Gastroscopy following a positive fecal occult blood test and negative colonoscopy: Systematic review and guideline


BACKGROUND: A sizeable number of individuals who participate in population-based colorectal cancer (CRC) screening programs and have a positive fecal occult blood test (FOBT) do not have an identifiable lesion found at colonoscopy to account for their positive FOBT screen.

OBJECTIVE: To evaluate the evidence and provide recommendations regarding the use of routine esophagogastroduodenoscopy (EGD) to detect upper gastrointestinal (UGI) cancers in patients participating in a population-based CRC screening program who are FOBT positive and colonoscopy negative.

METHODS: A systematic review was used to develop the evidentiary base and to inform the evidence-based recommendations provided.

RESULTS: Nine studies identified a group of patients who were FOBT positive and colonoscopy negative. Three studies found no cases of UGI cancer. Four studies reported cases of UGI cancer; three found UGI cancer in 1% or less of the population studied, and one study found one case of UGI cancer that represented 7% of their small subgroup of FOBT-positive/colonoscopy-negative patients. Two studies did not provide outcome information that could be specifically related to the FOBT-positive/colonoscopy-negative subgroup.

CONCLUSION: The current body of evidence is insufficient to recommend for or against routine EGD as a means of detecting gastric or esophageal cancers for patients who are FOBT positive/colonoscopy negative, in a population-based CRC screening program. The decision to perform EGD should be individualized and based on clinical judgement.

Key Words: Colonoscopy; Esophagogastroduodenoscopy; Fecal occult blood test; Mass screening; Systematic review; Upper gastrointestinal cancer

In Canada, colorectal cancer (CRC) is the fourth most commonly diagnosed cancer and the second leading cause of cancer death, with an estimated 21,500 new cases and 8900 deaths in 2008. In Ontario, approximately 8000 new cases and 3250 deaths occurred in 2008 (1). If CRC is detected early, the five-year survival rate is 93.2%, whereas the five-year survival rate for individuals with metastatic disease is only 8.1% (2). Most patients with early CRCs are asymptomatic and population screening has been shown to be an effective strategy in significantly reducing mortality rates (3-6). Toward this goal, a CRC screening program, ‘ColonCancerCheck’ (www.coloncancercheck.ca), was launched in the province of Ontario in 2008. Through this program, average-risk adults, defined as individuals who are asymptomatic, at least 50 years of age and without any first-degree relatives with a history of CRC, are screened with a fecal occult blood test (FOBT). Any individual with a positive FOBT is then referred to a specialist for colonoscopy.

La gastroscopie après une recherche positive de sang occulte dans les selles et une coloscopie négative : Une analyse systématique et des lignes directrices

HISTORIQUE : Bien des gens qui participent à un programme de dépistage du cancer colorectal (CCR) en population et obtiennent un résultat de recherche de sang occulte dans les selles (RSOS) positif n’ont pas de lésion identifiable à la coloscopie pour justifier ce dépistage positif.

OBJECTIF : Évaluer les données probantes et fournir des recommandations au sujet du recours systématique à l’esophagogastroduodénoscopie (OGD) pour déceler des cancers du transit oesogastroduodénal (TOGD) chez des patients qui participent à un programme de dépistage du CCR positif à une RSOS et négatif à la coloscopie.

MÉTHODOLOGIE : Une analyse systématique a permis d’établir une base de données probantes et d’étayer les recommandations probantes fournies.

RÉSULTATS : Neuf études portaient sur un groupe de patients positifs à une RSOS et négatifs à une coloscopie. Trois études n’ont décelé aucun cas de cancer du TOGD. Quatre études en ont déclaré des cas : trois ont décelé l’un de ces cancers chez 1 % ou moins de la population à l’étude et une en a découvert un cas, qui représentait 7 % du petit sous-groupe de patients positifs à la RSOS et négatifs à la coloscopie. Deux études ne contenaient pas d’information d’issues susceptible d’être reliée spécifiquement au sous-groupe positif à la RSOS et négatif à la coloscopie.

CONCLUSION : L’ensemble de données probantes actuel ne suffit pas pour recommander une OGD systématique afin de déceler les cancers gastriques ou œsophagiens chez les patients positifs à la RSOS et négatifs à la coloscopie dans le cadre d’un programme de dépistage du CCR en population. La décision d’exécuter une OGD doit être personnalisée et se fonder sur le jugement clinique.
A large number of individuals with a positive FOBT do not have an identifiable lesion found at colonoscopy to account for their positive FOBT screen (7). Results of a pilot CRC screening program in the United Kingdom (UK) indicated that 1.9% of those screened for CRC were FOBT positive, and 53% of these patients screened negative at colonoscopy (8). Therefore, approximately 1% of those who presented for CRC population screening were FOBT positive and colonoscopy negative in the UK experience. Results from a French CRC screening program (9) reported that 2.6% of those screened were FOBT positive, of which 37% were negative at colonoscopy. While some of these cases may be attributable to a false-positive FOBT, some may also be attributable to blood loss from upper gastrointestinal (UGI) or small bowel lesions including, but not limited to, possible malignancies – diagnosis of these lesions require further investigation. Presently, only some FOBT-positive/colonoscopy-negative patients are referred for UGI investigations and up to 5.9% of negative colonoscopy investigations may be owing to false-negative test results (10). Currently, there is a lack of consensus regarding whether UGI investigation by esophagogastroduodenoscopy (EGD) is routinely warranted in FOBT-positive/colonoscopy-negative cases. Given the limited endoscopy resources in Ontario and other jurisdictions, it is important to determine whether EGD is appropriate in this clinical context.

The aim of the present systematic review was to evaluate the evidence and provide recommendations concerning the use of routine EGD to detect UGI cancers in men and women participating in a population-based CRC screening program who have had a positive FOBT followed by colonoscopy without identifiable colonic lesions to account for their positive FOBT.

The intended audience for the present guidance document is health professionals, which may include gastroenterologists, family physicians, surgeons and other health care professionals involved in the screening, diagnosis, treatment and follow-up of individuals enrolled in a population-based CRC screening program.

METHODS

The present guideline, developed by Cancer Care Ontario’s (CCO) Program in Evidence-Based Care (PEBC), used the methods of the practice guidelines development cycle (11). The core methodology used to develop the evidentiary base was the systematic review. Evidence was selected by one author who was a methodologist (RC) and reviewed by two other authors (JA and EJI). The reference lists from these sources were also searched for additional trials.

The UGI Screening Working Panel consisted of several gastroenterologists, a family physician, a methodologist and a CCO representative from the CRC screening program.

The present systematic review is a convenient and up-to-date source of the best available evidence examining UGI endoscopic screening subsequent to a positive FOBT and negative colonoscopy. The body of evidence in the review, which forms the basis of the recommendations, is primarily comprised of prospective and retrospective cohort and cross-sectional studies that have evaluated the role of UGI investigation in FOBT-positive/colonoscopy-negative patients. The systematic review and companion recommendations are intended to promote evidence-based practice in Ontario. The PEBC is supported by the Ontario Ministry of Health and Long-Term Care through CCO. All work produced by the PEBC is editorially independent from its funding source.

Literature search strategy

The MEDLINE (1990 through May [week one] 2008) and EMBASE (1990 through week 20, 2008) databases were searched for relevant publications, using terms pertaining to colonoscopy, gastroscopy and gastrointestinal neoplasms. The reference lists of included studies were also searched. The start year of the search was 1990 because it is when evidence regarding screening began to appear in the literature. The full MEDLINE and EMBASE literature search strategies can be found in Appendixes 1 and 2, respectively.

Environmental scan

An environmental scan was conducted in May 2008, to locate published and unpublished documents outside the indexed literature. Documents pertaining to UGI screening for patients who were colonoscopy negative following a positive FOBT in a population-based CRC screening program from Canada and health care organizations in the United States, UK, Australia and New Zealand were searched. The complete list of resources used in the search are presented in Appendix 3.

Study selection criteria

Articles selected for inclusion in the present systematic review were English language reports involving human participants. They included practice guidelines, systematic reviews (with or without meta-analyses), and all publication types that examined the role of UGI screening in patients who had a negative colonoscopy following a positive FOBT. Articles such as letters, editorials, notes, case reports, commentaries and non-systematic reviews, were excluded.

If an EGD was not performed after a negative colonoscopy and patients were followed to determine a new occurrence of UGI cancer, the studies involved were included only if they reported cases of UGI cancers occurring within three years of the positive FOBT. Three years was chosen based on the mean sojourn time for CRC (the time between an undetectable preclinical screening and the clinical phase), reported to be 2.8 years in a Taiwanese study (12) and 2.6 years in a French study (13).

In theory, population screening should include only asymptomatic participants. However, in practice, some individuals presenting for screening are symptomatic, which realistically reflects medical practice. For this reason, articles relating to either symptomatic or asymptomatic patients were retained. At minimum, a group of FOBT-positive/colonoscopy-negative patients were required to be identified in the article.

Internal and external review

All documents produced in collaboration with the PEBC undergo rigorous internal review, including a full data audit as well as copyediting by staff not involved in the development of the document. Before submission of the present report (systematic review and companion guideline) to external review, the report was reviewed by an expert panel that consisted of a group of endoscopists from the Clinical Advisory Committee.
of CCO's Colorectal Cancer Screening Program. In addition, the report was reviewed and approved by the PEBC Report Approval Panel, which consists of two members: an expert methodologist and an oncologist with expertise in clinical and methodology issues.

The PEBC external review process is two pronged and includes a targeted peer review that is intended to obtain direct feedback on the draft report from a small number of specified content experts and a professional consultation that is intended to facilitate dissemination of the final guidance report to Ontario practitioners. Following endorsement by the PEBC Report Approval Panel, the present document was sent to four targeted peer reviewers from across Canada considered to be clinical and/or methodological experts on the topic; written comments were invited. Feedback was also obtained through a brief online survey of health care professionals who are the intended users of the guideline, namely, gastroenterologists, family physicians and surgeons. Written comments were invited from this group as well.

RESULTS

Literature search and environmental scan

The MEDLINE and EMBASE searches resulted in 1558 hits. A total of nine articles met selection criteria and were retained (7,14-21): five from MEDLINE, three from EMBASE and one from the reference list of included studies. The environmental scan did not yield any papers, documents or guidance pertaining to the use of UGI investigations in patients who were FOBT-positive/colonoscopy-negative.

Study characteristics and quality

Five studies that examined the occurrence of gastric cancer following a negative colonoscopy in patients who had positive FOBTs were identified (7,14-17). Of these, two (7,14) were studies in which patients were originally part of a population screening program for CRC. Participants in the Thomas and Hardcastle (14) study who were FOBT-positive/colonoscopy-negative, and were subsequently noted to be asymptomatic, underwent EGD. The outcomes for this small group of patients (n=14) were reported. Although Zappa et al (7) identified a large group of FOBT-positive/colonoscopy-negative patients who did not undergo EGD, they performed a follow-up using database linkage procedures. Because they reported the number of UGI cancers that occurred within a three-year follow-up period, this study was included in the present report. The remaining three studies (15-17) were prospective (15) and retrospective (16,17), in which all FOBT-positive/colonoscopy-negative patients were assessed by EGD for gastric cancer.

Four studies that examined the diagnosis of gastric cancer in patients who had same-day EGD and colonoscopy after a positive FOBT were identified; in these studies, a subgroup of colonoscopy-negative patients could be identified (18-21). Two studies collected data prospectively (18,21), and two studies collected data retrospectively (19,20). Participants in all four studies underwent bidirectional endoscopy. The order of endoscopy was either colonoscopy followed by EGD (20,21), EGD followed by colonoscopy (19) or as determined by institutional availability (18). Because the outcomes related to the subgroup of FOBT-positive/colonoscopy-negative patients were not reported separately in three (18,19,21) of these four studies, only limited information could be obtained from these articles. Table 1 provides a summary of each study.

Five studies used a guaiac FOBT only (15,16,18,20,21), two studies used both a guaiac and an immunochemical FOBT, with patients being tested with a guaiac test until 1995, and with an immunochemical test after 1995 in one study (7), and with one, the other, or both tests in some cases, in the other study (14). Two studies (17,19) did not report the type of FOBT used. Three studies (14,18,20) did not rehydrate samples, while the remaining studies did not indicate whether samples were rehydrated.

Measures of study quality included conflict of interest reporting and the identification of funding sources. Only Hisamuddin et al (20) reported specifically on authors' conflicts of interest and indicated they had no conflicts. No other papers reported on conflicts of interest. Information regarding funding sources was not reported in any of these studies.

Cancer outcomes

Nine studies identified a group of patients who were FOBT-positive/colonoscopy-negative (Table 2). Some studies were limited to patients who were either symptomatic (14) or asymptomatic (15,17), others included both symptomatic and asymptomatic patients (16,18,20), and several studies did not report whether patients were symptomatic (7,19,21). Chen et al (16) categorized their patients into four groups: asymptomatic, symptomatic, severely anemic (which included those who were both symptomatic and severely anemic) and ‘incomplete’, which was a group of patients with incomplete documentation with respect to anemia and/or symptoms. One study (20) did not provide a definition of ‘symptomatic’. In all other studies that included symptomatic patients, symptomatic was defined as including dyspepsia as well as a subset of the following: dysphagia, heartburn, abdominal pain, nausea, vomiting, weight loss and diarrhea (14,16,18). One study (20) reported that patients were required to have at least one positive window to be considered FOBT positive. No other studies reported how many windows were required to be positive for a patient to be considered FOBT positive.

Eight of the studies performed EGD on all FOBT-positive/colonoscopy-negative patients (14-21), whereas one study (7) conducted a large retrospective cohort study of patients who were followed through a cancer registry linkage. Rates of positive findings (not limited to cancers) at EGD ranged from a low of 13% (19) to a high of 43% (14). Of note, this latter study is a report of a very small subgroup (n=14) of symptomatic patients.

Three studies (15,16,20) found no cases of UGI cancer, defined as either gastric or esophageal cancer. Four studies reported cases of UGI cancer. Thomas and Hardcastle (14) found only one case of gastric cancer in their small subgroup of 14 symptomatic FOBT-positive/colonoscopy-negative patients. Bini et al (17) found five cases of UGI cancer (four gastric and one esophageal) in their study of 498 asymptomatic patients, and Zappa et al (7) found 14 cases of gastric cancer in 3555 patients between zero and 35 months following a positive FOBT (unknown if patients were symptomatic or asymptomatic). Zuckerman and Benitez (18) found one case of UGI cancer in their study of 74 asymptomatic and symptomatic FOBT-positive/colonoscopy-negative patients.
This represents 1% or less of the total population studied. The final two studies (19,21) did not provide outcome information that could be specifically related to the FOBT-positive/colonoscopy-negative subgroup. Therefore, no conclusion can be drawn regarding the UGI cancer yield in these studies.

**Probable UGI contributors to a positive FOBT**
Five studies reported outcomes other than UGI cancer outcomes (Table 2). Based on expert opinion, the panel divided these other noncancer outcomes into two groups: probable UGI contributors to a positive FOBT and probable UGI incidental finding. Probable UGI contributors include peptic ulcer disease (stomach, esophagus and duodenum), esophagitis, vascular malformations and large gastric polyps (larger than 1 cm in size). One study of symptomatic patients (14) reported no probable contributors. Among other symptomatic patients (including those with severe anemia), 11% to 21% had a probable UGI contributor to a positive FOBT (16), whereas 7% to 19% of asymptomatic patients had a probable contributor (15-17). The Hisamuddin et al (20) study involved both asymptomatic and symptomatic patients, and reported that 16% of this aggregate group had a probable contributor.

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**TABLE 1**
Summary of studies of fecal occult blood test-positive (FOBT-pos)/colonoscopy-negative (col-neg) patients or studies with an identifiable FOBT-pos/col-neg subgroup of patients

<table>
<thead>
<tr>
<th>Author (reference)</th>
<th>Study type</th>
<th>Study details</th>
<th>Type of FOBT</th>
<th>Guaiac sample hydration status</th>
<th>FOBT-pos/col-neg, patients, n</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thomas and Hardcastle (14)</td>
<td>Prospective</td>
<td>Asymptomatic patients randomly assigned to colorectal cancer screening with FOBT or control group (received no screening) FOBT-pos patients who underwent colonoscopy or flexible sigmoidoscopy combined with double-contrast barium enema EGD performed on FOBT-pos/col-neg patients who were subsequently deemed to be symptomatic</td>
<td>Guaiac and/or immunochemical</td>
<td>Not rehydrated</td>
<td>14</td>
</tr>
<tr>
<td>Hsia and al-Kawas (15)</td>
<td>Prospective</td>
<td>Asymptomatic FOBT-pos/col-neg patients referred for upper endoscopy</td>
<td>Guaiac</td>
<td>Not reported</td>
<td>70</td>
</tr>
<tr>
<td>Chen et al (16)</td>
<td>Retrospective</td>
<td>FOBT-pos/col-neg patients who were asymptomatic, symptomatic, anemic or had incomplete documentation and were referred for upper endoscopy</td>
<td>Guaiac</td>
<td>Not reported</td>
<td>211</td>
</tr>
<tr>
<td>Bini et al (17)</td>
<td>Retrospective</td>
<td>Asymptomatic FOBT-pos/col-neg patients who underwent upper endoscopy</td>
<td>Not reported</td>
<td>Not reported</td>
<td>498</td>
</tr>
<tr>
<td>Zappa et al (7)</td>
<td>Retrospective</td>
<td>Patients attending FOBT screening identified from a screening database FOBT-pos patients underwent colonoscopy Authors report on a FOBT-pos/col-neg subgroup EGD not performed but patients followed through databases and registries Authors report the number of upper gastrointestinal cancers in the first 3 years after a positive FOBT</td>
<td>Guaiac or immunochemical</td>
<td>Not reported</td>
<td>3555</td>
</tr>
<tr>
<td>Hisamuddin et al (20)</td>
<td>Retrospective</td>
<td>FOBT-pos patients who underwent same-day bidirectional endoscopy Sequence was colonoscopy followed by EGD Authors report outcomes from an FOBT-pos/col-neg subgroup</td>
<td>Guaiac</td>
<td>Not rehydrated</td>
<td>70</td>
</tr>
<tr>
<td>Zuckerman and Benitez (18)</td>
<td>Prospective</td>
<td>FOBT-pos or iron-deficiency anemia patients who underwent bidirectional endoscopy Sequence dependent on scheduling and could be separated by up to 14 days Authors identify a group of FOBT-pos/col-neg patients but do not report separate outcomes for this subgroup</td>
<td>Guaiac</td>
<td>Not rehydrated</td>
<td>74</td>
</tr>
<tr>
<td>Ali et al (19)</td>
<td>Retrospective</td>
<td>FOBT-pos patients underwent same-day bidirectional endoscopy Sequence was EGD followed by colonoscopy in 85% of cases Authors identify a group of FOBT-pos/col-neg patients but do not report separate outcomes for this subgroup</td>
<td>Not reported</td>
<td>Not reported</td>
<td>125</td>
</tr>
<tr>
<td>Stray and Weberg (21)</td>
<td>Prospective</td>
<td>FOBT-pos patients with or without iron-deficiency anemia, solely iron-deficiency anemia or solely iron-deficiency and underwent same-day bidirectional endoscopy Sequence was colonoscopy followed by EGD Authors identify a group of FOBT-pos/col-neg patients but do not report separate outcomes for this subgroup</td>
<td>Guaiac</td>
<td>Not reported</td>
<td>146</td>
</tr>
</tbody>
</table>

EGD Esophagogastroduodenoscopy
None of the remaining four studies (7,18,19,21) reported on noncancer outcomes separately for their FOBT-positive/colonoscopy-negative subgroups.

**Probable UGI incidental findings**

Probable UGI incidental findings were defined by the panel as lesions not likely to contribute to a positive FOBT. These include Barrett’s esophagus, gastric and duodenal erosions, duodenitis, jejunitis, esophageal and gastric varices, esophageal stricture, duodenal adenoma, nonerosive esophagitis, benign gastroduodenal disease and small gastric polyps (<1 cm). Among asymptomatic patients (including those with severe anemia in one study), 24% to 36% had a probable UGI incidental finding at EGD (14,16), while 10% to 36% of asymptomatic patients had a probable incidental finding (15-17). Hisamuddin et al (20) had both asymptomatic and symptomatic patients in their study and reported that 20% of this aggregate group had a probable incidental finding.

None of the other four studies (7,18,19,21) reported on noncancerous outcomes separately for their FOBT-positive/colonoscopy-negative subgroups.

**Effect of anemia**

There were few data from these studies relating to anemia as a possible predictor of EGD results (in addition to positive FOBT and negative colonoscopy). Furthermore, the outcomes reported differed among studies and could not be combined.

**Effect of nonsteroidal anti-inflammatory drug use**

There were few data from these studies relating to the investigation of acetylsalicylic acid or nonsteroidal anti-inflammatory drug (NSAID) use as a possible predictor of EGD results (in addition to positive FOBT and negative colonoscopy). Furthermore, the outcomes reported differed among studies and could not be combined.

**Development of recommendations; internal and external review**

Once the data were collected and collated, the panel reconvened to formulate the guideline recommendation(s) based on the available evidence. At this point, the document was sent for internal review followed by external review as described in the Methods section.

All feedback received through the internal and external review process was discussed by the panel and changes to the document were made, where appropriate. The main comment made by several reviewers was that a 1% yield for the detection of UGI malignancy may make it worth performing EGD given that the detection rate of CRC in CRC screening programs would be similar. The panel believed that this warranted comment and, consequently, a revision to the Discussion section was made.

**DISCUSSION**

The current management of patients who undergo screening for CRC and test FOBT positive and colonoscopy negative is inconsistent. As a greater proportion of the population complies with guidelines for CRC screening, there will be an increasing and perhaps substantial number of such patients who fall into this category. However, there are relatively few studies that used sound study design and appropriate outcome assessment to fully address whether EGD is warranted in this situation. The evidence base compiled for the present document consists of four prospective and five retrospective studies. Two of these studies (19,21) only identified a group of FOBT-positive/colonoscopy-negative patients with a positive EGD, but provided no information regarding the endoscopic findings, making the results uninterpretable. The current document does not address the entire issue of how to manage FOBT-positive/colonoscopy-negative patients, but examines whether EGD is warranted to detect UGI cancer in this group of patients.
In the remaining seven studies, the prevalence of UGI cancer was very low. Three studies (15,16,20) found no UGI cancers. Three studies (7,17,18) observed UGI cancer rates of 1% or lower, and one study (14) noted a UGI cancer rate of 7%, representing one case out of 14 symptomatic patients. It should be noted that while a 1% yield of UGI cancer may seem comparable to the yield of colon cancers detected at screening colonoscopy using FOBT, the quality of the available studies evaluating EGD in FOBT-positive/colonoscopy-negative patients are not as scientifically rigorous as those evaluating the use of FOBT in CRC screening. The gastroscopy studies contain heterogeneous samples that are not truly representative of a screening population and the number of patients evaluated is smaller than that used in the landmark CRC screening studies that established the benefit of FOBT.

Diagnostic findings at EGD other than UGI cancer were more prevalent. Overall, a probable UGI contributor to the positive FOBT was reported in 7% to 21% of FOBT-positive/colonoscopy-negative patients. Probable contributors were reported to be peptic ulcer disease, esophagitis, vascular malformations and gastric polyps (greater than 1 cm in size). It is not known if gastric polyps greater than 1 cm in size cause a positive FOBT. These larger gastric polyps were only reported in two studies (17,20), and their inclusion as a probable UGI contributor to a positive FOBT did not change any conclusions of the present report. The proportion of cases with findings that were reported as likely to be incidental (and unlikely to account for a positive FOBT) occurred in 10% to 36% of patients. These incidental findings included Barrett’s esophagus, gastric and duodenal erosions, gastritis, duodenitis, jejunitis, esophageal and gastric varices, esophageal stricture, duodenal adenoma, benign gastroduodenal disease and gastric polyps. The variability in the descriptions of UGI lesions other than cancer is likely the result of the variations between studies in defining what constitutes a positive EGD.

There were very few data regarding EGD results and the presence or absence of anemia, and even fewer data regarding EGD and NSAID use. Moreover, the papers that did report on these variables all disclosed different outcomes. Because most patients who undergo screening are older than 50 years of age, many will be taking acetylsalicylic acid for cardiovascular disease prevention and/or NSAIDs for arthritis and analgesia. In addition, anemic patients are unique and should be considered symptomatic and, therefore, do not fall under the auspices of screening programs. Anemia is a ‘red-flag’ sign that requires further investigation. These two groups of patients would benefit from further study as separate subgroups.

The body of literature examining the controversial issue of performing routine EGD in FOBT-positive/colonoscopy-negative patients is sparse. The data gathered from the studies in the present systematic review suggest that the number of UGI cancers found in FOBT-positive/colonoscopy-negative patients is small – in the order of 1% or less – although the rate of other UGI findings was higher. However, although the risk associated with EGD is small (approximately 0.03% for perforation [22,23]), other factors related to cost and endoscopic resources are significant, considering the large numbers of patients who will emerge from CRC screening programs with a positive FOBT and a negative colonoscopy. Performing routine EGD in these patients would significantly add to the cost of screening programs while potentially adding little value with respect to the effectiveness of screening for UGI cancer.

**CONCLUSION**

After examining the available evidence, the panel provided the following recommendation:

The current body of evidence is insufficient to recommend for or against routine EGD as a means of detecting gastric or esophageal cancers for patients who are FOBT-positive/colonoscopy-negative in a population-based CRC screening program. The decision to perform EGD should be individualized and based on clinical judgement.

The key evidence supporting this recommendation is as follows:

- Four prospective (14,15,18,21) and five retrospective (7,16,17,19,20) studies of patients who were FOBT-positive/colonoscopy-negative and underwent EGD. Of these, two studies (19,21) reported positive EGD but no information about endoscopic findings, and several studies did not document the presence of anemia, UGI symptoms or the use of NSAIDs.

- Based on this limited evidence, EGD had a low yield for UGI cancer, generally 1% or less, even in symptomatic or severely anemic patients. The yield for detecting nonmalignant findings potentially contributing to a positive FOBT was 11% to 21%, while the yield for incidental findings unlikely contributing to a positive FOBT was 10% to 36%. There were very few data regarding EGD results in the context of anemia or NSAID use.

A recommendation regarding the use of EGD for the detection of noncancerous pathology is not provided because it is beyond the scope of the present review.

Furthermore, adequately powered studies are needed to investigate the incidence of gastric or esophageal cancer in patients enrolled in a population-based CRC screening program who are FOBT positive and colonoscopy negative.

**CONFLICTS OF INTEREST:** All authors declare no conflicts of interest.

**APPENDIX 1**

**MEDLINE search strategy**

**Search 1**

1. Colorectal neoplasms/di
2. exp Colonoscopy/
3. 1 or 2
4. Digestive System Diseases/di
5. Gastrointestinal neoplasms/di
6. Gastrointestinal diseases/di
7. Stomach ulcer/di
8. Stomach neoplasms/di
9. Peptic ulcer/di
10. Peptic ulcer hemorrhage/di
11. Liver disease/di
12. or/4–11
13. exp Mass Screening/
14. 3 and 12 and 13
15. limit 18 to english language
16. limit 19 to yr = “1990–2008”
EGD after positive FOBT and negative colonoscopy

APPENDIX 2
EMBASE search strategy

Search 1
1. Colorectal cancer/di
2. exp COLONOSCOPY/
3. exp Cancer Screening/
4. exp Occult Blood/
5. exp GASTROSCOPY/
6. exp ESOPHAGOGASTRODUODENOSCOPY/
7. exp Gastrointestinal Endoscopy/
8. Digestive system cancer/di
9. Stomach ulcer/di
10. Esophagus cancer/di
11. Liver disease/di
12. Peptic ulcer/di
13. or/5–14
14. 1 and 2 and 3 and 4 and 15
15. limit 16 to English language

Search 2
1. Colorectal cancer/di
2. exp COLONOSCOPY/
3. 1 or 2
4. Digestive System Cancer/di
5. gastrointestinal disease/di
6. stomach ulcer/di
7. stomach cancer/di
8. upper gastrointestinal bleeding/di
9. esophagus cancer/di
10. peptic ulcer/di
11. liver disease/di
12. or/4–11
13. 3 and 12
14. limit 13 to English language
15. limit 14 to yr = “1990–2008”

Search 3
1. exp GASTROSCOPY/
2. exp ESOPHAGOGASTRODUODENOSCOPY/
3. exp Gastrointestinal Endoscopy/
4. 1 or 2 or 3
5. Digestive System Cancer/di
6. Gastrointestinal disease/di
7. stomach ulcer/di
8. stomach cancer/di
9. Upper gastrointestinal bleeding/di
10. Esophagus cancer/di
11. Peptic ulcer/di
12. Liver disease/di
13. or/5–12
14. exp Mass Screening/
15. exp Cancer Screening/
16. 14 or 15
17. 4 and 13 and 16
18. limit 17 to English language
19. limit 18 to yr = “1990–2008”

APPENDIX 3
Environmental scan (National Guideline Clearinghouse)

International guideline developers:
- National Institute for Health Excellence (NICE) (United Kingdom [UK]) – NICE guidance
- The Scottish Intercollegiate Guidelines Network (SIGN) (UK) – SIGN guidelines
- American Society of Clinical Oncology (ASCO) (United States) – ASCO guidelines
- National Comprehensive Cancer Network (NCCN) (United States) – NCCN home (consensus based)
- National Health and Medical Research Council (Australia) – Cancer guidelines
- New Zealand Guidelines Group – Guidelines
- Canadian provincial cancer agencies:
  - BC Cancer Agency – Cancer management guidelines
  - Alberta Cancer Board – Treatment guidelines
  - Saskatchewan Cancer Agency – Follow-up guidelines
  - Cancer Care Manitoba (CCM) – CCM home
  - Cancer Care Nova Scotia – Guidelines

National cancer agencies (UK, Australia, New Zealand):
- New Zealand Cancer Control Trust
- The Cancer Council Australia
- National Cancer Control Initiative (Australia)
- The Collaboration for Cancer Outcomes Research and Evaluation (Australia)
- State Government of Victoria, Australia
- Peter MacCallum Cancer Centre (Australia)
- Medical Oncology Group of Australia
- Cancer UK
REFERENCES


