Patient activation increases colorectal cancer screening rates: A randomized trial among low-income minority patients

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Abstract

Background: Colorectal cancer (CRC) screening rates remain low among low-income and minority populations. The purpose of this study was to determine if providing patients with screening information, activating them to ask for a screening test, and telephone barriers counseling improves CRC screening rates compared to providing screening information only.

Methods: Patients were randomized to CRC screening information plus patient activation and barriers counseling (n=138) or CRC screening information (n=132). Barriers counseling was attempted among activated patients if screening was not completed after one month. CRC screening test completion was determined by medical record review at two months after the medical visit. Logistic regression was used to determine whether activated patients were more likely to complete CRC screening, after adjustment for confounding factors (e.g. demographic characteristics, CRC knowledge).

Results: Patients were African American (72.2%), female (63.7%), had annual household incomes <$20,000 (60.7%), no health insurance (57.0%), and limited health literacy skills (53.7%). In adjusted analyses, more patients randomized to the activation group completed a screening test (19.6% vs. 9.9%; OR=2.35, 95%CI: 1.14, 5.56; p=0.020). Additionally, more activated patients reported discussing screening with their provider (54.4% vs. 27.5%, OR=3.29, 95%CI: 1.95, 5.56; p<0.001) and had more screening tests ordered (39.1% vs. 17.6%; OR=3.40, 95%CI: 1.88, 6.15; p<0.001) compared to those in the control group.
Conclusion: Patient activation increased CRC screening rates among low-income minority patients.

Impact: Innovative strategies are still needed to increase CRC screening discussions, motivate providers to recommend screening to patients, as well as assist patients to complete ordered screening tests.
Introduction

Colorectal cancer (CRC) incidence and mortality rates have decreased over the past two decades in the United States (U.S.) due to increased screening rates and advances in treatment [1-4]. Nonetheless, CRC remains a leading cause of cancer mortality in the U.S. Certain segments of the population, namely African Americans, have not benefitted equally from screening and still have elevated CRC incidence and mortality rates [1, 2]. Reasons for CRC disparities are numerous, complex, and occur at multiple levels (patient, provider, health system, society) [5, 6]. One reason for CRC disparities is that lower CRC screening rates occur among African Americans and among lower socioeconomic (SES) populations [1, 2, 7-9].

Since CRC screening tests are available and CRC screening has been shown to be cost-effective [10-12], the increased CRC mortality rates in these populations indicate the need for programs to increase the use of these tests to help reduce disparities. Over a decade of cancer behavioral research has provided insight to barriers to and facilitators for CRC screening [5, 13, 14]. In the past, interventions directed at the individual patient-level have usually provided CRC and CRC screening information to patients in expectation that increased knowledge would improve CRC screening rates. Prior research also suggests that a healthcare provider’s recommendation to undergo screening has been one of the strongest predictors of an individual completing a CRC screening test [13, 14].

Previously, to improve patient-provider discussions about health-related issues, communication training has mostly centered on the physician half of the patient-provider dyad with little attention given to improving patients’ communication skills [15]. Existing
literature on patient communication skills training, however, supports its value in improving patients’ participation in medical interviews, recall of treatment information and recommendations, and patient outcomes [16-19]. The goal of this study was to evaluate if average-risk patients provided with CRC screening information, activated to ask their healthcare provider for a CRC screening test, and given telephone barriers counseling would complete more CRC screening tests compared to patients provided with CRC screening information only. In addition, secondary outcomes were to evaluate if activated patients also: (a) demonstrate greater information seeking about CRC screening; and (b) have more CRC screening tests ordered by their providers compared to patients provided with CRC screening information only.

**Materials and Methods**

**Setting and Study Participants**

The study was conducted from November 2007 to May 2010 in one federally qualified health center (FQHC) that serves a mostly minority and low SES population in Columbus, Ohio. On average, the health center addresses the medical needs of approximately 6,000 patients annually, 30% of patients are 50+ years old, and 54% are African American. Healthcare providers at the center were aware that two CRC screening educational programs were being tested; however, they were not aware of the purpose of the study.

To be eligible for this study, men and women had to be age 50 or older, average-risk for CRC, not within CRC screening guidelines, able to speak and understand English, and have a working telephone. Additionally, patients had to have a scheduled appointment with a provider for a non-acute medical reason, and be able to come to the...
health center one hour prior to their scheduled appointment. Eligibility of patients was determined after medical record review and a brief telephone screening interview. Informed consent procedures and study protocols were approved by the Institution Review Board of The Ohio State University.

**Randomization and Intervention Design**

After signing a consent form, HIPAA form, and a medical record release form, patients completed a face-to-face baseline interview conducted by a research assistant (Figure 1). Patients were then randomized into the intervention arm (patient activation plus CRC screening information and barriers counseling) or control arm (CRC screening information only) via a computerized permuted randomization using a block size of eight. A second research assistant delivered the intervention to the patients.

The intervention group watched a 12 minute video entitled “Ask your doctor about colon cancer screening,” received a brochure that supplemented the video and focused on asking their provider for a CRC screening test, and received a second brochure on tips to prevent CRC (e.g. the importance of daily exercise). A description of the video content and production has been previously reported [20]. The intervention was based on the Protection Motivation Theory (PMT) [21, 22]. According to the PMT, the contradictory impact of threatening information (threat appraisal) followed by coping appraisal influences an individual’s decision to react to health information. Additionally, the intervention included the PACE (Presenting information, Asking questions, Checking for understanding, Expressing concerns) communication system which focused communication training for the patient to ask their healthcare provider about
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The control group watched a 10 minute video entitled “Colon cancer screening.” The video for the participants in the control group was the same as shown to the intervention group except the patient activation section was not included in this video. Additionally, the participants in the control group received the brochure focused on tips to prevent CRC. Following the educational session, all patients completed a brief face-to-face interview to document changes in CRC and CRC screening knowledge, attitudes, and intention to complete CRC screening. Subsequent medical visits were audio taped, if the patient and provider agreed and consented to taping.

Following the medical visit and prior to leaving the health center, patients completed a short face-to-face interview that addressed: if CRC screening was discussed with the provider; who initiated the CRC screening discussion if it occurred; and what CRC screening test was ordered or why a screening test was not ordered. All patients received a $25 gift card in appreciation of their time. Medical record reviews were conducted to collect information about any CRC screening test ordered by the provider and completion of any CRC screening test on all patients as soon as the medical chart became available after the visit, and at one and two months following the medical visit.

One month after the medical visit, if a patient in the intervention group had a CRC screening test ordered and did not complete the test, telephone barriers counseling to address patient identified CRC screening barriers was conducted. If a patient in the intervention group did not have a screening test ordered, telephone
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barriers counseling focused on activating them to ask for a CRC screening test by calling their provider or asking their provider at their next medical visit. Several attempts were made to contact each activated patient on different days and times. In addition, calls were made to patients who completed a fecal occult blood test (FOBT) to assess what components of the intervention motivated them to complete the screening test.

**Measures**

Baseline information collected was based on the constructs included in PMT including CRC susceptibility, self- and response-efficacy for CRC screening, etc. Survey items included: demographic characteristics (age, gender, race, ethnicity, marital status, education, employment status, annual household income, and health insurance); medical history; past cancer screening behaviors; CRC and CRC screening knowledge (10 true and false questions), CRC screening attitudes, barriers, and intention measured by a validated instrument (Lickert scale: strongly agree to strongly disagree) [23, 24]; health literacy (REALM) [25]; and one item measuring shared decision making preference [26].

The primary outcome in this study was whether or not a participant completed a CRC screening test by review of the medical record and laboratory log book at two months following randomization. Two months was considered an adequate follow-up period for this study because the CRC screening test recommended most frequently at the health center was the FOBT. Secondary outcomes were whether patients discussed CRC screening with their providers based on self-report in the post medical
visit interview, and whether patients had CRC tests ordered as found in the medical record review.

**Statistical analysis**

Descriptive statistics were used to provide overall characteristics by treatment arm and to ensure balancing of covariates after randomization. An intention-to-treat analysis, based on random assignment to the intervention or control arm, was used to determine the effect of the intervention on completing a CRC screening test in the two months following the medical visit. Logistic regression models were constructed to evaluate the intervention effect on the primary outcome (completion of CRC screening), and two secondary outcomes defined *a priori* (a CRC screening test ordered, and a patient-provider CRC screening discussion) and to control confounding by factors measured at baseline. Confounding was controlled by inclusion in the regression model when removal of the confounding factor from the model resulted in at least a 10% change in the intervention effect. For logistic regression analyses, likelihood ratio chi-square tests were used to determine improved statistical fit. All statistical analyses were conducted using SAS version 9.2 (SAS Institute Inc, Cary, NC).

**Results**

**Study Participants**

Figure 2 displays the number of patients selected, assessed for eligibility, accrued, randomized, and assessed for the primary outcome. The main reasons patients were ineligible were: within screening guidelines (n=291), high risk (CRC personal or family history; n=201), did not speak English (n=172), medical reasons
(n=121), medical appointment rescheduled (n=18), or a CRC screening test was recently ordered by medical record review (n=11). There were 148 patients who refused to participate because they were not able to come to the health center early (n=66), were not interested (n=47), provided no reason (n=21), or had no time (n=14). The patients who refused participation in the study were older (mean=60.5 years) and more were males (39.6%) than the participants (mean age=56.0 years, 36.4% males; p<0.05).

Among the 331 patients who agreed to participate, 47 did not arrive at the health center in time to participate in the study. Of the 284 patients who met all eligibility criteria and consented, 141 were randomized to the patient activation intervention group and 143 patients were randomized to the education only group. Fourteen patients were found to be ineligible after randomization (intervention group: one patient was high risk for CRC and not eligible for the FOBT, and two patients were found to be within CRC screening guidelines; control group: seven patients did not have a medical visit because of co-pay issues, three patients were determined to be high-risk patients not eligible for the FOBT, and one patient was randomized previously). The overall response rate was 58.1% (270/465).

Baseline characteristics of intervention and control groups are shown in Table 1. Participants (n=270) were predominantly female (63.7%), African American (72.2%), had a high school education (72.6%), had annual household incomes <$20,000 (60.7%), no health insurance (57.0%), and limited health literacy skills (53.7%). Although most participants were not married (85.9%), more participants in the control arm of the study (17.4%) were married/living together compared to participants in the
Patient activation increases colorectal cancer screening intervention arm (10.9%; p<0.05). There were no other statistically significant differences in characteristics at baseline between the intervention and control groups. Among the 270 patients, 83 (30.7%) reported having 2 or more co-morbidities, 65 (24.1%) reported having completed CRC screening in the past, and 206 (76.3%) preferred some degree of shared decision making. There were no significant differences between participant groups for baseline CRC screening knowledge and CRC screening attitudes and beliefs (Table 2).

Intervention effectiveness

Medical record review pertaining to CRC screening status was completed for all 270 patients. Overall, CRC screening completion was documented in 40 patients (27 in the intervention group and 13 in the control group). Thirty-five patients completed FOBT tests and five patients completed a colonoscopy within the two month follow-up period.

Logistic regression modeling (Table 3) showed that patients randomized to the intervention arm were 2.35 (95% CI: 1.14, 5.56) times as likely to complete CRC screening based on medical record review at 2 months after the medical visit. Additionally, patients in the intervention arm: were more likely to self-report discussing CRC screening with their provider (OR=3.29; 95% CI: 1.95, 5.56); and had more CRC screening tests ordered as determined by medical record review (OR=3.40; 95% CI: 1.88, 6.15). There was no evidence of confounding or effect modification by any of the pre-specified variables collected at baseline.
**Process evaluation**

Process evaluation documented that: 1) two patients in the intervention group watched most (90%) but not the entire video because the health care provider requested to see the patient (post intervention survey was completed); 2) five patients in the intervention group completed the education session but had it interrupted for a phone call, to use the rest room, or because a nurse needed to talk to the patient; 3) all patients but one in the intervention group received the brochures; 4) among the 54 patients in the intervention group who had a CRC screening test ordered: 18 patients returned a FOBT within a month and did not need telephone barriers counseling; 12 patients could not be reached for barriers counseling; and we were able to contact 24 patients for telephone barriers counseling at one month after their medical visit (one person completed the FOBT and two patients completed their scheduled colonoscopy); 5) among the 84 patients in the intervention group who did not have a CRC screening test ordered: 2 patients returned a FOBT within a month and did not need barriers counseling; 25 patients could not be reached for telephone barriers counseling; and we reached 57 patients at one month and reminded them to call the doctor’s office or to ask their provider for a CRC screening test at their next medical visit (one patient completed screening after being activated during the one month telephone counseling call); and 6) 43.5% (103/237) of patients agreed to have their medical visit audio taped (near the end of the study one provider (33 patients) refused to have medical visits recorded).

The most frequent CRC screening barriers reported by the participants who had a CRC screening test ordered were: dealing with other medical issues, keep putting it off, too busy, or waiting for a scheduled colonoscopy. Among patients who did not have
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a CRC screening test ordered, the most frequent comment made was that providers did not mention CRC screening to them. Additionally, among patients reached after completing the FOBT, most patients stated that the educational video made them realize how important CRC screening was and that the video showed them how to complete the FOBT.

**Discussion**

Improving colorectal cancer screening rates among minority and low SES populations is critical to reduce CRC disparities in the United States. Previous studies have shown that a provider’s recommendation for a CRC screening test is the strongest facilitator to get patients to complete a CRC screening test [13, 14, 27]. More recently, however, studies have shown that patient-provider discussions about CRC screening often do not occur or patient-provider conversations about CRC screening are limited in the information that is exchanged (e.g. lack discussion of different screening test options) [28-30].

In this randomized trial, we tested the efficacy of providing CRC screening information, activating patients to discuss CRC screening with their provider, and telephone barriers counseling to improve CRC screening rates among average-risk patients in need of a CRC screening test who were recruited from one FQHC. The intervention was hypothesized to improve CRC screening knowledge and attitudes, and empower patients to initiate CRC screening discussions with providers. Subsequent to more patient-provider CRC screening discussions, there would be an increase in CRC screening tests ordered, and thus, an increase in CRC screening test completion rates. The impact of the intervention on CRC screening knowledge, barriers, attitudes, and
Patient activation increases colorectal cancer screening intention is the topic of another manuscript, as is a content analysis of the CRC screening discussions.

Overall, the results demonstrate an increase in CRC screening completion rates among activated patients by medical record review. Self-report of CRC screening discussions and CRC screening tests ordered by medical record review were also found among the patients who received the patient activation intervention. In this study, the control group received CRC screening information, thus the intervention effect, although significant, may have been greater if we used a true control group. In this study, a true control group was not used because we thought it was unethical not to provide minority and low-income patients information about CRC screening.

Telephone CRC screening barriers counseling among this population was not successful due to the fact that only 54 (39.1%) patients in the intervention group received a CRC screening recommendation, 37 (31.4%) of the 118 patients needing CRC screening barriers counseling were not reached, and just 4 (4.9%) of the 81 patients completed screening after telephone barriers counseling (two patients completed FOBT and two patients completed scheduled colonoscopy). The main barriers reported by patients (other medical issues, too busy, etc.) in this study are similar to previous reports of patient-level CRC screening barriers [14, 31, 32]. In addition, it must be noted that even when CRC screening was reported as being discussed during a medical visit (n=111), a CRC screening test was not ordered for 34 (30.6%) patients. Reasons for not ordering a CRC screening test following a CRC screening discussion in this study include forgetfulness, concentration on a different
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medical issue, and not ordering a FOBT for patients without health insurance and who could not afford a colonoscopy.

Although the difference between study arms was significant in this study, the findings also identified that patient-provider CRC screening discussions may not always lead to a CRC screening test being ordered and a test being ordered may not always lead to completion of the recommended CRC screening test. Only 14.8% (40/270) of the patients in this study completed CRC screening. This is similar to national data which reports a 19.5% CRC screening rate among individuals without health insurance [1].

Even though CRC screening rates have increased in the past decade, there remains a trend of lower CRC screening rates within recommended guidelines among minority and underserved populations [9]. Since health care providers are aware of the importance of screening to reduce CRC mortality, there appear to be problems along the steps (encounters) and interfaces (transfer of information) of the screening process [5, 33]. Innovative strategies at the patient, provider, and system levels to improve CRC screening rates among minority and low SES populations are still needed.

In this study, the intervention focused on activating patients to ask their health care provider for a CRC screening test. The results of this study are similar to other patient-level interventions reported recently that used different content, intensity, tailoring, and delivery channels to increase CRC screening rates [34-40]. Overall, modest increases in CRC screening have been documented among patients randomized to a CRC screening intervention delivered via brochures, physician letters, videos, DVDs, decision aids, automated telephone calls, motivational interviewing, or
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patient navigation. For example, Miller and colleagues randomized predominantly minority and low-income patients to a CRC screening decision aid that encouraged patients to discuss CRC screening with their provider vs. a control group [34]. Patients randomized to the CRC screening intervention had more screening tests ordered (30% vs. 21%) and completed (19% vs. 14%) than the patients in the control group, although the differences were not statistically significant.

Even though these interventions have increased CRC screening, they have not improved screening rates to the level of other cancer screening rates (e.g. breast cancer), especially among minority and low-income populations. It is time to recognize that interventions focused only at the patient-level have limited improvements in CRC screening rates. The results of our study and others support evidence that future interventions to increase CRC screening rates need to consider contextual factors at multiple levels [41, 42].

Our study should be interpreted with several limitations. First, we were unable to contact many potentially eligible patients and many patients were not able to arrive one hour early for their appointments to participate in the study. To minimize this problem, we tried to contact patients numerous times on different days and times. Still, the refusal rate for this study was 42%. Although these issues may have caused a selection bias, the internal validity of this study was likely protected by randomization and statistical control for potential confounding by measured baseline characteristics. The generalizability of the results is limited by conducting the study in one FQHC. It is possible that the minority and low-income patients using this health center could be significantly different than other minority and low-income patients using other FQHCs. In
addition, patient agreement for audio taping their medical visits varied by the research assistant (24-83%) and we were not able to contact almost one third of participants in the intervention group for CRC screening barriers counseling, largely because of disconnected telephone numbers or unanswered calls. Finally, we believe a longer follow-up time is needed for CRC screening intervention studies because six additional activated patients who had a colonoscopy ordered at the time of the medical visit completed the test within six months instead of two months, thus outside of the follow-up date set *a priori* in this study.

In spite of limitations, this study was able to recruit a mostly African American and low SES population in need of CRC screening. Additionally, we captured information on participant characteristics, such as health literacy, to assess factors associated with both the intervention and outcome. Audio-taped medical visits provided insight into why screening tests were not recommended in several cases. Furthermore, CRC screening completion was determined by medical record and laboratory log book review. Results, however, may not be generalizable to all FQHCs and to populations outside of Ohio.

In conclusion, activating patients to ask healthcare providers for CRC screening tests improves CRC screening rates compared to providing patients with CRC screening information only. Since the addition of activating patients was successful among minority and low-income patients, this study speaks to the added importance of activating patients in the CRC screening process. Future studies to improve CRC screening among minority and low SES populations should include innovative strategies to motivate providers to recommend screening tests to patients, as well as to assist
patients to complete ordered CRC screening tests (but not telephone barriers counseling).

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The authors would like to acknowledge support from the following grants: K07 CA107079 (MLK); and P30 CA016058 (Behavioral Measurement Shared Resource at The Ohio State University).
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References


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Table 1. Baseline participant demographics by treatment arm (N=270)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>CRC screening information plus patient activation n=138</th>
<th>CRC screening information only n=132</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (median, in years)</td>
<td>55.7</td>
<td>56.3</td>
</tr>
<tr>
<td>Gender (% female)</td>
<td>92 (66.7%)</td>
<td>80 (60.6%)</td>
</tr>
<tr>
<td>Race (% African-American)</td>
<td>104 (75.4%)</td>
<td>91 (68.9%)</td>
</tr>
<tr>
<td>Marital Status* (% married/living as married)</td>
<td>15 (10.9%)</td>
<td>23 (17.4%)</td>
</tr>
<tr>
<td>Education (% less than high school)</td>
<td>40 (29.0%)</td>
<td>34 (25.8%)</td>
</tr>
<tr>
<td>Household Income (% with income &lt; $20,000)</td>
<td>87 (63.0%)</td>
<td>77 (58.3%)</td>
</tr>
<tr>
<td>Health Insurance (% with no insurance)</td>
<td>82 (59.4%)</td>
<td>72 (54.5%)</td>
</tr>
<tr>
<td>Health Literacy: &lt; high school reading level (% with Realm Score &lt; 60)</td>
<td>80 (58.0%)</td>
<td>65 (49.2%)</td>
</tr>
<tr>
<td>Co-morbid conditions (% with 2+ conditions)</td>
<td>46 (33.3%)</td>
<td>37 (28.0%)</td>
</tr>
<tr>
<td>Previous CRC screening (% reporting previous test completion)</td>
<td>34 (24.6%)</td>
<td>31 (23.5%)</td>
</tr>
<tr>
<td>Shared decision making (% prefer involvement with medical decisions)</td>
<td>101 (73.2%)</td>
<td>105 (79.6%)</td>
</tr>
</tbody>
</table>

CRC = colorectal cancer
*p<0.05
Table 2. Baseline colorectal cancer (CRC) screening characteristics of participants by treatment arm (N=270)

<table>
<thead>
<tr>
<th></th>
<th>CRC screening information plus patient activation (n=138)</th>
<th>CRC screening information only (n=132)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRC screening knowledge*</td>
<td>6.1 ± 1.8</td>
<td>6.3 ± 1.7</td>
</tr>
<tr>
<td>(mean ± SD)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CRC screening attitudes and beliefs**</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(mean ± SD)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salience and Coherence</td>
<td>2.9 ± 0.9</td>
<td>2.8 ± 0.8</td>
</tr>
<tr>
<td>Self-Efficacy</td>
<td>2.5 ± 0.8</td>
<td>2.5 ± 0.8</td>
</tr>
<tr>
<td>Perceived Susceptibility</td>
<td>2.5 ± 0.7</td>
<td>2.3 ± 0.8</td>
</tr>
<tr>
<td>Worries and Fears (including barriers)</td>
<td>3.0 ± 1.0</td>
<td>2.9 ± 0.9</td>
</tr>
<tr>
<td>Intention</td>
<td>2.4 ± 0.8</td>
<td>2.6 ± 0.7</td>
</tr>
</tbody>
</table>

*CRC screening knowledge (correct number from 10 true/false questions)
**CRC screening attitudes and beliefs (strongly agree=1 and strongly disagree=4)[33]
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Table 3. Odds ratios and 95% confidence intervals from adjusted logistic regression models for discussing CRC screening with a healthcare provider, having a CRC screening test ordered, and completing CRC screening

<table>
<thead>
<tr>
<th>CRC screening</th>
<th>CRC screening information plus patient activation (n=138)</th>
<th>CRC screening information only (n=132)</th>
<th>OR (95% CI)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Discussion*</td>
<td>75 (54.4)</td>
<td>36 (27.5)</td>
<td>3.29 (1.95, 5.56)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Ordered**</td>
<td>54 (39.1)</td>
<td>23 (17.6)</td>
<td>3.40 (1.88, 6.15)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Completion**</td>
<td>27 (19.6)</td>
<td>13 (9.9)</td>
<td>2.35 (1.14, 5.56)</td>
<td>0.020</td>
</tr>
</tbody>
</table>

*Self-report; **Medical Record Review; adjusted for age, gender, race, and provider
Figure 1. Study design

Eligible patients
Consent forms signed
Baseline survey

Randomization

Intervention Group
CRC screening information
plus patient activation

Control Group
CRC screening
information only

Post intervention survey

Medical visit
(Audio recording if consented)

Post medical visit survey
and medical record review

1 month telephone
barriers call
(if needed among
activated patients)

1 month medical record/laboratory log review

2 month medical record/laboratory log review
Figure 2. Study flow of participants, CONSORT diagram

1,597 Medical Records Reviewed

814 Patients Ineligible
1) Within screening guidelines=291
2) High risk = 201
   (CRC personal or family history, polyps)
3) Not English speaking=172
4) Medical reason=121
   (blindness, deafness, cognitive limitations)
5) Medical appointment rescheduled=18
6) CRC screening test recently ordered=11

783 Patients Eligible
1) Contacted and agreed to participate=331
2) Contact not attempted=209
   (another patient scheduled at that time/staff not available)
3) Refused=148
   (cannot arrive early=66; not interested=47; no reason=21; no time=14)
4) Not able to contact = 95
   (No answer=53; number d/c or wrong=28; lives in shelter=14)
5) Contact not attempted=209
   (another patient scheduled at that time/staff not available)
6) Refused=148
   (cannot arrive early=66; not interested=47; no reason=21; no time=14)

331 Patients agreed to participate
   (No show=47)

284 Participants Randomized

138 Randomized to CRC screening information plus activation
132 Randomized to CRC screening information

270 Analysis: MRR at 2 months
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