Implementation of a mailed faecal immunochemical test programme for colorectal cancer screening among Veterans

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ABSTRACT

Background Screening for colorectal cancer (CRC) with faecal immunochemical test (FIT) is effective at reducing CRC mortality. Unfortunately, the COVID-19 pandemic has been associated with deferred care, especially screening for CRC.

Aim We sought to develop a mailed FIT programme (MFP) to increase CRC screening and make recommendations for adoption across the Veterans Health Administration (VHA) and for other large healthcare systems.

Setting 2 regional VA medical centres in California and Washington state.

Participants 5667 average risk veterans aged 50–75 overdue or due within 90 days for CRC screening.

Programme description A multidisciplinary implementation team collaborated to mail an FIT kit to eligible veterans. Both sites mailed a primer postcard, and one site added an automated reminder call.

Programme evaluation We monitored FIT return and positivity rate, as well as impact of the programme on clinical staff. 34% of FIT kits were returned within 90 days and 7.8% were abnormal.

Discussion We successfully implemented a population-based MFP at multiple regional VA sites and recommend that these efforts be spread across VA. Our model of regional leadership, facility champions and using centralised resources can be adaptable to other large healthcare systems. MFPs support catch-up from disrupted care by addressing access to CRC screening, unburden primary care visits and conserve limited procedural resources.

INTRODUCTION

Colorectal cancer (CRC) is the second leading cause of cancer death in the USA. Screening reduces CRC mortality and incidence. The faecal immunochemical test (FIT) is one of several recommended CRC screening options and can be performed at home.3 4 The standard of care for CRC screening in the Veterans Health Administration (VA) includes identifying individuals due for screening through an electronic health record (EHR) reminder during a primary care visit. Providers determine if the veteran is appropriate for screening and provide a FIT kit during the clinic visit, or order an alternative screening test (eg, colonoscopy) as clinically indicated.

At the start of the COVID-19 pandemic, the VA issued guidance to defer all non-urgent and elective procedures, including screening colonoscopy, and FIT was designated as the preferred CRC screening option.6 At the same time, a dramatic shift of primary care visits to telehealth modalities disrupted traditional CRC screening workflows.5 The USA as a whole experienced a decline in CRC diagnosis and screening during the pandemic.6,7 In response, the VA sought to identify an alternative approach to CRC screening.

Based on the urgent need to increase CRC screening, our teams in Washington state and California collaborated on a
quality improvement initiative to establish a mailed FIT programme (MFP) in the VA. MFPs are a population health approach used by healthcare systems to increase CRC screening rates. These programmes proactively target an entire population of patients who are due for CRC screening, but do not rely on a traditional primary care visit. This helps unburden primary care teams who are responsible for numerous preventative health recommendations. MFPs have demonstrated increased screening rates across various populations including low-income and racially diverse groups, and among rural veterans.

In this report, we describe the process for establishing an MFP at multiple regions in the VA. We describe initial results from large-scale implementation at pilot sites and make recommendations on ways to implement MFPs at VAs across the country.

Setting and participants
We implemented an MFP at two regional pilot sites: VA Central California Health Care System and VA Puget Sound Health Care System. VA Central California serves approximately 26,000 veterans across four sites of care and VA Puget Sound serves over 100,000 veterans across eight sites of care in Washington state. The MFP teams consisted of a clinical lead, programme manager and analysts who partnered with supply chain staff, laboratory staff and contracted mailing and logistics entities. The Central California team operated at the regional office level and the Puget Sound team was a local facility-based team. The pilot programme had the support of regional and national primary care and gastroenterology leadership.

Eligible veterans included individuals aged 50–75 years with at least one outpatient visit within the past 2 years (Puget Sound, n=20,090; Central California, n=12,167). The VA uses an EHR-based clinical reminder that prompts average risk screening for age-appropriate individuals unless prior screening (eg, prior colonoscopy, sigmoidoscopy or FIT) indicated that other action was appropriate (eg, repeat colonoscopy). This clinical reminder formed the basis of the average risk cohort identified for the MFP (Puget Sound, n=14,543; Central California, n=9,186). Veterans were excluded if they were up to date with appropriate CRC screening (Puget Sound, n=9,928; Central California, n=6,196), scheduled for upcoming colonoscopy or sigmoidoscopy within 90 days (Puget Sound, n=36; Central California, n=n/a), enrolled in hospice (Puget Sound, n=29; Central California, n=11), lacked a mailing address (Puget Sound, n=167; Central California, n=2) or were newly started on clopidogrel within the past 6 months (Puget Sound, n=140; Central California, n=67). We defined a new clopidogrel start as a prescription filled within the last 180 days, but not the prior 180–365 days. A total of 4887 individuals at Puget Sound and 2912 at Central California were eligible for inclusion at the start of the programme. These inclusion and exclusion criteria were reapplied to eligible patients before each subsequent round of approximately 500 FIT kit mailings, capped by the availability of laboratory processing.

Programme description
Our MFP (figure 1) incorporated best practices from other reported MFPs, including a primer postcard mailed before the FIT kit and an automated reminder call. VA Central California followed this protocol for the pilot. To gain further insight into what components are necessary for an MFP, the VA Puget Sound team designed randomised controlled trials of programme elements. The data presented include the first five rounds of the VA Puget Sound pilot, during which primer postcards were sent to half of patients and no reminder calls were sent. The data presented include the first five rounds at Central California, which were conducted over a period of 3 months.

We used centralised contractors for printing, labelling and mailing FIT kits (US Government Printing Office and Western States Network Consortium Regional Reproduction Center). A primer postcard was mailed to participants introducing FIT for CRC screening (online supplemental figure 1). Two weeks later, veterans were mailed prelabelled FIT kits, instructions and a return envelope with prepaid postage. Veterans were instructed to collect a stool sample, record the date of collection and return the kit as soon as possible.

Returned FIT kits were analysed by laboratory staff, with results recorded in the EHR and alerted to the
primary care provider (PCP). Tests received without a collection date or 15 days after the collection date have a comment to alert the PCP, as FIT kits processed >15 days after collection have a high false-negative rate. VA Central California patients who had not returned their kits within 2 weeks received an automated phone call reminder (AudioCARE Communicator, Wayne, Pennsylvania).

The MFP teams monitored for kits returned to sender and FIT sample laboratory errors. We communicated with contractors, laboratory staff and clinical teams via secure email and through staff meetings to identify and address any issue with the MFP.

The implementation of the MFP was operational and designated as a non-research activity.

**Patient and public involvement**

Prior to launch of the MFP, a primary care clinic advisory board that comprised patients provided feedback on the programme and mailed material at VA Puget Sound. Patients were not involved in the design, delivery or dissemination of the MFP.

**Programme monitoring**

Our MFP pilot was conducted between March and July 2021 at the VA Central California and between July and October 2021 at VA Puget Sound. Table 1 shows the demographic data for the cohort included in the MFP intervention.

To capture routine care at each site, we measured the number of ordered and resulted FITs before implementation. In order to avoid overlap with the MFP cohort, preimplementation FIT data were from 3 to 6 months prior to the MFPs (ie, September through December for Central California, and January through April for Puget Sound). We implemented the programme and analysed the total return and positivity rates at 90 days after mailing the kits. FIT results were limited to one result per patient during the evaluation period and excluded kits returned before the mailed date. Table 2 shows the mean 90-day FIT return rates before and after MFP implementation at each site. During 4 months of routine care prior to the MFP implementation, 1259 FITs were ordered at Central California with a 39% (n=492) 90-day return rate, and 1510 FITs were ordered at Puget Sound with a 48% (n=725) 90-day return rate. After implementation of the MFP, 3635 FITs were mailed and the 90-day return rate was 35% (n=1277) at Central California, and 2022 FITs were mailed and 90-day return rate was 33% (n=672) at Puget Sound.

The Central California programme included automated reminder calls sent 2–4 weeks after the mailed FIT. Of the 3635 individuals included in the MFP, 3513 (96%) patients were successfully called by the automated phone system, of which 2989 patients (43.5%) were reached (ie, call answered or voice mail).

### Table 1 Mailed FIT programme cohort demographics

|                       | Central California (n=3635) | Puget Sound (n=2022) |
|-----------------------|---------------------------|----------------------|
| **Age**               |                           |                      |
| 50–64                 | 1552 (43)                 | 1011 (50)            |
| 65–75                 | 2083 (57)                 | 1011 (50)            |
| **Sex**               |                           |                      |
| Male                  | 3415 (94)                 | 1806 (89)            |
| Female                | 220 (6)                   | 216 (11)             |
| **Race/ethnicity**    |                           |                      |
| Non-Hispanic white    | 2046 (56)                 | 1354 (67)            |
| Hispanic              | 852 (23)                  | 79 (3.9)             |
| Non-Hispanic black    | 301 (8.3)                 | 359 (18)             |
| Asian/Pacific Islander/Native Hawaiian | 106 (2.9) | 101 (5.0) |
| Multirace             | 54 (1.5)                  | 44 (2.2)             |
| American Indian/ Alaska Native | 34 (0.9) | 23 (1.1) |
| Other                 | 18 (0.5)                  | 3 (0.1)              |
| **Rurality**          |                           |                      |
| Urban                 | 2673 (74)                 | 1521 (75)            |
| Rural                 | 962 (26)                  | 454 (22)             |
| **Prior FIT completion in the VA** |                  |                      |
| Completed ≥1 FIT†     | 668 (18)                  | 1046 (52)            |
| Never completed       | 2967 (82)                 | 976 (48)             |

*Numbers may not add to totals and percentages to 100% due to missing data (ie, unknown response).
†Completed ≥1 FIT kit within the past 5 years.
FIT, faecal immunochemical test; VA, Veterans Health Administration.

**DISCUSSION**

We successfully implemented an MFP across two regional pilot sites to improve access to CRC screening for veterans. Our preliminary results found a veteran patient population with an average 34% FIT completion rate within 90 days across both sites. The return rate was slightly higher at the Central California VA than the Puget Sound VA (35% vs 33%), which may be attributable to the addition of an automated reminder call in the Central California programme. Despite baseline practice variation in use of FIT, with more patients having completed a prior FIT at the Puget Sound than the Central California VA (52% vs 18%), the MFP return rate was similar at both sites. Additionally, our MFP return rate was comparable to that seen in other MFPs, which range from 26% to 59%.10 Though FIT return rates were lower with the MFP compared to the MFP cohort, the return rate was significantly higher than the return rate seen in other MFPs, which range from 26% to 59%.10

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than the preimplementation FIT screening, the MFP has several added benefits. The MFP: (1) systematically reaches more patients in a shorter time period, (2) saves time for busy primary care clinics, which carry the burden of preventative screening efforts, (3) includes individuals who may not seek in-person healthcare visits and (4) reduces demand for colonoscopies as the primary screening strategy, which is especially valuable given the deferral of procedures due to the COVID-19 pandemic.

VA strives to be an ever-evolving learning health system (LHS), through leveraging research and a culture of innovation. An LHS approach promotes continuous, timely learning to improve the outcomes of individuals, populations and healthcare organisations. We adapted an LHS framework to describe the key components of our MFP, as well as our success, challenges and lessons learnt (Box 1).

At the individual level, we increased screening access and efficiently screened numerous veterans. However, similar to other MFPs, the majority still did not return their FIT kit. MFPs that incorporated advanced notifications and reminders have shown modest effects on increasing FIT completion rates. More work is needed to understand which strategies may increase MFP screening participation among veterans. In addition, we identified some logistical challenges. Despite incorporating best practices from other programmes, including a prelabelled kit, many veterans neglected to date their FIT kit, which can lead to unclear results and repeat testing.

At the team level, primary care and laboratory staff were familiar with processes for FIT. This enabled quicker programme implementation as no significant training was needed. Given that the programme is run by a separate dedicated team and not the individual PCP team, less workload fell onto primary care staff. While the MFP reduced primary care workload, it increased laboratory staff work due to manual FIT order entry. Work is underway to transition to a batch FIT ordering process. Early pilots used clinic staff for printing, labelling and mailing; however, we quickly determined that this was not sustainable for a large-scale effort. Moreover, we found it was important to establish clear communication channels between the MFP implementation team and front-line staff. This allowed staff to raise issues with the MFP, and to clarify workflows for in-clinic versus population health CRC screening. The communication channels with front-line staff were also essential for quality assurance. Early in the Puget Sound pilot, staff reported FIT results for some veterans who were not average risk, which identified an error in cohort identification that was quickly corrected.

| Intervention phase | Total ordered/mailed | Total returned 90 days n (%) | Total positive n (%) |
|--------------------|---------------------|-----------------------------|---------------------|
| Central California |                     |                             |                     |
| Premailed FIT      | 1259                | 492 (39)                    | 49 (9.9)            |
| Mailed FIT         | 3635                | 1277 (35)                   | 88 (6.9)            |
| Puget Sound        |                     |                             |                     |
| Premailed FIT      | 1510                | 725 (48)                    | 87 (12)             |
| Mailed FIT         | 2022                | 672 (33)                    | 58 (8.6)            |

To capture routine care, we identified ordered and returned FIT kits from 3 to 6 months prior to the implementation of the MFPs (ie, premailed FIT above). 90-day return rates are shown for the pre-MFP and post-MFP implementation.

FIT, faecal immunochemical test; MFP, mailed FIT programme.
At the organisational and operating level, primary care, gastroenterology and laboratory leadership were supportive of the MFP. Our MFPs were quickly implemented in response to reduced CRC screening due to the COVID-19 pandemic. However, we identified key steps to address with leaders. First, it was necessary for leadership to budget time and resources to create an MFP team—rather than use internal staff. Teams need protected time to manage the MFP, including working with logistics, contractors and health administration systems. Second, it was important for the MFP team to have experienced analysts. This allowed for continuous monitoring of outcomes and adaptation of the programme during implementation. Regional MFP teams should work in partnership with facility leadership, identify site champions and ensure local workflows allow proper follow-up of FIT results. Finally, implementation of the MFP required special agreements with leadership in several divisions. We had to have special agreements with laboratory staff to enter FIT orders. In addition, it was vital to have discussions with gastroenterology to address priority colonoscopy for veterans with a positive test result.

Given the known benefits of the MFP and the successes our team had in adapting the programme for the VA setting, we believe our pilot provides a blueprint for other VA primary care sites to implement MFP. Throughout this project, our team considered how our programme can be both implemented and sustained as a model that could be expanded nationally in the VA. At each step, we attempted to build a sustainable model, customised to the individual, team and organisational and operational factors, needed to develop an MFP. Moreover, we find that with leadership to champion the programme, appropriate staff for implementation, and funding support, this model could be sustainable for continuous operations of the MFP in the VA.

Limitations
We acknowledge several limitations. Importantly, we cannot account for the exact time and effort spent by primary care team members on screening efforts that may have occurred concurrently to the MFP efforts. Routine care continued throughout the MFP, and it is possible that veterans received kits and/or reminders from their clinical team at visits in addition to those from the MFP. Second, this pilot was conducted among veterans enrolled at the VA, a majority of whom are men, and results may not generalise to populations outside the VA. However, VA cares for over 6 million veterans in primary care and improving CRC screening among veterans is key priority.

One concern raised is that overscreening may happen when implementing an MFP. Overall, using an MFP results in only a small number of individuals who get duplicative screening. Overscreening can also happen in general practice when screening records are not up to date and unnecessary testing is performed.

CONCLUSIONS
We successfully implemented a large-scale population-based MFP at multiple regional VA sites. Our preliminary results show 34% of veterans completed CRC screening, and we were able to invite more patients to complete screening than through the usual care process. With pandemic-related care disruptions, it is more important than ever that healthcare systems seek new ways to provide needed preventive services. VA is primed for the opportunity to create a national MFP. Additional work is needed to determine if the MFP implementation will be as successful when spread to additional sites, and if the same model is viable. Further evaluations will assess the impact of the MFP on disparities and seek to test specific programmatic features that can enhance CRC screening rates among veterans.

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Contributors
All authors played a role in fulfilling all four criteria: (1) design, data, interpretation; (2) drafting and revising for intellectual content; (3, 4) final approval and will stand by the work.

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