Screening for Colorectal Cancer
US Preventive Services Task Force Recommendation Statement

US Preventive Services Task Force

IMPORTANCE Colorectal cancer is the third leading cause of cancer death for both men and women, with an estimated 52,980 persons in the US projected to die of colorectal cancer in 2021. Colorectal cancer is most frequently diagnosed among persons aged 65 to 74 years. It is estimated that 10.5% of new colorectal cancer cases occur in persons younger than 50 years. Incidence of colorectal cancer (specifically adenocarcinoma) in adults aged 40 to 49 years has increased by almost 15% from 2000-2002 to 2014-2016. In 2016, 26% of eligible adults in the US had never been screened for colorectal cancer and in 2018, 31% were not up to date with screening.

OBJECTIVE To update its 2016 recommendation, the US Preventive Services Task Force (USPSTF) commissioned a systematic review to evaluate the benefits and harms of screening for colorectal cancer in adults 40 years or older. The review also examined whether these findings varied by age, sex, or race/ethnicity. In addition, as in 2016, the USPSTF commissioned a report from the Cancer Intervention and Surveillance Modeling Network Colorectal Cancer Working Group to provide information from comparative modeling on how estimated life-years gained, colorectal cancer cases averted, and colorectal cancer deaths averted vary by different starting and stopping ages for various screening strategies.

POPULATION Asymptomatic adults 45 years or older at average risk of colorectal cancer (ie, no prior diagnosis of colorectal cancer, adenomatous polyps, or inflammatory bowel disease; no personal diagnosis or family history of known genetic disorders that predispose them to a high lifetime risk of colorectal cancer [such as Lynch syndrome or familial adenomatous polyposis]).

EVIDENCE ASSESSMENT The USPSTF concludes with high certainty that screening for colorectal cancer in adults aged 50 to 75 years has substantial net benefit. The USPSTF concludes with moderate certainty that screening for colorectal cancer in adults aged 45 to 49 years has moderate net benefit. The USPSTF concludes with moderate certainty that screening for colorectal cancer in adults aged 76 to 85 years who have been previously screened has small net benefit. Adults who have never been screened for colorectal cancer are more likely to benefit.

RECOMMENDATION The USPSTF recommends screening for colorectal cancer in all adults aged 50 to 75 years. (A recommendation) The USPSTF recommends screening for colorectal cancer in adults aged 45 to 49 years. (B recommendation) The USPSTF recommends that clinicians selectively offer screening for colorectal cancer in adults aged 76 to 85 years. Evidence indicates that the net benefit of screening all persons in this age group is small. In determining whether this service is appropriate in individual cases, patients and clinicians should consider the patient’s overall health, prior screening history, and preferences. (C recommendation)


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Summary of Recommendations

| Adults aged 50 to 75 years | The USPSTF recommends screening for colorectal cancer in all adults aged 50 to 75 years. | A |
| Adults aged 45 to 49 years | The USPSTF recommends screening for colorectal cancer in adults aged 45 to 49 years. | B |
| Adults aged 76 to 85 years | The USPSTF recommends that clinicians selectively offer screening for colorectal cancer in adults aged 76 to 85 years. Evidence indicates that the net benefit of screening all persons in this age group is small. In determining whether this service is appropriate in individual cases, patients and clinicians should consider the patient’s overall health, prior screening history, and preferences. | C |

See the Summary of Recommendation figure.

Importance

Colorectal cancer is the third leading cause of cancer death for both men and women, with an estimated 52,980 persons in the US projected to die of colorectal cancer in 2021. It is most frequently diagnosed among persons aged 65 to 74 years. It is estimated that 10.5% of new colorectal cancer cases occur in persons younger than 50 years. Incidence of colorectal cancer (specifically adenocarcinoma) in adults aged 40 to 49 years has increased by almost 15% from 2000-2002 to 2014-2016. In 2016, 25.6% of eligible adults in the US had never been screened for colorectal cancer and in 2018, 31.2% were not up to date with screening.

USPSTF Assessment of Magnitude of Net Benefit

The US Preventive Services Task Force (USPSTF) concludes with high certainty that screening for colorectal cancer in adults aged 50 to 75 years has substantial net benefit. The USPSTF concludes with moderate certainty that screening for colorectal cancer in adults aged 45 to 49 years has moderate net benefit. The USPSTF concludes with moderate certainty that screening for colorectal cancer in adults aged 76 to 85 years who have been previously screened has small net benefit. Adults who have never been screened for colorectal cancer are more likely to benefit.

Assessment of Risk

Age is one of the most important risk factors for colorectal cancer, with incidence rates increasing with age and nearly 94% of new cases of colorectal cancer occurring in adults 45 years or older. Rates of colorectal cancer incidence are higher in Black adults and American Indian and Alaskan Native adults, persons with a family history of colorectal cancer (even in the absence of any known inherited syndrome such as Lynch syndrome or familial adenomatous polyposis), men, and persons with other risk factors (such as obesity, diabetes, long-term smoking, and unhealthy alcohol use). However, all adults 45 years or older should be offered screening, even if these risk factors are absent.

Screening Tests

The risks and benefits of different screening tests vary. See Table 1 for characteristics of recommended screening strategies, which may include combinations of screening tests. Because of limited available evidence, the USPSTF recommendation does not include serum tests, urine tests, or capsule endoscopy for colorectal cancer screening. Recommended stool-based and direct visualization screening tests are described below.

Stool-Based Tests

Stool-based tests include the high-sensitivity guaiac fecal occult blood test (gFOBT), fecal immunochemical test (FIT), and stool DNA test. Both high-sensitivity gFOBT and FIT detect blood in the stool; however, they use different methods. High-sensitivity gFOBT is based on chemical detection of blood, while FIT uses antibodies to detect blood. Stool DNA tests detect DNA biomarkers for cancer in cells shed from the lining of the colon and rectum into stool.
Table 1. Characteristics of Recommended Colorectal Cancer Screening Strategies

<table>
<thead>
<tr>
<th>Screening method</th>
<th>Frequency</th>
<th>Evidence of efficacy</th>
<th>Other considerations</th>
</tr>
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</table>
| High-sensitivity gFOBT | Every year | • Evidence from RCTs that gFOBT reduces colorectal cancer mortality  
  • Harms from screening with gFOBT arise from colonoscopy to follow up abnormal gFOBT results  
  • Requires dietary restrictions and 3 stool samples  
  • Does not require bowel preparation, anesthesia or sedation, or transportation to and from the screening examination (test is performed at home) | |
| FIT | Every year | • Evidence from 1 large cohort study that screening with FIT reduces colorectal cancer mortality  
  • Certain types of FIT have improved accuracy compared to gFOBT and HSgFOBT (20 μg hemoglobin per gram of feces threshold was used in the CISNET modeling)  
  • Harms from screening with FIT arise from colonoscopy to follow up abnormal FIT results  
  • Can be done with a single stool sample  
  • Requires good adherence over multiple rounds of testing  
  • Does not require bowel preparation, anesthesia or sedation, or transportation to and from the screening examination (test is performed at home) | |
| sDNA-FIT | Every 1 to 3 y | • Improved sensitivity compared with FIT per 1-time application of screening test  
  • Specificity is lower than that of FIT, resulting in more false-positive results, more follow-up colonoscopies, and more associated adverse events per sDNA-FIT screening test compared with per FIT test  
  • Modeling suggests that screening every 3 y does not provide a favorable (ie, efficient) balance of benefits and harms compared with other stool-based screening options (ie, annual FIT or sDNA-FIT every 1 or 2 y)  
  • Insufficient evidence about appropriate longitudinal follow-up of abnormal findings after a negative follow-up colonoscopy  
  • No direct evidence evaluating the effect of sDNA-FIT on colorectal cancer mortality  
  • Harms from screening with sDNA-FIT arise from colonoscopy to follow up abnormal sDNA-FIT results  
  • Can be done with a single stool sample but involves collecting an entire bowel movement  
  • Requires good adherence over multiple rounds of testing  
  • Does not require bowel preparation, anesthesia or sedation, or transportation to and from the screening examination (test is performed at home) | |
| Colonoscopy | Every 10 y | • Evidence from cohort studies that colonoscopy reduces colorectal cancer mortality  
  • Harms from colonoscopy include bleeding and perforation, which both increase with age  
  • Screening and follow-up of positive results can be performed during the same examination  
  • Requires less frequent screening  
  • Requires bowel preparation, anesthesia or sedation, and transportation to and from the screening examination | |
| CT colonography | Every 5 y | • Evidence available that CT colonography has reasonable accuracy to detect colorectal cancer and adenomas  
  • No direct evidence evaluating effect of CT colonography on colorectal cancer mortality  
  • Limited evidence about the potential benefits or harms of possible evaluation and treatment of incidental extracolonic findings, which are common. Extracolonic findings detected in 1.3% to 11.4% of examinations; <3% required medical or surgical treatment  
  • Additional harms from screening with CT colonography arise from colonoscopy to follow up abnormal CT colonography results  
  • Requires bowel preparation  
  • Does not require anesthesia or sedation or transportation to and from the screening examination | |
| Flexible sigmoidoscopy | Every 5 y | • Evidence from RCTs that flexible sigmoidoscopy reduces colorectal cancer mortality  
  • Risk of bleeding and perforation but less than risk with colonoscopy  
  • Modeling suggests that it provides fewer life-years gained alone than when combined with FIT or in comparison to other strategies  
  • Additional harms may arise from colonoscopy to follow up abnormal flexible sigmoidoscopy results  
  • Test availability has declined in the US but may be available in some communities where colonoscopy is less available | |
| Flexible sigmoidoscopy with FIT | Flexible sigmoidoscopy every 10 y plus FIT every year | • Evidence from RCTs that flexible sigmoidoscopy + FIT reduces colorectal cancer mortality  
  • Modeling suggests combination testing provides benefits similar to those of colonoscopy, with fewer complications  
  • Risk of bleeding and perforation from flexible sigmoidoscopy but less than risk with colonoscopy  
  • Additional potential harms from colonoscopy to follow up abnormal flexible sigmoidoscopy or FIT results  
  • Flexible sigmoidoscopy availability has declined in the US but may be available in some communities where colonoscopy is less available  
  • Screening with FIT requires good adherence over multiple rounds of testing | |

Abbreviations: CISNET, Cancer Intervention and Surveillance Modeling Network; CT, computed tomography; FIT, fecal immunochemical test; gFOBT, guaiac fecal occult blood test; RCT, randomized clinical trial; sDNA-FIT, stool DNA test with fecal immunochemical test.

a To achieve the benefits of screening, abnormal results from stool-based tests, CT colonography, and flexible sigmoidoscopy should be followed up with colonoscopy.

b Applies to persons with negative findings (including hyperplastic polyps) and is not intended for persons in surveillance programs. Evidence of efficacy is not informative of screening frequency, with the exception of gFOBT and flexible sigmoidoscopy alone.

c As stated by the manufacturer.
Currently, the only stool DNA test approved by the US Food and Drug Administration is a multitarget stool DNA test that also includes a FIT component, referred to as sDNA-FIT in this recommendation. When stool-based tests reveal abnormal results, follow-up with colonoscopy is needed for further evaluation. Among the stool-based tests, screening with annual FIT or annual sDNA-FIT provides an estimated greater life-years gained than annual high-sensitivity gFOBT or sDNA-FIT every 3 years.12,13 Additionally, modeling estimates that screening with sDNA-FIT annually would result in more colonoscopies than annual screening with FIT.12,13 However, sDNA-FIT every 1 to 3 years is estimated to provide a reasonable balance of life years gained per estimated follow-up colonoscopy compared with no screening. Currently, there is uncertainty around the accuracy of high-sensitivity gFOBT to detect colorectal cancer and advanced adenomas, although it is likely lower than the accuracy of FIT and sDNA-FIT, and high-sensitivity gFOBT is more difficult for patients to administer.9,10 However, randomized trials demonstrate direct evidence of decreased deaths from colorectal cancer when screening with non-high-sensitivity gFOBT is performed.9,10

**Direct Visualization Tests**
Direct visualization tests to screen for colorectal cancer include colonoscopy, CT colonography, and flexible sigmoidoscopy. All 3 screening tests visualize the inside of the colon and rectum, although flexible sigmoidoscopy can only visualize the rectum, sigmoid colon, and descending colon, while colonoscopy and CT colonography can generally visualize the entire colon. For colonoscopy and flexible sigmoidoscopy, a camera is used to visualize the inside of the colon, while CT colonography uses x-ray images. When abnormal results are found on flexible sigmoidoscopy or CT colonography, follow-up with colonoscopy is needed for further evaluation. Among the direct visualization tests, a colonoscopy every 10 years or CT colonography every 5 years have greater estimated life-years gained than flexible sigmoidoscopy every 5 years.12,13 Unlike colonoscopy and flexible sigmoidoscopy, CT colonography may reveal extracolonic
Table 2. Summary of USPSTF Rationale

<table>
<thead>
<tr>
<th>Rationale</th>
<th>Adults aged 45-49 y</th>
<th>Adults aged 50-75 y</th>
<th>Adults 76 y or older</th>
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<tbody>
<tr>
<td>Detection</td>
<td>The USPSTF found adequate evidence that screening for colorectal cancer in adults aged 45 to 49 y provides a moderate benefit in terms of reducing colorectal cancer mortality and increasing life-years gained.</td>
<td>The USPSTF found convincing evidence that screening for colorectal cancer with several different methods can accurately detect early-stage colorectal cancer and adenomatous polyps.</td>
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</tr>
<tr>
<td>Benefits of early detection and intervention and treatment</td>
<td>The USPSTF found adequate evidence that screening for colorectal cancer with stool tests, colonoscopy, CT colonography, or flexible sigmoidoscopy in adults aged 50 to 75 y provides a substantial benefit in reducing colorectal cancer mortality and increasing life-years gained.</td>
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<td>The USPSTF found adequate evidence that routine screening for colorectal cancer with stool tests, colonoscopy, CT colonography, or flexible sigmoidoscopy in adults aged 76 to 85 y provides a small to moderate benefit in reducing colorectal cancer mortality and increasing life-years gained.</td>
</tr>
<tr>
<td>Harms of early detection and intervention and treatment</td>
<td>The USPSTF found adequate evidence that the harms of screening for colorectal cancer in adults aged 45 to 49 y are small. The majority of harms result from the use of colonoscopy (such as bleeding and perforation), either as the screening test or as follow-up for positive findings detected by other screening tests.</td>
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<tr>
<td>USPSTF assessment</td>
<td>The USPSTF concludes with moderate certainty that there is a moderate net benefit of starting screening for colorectal cancer in adults aged 45 to 49 y.</td>
<td>The USPSTF concludes with high certainty that there is a substantial net benefit of screening for colorectal cancer in adults aged 50 to 75 y.</td>
<td>The USPSTF concludes with moderate certainty that there is a small net benefit of screening for colorectal cancer in adults aged 76 to 85 y who have been previously screened.</td>
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Abbreviations: CT, computed tomography; gFOBT, guaiac fecal occult blood test; USPSTF, US Preventive Services Task Force.

findings that require additional workup, which could lead to other potential benefits or harms.930

Starting and Stopping Ages

The USPSTF recommends offering colorectal cancer screening starting at age 45 years. Although the absolute risk of developing colorectal cancer is much lower in adults younger than 50 years (20.0 new colorectal cancer cases per 100 000 persons aged 40 to 49 years, 47.8 new cases per 100 000 persons aged 50 to 59 years, and 105.2 new cases per 100 000 persons 60 years or older16), age-period cohort analysis indicates a recent trend for increasing risk of colorectal cancer in birth cohorts of adults younger than 50 years.15 The benefit of reducing colorectal cancer deaths by screening for colorectal cancer in adults 50 years or older is well established through trial data. Some of these trials16-18 also included adults younger than 50 years, although results are not reported separately for younger age groups. Additionally, modeling performed by the Cancer Intervention and Surveillance Modeling Network (CISNET) suggests that starting colorectal cancer screening at age 45 years may moderately increase life-years gained and decrease colorectal cancer cases and deaths compared with beginning screening at age 50 years.16,18

In adults aged 76 to 85 years, the age at which the balance of benefits and harms of colorectal cancer screening becomes less favorable and screening should be stopped varies based on a patient’s health status (eg, life expectancy, comorbid conditions), prior screening status, and individual preferences.19 Limited evidence suggests that harms from colonoscopy, such as perforation and bleeding, and extracolonic findings on CT colonography increase with age.930 Modeling studies estimate that generally, few additional life-years are gained when screening is extended past age 75 years among average-risk adults who have previously received adequate screening.12,13 In adults 86 years or older, evidence on benefits and harms of colorectal cancer screening is lacking, and competing causes of mortality likely preclude any survival benefit that would outweigh the harms of screening.

Screening Intervals

Recommended intervals for colorectal cancer screening tests include:

- High-sensitivity gFOBT or FIT every year
- sDNA-FIT every 1 to 3 years
- CT colonography every 5 years
- Flexible sigmoidoscopy every 5 years
- Flexible sigmoidoscopy every 10 years + FIT every year
- Colonoscopy screening every 10 years

Treatment or Interventions

Localized cancer is generally treated with surgical resection.20 Depending on cancer location and stage/progression, additional
Screening for Colorectal Cancer in Black Adults

Colorectal Cancer Burden

Black adults have the highest incidence of and mortality from colorectal cancer compared with other races/ethnicities. From 2013 to 2017, incidence rates for colorectal cancer were 43.6 cases per 100,000 Black adults, 39.0 cases per 100,000 American Indian/Alaska Native adults, 37.8 cases per 100,000 White adults, 33.7 cases per 100,000 Hispanic/Latino adults, and 31.8 cases per 100,000 Asian/Pacific Islander adults. Colorectal cancer death rates in 2014 to 2018 were 18.0 deaths per 100,000 Black adults, 15.1 deaths per 100,000 American Indian/Alaska Native adults, 13.6 deaths per 100,000 non-White Hispanic White adults, 10.9 deaths per 100,000 Hispanic/Latino adults, and 9.4 deaths per 100,000 Asian/Pacific Islander adults.

The causes for these health disparities are complex; recent evidence points to inequities in the access to and utilization and quality of colorectal cancer screening and treatment as the primary driver for this health disparity rather than genetic differences. The recent trend for increasing colorectal cancer incidence in adults younger than 50 years has been observed in White and Hispanic/Latino adults but not Black or Asian/Pacific Islander adults. However, despite these trends, Black adults across all age groups, including those younger than 50 years, continue to have a higher incidence of and mortality from colorectal cancer than White adults.

Available Evidence

The USPSTF sought evidence on the potential benefits and harms of colorectal cancer screening in Black adults; however, little empirical evidence was identified. Although some studies on the effectiveness of colorectal cancer screening included non-White participants, no studies reported results of screening by race/ethnicity. Few studies on screening accuracy reported findings by race; however, those studies that did generally found no difference in accuracy to detect colorectal cancer in Black adults compared with White adults for FIT (in 1 study) or sDNA-FIT (in 1 study). The 4 studies of screening colonoscopy that reported harms by race/ethnicity had inconsistent findings. No other studies on harms reported results by race/ethnicity. Modeling studies that assume perfect adherence to screening and no racial differences in screening accuracy or natural history of colorectal cancer (ie, no biological differences in the risks of adenoma onset and progression to colorectal cancer), but lower relative colorectal cancer survival rates and increased all-cause mortality in Black adults vs White adults, estimate similar life-years gained from screening Black adults and White adults and a similar balance of the benefits and harms for each screening strategy.

Advising Black Adults

Based on the limited available empirical evidence, the USPSTF is not able to make a separate, specific recommendation on colorectal cancer screening in Black adults. Results from CISNET modeling also do not support different screening strategies by race. Other organizations such as the US Multi-Society Task Force recommend starting screening in Black adults at age 45 years while starting screening at age 50 years for persons of other races. The current USPSTF statement recommends starting screening for everyone at age 45 years, including Black adults.
usually used during colonoscopy; hence, assistance with transportation home and recovery time after colonoscopy is required. Abnormal findings identified by flexible sigmoidoscopy or CT colonography screening require follow-up colonoscopy for screening benefits to be achieved.

Additional Tools and Resources
The National Cancer Institute and the Centers for Disease Control and Prevention have developed patient and clinician guides on screening for colorectal cancer:

- Colorectal Cancer Screening (PDQ)—Health Professional Version https://www.cancer.gov/types/colorectal/hp/colorectal-screening-pdq
- Colorectal Cancer Screening Tests https://www.cdc.gov/cancer/colorectal/basic_info/screening/tests.htm

The Community Preventive Services Task Force has also issued recommendations on interventions to increase colorectal cancer screening at https://www.thecommunityguide.org/content/task-force-findings-cancer-prevention-and-control.

Other Related USPSTF Recommendations
The USPSTF has a recommendation statement on aspirin use for the primary prevention of cardiovascular disease and colorectal cancer in average-risk adults (available at https://uspreventiveservicestaskforce.org).37

Update of Previous USPSTF Recommendation
This final recommendation replaces the 2016 USPSTF recommendation on screening for colorectal cancer. In 2016, the USPSTF recommended screening for colorectal cancer starting at age 50 years and continuing until age 75 years (A recommendation). In addition, the USPSTF concluded that the decision to screen for colorectal cancer in adults aged 76 to 85 years should be an individual one, taking into account the patient’s overall health and prior screening history (C recommendation) and that screening should be discontinued after age 85 years.

In the current recommendation, while continuing to recommend colorectal cancer screening in adults aged 50 to 75 years (A recommendation), the USPSTF now recommends offering screening starting at age 45 years (B recommendation). As it did in 2016, the USPSTF continues to conclude that screening in adults aged 76 to 85 years should be an individual decision (C recommendation) and screening should be discontinued after age 85 years.

Supporting Evidence
Scope of Review
To update its 2016 recommendation, the USPSTF commissioned a systematic review9,10 to evaluate the benefits and harms of screening for colorectal cancer in adults 40 years or older. As in 2016, the USPSTF reviewed the evidence on (1) the effectiveness and comparative effectiveness of screening strategies to reduce colorectal cancer incidence, colorectal cancer mortality, or both; (2) the accuracy of various screening tests to detect colorectal cancer, advanced adenomas, or adenomatous polyps based on size; and (3) the serious harms of different screening tests. The review also examined whether these findings varied by age, sex, or race/ethnicity.

In addition, as in 2016, the USPSTF commissioned a report from the CISNET Colorectal Cancer Working Group12,13 to provide information from comparative modeling on how estimated life-years gained, colorectal cancer cases averted, and colorectal cancer deaths averted as well as colonoscopy burden and harms vary by different starting and stopping ages for various screening strategies. New analyses included in the current modeling for the USPSTF that were not performed in the models commissioned by the USPSTF in 2016 included analyses with elevated risk scenarios to reflect recent population trends in colorectal cancer incidence95 and analyses by race.12,13

Accuracy of Screening Tests
The USPSTF focused on reviewing evidence that reported accuracy of screening tests compared with colonoscopy as the reference standard. Colonoscopy accuracy is reported with a reference standard of either repeat colonoscopy or CT colonography-enhanced colonoscopy. The following accuracy results reflect accuracy after only a single application of the test rather than a program of repeated screenings.

Stool-Based Tests
Evidence on accuracy of high-sensitivity gFOBT to detect colorectal cancer and advanced adenomas compared with a colonoscopy reference standard was reported in 2 studies (n = 3503).9,10 Reported sensitivity to detect colorectal cancer ranged from 0.50 to 0.75 (95% CI, 0.09-1.0) and reported specificity ranged from 0.96 to 0.98 (95% CI, 0.95-0.99). Sensitivity for detecting advanced adenomas was lower, ranging from 0.06 to 0.17 (95% CI, 0.02-0.23), while specificity was similar (0.96 to 0.99 [95% CI, 0.96-0.99]).9,10 A larger evidence base was available on the accuracy of FIT, with the most evidence available on the OC-Sensor family of FITs (13 studies; n = 44 887).9,10 Using the threshold recommended by the manufacturer (20 μg hemoglobin per gram of stool), the pooled sensitivity for detection of colorectal cancer was 0.74 (95% CI, 0.64-0.83; 9 studies; n = 34 352) and pooled specificity was 0.94 (95% CI, 0.93-0.96; 9 studies; n = 34 352). Similar to high-sensitivity gFOBT, sensitivity for detecting advanced adenomas was lower while specificity was similar; pooled sensitivity was 0.23 (95% CI, 0.20-0.25) and pooled specificity was 0.96 (95% CI, 0.95-0.97).9,10 Accuracy estimates of 9 other types of FIT were similar but were generally reported only in single studies. In 4 studies (n = 12 424) reporting the accuracy of sDNA-FIT,9,10 pooled sensitivity for colorectal cancer detection was 0.93 (95% CI, 0.87-1.0) and pooled specificity was 0.84 (95% CI, 0.84-0.86), with a lower pooled sensitivity for detecting advanced adenomas (0.43 [95% CI, 0.40-0.46]) but higher pooled specificity (0.89 [95% CI, 0.86-0.92]).9,10 Ten of the accuracy studies on FIT also reported results by age strata and generally found no significant difference; 2 reported stratified analyses for individuals younger than 50 years. Two studies suggested lower specificity for colorectal cancer detection in adults 70 years or older; a single study on sDNA-FIT suggested decreasing specificity with increasing age.9,10
Direct Visualization Tests
Colonoscopy was evaluated in 4 studies (n = 4821) on accuracy, with 3 studies (n = 2290) determining missed cases of colorectal cancer by follow-up CT colonography-enhanced colonoscopy or CT colonography and repeat colonoscopy for discrepant findings. In all 4 studies, sensitivity for detection of adenomas measuring 10 mm or larger ranged from 0.89 (95% CI, 0.78-0.96) to 0.95 (95% CI, 0.74-0.99); specificity was reported in a single study as 0.89 (95% CI, 0.86-0.91). Two of the studies on colonoscopy accuracy included patients younger than 50 years, although results in this age group were not reported separately.

Seven studies (n = 5328) reported on accuracy of CT colonography. The studies were heterogeneous in study design, population, imaging technique, and reader experience or protocol. Sensitivity for colorectal cancer detection was reported in 6 of the studies and ranged from 0.86 to 1.0 (95% CI range, 0.21-1.0); specificity was not reported. Pooled sensitivity for detection of adenomas measuring 10 mm or larger was 0.89 (95% CI, 0.83-0.96) and pooled specificity was 0.94 (95% CI, 0.89-1.0). One study reported CT colonography accuracy by age and suggested that sensitivity was lower in adults 65 years or older; however, this finding was not statistically significant.

The USPSTF did not identify any studies that reported on the accuracy of flexible sigmoidoscopy using colonoscopy as the reference standard.

Benefits of Early Detection and Treatment
Direct evidence on the benefits of colorectal cancer screening to decrease colorectal cancer mortality are available from randomized clinical trials (RCTs) on gFOBT and flexible sigmoidoscopy as well as from cohort studies on FIT and colonoscopy. Pooled results from 4 RCTs (n = 458 002) on flexible sigmoidoscopy compared with no screening show a significant decrease in colorectal cancer mortality (mortality rate ratio, 0.74 [95% CI, 0.68-0.80]) over 11 to 17 years of follow-up. Most studies reported outcomes after a single round of screening, although the 1 trial conducted in the US, the Prostate, Lung, Colorectal and Ovarian Cancer Screening Trial, evaluated 2 rounds of screening. Decreased mortality with flexible sigmoidoscopy screening was consistently reported across the 4 trials. None of the trials included persons younger than 50 years.

Trials that report on colorectal cancer outcomes with high-sensitivity gFOBT screening are currently lacking, although several older trials report decreased colorectal cancer mortality with Hemoccult II screening (an older gFOBT no longer commonly used). After 2 to 9 rounds of biennial gFOBT screening, colorectal cancer mortality was found to be lower at 11 to 30 years of follow-up (relative risk range, 0.78 [95% CI, 0.65-0.93] to 0.91 [95% CI, 0.84-0.98]). Participants younger than 50 years were included in 3 trials, although results for that age group were not reported separately.

Two prospective cohort studies (n = 436 927) in US-based populations reported on colorectal cancer outcomes after colonoscopy screening. One study among health professionals found that after 22 years of follow-up, colorectal mortality was lower in persons who reported receiving at least 1 colonoscopy (adjusted hazard ratio, 0.32 [95% CI, 0.24-0.45]), although findings were no longer significant after 5 years for adults with a first-degree relative with colorectal cancer. This study included persons younger than 50 years, although results for this age group were not reported separately. Another cohort study among Medicare beneficiaries reported that the risk of colorectal cancer was significantly lower in adults aged 70 to 74 years (but not aged 75 to 79 years) 8 years after receiving a screening colonoscopy (standardized risk, 0.42% [95% CI, 0.24%-0.63%]). One large, prospective cohort study (n = 5 417 699) from Taiwan reported on colorectal cancer mortality after introduction of a nationwide screening program with FIT in adults aged 50 to 69 years. After 1 to 3 rounds of biennial FIT screening, lower colorectal cancer mortality was found at 6 years of follow-up (adjusted relative risk, 0.90 [95% CI, 0.84-0.95]).

The CISNET modeling study commissioned for this review estimated the number of life-years gained, colorectal cancer cases and deaths averted, lifetime colonoscopies required (as a proxy measure for the burden of screening), and resulting harms from colonoscopy (ie, gastrointestinal and cardiovascular events) for various screening strategies. These strategies varied in the screening modality, the age at which to start and stop screening, and the frequency of screening. The USPSTF focused on findings from models that assumed an elevated population risk of colorectal cancer. These models were thought to better capture the currently observed epidemiologic trend of increasing incidence in adults younger than 50 years, which is thought to reflect cohort effects, with younger birth cohorts at greater risk for colorectal cancer than older cohorts. The USPSTF focused on estimated life-years gained (compared with no screening) as the primary measure of the benefit of screening. Given this elevated population risk assumption, as well as assuming 100% adherence, the USPSTF determined that beginning screening at age 45 years and continuing to the age of 75 years, for the following screening strategies, yielded a reasonable balance of benefits (life-years gained) and burdens or harms (number of colonoscopies): annual FIT, sDNA-FIT every 1 to 3 years, CT colonography or flexible sigmoidoscopy every 5 years, colonoscopy every 10 years, or flexible sigmoidoscopy every 10 years with annual FIT (Figure 2 and Figure 3). Modeling estimates that screening with sDNA-FIT annually results in additional colonoscopy burden compared with annual FIT screening (approximately 850 more subsequent follow-up and surveillance colonoscopies needed per 1000 adults screened with annual sDNA-FIT). Screening with sDNA-FIT every 2 years is estimated to result in approximately 300 more subsequent follow-up and surveillance colonoscopies per 1000 adults screened compared with annual FIT. In modeling analyses, performing sDNA-FIT every 3 years or high-sensitivity gFOBT annually (also included in Figure 2 and Figure 3) did not provide an efficient balance of the estimated lifetime number of colonoscopies vs the estimated life-years gained, compared with other options for stool-based screening. However, sDNA-FIT every 3 years or high-sensitivity gFOBT annually is still estimated to provide a reasonable balance of benefit in life-years gained and harms compared with no screening. Additionally, there is greater uncertainty in the model predictions for high-sensitivity gFOBT strategies, given the underlying uncertainty around the sensitivity and specificity of high-sensitivity gFOBT to detect adenomas and colorectal cancer. Based on averaging estimates across the 3 CISNET models, if screening were performed from ages 45 to 75 years with one of the USPSTF recommended strategies, an estimated 286 to 337 life-years would be gained, an estimated 42 to 61 cases of colorectal cancer would be averted, and an estimated 24 to 28 colorectal cancer deaths would be averted, per 1000 adults screened, depending on...
the specific strategy used (Figure 2). This finding translates to an estimated 104 to 123 days of life gained per person screened. Lowering the starting age of screening from age 50 years to age 45 years results in an estimated additional 2 to 3 cases of colorectal cancer being averted, an estimated 1 additional colorectal cancer death averted, and an estimated 22 to 27 additional life-years gained per 1000 adults (ie, 8 to 10 additional days of life gained per person screened) (Figure 2).

**Harms of Screening and Treatment**

No studies reported on harms from stool-based tests. The primary harms from stool-based screening tests are thought to come

**Figure 2. Benefits of Colorectal Cancer Screening**

**A** Benefit: Estimated life-years gained per 1000 individuals screened

<table>
<thead>
<tr>
<th>Screening modality and frequency</th>
<th>Mean life-years gained if start screening</th>
<th>Additional life years gained if start screening at age 45 y</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIT every year</td>
<td>292</td>
<td>26</td>
</tr>
<tr>
<td>HSgFOBT every year, d</td>
<td>272</td>
<td>26</td>
</tr>
<tr>
<td>sDNA-FIT every year</td>
<td>307</td>
<td>26</td>
</tr>
<tr>
<td>sDNA-FIT every 3 y</td>
<td>278</td>
<td>25</td>
</tr>
<tr>
<td>Direct visualization tests</td>
<td></td>
<td></td>
</tr>
<tr>
<td>COL every 10 y</td>
<td>310</td>
<td>27</td>
</tr>
<tr>
<td>CT colonography every 5 y</td>
<td>293</td>
<td>24</td>
</tr>
<tr>
<td>Flexible SIG every 5 y</td>
<td>264</td>
<td>22</td>
</tr>
<tr>
<td>Flexible SIG every 10 y plus FIT every year</td>
<td>306</td>
<td>26</td>
</tr>
</tbody>
</table>

**B** Benefit: Estimated No. of CRC cases averted per 1000 individuals screened

<table>
<thead>
<tr>
<th>Screening modality and frequency</th>
<th>Mean CRC cases averted if start screening</th>
<th>Additional CRC cases averted if start screening at age 45 y</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIT every year</td>
<td>47</td>
<td>3</td>
</tr>
<tr>
<td>HSgFOBT every year, d</td>
<td>39</td>
<td>3</td>
</tr>
<tr>
<td>sDNA-FIT every year</td>
<td>54</td>
<td>3</td>
</tr>
<tr>
<td>sDNA-FIT every 3 y</td>
<td>44</td>
<td>3</td>
</tr>
<tr>
<td>Direct visualization tests</td>
<td></td>
<td></td>
</tr>
<tr>
<td>COL every 10 y</td>
<td>58</td>
<td>3</td>
</tr>
<tr>
<td>CT colonography every 5 y</td>
<td>53</td>
<td>2</td>
</tr>
<tr>
<td>Flexible SIG every 5 y</td>
<td>49</td>
<td>2</td>
</tr>
<tr>
<td>Flexible SIG every 10 y plus FIT every year</td>
<td>54</td>
<td>3</td>
</tr>
</tbody>
</table>

**C** Benefit: Estimated No. of CRC deaths averted per 1000 individuals screened

<table>
<thead>
<tr>
<th>Screening modality and frequency</th>
<th>Mean CRC deaths averted if start screening</th>
<th>Additional CRC deaths averted if start screening at age 45 y</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIT every year</td>
<td>25</td>
<td>1</td>
</tr>
<tr>
<td>HSgFOBT every year, d</td>
<td>23</td>
<td>1</td>
</tr>
<tr>
<td>sDNA-FIT every year</td>
<td>27</td>
<td>1</td>
</tr>
<tr>
<td>sDNA-FIT every 3 y</td>
<td>24</td>
<td>1</td>
</tr>
<tr>
<td>Direct visualization tests</td>
<td></td>
<td></td>
</tr>
<tr>
<td>COL every 10 y</td>
<td>27</td>
<td>1</td>
</tr>
<tr>
<td>CT colonography every 5 y</td>
<td>26</td>
<td>0.9</td>
</tr>
<tr>
<td>Flexible SIG every 5 y</td>
<td>23</td>
<td>0.9</td>
</tr>
<tr>
<td>Flexible SIG every 10 y plus FIT every year</td>
<td>26</td>
<td>1</td>
</tr>
</tbody>
</table>

CRC indicates colorectal cancer; CT, computed tomography; FIT, fecal immunochemical test (with positivity cutoff of 20 μg of hemoglobin per gram of feces); HSgFOBT, high-sensitivity guaiac fecal occult blood test; sDNA-FIT, stool DNA tests with FIT (multitarget stool DNA test); SIG, sigmoidoscopy; COL, colonoscopy.

* Outcomes are expressed per 1000 40-year-olds who start screening at age 45 or at age 50.

* Mean estimate across the 3 Cancer Intervention and Surveillance Modeling Network colorectal cancer models. See modeling report12,13 for additional details and model-specific estimates.

* Because of imprecision in sensitivity and specificity, there is considerable uncertainty in model predictions for HSgFOBT strategies. See modeling report12 for more information.

* Compared with other options for stool-based screening, these strategies do not provide an efficient balance of the benefits (life-years gained) vs harms and burden (ie, lifetime number of colonoscopies) of screening. See modeling report12,13 for more information.
from false-positive and false-negative results and from harms of workup of positive screening results, such as colonoscopy. Serious harms from colonoscopy to follow-up positive screening results are estimated to be 17.5 serious bleeding events (95% CI, 7.6-27.5; 11 studies; n = 78 793) and 5.4 perforations (95% CI, 3.4 to 7.4; 12 studies; n = 341 922) per 10 000 colonoscopies.9,10

Harms from screening colonoscopy have been reported in 67 observational studies (n = 27 746 669).9 Rates of serious bleeding events and perforations are lower with screening colonoscopy than with colonoscopy performed following positive stool-based screening test results (presumably because of fewer biopsies and adenoma removals), with 14.6 major bleeding events per 10 000 colonoscopies (95% CI, 9.4-19.9; 20 studies; n = 5 172 508) and 3.1 perforations per 10 000 colonoscopies (95% CI, 2.3-4.0; 26 studies; n = 5 272 600).9,10 If sedation is used during colonoscopy, cardiopulmonary events may rarely occur, although the precise frequency of occurrence is not known. No higher risk of other serious harms with screening colonoscopy was seen in 4 cohort studies.
Other serious reported harms include infection and other gastrointestinal events (besides bleeding and perforation). Twenty-three studies reported on differences in harms by age, and 21 studies included persons younger than 50 years. Overall, all harms were estimated as rare. Overall findings indicated increasing risk of bleeding and perforation with increasing age. Bowel preparation for colonoscopy, flexible sigmoidoscopy, and CT colonography may lead to dehydration or electrolyte imbalances, particularly in older adults or persons with comorbid conditions; accurate estimates of the rates of these events are not available.

Harms from flexible sigmoidoscopy were reported in 18 studies (n = 395,077). Rates of serious harms were 0.5 bleeding events per 10,000 sigmoidoscopies (95% CI, 0.1-1.3; 10 studies; n = 179,854) and 0.2 perforations per 10,000 sigmoidoscopies (95% CI, 0.1-0.4; 11 studies; n = 359,679). No studies included persons younger than 50 years and no subgroup analyses on harms by age were reported. Rates of harms from colonoscopy following abnormal flexible sigmoidoscopy results include 20.7 major bleeding events per 10,000 colonoscopies (95% CI, 8.2-33.2; 4 studies; n = 5790) and 12.0 perforations per 10,000 colonoscopies (95% CI, 7.5-16.5; 4 studies; n = 23,022).

Harms from CT colonography are uncommon (19 studies; n = 90,133), and the reported radiation dose for CT colonography ranges from 0.8 to 5.3 mSv (compared with an average annual background radiation dose of 3.0 mSv per person in the US). Accurate estimates of rates of serious harms from colonoscopy following abnormal CT colonography results are not available. Extracolonic findings on CT colonography are common. Based on 27 studies that included 48,235 participants, 1.3% to 11.4% of examinations identified extracolonic findings that required workup. Three percent or less of individuals with extracolonic findings required definitive medical or surgical treatment for an incidental finding. A few studies suggest that extracolonic findings may be more common in older age groups. Long-term clinical follow-up of extracolonic findings was reported in few studies, making it difficult to know whether it represents a benefit or harm of CT colonography.

Potential harms of colorectal cancer screening include possible overdetection of adenomas not destined to become cancer; however, no studies directly assessing the health effects of these harms were identified.

Based on the available empirical evidence, harms from colonoscopy (either a screening colonoscopy, follow-up colonoscopy after a positive screening result from other methods, or surveillance colonoscopy in persons in whom adenomas have previously been detected) were considered to be the main source of colorectal cancer screening harms in the CIRNet modeling study. Thus, harms were quantified as the lifetime number of colorectal cancer complications associated with screening, and the lifetime number of colonoscopies was used as a proxy for the burden of screening. Based on averaging estimates across the 3 models, if screening were performed from ages 45 to 75 years with 1 of the USPSTF recommended strategies, an estimated 1535 to 4248 colonoscopy procedures and 10 to 16 colonoscopy complications would be expected over the lifetime of 1000 screened adults (ie, 1.5 to 4.2 colonoscopies per person over the lifetime and complications estimated as occurring in 1 in every 63 to 102 adults screened from ages 45 to 75 years).

Response to Public Comments

A draft version of this recommendation statement was posted for public comment on the USPSTF website from October 27, 2020, to November 23, 2020. Many comments were received on the USPSTF’s new B recommendation to screen adults aged 45 to 49 years; some supported the new recommendation, others requested that screening begin at an even younger age, and still others disagreed with starting screening before age 50 years. The USPSTF appreciates the various perspectives that were shared. Although future research could further strengthen the USPSTF’s understanding about the benefits and harms of colorectal cancer screening in adults aged 45 to 49 years, based on the USPSTF’s assessment of the available empirical, modeling, and epidemiologic data, the USPSTF finds adequate evidence that screening this age group provides a moderate net benefit. Several comments requested that colonoscopy to follow up an abnormal noncolonoscopy screening test result be considered part of screening. The USPSTF recognizes that the benefits of screening can only be fully achieved when follow-up of abnormal screening test results is performed. The USPSTF added language to the Practice Considerations section to clarify this.

Several comments also requested clarification about how frequently sDNA-FIT is being recommended. The USPSTF has clarified that screening every 1 to 3 years with sDNA-FIT would be reasonable. Comments were also received requesting that the USPSTF provide a tiered or ranked list of screening strategies. Because no direct evidence compares different screening tests, and because local resources or patient factors may influence feasibility of different screening strategies, the USPSTF is unable to determine which tests are unequivocally “better” or “worse.” In Table 1, the USPSTF describes the available empirical and modeling evidence on the benefits and harms of each screening strategy and also highlights additional considerations that may help an individual patient and clinician select a specific screening strategy. Comments also requested that persons with a personal or family history of Lynch syndrome be added to the recommendation. Persons who have hereditary cancer syndromes such as Lynch syndrome are at very high risk for colorectal cancer and may need screening strategies that go beyond the evidence that the USPSTF reviewed. Persons with a personal or family history of Lynch syndrome should speak with their health care professional about appropriate screening options.

Research Needs and Gaps

Although the benefits of screening for colorectal cancer are well established, the following important evidence gaps that need to be addressed by additional research persist.

• Randomized trials that directly compare the effectiveness of different colorectal cancer screening strategies (including hybrid strategies that switch between modalities over time) to reduce colorectal cancer mortality are needed.

• Studies are needed on screening effectiveness in adults younger than 50 years and whether screening strategies should be tailored in these populations.

• More research is needed to understand the factors that contribute to increased colorectal cancer incidence and mortality in Black adults, such as access to and availability of care and characteristics of systems providing health care. Once these factors are identified, more research is needed to test interventions designed to mitigate these differences for Black adults.
The US Preventive Services Task Force (USPSTF) makes recommendations about the effectiveness of specific preventive care services for patients without obvious related signs or symptoms. It bases its recommendations on the evidence both benefits and harms of the service and an assessment of the balance. The USPSTF does not consider the costs of providing a service in this assessment. The USPSTF recognizes that clinical decisions involve more considerations than evidence alone. Clinicians should understand the evidence but individualize decision-making to the specific patient or situation. Similarly, the USPSTF notes that policy and coverage decisions involve considerations in addition to the evidence of clinical benefits and harms.

REFERENCES

6. Joseph DA, King JB, Dowling NF, Thomas CC, Richardson LC. Vital signs: colorectal cancer