Colorectal cancer screening: A guide to the guidelines

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Convincing evidence of the effectiveness of fecal occult blood testing and sigmoidoscopy in reducing mortality from colorectal cancer has led a number of groups to publish statements or guidelines that endorse colorectal cancer screening (1,2). These published statements, in conjunction with the new coverage for colorectal cancer screening for Medicare beneficiaries in the United States, have combined to make colorectal cancer screening the standard of care for medical practice in the United States in 1998. Hopefully, private insurers will feel compelled by these steps to provide coverage for screening. The percentage of eligible screenees in the United States who have been screened is still low. Major education efforts from the Centers for Diseases Control, the Digestive Disease National Coalition, individual gastro-intestinal societies and patient advocacy groups are being undertaken to inform the public and primary care physicians about the importance of undergoing and carrying out colorectal cancer screening programs. Education combined with acceptable reimbursement are major factors that will drive increased use of screening. Malpractice litigation will be an additional factor that will increase physician compliance with offering screening and stimulate payers to provide coverage for screening.

Although the issue of whether screening works is considered settled, the issue of how screening should be done is evolving. Rigorous evidence of effectiveness exists only for the two most recent guidelines for colorectal cancer screening are those of the Agency for Healthcare Policy and Research, and the American Cancer Society. The guidelines are similar in many regards and reflect current literature, consensus opinion and compromise between members of multidisciplinary panels. The emphasis of both guidelines is to increase the options available for colorectal cancer screening. Increasing choice should expand the attractiveness of colorectal cancer screening to more patients and physicians, and the development of guidelines should help compel payers to provide reimbursement for colorectal cancer screening. These guidelines are summarized and evaluated as they pertain to colorectal cancer screening.

Key Words: Colorectal cancer; Endoscopy; Fecal occult blood testing; Sigmoidoscopy
TABLE 1
Options for screening average-risk persons: Agency for Health Care Policy and Research, and American Cancer Society guidelines

<table>
<thead>
<tr>
<th>Method</th>
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<tbody>
<tr>
<td>Annual fecal occult blood test</td>
</tr>
<tr>
<td>Flexible sigmoidoscopy every 15 years</td>
</tr>
<tr>
<td>Annual fecal occult blood test plus</td>
</tr>
<tr>
<td>flexible sigmoidoscopy every five years</td>
</tr>
<tr>
<td>Double contrast barium enema every 5 to</td>
</tr>
<tr>
<td>10 years</td>
</tr>
<tr>
<td>Colonoscopy every 10 years</td>
</tr>
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*American Cancer Society guidelines do not include fecal occult blood test by itself*

The average-risk population is defined as age 50 years or older and lacking other known risk factors. The guidelines from the American Health Care Policy and Research (AHCPR) (1) and the American Cancer Society (ACS) as they pertain to screening (2). It is no accident that these guidelines are very similar. Clearly, the AHCPR guidelines significantly influenced the ACS guidelines. The present article serves to familiarize readers with the content of the guidelines and to point out for the reader specific strengths and weaknesses of the guidelines as perceived by the author.

SUPPORT SCREENING OPTIONS

The philosophy of the two most recently published major guidelines is to create a menu of choices for patients, physicians and payers to choose from. Clearly these choices are not equal from the perspectives of effectiveness, cost or risk; these differences provide the rationale for creating a menu of options.

The present review compares and contrasts the new guidelines from the Agency for Health Care Policy and Research (AHCPR) (1) and the American Cancer Society (ACS) as they pertain to screening (2). It is no accident that these guidelines are very similar. Clearly, the AHCPR guidelines significantly influenced the ACS guidelines. The present article serves to familiarize readers with the content of the guidelines and to point out for the reader specific strengths and weaknesses of the guidelines as perceived by the author.

**AVERAGE-RISK SCREENING: EVIDENCE TO SUPPORT SCREENING OPTIONS**

The average-risk population is defined as age 50 years or older and lacking other known risk factors. The guidelines from the AHCPR and ACS present a menu of options for average-risk screening (Table 1). The options are identical, except that the ACS declined to recommend fecal occult blood testing by itself, citing the relatively low mortality reductions that have been associated with its use. Table 2 shows the categories of data that support the screening modalities listed in the menus. For example, evidence supporting the use of fecal occult blood testing has been published in three randomized, controlled trials (5-7) and in a case-control study (9), and a large amount of cross-sectional data have been collected. Cross-sectional studies provide prevalence data, useful for describing the expected yield of a screening modality. For sigmoidoscopy, there are no data from randomized, controlled trials, but results from two strong case-control studies support its effectiveness (10,11). These studies were largely performed using rigid sigmoidoscopy but have been widely extrapolated to flexible sigmoidoscopy. Cross-sectional data obtained by using flexible sigmoidoscopy from well over 10,000 patients have accumulated (12). Thus, the single time yield of screening flexible sigmoidoscopy in average-risk persons is well understood.

For colonoscopy, data from randomized, controlled trials or case-control studies are not available. A large amount of
cross-sectional data have accumulated, including data from five studies in average-risk persons (13–17), more than 10 studies (12,18-28) in persons with a positive family history of cancer that does not meet the criteria for hereditary non-polyposis colon cancer (HNPCC) and a number more in persons with a history of HNPCC (29–33). For DCBE, there are essentially no available data, although the original screening colonoscopy study in average-risk persons included 90 patients who also underwent DCBE (14). In one study of 738 persons using single contrast barium enema, the yield in screening populations was dramatically lower than has been detected using colonoscopy (34).

OBSERVED INTERVALS
Data from a randomized, controlled trial of fecal occult blood testing using rehydrated slides showed effectiveness at one year (5). In two studies that did not use rehydration, fecal occult blood testing at an interval of two years produced mortality reductions of 15% and 18% (6,7). The recommended interval in both guidelines was one year (1,2).

With flexible sigmoidoscopy, the recommended interval is five years. The reason why the United States Federal Government provides coverage every four years is uncertain but may reflect previous guidelines that recommended flexible sigmoidoscopy at three- to five-year intervals, and the need to provide coverage at a single designated time point that was arrived at by averaging the three- to five-year interval. The longest observed interval in which a group of average-risk asymptomatic individuals who were initially screened with flexible sigmoidoscopy and found to be negative, and then rescreened at a later date is 3.4 years (35). Collectively, these data support a very low yield of repeat screening at both one and three years, and support a move to a five-year interval. Evidence to support a longer interval also arises from the Kaiser case-control study of sigmoidoscopy, in which the protective effect of a sigmoidoscopy appeared to last for at least 10 years (10). The Prostate, Lung, Colorectal and Ovarian (PLCO) Cancer Screening Trial in the United States is about to begin accumulating observational data on flexible sigmoidoscopy at five years, which may support a move to even longer intervals.

INTERVALS FOR DCBE AND COLONOSCOPY
For DCBE, the recommended interval in the guidelines is five to 10 years. However, there are no observational data to support this because even the initial yield with barium enema in a screening population has been determined only in studies using single contrast barium enema and in no studies using DCBE. The intervals are based on estimates of sensitivity.

For colonoscopy, the recommended interval is 10 years. It is of interest that the interval for flexible sigmoidoscopy is five years and the interval for colonoscopy is 10 years because the strategies are fundamentally similar (endoscopic screening). The shorter interval for flexible sigmoidoscopy may reflect its lower cost. The longest observed interval in a screening population of individuals who were initially colonosced and were negative, and then recolonosced is 5.5 years (36). The incidence of new adenomas was 27%, but the incidence of ‘advanced’ adenomas (those with villous or tubulovillous histology, high-grade dysplasia or size greater than 1 cm) was less than 1%. These data, plus those from the Kaiser case-control study (10) showing that the protective effect of a sigmoidoscopy is at least 10 years, helped lend support for the 10-year interval for colonoscopy.

EFFECTIVENESS OF INDIVIDUAL OPTIONS: FECAL OCCULT BLOOD TEST
In randomized, controlled trials, fecal occult blood testing on an annual to a semiannual basis reduced mortality by 15% to 33%. Thus, ironically, while fecal occult blood testing provided the most rigorous evidence of the effectiveness of colorectal cancer screening, its overall impact was only modest. Fecal occult blood testing has the lowest up front cost of any screening method, but the downstream costs associated with evaluation of positive fecal occult blood tests and the cancer care costs associated with this strategy are substantial (37). A recent guideline made suggestions for the use of fecal occult blood tests in clinical practice (Table 3) (38). The negative predictive value of colonoscopy for cancer is very high. Thus, after a negative colonoscopy of the cecum by a competent endoscopist, with a good bowel preparation, all colorectal cancer screening should be stopped for at least five years.

FLEXIBLE SIGMOIDOSCOPY
The advantages of flexible sigmoidoscopy are its low cost, low risk, high accuracy over the area of colon examined, no need for sedation, easy preparation (enemas only in some practices) and a proven long protective effect (10). The only disadvantage is that it fails to examine the entire colon. The flexible sigmoidoscope reaches 50% to 60% of colon cancers in adenomas. Forty per cent of all colorectal cancers arise proximal to the splenic flexure (39), an area of the colon that is typically outside the reach of 60 cm flexible sigmoidoscopes. Only 25% of persons with cancer proximal to the splenic flexure have an adenomatous polyp distal to the splenic flexure. Thus, at least 30% of all persons undergoing screening flexible sigmoidoscopy have a normal examination (40). The overall effectiveness of combining fecal occult blood testing with flexible sigmoidoscopy remains uncertain, though estimates of mortality reduction of around 80% seem reasonable (37).
TABLE 4
Recommmendations for screening persons with a positive family history less than hereditary nonpolyposis colon cancer

<table>
<thead>
<tr>
<th>Family history</th>
<th>Recommendation</th>
</tr>
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<tbody>
<tr>
<td>First degree relative with cancer or adenoma diagnosed at age younger than 60 years, or two or more first degree relatives with cancer and/or adenomas</td>
<td>Colonoscopy every three to five years beginning at age 40 years, or 10 years younger than youngest affected relative, whichever comes first</td>
</tr>
<tr>
<td>Single first degree relative with cancer or adenoma diagnosed at age 60 years or older</td>
<td>Same options as average-risk, but begin screening at age 40 years</td>
</tr>
</tbody>
</table>

DCBE
There are no data on the sensitivity of DCBE in a screening population. Data on sensitivity in symptomatic populations are quite mixed (41) and cannot be extrapolated to screening populations, which have a dramatically different spectrum of disease. The Office of Technology Assessment, an independent panel, estimated that the sensitivity of DCBE for large polyps and cancer in a screening population might be 70%, although they included a calculation at 50% in their cost effectiveness model (41). The sensitivity may be in the range of 50%, as was demonstrated in the National Polyp Study (42). This study is perhaps the best prospective blinded evaluation of the sensitivity of DCBE ever performed. Although the study was of a surveillance population, its disease spectrum and prevalence were very similar to those of a screening population. In that study, DCBE missed 50% of all adenomas greater than 1 cm in size. A particular difficulty with DCBE is that uncertainty regarding its sensitivity and the complete lack of observational data make determining intervals quite difficult. The recommended interval in the guidelines of every five to 10 years is a guess at the safe intervals, likely based on over-inflated estimates of sensitivity (1). In particular, the longer interval of 10 years seems ill advised. The specificity of barium enema in clinical practice is 85% to 90%, so that short intervals result in a substantial portion of the screened population undergoing colonoscopy in any case.

Colonoscopy can now be performed by experts in many cases without sedation (46,47), although the practice is widespread in some countries (48) but not in the United States. Perforation from diagnostic colonoscopy has been less than one in 3000 in studies published in the 1990s (5,49,50). Complications associated with polypectomy are inevitable during colonoscopy and can be only partly reduced by proper technique. Because colonoscopy detects more polyps than other modalities, there will continue to be a higher complication rate associated with its use. However, to the extent that any initial strategy identifies colon polyps, it leads to colonoscopy. Therefore, to the extent that other strategies effectively identify polyps, they do not avoid the complications of colonoscopy and polypectomy.

FAMILY HISTORY OF COLORECTAL NEOPLASIA LESS THAN HNPCC
Persons with even one first degree relative with colorectal cancer incur about a twofold increased lifetime risk of developing colorectal cancer (51-54). Both the risk of colorectal cancer and the prevalence of adenomas at screening appear to be increased in persons with multiple first degree relatives with either cancer or adenoma (55), or persons with relatives diagnosed at a young age (younger than 60 years) with cancer or adenoma. The AHCPR guidelines recommend that persons with a first degree relative who has colorectal cancer or with adenomatous polyps at age younger than 60 years should be offered the same menu of options as average-risk persons but that screening should begin at age 40 years. The basis for this recommendation is that the incidence of colorectal cancer in such persons at age 40 years appears to be similar to the average-risk population risk at age 50 years (1). The guidelines state that “if the close relative was diagnosed with colorectal cancer before the age of 55 years or with an adenomatous polyp before age 60 years, special efforts should be made to assure that screening takes place”. The guidelines of the ACS differ in some regards. In persons with a single first degree relative with cancer or adenomatous polyps diagnosed after age 60 years, or with relatives (ie, second or higher degree relatives) with colorectal cancer, the recommendation is that average-risk options be offered to the patient and that the physician may consider screening before the age of 50 years. If however, there is colorectal cancer or adenomas in a first degree relative younger than 60 years of age or in two or more first degree relatives, the ACS recommends total colon evaluation (TCE). TCE comprises either colonoscopy or DCBE, and the recommended interval of examination is five years. The ACS guidelines are preferred because they provide for TCE in patients with a higher risk family history. The unfortunate aspect of the ACS guidelines is the presentation of colonoscopy and DCBE as equivalent, which is hardly the case (36). In addition, there are no data on the use of DCBE in this population, whereas at least cross-sectional studies are available for colonoscopy (see above). Furthermore, the ACS guidelines state that DCBE can be performed without flexible sigmoidoscopy, despite clear evidence in the literature of major deficiencies of

Colorectal Screening in Average-Risk Individuals

Table 4 outlines the recommendations for screening persons with a positive family history less than hereditary nonpolyposis colon cancer. It categorizes the family history into three groups: single first degree relative with cancer or adenomas at age 60 years or older, or more first degree relatives with cancer and/or adenomas; single first degree relative with cancer or adenomas diagnosed at age 60 years or older; and first degree relative with cancer or adenomas diagnosed at age younger than 60 years, or two or more first degree relatives with cancer and/or adenomas.

The ACS guidelines recommended total colon evaluation (TCE) comprising either colonoscopy or DCBE with an interval of every five to 10 years. The AHCPR guidelines recommended that persons with a first degree relative with colorectal cancer or with adenomatous polyps at age younger than 60 years should be offered the same menu of options as average-risk persons but that screening should begin at age 40 years. The basis for this recommendation is that the incidence of colorectal cancer in such persons at age 40 years appears to be similar to the average-risk population risk at age 50 years (1). The guidelines state that “if the close relative was diagnosed with colorectal cancer before the age of 55 years or with an adenomatous polyp before age 60 years, special efforts should be made to assure that screening takes place”. The guidelines of the ACS differ in some regards. In persons with a single first degree relative with cancer or adenomatous polyps diagnosed after age 60 years, or with relatives (ie, second or higher degree relatives) with colorectal cancer, the recommendation is that average-risk options be offered to the patient and that the physician may consider screening before the age of 50 years. If however, there is colorectal cancer or adenomas in a first degree relative younger than 60 years of age or in two or more first degree relatives, the ACS recommends total colon evaluation (TCE). TCE comprises either colonoscopy or DCBE, and the recommended interval of examination is five years. The ACS guidelines are preferred because they provide for TCE in patients with a higher risk family history. The unfortunate aspect of the ACS guidelines is the presentation of colonoscopy and DCBE as equivalent, which is hardly the case (36). In addition, there are no data on the use of DCBE in this population, whereas at least cross-sectional studies are available for colonoscopy (see above). Furthermore, the ACS guidelines state that DCBE can be performed without flexible sigmoidoscopy, despite clear evidence in the literature of major deficiencies of...
DCBE in the rectosigmoid colon (57-59). Table 4 shows results from my own practice with regard to persons with a positive family history.

HNPPC

HNPPC is characterized by the development of colorectal cancer at an early age, with a tendency toward right-sided lesions. The syndrome has autosomal dominant inheritance and results from germline mutations in one of four mismatch repair genes. Cancers also occur in a number of other organs, particularly the female genital tract. Identification of affected kindreds can be difficult because there is no absolute definition. A history with high predictive value for a mismatch repair mutation is shown in Table 5. However, lesser histories of colorectal cancer may be present, particularly if cancers have occurred in other organs. Therefore, taking a detailed family history is necessary to identify adequately HNPPC kindreds. Traditional screening is by colonoscopy every two years beginning at age 20 years and continuing until age 40 years, and annually thereafter. This is the only group other than persons with ulcerative colitis for whom annual colonoscopy is still appropriate. Genetic testing is available and is typically effective in about 50% of kindreds (60). When a family history is identified, genetic testing should be offered in the context of genetic counselling.

FAMILIAL ADENOMATOUS POLYPOSIS

Familial adenomatous polyposis accounts for less than 1% of the colorectal cancer burden in the United States. A detailed description of this syndrome is not provided here. When a proband is identified, genetic testing should be offered in conjunction with a genetic counsellor. The genetic test is positive in about 80% of probands (61). It can then be used with complete accuracy to identify family members who are affected. Family members who test positive or all family members in a kindred in whom the genetic test is negative should undergo flexible sigmoidoscopy on a yearly basis beginning at puberty. If the proband's genetic test is positive, family members with a negative genetic test can be screened according to average-risk guidelines.

CONCLUSIONS

For average-risk individuals, the current guidelines create a menu of options that differ with respect to effectiveness, up front costs and risk. The choice of test depends on the relative importance placed on these factors by patients, physicians and payers.

Persons with certain family histories of either colorectal cancer or adenomas are at substantially increased risk, and should be screened with colonoscopy at an earlier age and more frequent intervals than the average-risk person. Genetic testing is a consideration only when individuals are identified who have either the familial adenomatous polyposis phenotype or a combination of personal and family history with high predictive value for a mismatch repair mutation (HNPPC).

REFERENCES

42. Zauber A. Presentation at Digestive Disease Week, New Orleans, May 1994.
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