM</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>86/M</td>
<td>Iron deficiency anemia</td>
<td>76/3</td>
<td>20</td>
<td>Tumour, fresh blood</td>
<td>Tumour, fresh blood</td>
<td>Fresh blood</td>
<td>Fresh blood</td>
<td>Negative</td>
<td>–</td>
</tr>
<tr>
<td>7</td>
<td>81/F</td>
<td>Iron deficiency anemia</td>
<td>110/27</td>
<td>31</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>3 AVM</td>
<td>–</td>
</tr>
<tr>
<td>8</td>
<td>73/M</td>
<td>Melena</td>
<td>85/10</td>
<td>93</td>
<td>Negative, 2 small polyps</td>
<td>Negative, 1 small polyp</td>
<td>Negative, 2 small polyps</td>
<td>Negative, 1 small polyp</td>
<td>Negative, Myelodysplasia</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>75/F</td>
<td>Melena</td>
<td>120/10</td>
<td>68</td>
<td>3 AVM</td>
<td>4 AVM</td>
<td>1 AVM</td>
<td>1 AVM</td>
<td>1 AVM</td>
<td>Distal AVM not seen in CE2 due to fluid in the fundus</td>
</tr>
<tr>
<td>10</td>
<td>68/M</td>
<td>Iron deficiency anemia</td>
<td>130/22</td>
<td>1</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Negative</td>
<td>Cameron lesions in the small bowel</td>
</tr>
</tbody>
</table>

*Age is presented in years. A VM Angiodysplasia; F Female; Hb Hemoglobin; M Male

### Table 3

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Positive findings, % (patients, n)</th>
<th>Type of finding (patients, n)</th>
<th>Total lesions found, n</th>
</tr>
</thead>
<tbody>
<tr>
<td>CE1</td>
<td>40 (4)</td>
<td>AVM (3), tumour (1)</td>
<td>16</td>
</tr>
<tr>
<td>CE2</td>
<td>50 (5)</td>
<td>AVM (4), tumour (1)</td>
<td>14</td>
</tr>
<tr>
<td>CE1 + CE2</td>
<td>50 (5)</td>
<td>AVM (4), tumour (1)</td>
<td>18</td>
</tr>
<tr>
<td>PE</td>
<td>40 (4)</td>
<td>AVM (4)</td>
<td>8</td>
</tr>
<tr>
<td>CE1 + CE2 + PE</td>
<td>60 (6)</td>
<td>AVM (5), tumour (1)</td>
<td>21</td>
</tr>
</tbody>
</table>

AVM Angiodysplasia, CE1 First capsule endoscopy; CE2 Second capsule endoscopy; PE Push enteroscopy

### Table 4

#### Kappa analysis (measure of agreement) for the significant and nonsignificant findings between the two investigators and between the first capsule endoscopy (CE1) and the second capsule endoscopy (CE2)

<table>
<thead>
<tr>
<th>Finding</th>
<th>Investigator 1 versus investigator 2</th>
<th>CE1 versus CE2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>k</td>
<td>ASE</td>
</tr>
<tr>
<td>AVMs*</td>
<td>0.883</td>
<td>0.113</td>
</tr>
<tr>
<td>Tumour*</td>
<td>0.642</td>
<td>0.326</td>
</tr>
<tr>
<td>Fresh blood*</td>
<td>1.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Melena*</td>
<td>1.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Indeterminate red lesions†</td>
<td>0.374</td>
<td>0.207</td>
</tr>
<tr>
<td>Tiny red spots‡</td>
<td>0.240</td>
<td>0.190</td>
</tr>
<tr>
<td>Phlebectasias§</td>
<td>0.578</td>
<td>0.203</td>
</tr>
<tr>
<td>Lymphangiectasias§</td>
<td>0.424</td>
<td>0.159</td>
</tr>
<tr>
<td>Small polyps†</td>
<td>1.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Lymphoid nodules‡</td>
<td>1.000</td>
<td>0.000</td>
</tr>
</tbody>
</table>

Agreement was poor if the kappa value (k) ≤ 0.20; fair if 0.21 ≤ k ≤ 0.40; moderate if 0.41 ≤ k ≤ 0.60; substantial if 0.61 ≤ k ≤ 0.80, and good if k>0.80. *Significant finding; †Nonsignificant finding; §No statistics were computed because CE2 had a constant negative finding (no variance). ASE Asymptotic standard error; AVM Angiodysplasia
was diagnosed as a tumour (Figure 6). The patient underwent a computed tomography scan, which showed a focal irregular mass with thickening of the wall of the small bowel for approximately 5 cm to 6 cm, with an irregular but patent lumen. There was no evidence of a small bowel obstruction proximal to the mass. There were also a few mildly prominent mesenteric nodes in the vicinity, which measured up to 9 mm. Unfortunately, the patient died as the result of a heart attack before undergoing the operation.

In the other five patients, CE1 did not demonstrate a definite source of bleeding, but only some prominent submucosal veins, lymphangiectasias (n=4) and some small duodenal polyps (n=1). In one patient, on CE1, the M2A capsule remained in the stomach for the whole duration of the test. Gastric findings included a few antral erosions, a gastric scar and mild erythema. Limited endoscopic views of the colonic mucosa were obtained in six patients, while two others had changed blood (ie, melena) and one other had a large amount of stool that did not allow any view of the colonic mucosa. CE2 gave results similar to those of CE1 (Figure 7). One of the investigators identified an AVM on one more patient (patient 7) on CE2 and this finding was confirmed by PE. In the diabetic patient (patient 5), the first capsule failed to get through the pylorus and the second capsule got through the pylorus 3 h 41 min after ingestion, but failed to demonstrate any bleeding sites.

Therapy
PE identified a potential small bowel bleeding site in four of the 10 patients. All four patients had AVMs. The first of
them had three AVMs that were treated with APC, while the respective capsule studies had also shown AVMs on that patient. The second patient had one large jejunal AVM that was treated with APC, and that lesion was not identified by CE (however, the first capsule stayed in the stomach and in the second CE study, the view was limited because of fluid). The third patient had AVMs detected by PE (also found in one of the CE studies) and the fourth patient had a small AVM detected by PE (while AVMs were also identified by CE studies on that patient). Both of those patients received treatment with APC. In total, more significant findings were identified by CE (n=18) than by PE (n=8) (Table 3).

Four patients had negative findings in both CE studies and PE examination. The first patient had diverticulosis and a history of cecal AVM, the second patient had chronic, intermittent blood loss from a stoma site, as proven by direct magnification inspection of the stoma during retrograde endoscopy, the third patient had Cameron gastric lesions and the fourth patient was found to have myelodysplastic syndrome as the cause of his chronic anemia.

The course of the disease was improved in the four patients who received APC treatment during PE for their AVMs. For the four patients with negative findings by CE and PE, the exclusion of a small bowel disease led to improved management of their underlying GI disease or other disease. The patient with AVMs found by CE that were not found by PE (patient 1) underwent intraoperative enteroscopy with cauteryization of some AVMs and the requirements for blood transfusions were reduced.

**DISCUSSION**

Evaluation of COGB is still one of the challenges in gastroenterology. It is estimated that in 5% to 10% of patients with COGB, the bleeding source cannot be identified by standard endoscopic techniques (ie, upper endoscopy and colonoscopy) (1,2,12). For the diagnosis of COGB, it is suggested that a second upper endoscopy or “second opinion endoscopy” be performed, due to the high frequency of lesions that are overlooked at the initial endoscopy (1). Among these lesions are large hiatal hernias with Cameron lesions (underlying erosions or ulcers), peptic ulcer disease and vascular ectasia in the upper GI tract. Small bowel follow-through or enterolysis, nuclear red blood cell scan and angiography can locate the bleeding site in only up to 10% to 20% of patients with COGB (13-15). PE, along with CE, where available, is now the standard approach for the evaluation of COGB (2,16,17). Intraoperative enteroscopy is usually performed when patients have transfusion-dependent bleeding from a source that cannot be located despite extensive diagnostic evaluation and when the risks of continued bleeding are judged to outweigh the risks of laparotomy. In general, the overall diagnostic yield of PE in identifying potential bleeding lesions in patients with COGB lies in the range of 38% to 75%, whereas that of intraoperative enteroscopy ranges from 70% to 100% (2,13,14,16,18,19).

The present study showed that CE has good reproducibility and a satisfactory interobserver agreement rate. A second capsule, administered the next day, led to an identification of the same lesions as the first capsule for the great majority of lesions, with few exceptions. Good bowel preparation and passage of the capsule through the entire small bowel were the most important factors for identification of the same lesions by CE1 and CE2.

In an experimental study (3) that preceded human studies, CE was compared with PE in detecting small bowel lesions in an animal model that was developed specifically for this purpose. Overall, the CE sensitivity in detecting lesions randomly sewn in the full length of the small bowel was 64% compared with 57% for PE. PE had a sensitivity of 94% in identifying beads within its range, compared with an overall sensitivity of 53% for CE within the same range. In another short report (5), CE provided good views and successfully imaged small bowel pathological features in four patients with obscure or uncontrolled GI bleeding.

Some prospective trials (9,10,20-27) compared CE with PE for the evaluation of COGB. Ell et al (10) included 32 patients in their study, and the diagnostic workup included small bowel enterolysis, angiography, scintigraphy, PE and CE. Enterolysis did not provide any diagnostic clue, while red blood cell scintigraphy was positive in one patient and celiac and mesenteric angiography was positive in four patients. PE provided relevant pathological findings in 38% of the patients and identified a clear bleeding source in 28% (n=9) of the 32 patients. CE provided definitive evidence of a bleeding source in 21 of 32 patients (66%), and the difference from PE was statistically significant. A study by Lewis and Swain (9) also compared CE with PE for the evaluation of suspected SI bleeding in 20 patients. The yield of PE in the evaluation of obscure bleeding was 30% (n=6) and the yield of CE was 55% (n=11). The difference in yield between PE and CE did not reach significance. In another study (20), CE found a distal source of bleeding in five of 14 patients who had normal PE. In a study by Mylonaki et al (21), CE was more effective than PE in the evaluation of COGB (68% versus 32%) and led to an alteration in therapy in 66% of patients with positive findings. Other studies (22-26) confirmed those findings, but also showed that CE did not change the management of patients with indeterminate lesions, such as red spots or slight erythema, while it improved the clinical course in patients with significant findings such as angiectasias or focal ulcers. The studies described above showed that CE is an invaluable tool for the investigation of COGB. Other authors (27) stressed the efficacy of CE for the diagnosis of small bowel lesions because they found significant lesions in 62.9% of patients and identified the bleeding source in 75% of patients with iron deficiency anemia of obscure origin.

In a prospective study (11,28) that compared CE with small bowel follow-through in 20 patients, CE was significantly more sensitive for the detection of small bowel diseases and of the potential SI bleeding source. Of interest, two case reports (29,30) showed that CE identified a bleeding Meckel diverticulum after an extensive, nonconclusive diagnostic workup, including Meckel scintigraphy, in one of the two cases. It is clear that wireless CE is already a first-line tool for the investigation of small bowel diseases, but there is still more to investigate surrounding the fine details and findings of the procedure (24,31-34). From this perspective, the present study confirmed the high reproducibility rate of consecutive CE studies and used a clear terminology for the identified lesions.

Delvaux et al (25) studied 44 patients who underwent CE as the initial investigation of the small bowel when the gastroscopy and colonoscopy findings were normal. Further management decisions were based on CE results. After 12 months, follow-up data were obtained from all patients and referring physicians. CE detected an intestinal lesion in 18 patients
for surgical laparotomy (43). Despite all the advantages of entrapped CE capsules from the small bowel without the need for enteroscopy has been an effective method for the extraction of Roux-en-Y anastomosis (42). Finally, double-balloon enteroscopy can be used to insert the endoscope in parts of the small bowel that have altered anatomy as the result of surgical procedures (e.g., the afferent limb of a COGB (by the anterograde or retrograde approach, or both). This method also makes the treatment of many of the lesions that are identified by CE feasible, obviating the need for intra-operative enteroscopy and laparotomy (39,40). It can also identify some of the lesions that are rarely missed by CE and the two methods can be considered complementary (41). Double-balloon enteroscopy can be used to insert the endoscope in parts of the small bowel that have altered anatomy as the result of surgical procedures (e.g., the afferent limb of a Roux-en-Y anastomosis) (42). Finally, double-balloon enteroscopy has been an effective method for the extraction of entrapped CE capsules from the small bowel without the need for surgical laparotomy (43). Despite all the advantages of double-balloon enteroscopy, the technique also has some limitations: it is not widely available, it is very time consuming and has increased costs.

In the present study, both CE and PE were negative in four of the 10 patients, so in combination with the patients’ relative history and relative findings, a bleeding source from the small bowel was excluded, as described above. For the six remaining patients with positive findings, CE identified one or more bleeding sources in five patients. The patient with a small bowel tumour had a regional transit abnormality of the M2A capsule at the area of the tumour, as was previously described by our group of researchers (32,33). PE identified potential small bowel bleeding sources in four of the 10 patients. The most important finding of the present study was the high reproducibility of CE and the high degree of agreement between the investigators that reviewed the capsule videos, with few exceptions. In addition, CE was better tolerated by patients than was PE, as was demonstrated by the subjective questionnaire completed by the patients (Table 1). No patient experienced pain or discomfort during CE and all patients rated the procedure very highly. The capsule was comfortable to swallow and the patients were willing to repeat the test, if necessary.

**CONCLUSIONS**

The present study provided further evidence regarding the value and reproducibility of CE during the evaluation of COGB. It is likely that CE, if available, should be the first test chosen for the investigation of suspected SI bleeding because it is easy, reliable and reproducible. In our opinion, PE should accompany CE for the complete investigation of the upper small bowel in patients with transfusion-dependent anemia and visible bleeding because it provides the option of therapeutic intervention. CE should be repeated if the capsule fails to reach the cecum during the recording period and in cases in which the view is limited due to the presence of fluid or food residue.

**ACKNOWLEDGEMENTS:** This study was sponsored by Given Imaging Ltd. Dr Christodoulou has received a scholarship/grant from the Greek Association of Gastroenterology for postgraduate training at St Michael’s Hospital, Toronto, Ontario.

**REFERENCES**