ABSTRACT

Objective: To investigate the diagnostic activity in general practice and the cumulative incidence of colorectal cancer (CRC) in individuals invited to the Danish national screening programme for CRC.

Design: A historical population-based cohort study.

Setting: The Danish CRC screening programme and general practice.

Subjects: The 376,198 individuals invited to the Danish CRC screening programme from 1 March to 31 December 2014.

Main outcome Measures: The diagnostic activity (consultations and haemoglobin measures) in general practice in the year preceding the screening invitation and the cumulated incidence of CRC in the year following the screening invitation.

Results: Screening participants had significantly higher diagnostic activity than non-participants. Individuals with a positive faecal immunochemical test (FIT) had higher diagnostic activity compared to individuals with a negative FIT, and a small increase in the months leading up to the invitation. Individuals with a screen-detected CRC had lower diagnostic activity than individuals with no CRC. In total, 308 (25.3%) of CRCs diagnosed in the invited population were diagnosed outside the screening programme. Non-participants with CRC more often had low socio-economic status, high comorbidity and stage IV CRC than participants with CRC.

Conclusions: There was a tendency that participants and those with a positive FIT had a higher diagnostic activity the year before the screening. This was not seen for those with CRC detected through screening. CRC must still be diagnosed in general practice in the invited population and non-participants are of special interest as they have higher risk of late stage CRC.

KEY POINTS

- Current awareness: Individuals with colorectal cancer (CRC) in screening may be symptomatic and CRC may still occur outside screening in the invited population.
- Most important points: The majority of individuals with CRC in screening cannot be expected to be diagnosed on symptomatic presentation in general practice.
- GPs have to be aware that CRC still occurs outside screening in the invited population.
- Non-participants with CRC are often deprived and have late stage CRC.

Introduction

Colorectal cancer (CRC) is the third most common cancer worldwide and one of the main reasons for cancer-related death [1]. During decades, Denmark has had poorer CRC survival than other Nordic countries. This may partly be explained by late stage of disease at diagnosis [2]. In an attempt to identify cancer at earlier stages and thereby improve the CRC prognosis, a screening programme based on the use of the faecal immunochemical test (FIT) was...
implemented in March 2014 for individuals aged 50-74 years [3].

For individuals participating in the screening, recent questionnaire studies have indicated that lower gastrointestinal (GI) symptoms are prevalent in up to 70-80% of the individuals with a positive faecal test [4–6]. If this is true, these individuals may be detected before entering the screening programme through symptomatic presentation in general practice if the general practitioner (GP) had access to the FIT and thus, providing the opportunity to detect the CRC earlier.

For individuals invited to, but not participating in the screening, studies indicate that general practice will play a key role in the detection of CRC [7]. However, so far, no study has investigated the amount of CRCs occurring outside the Danish screening programme in the invited population, but for GPs this is of special interest since non-participants diagnosed with CRC may have low socio-economic status (SES) and poor CRC prognosis [8–10].

Thus, the aim of this study was to investigate the diagnostic activity in general practice and the cumulative incidence of CRC in individuals invited to the Danish national screening programme for CRC.

**Material and methods**

**Study design**

We conducted a historical population-based cohort study using registry data to assess the diagnostic activity in general practice and the cumulative incidence of CRC in individuals invited to the Danish national screening programme for CRC.

**Setting**

From March 2014, CRC screening is being implemented over a 4-year period in Denmark and will be offered biennially after full implementation. The screening is free of charge and uses the FIT as a first line test, which is followed by colonoscopy (or alternatively CT-colonography) if the FIT is positive (≥100μg/L).

All citizens aged 50-74 years are invited to participate in the screening. Citizens participate by performing a FIT received in an invitational letter. The order in which citizens are invited is random and determined on the basis of their month of birth. However, citizens turning 50 or 74 years during the prevalence screening must receive a screening invitation before that particular birthday.

General practice is not involved in any part of the Danish screening programme for CRC. In Denmark, access to general practice is free of charge at the point of care and almost 99% of Danish citizens are listed with a general practice [11].

**Study population**

All individuals invited to screening for CRC from 1 March to 31 December 2014 were eligible for inclusion. To ensure complete follow-up, we excluded individuals who were not listed with a GP, had lived outside Denmark at some point during the year preceding the screening invitation or had died within one month after receiving the invitation. Furthermore, to ensure a homogeneous population, we restricted the analyses to individuals without a previous diagnosis of colorectal disease (CRC, Crohn’s disease, ulcerative colitis, familial adenomatous polyposis (FAP), hereditary nonpolyposis colorectal cancer (HNPCC) or adenomas followed by regular colonoscopy).

**Outcome measures**

**Diagnostic activity in general practice:** Daytime face-to-face consultations (incl. home visits) and point-of-care haemoglobin measurements in general practice (photometric analysis) were used as proxies for diagnostic activity in general practice related to gastrointestinal symptoms and signs, and assessed in the year preceding the screening invitation. The outcome was compared for participants vs. non-participants, individuals with positive FIT vs. individuals with negative FIT and CRC cases vs. non-CRC cases.

**Cumulative incidence of CRC:** Number of CRC diagnoses among individuals invited to screening in the year following the screening invitation. The CRC incidence was stratified into three subgroups: participants diagnosed in the screening, participants diagnosed outside the screening and non-participants.

**Data collection**

Data on diagnostic activity was collected from the Danish National Health Service Register until three years preceding the screening invitation to allow for extension of the study period if differences were present for more than one year [12]. The database holds no information on results of laboratory analyses.

Data on CRC (ICD-10: DC180-9 and DC200-9) and UICC stages was collected from the Danish Cancer Registry from the day of invitation until one year following the screening invitation [13,14].

The Danish Colorectal Cancer Screening Database (DCCSD) provided information on the screening
invitation, FIT results, performed colonoscopy or CT-colonography and screen-detected CRCs [15].

The Danish National Patient Register (NPR) was used to identify previous diagnoses of colorectal disease (Crohn’s disease (ICD-10: DK500-9), ulcerative colitis (ICD-10: DK510-9), FAP (ICD-10: DD126F), HNPCC (ICD-10: DC188A) and adenomas followed by regular colonoscopy (ICD-10: DZ018B) [16]. Furthermore, the NPR was used to identify diagnoses for generating the Charlson Comorbidity Index, which was categorised into “low” (CCI score = 0), “moderate” (CCI score = 1-2) and “severe” (CCI score ≥3) (CCI) [17,18].

The Danish National Prescription Registry was used to collect information on prescriptions for medications against haemorrhoids (ATC: C05A) or drugs with anticoagulatory effect (NSAIDs (ATC: M01A), acetylsalicylic acids (ATC: B01AC06, N02BA01, N02BA51) and anticoagulants (ATC: B01A)) as these covariates were considered potential confounders for the outcomes [19].

Statistics Denmark provided data on vital status, socio-economic characteristics and demographic factors [20]. Marital status was dichotomised into living with a partner (“married/cohabitating”) or living alone (“alone”). Country of origin was categorised into “Danish”, “immigrant from a western country” or “immigrant from a non-western country”. Labour market affiliation was categorised into “working”, “unemployed” or “retirement pension”. Educational level was categorised into “basic” (<10 years), “medium” (10-15 years) and “high” (>15 years) [21].

All data were linked by the civil registration number [22].

**Statistical analysis**

The index date was defined as the date the individual was invited to screening for CRC.

The diagnostic activity in general practice was investigated in the year preceding the screening invitation by estimating the consultation rates and haemoglobin measurements rates. This was done in intervals of three months. The diagnostic activity was compared between subgroups by estimating incidence rate ratios (IRRs) using negative binomial regression models with cluster robust variance estimation to account for heterogeneity between individuals. Estimates of the IRRs were adjusted for age, gender, marital status, country of origin, level of education, labour market affiliation, comorbidity, prescriptions of medicine against haemorrhoids and prescriptions of medicine with an anticoagulatory effect. The cumulative incidence of CRC among individuals invited to screening was investigated in three subgroups to allow assessment of the number of CRCs diagnosed outside the screening programme. CRCs diagnosed among participants outside the screening programme were defined as; a CRC diagnosis after a negative FIT in the screening, or a CRC diagnosis registered in the DCR for an individual with a positive FIT in the screening, but no registered CRC diagnosis in the DCCSD.

In addition, we compared participants’ and non-participants’ risk of being diagnosed with a stage IV CRC by using a Poisson regression model with time at risk as exposure while taking into account the competing risk of getting a lower stage cancer.

All analyses were performed on the server of Statistics Denmark using Stata 14.0.

**Ethical approvals**

The study was approved by the Danish Data Protection Agency (j.no. 2014-41-3143).

**Results**

**Diagnostic activity in the year preceding the screening invitation**

In total, 390,552 individuals were invited to screening from 1 March 2014 to 31 December 2014. After exclusions, 376,198 individuals were included in the analyses (Figure 1). Among the 245,299 (65%) participants, 16,206 (6.6%) had a positive FIT result and 907 (6.1%) of individuals who had a colonoscopy or CT-colonography performed were diagnosed with CRC (Figure 1).

Participants vs. non-participants: Participants in screening were more often females and had higher SES and less comorbidity than non-participants (Table 1). During the year preceding the screening invitation, participants had significantly higher consultation rates and haemoglobin measurements than non-participants (Figure 2(A)). This was also seen when extending the study period to three years (results not shown).

Individuals with positive FIT vs. individuals with negative FIT: More males than females had a positive FIT in screening. In addition, individuals with a positive FIT had lower SES, higher CCI score and more often received medicine with an anticoagulatory effect (Table 1). Higher consultation rates were seen for individuals with a positive FIT compared to individuals with a negative FIT, but also a small insignificant increase in the last three months preceding the screening invitation compared with the three-month
period earlier (Figure 2(B)). The rate of haemoglobin measurements was generally higher for individuals with positive FIT, also with an insignificant increase in the last three months.

**Individuals with colorectal cancer vs. individuals with no colorectal cancer:** Individuals diagnosed with a screen-detected CRC were more often males and of higher age than individuals with no CRC (Table 1). The consultation rates were lower for individuals who were diagnosed with CRC compared to individuals without CRC (Figure 2(C)). This did not change if the rates were stratified for CRC stages and was seen constantly if the study period was extended to three years (data not shown). There were no significant differences in the rates of haemoglobin measurements.

**Cumulative incidence of CRC in the year following the screening invitation**

One year after the screening invitation, 239 (0.18%) non-participants and 976 (0.40%) participants had
been diagnosed with CRC. Among participants, 69 CRC incidents were diagnosed outside the screening programme (Figure 3) implying that in total; 308 (25.3%) of the CRCs diagnosed within the first year after the invitation were identified outside the screening programme.

The characteristics of individuals diagnosed with CRC in the year following the screening invitation are shown in Table 2. Non-participants with CRC were more likely to have low SES and to have higher CCI score compared to participants. The overall risk of being diagnosed with a stage IV CRC was 0.021% (95% confidence interval (95%CI): 0.015; 0.027) for participants and 0.035% (95%CI: 0.026; 0.047) for non-participants, corresponding to a 67% increased risk of late stage (data not shown).

**Table 1. Characteristics of individuals included in the analyses on diagnostic activity.**

|                      | Participants | Non-participants | Positive FIT | Negative FIT | Colorectal cancer | No colorectal cancer |
|----------------------|--------------|------------------|--------------|--------------|-------------------|---------------------|
| Total                | 245,299      | 130,899          | 16,206       | 229,093      | 907               | 13,955              |
| Gender†              |              |                  |              |              |                   |                     |
| Male                 | 113,632      | 46.3             | 71,912       | 53.7         | 9,083             | 56.1                |
| Female               | 131,663      | 53.7             | 58,978       | 45.1         | 7,123             | 43.9                |
| Age (years)          |              |                  |              |              |                   |                     |
| 50-54                | 83,138       | 33.9             | 54,306       | 41.5         | 3,467             | 21.4                |
| 55-59                | 35,887       | 14.6             | 18,727       | 14.3         | 1,910             | 11.8                |
| 60-64                | 35,599       | 14.5             | 16,003       | 12.2         | 2,392             | 14.7                |
| 65-69                | 39,734       | 16.2             | 15,903       | 12.2         | 3,254             | 20.1                |
| 70-74                | 50,941       | 20.8             | 25,960       | 19.8         | 5,183             | 32.0                |
| Country of origin    |              |                  |              |              |                   |                     |
| Danish               | 229,909      | 93.7             | 118,402      | 90.5         | 15,360            | 94.8                |
| Immigrant (western)  | 7,569        | 3.1              | 5,486        | 4.2          | 468               | 2.9                 |
| Immigrant (non-western) | 7,821    | 3.2              | 7,011        | 5.3          | 378               | 2.3                 |
| Educational level     |              |                  |              |              |                   |                     |
| Basic                | 65,204       | 26.6             | 46,202       | 35.3         | 5,048             | 31.1                |
| Medium               | 121,980      | 49.7             | 59,217       | 45.2         | 8,273             | 51.1                |
| High                 | 58,115       | 23.7             | 25,480       | 19.5         | 2,885             | 17.8                |
| Labour marked affiliation† |        |                  |              |              |                   |                     |
| Working              | 136,066      | 55.5             | 66,792       | 51.0         | 6,576             | 40.6                |
| Unemployed           | 37,298       | 15.2             | 29,619       | 22.6         | 2,691             | 16.6                |
| Pension              | 71,931       | 29.3             | 34,479       | 26.4         | 6,939             | 42.8                |
| Marital status†      |              |                  |              |              |                   |                     |
| Married/cohabiting   | 184,196      | 75.1             | 77,840       | 59.5         | 11,526            | 71.1                |
| Single/living alone  | 61,099       | 24.9             | 53,050       | 40.5         | 4,680             | 28.9                |
| Charlson Comorbidity Index |           |                  |              |              |                   |                     |
| Low                  | 186,198      | 75.9             | 95,442       | 72.9         | 10,591            | 65.4                |
| Moderate             | 47,443       | 19.3             | 26,000       | 19.9         | 4,109             | 25.3                |
| Severe               | 11,658       | 4.8              | 9,457        | 7.2          | 1,506             | 9.3                 |
| Prescription for medicine against haemorrhoids No | 236,184 | 96.3 | 127,613 | 97.5 | 15,521 | 95.8 | 220,663 | 96.3 | 870 | 95.9 | 13,396 | 96.0 |
| Yes                  | 9,115        | 3.7              | 3,286        | 2.5          | 685               | 4.2                 |
| Prescription for NSAID No | 194,067 | 79.1 | 105,945 | 80.9 | 12,479 | 77.0 | 181,588 | 79.3 | 743 | 81.9 | 10,691 | 76.6 |
| Yes                  | 51,232       | 20.9             | 24,954       | 19.1         | 3,727             | 23.0                |
| Prescription for acetylsalicylic acids No | 214,396 | 87.4 | 114,097 | 87.2 | 12,942 | 79.9 | 201,454 | 87.9 | 729 | 80.4 | 11,214 | 80.4 |
| Yes                  | 30,903       | 12.6             | 16,802       | 12.8         | 3,264             | 20.1                |
| Prescription for anticoagulant drugs No | 228,729 | 93.2 | 120,686 | 92.2 | 13,955 | 86.1 | 214,774 | 93.8 | 803 | 88.5 | 12,091 | 86.6 |
| Yes                  | 16,570       | 6.8              | 10,213       | 7.8          | 2,251             | 13.9                |

†Information on gender, labour marked affiliation and marital status were missing for 13 individuals.

Discussion

Principal findings

Compared to non-participants, screening-participants had higher SES and a higher diagnostic general practice activity in the year preceding the screening. Individuals with a positive FIT had lower SES, more comorbidity and a higher diagnostic activity compared to individuals with a negative FIT. This group also showed a small increase in the three months preceding screening. Individuals diagnosed with CRC in the screening had lower diagnostic activity than individuals with no CRC.

As expected, the screening helped identify cases of CRC. However, approx. 25% of the CRCs diagnosed in the year following the screening invitation were found...
Figure 2. (A-C) Mean rates of daytime face-to-face consultations and haemoglobin measurements for subgroups in the screening for CRC. Estimates are for 3-month intervals, 12 months preceding invitation to screening. The upper graph illustrates the unadjusted average consultation rate for individuals in subgroups. The lower graph illustrates the IRRs for comparison of subgroups adjusted for age, gender, country of origin, educational level, labour market affiliation, marital status, CCI score, prescriptions of medicine against haemorrhoids, NSAIDs, acetylsalicylic acids and anticoagulant drugs.
Figure 3. Cumulative incidence of CRC among individuals invited for screening participation, one year following invitation. Incidence was stratified for screening participation. In total, 907 CRCs were diagnosed in the screening (red curve) and 308 CRCs were diagnosed outside the screening (blue and green curves).

Table 2. Characteristics for individuals diagnosed with CRC in the year following the screening invitation.

|                        | Participants diagnosed with CRC in the screening | Participants diagnosed with CRC outside the screening | Non-participants diagnosed with CRC |
|------------------------|--------------------------------------------------|------------------------------------------------------|------------------------------------|
|                        | n       | %        | n       | %        | n       | %        |
| Total                  | 907     |          | 69      |          | 239     |          |
| Gender                 |         |          |         |          |         |          |
| Male                   | 538     | 59.3     | 35      | 50.7     | 147     | 61.5     |
| Female                 | 369     | 40.7     | 34      | 49.3     | 92      | 38.8     |
| Age (years)            |         |          |         |          |         |          |
| 50-54                  | 87      | 9.6      | 10      | 14.5     | 28      | 11.7     |
| 55-59                  | 77      | 8.5      | 0       | –        | 14      | 5.9      |
| 60-64                  | 126     | 13.9     | 9       | 13.0     | 47      | 19.7     |
| 65-69                  | 198     | 21.8     | 22      | 31.9     | 38      | 15.9     |
| 70-74                  | 419     | 46.2     | 28      | 40.6     | 112     | 46.8     |
| Country of origin      |         |          |         |          |         |          |
| Danish                 | 881     | 97.1     | 66      | 95.7     | 230     | 96.2     |
| Immigrant (western)    | 19      | 2.1      | 3       | 4.3      | NA      | –        |
| Immigrant (non-western)| 7       | 0.8      | 0       | –        | NA      | –        |
| Educational level      |         |          |         |          |         |          |
| Basic                  | 295     | 32.5     | 24      | 34.8     | 84      | 35.1     |
| Medium                 | 467     | 51.5     | 35      | 50.7     | 109     | 45.6     |
| High                   | 145     | 16.0     | 10      | 14.5     | 46      | 19.3     |
| Labour market affiliation|         |          |         |          |         |          |
| Working                | 310     | 34.2     | 21      | 36.2     | 75      | 31.4     |
| Unemployed             | 88      | 9.7      | 8       | 13.8     | 41      | 17.1     |
| Pension                | 509     | 56.1     | 29      | 50.0     | 123     | 51.5     |
| Marital status         |         |          |         |          |         |          |
| Married/cohabiting     | 665     | 73.3     | 53      | 76.8     | 141     | 59.0     |
| Single/living alone    | 242     | 26.7     | 16      | 23.2     | 98      | 41.0     |
| Charlson Comorbidity Index|       |          |         |          |         |          |
| Low                    | 614     | 67.7     | 45      | 65.2     | 144     | 60.3     |
| Moderate               | 226     | 24.9     | 17      | 24.6     | 60      | 25.1     |
| Severe                 | 67      | 7.4      | 7       | 10.1     | 35      | 14.6     |
| UICC stage             |         |          |         |          |         |          |
| I                      | 246     | 27.1     | 24      | 10.0     | 10      | 14.5     |
| II                     | 160     | 17.6     | 40      | 16.7     | 12      | 17.4     |
| III                    | 171     | 18.9     | 51      | 21.3     | 13      | 18.8     |
| IV                     | 51      | 5.6      | 46      | 19.3     | 13      | 18.9     |
| Unknown                | 279     | 30.8     | 78      | 32.7     | 21      | 30.4     |
outside the screening programme. Non-participants with CRC were more likely to have low SES and more comorbidity and had a 67% increased risk of being diagnosed with a stage IV CRC.

**Strengths and weaknesses**

A major strength of the study was the large study population. This, together with analysing the diagnostic activity in three-month intervals, ensured a high statistical power for these analyses. However, using three-month intervals might have hidden more abrupt changes in the activity. Therefore, we also performed the analyses with monthly intervals, but this did not change the overall findings and estimates.

Using daytime face-to-face consultations and hemoglobin measurements as proxies for presentation of gastrointestinal symptoms and signs in general practice has both advantages and disadvantages. The approach has previously been used in similar studies and is therefore a well-investigated and accepted method [23,24]. These previous studies have shown that CRC patients visit their GP more than the average population in the time leading up to the diagnosis. Using the Danish National Health Service Register for collecting information on general practice consultations enabled us to leave out consultations with preventive focus as these have specific codes in the register. Therefore consultations in this study can be considered as “new events” of symptoms and disease [12]. However, some contacts may have been due to other symptoms than GI-symptoms.

In general, we consider the internal validity of the study to be high. Information bias was diminished by using Danish registers as the primary data source. These databases contain data collected prospectively and independently of this study. However, the DCCSD is a rather newly established database and a recently published paper has shown a sensitivity for CRC of 72%. Therefore, some screen-detected CRC diagnoses may have been missed and overestimated the amount of CRC found outside screening [15]. Selection bias was minimal due to the random order in which individuals were invited to screening and that all citizens are invited regardless of SES and morbidity. However, we cannot rule out that some selection occurred in relation to who chose to participate in the screening. Furthermore, some of the difference found between subgroups regarding SES and comorbidity may be related to differences in age between the groups. Finally, we cannot rule out the presence of residual confounding from comorbidity and medicine with anticoagulatory effect. Firstly, the CCI score accounts for only 17 comorbid conditions and are based on diagnoses from secondary care. Therefore, it may leave out additional comorbidity that could be a reason for increased health-care seeking in general practice. However, we chose to use the CCI score since it is regarded among the best methods to measure comorbidity and that it was not possible to collect diagnoses registered in primary care since the Danish authorities made the existing database inaccessible for researchers in 2015. Secondly, we did not have the possibility of collecting information on over-the-counter sale of drugs with anticoagulatory effect (primarily NSAIDs) and information on diagnoses from general practice. This residual confounding may therefore account for some of the difference between subgroups.

The reported results on the incidence of CRC outside the screening programme are generalizable to other countries using FIT-based screening, but are dependent on participation rate and the screened age group. Furthermore, the results on diagnostic activity are generalizable to health care settings similar to the Danish.

**Comparison with other research**

From the present results it seems that the majority of individuals with a screen-detected CRC are not symptomatic as indicated by similar use of general practice. Thus, it seems unlikely that these individuals may be diagnosed earlier in general practice. However, previous questionnaire studies have indicated that approx. 75-80% of individuals with a positive FIT in a screening programme have lower GI symptoms [4–6]. An explanation for these findings could be recall bias among individuals with a screen-detected CRC. Another explanation could be that the symptoms the patient experience are not serious enough for the patient to consult a GP. Nevertheless, our findings represent a mean for all patients. This could imply that a proportion of the population attend the GP more often before attending the screening due to gastrointestinal symptoms. However, in this study, we cannot identify who or how many this concerns.

In the analyses comparing the diagnostic activity for participants vs. non-participants, we found that participants in screening generally had higher diagnostic activity in the years preceding screening invitation. This is supported by the literature which shows that low SES and a low use of health care services are known to be related to non-participation in screening [8]. We therefore consider the difference a consequence of another health care seeking behaviour and
a lower threshold for acting on symptoms, rather than a true difference in symptom prevalence [25].

In line with the literature, we found that CRC occurs in general practice in the screened age group [7,26]. It has been shown that individuals diagnosed with CRC in general practice attend their GP more in the year preceding the diagnosis than the background population [24]. This may indicate a diagnostic window to detect CRC earlier. Recent studies have suggested that the FIT can be used outside screening in general practice to detect CRC in symptomatic individuals [27–30]. In consideration of this, the FIT may be used in this diagnostic window to reduce the time to diagnosis. In relation to our findings, the majority of screening non-participants were individuals with low SES and high levels of comorbidity. We also saw that this population had higher frequency of positive FITs when they participated in the screening and therefore, the FIT may be a useful diagnostic tool in general practice to detect CRC in this population. However, further research is needed on this topic.

In conclusion, we did not find evidence to support that the majority of individuals with a positive FIT or CRC diagnosed in the screening programme were symptomatic. However, CRC will still occur in general practice in the screened population, and special notice has to be paid on non-participants since they are often deprived and have higher risk of late stage CRC.

Acknowledgements
The authors would like to thank datamanager Kaare Rud Flarup who has assisted in the handling of the data.

Declarations of interest
No potential conflict of interest was reported by the author(s).

Funding
This study was funded by the Committee for Quality Improvement and Continuing Medical Education (KEU) of the Central Denmark Region and the Danish Cancer Society. None of the funding bodies have been involved in designing the study or writing the article, or in the collection, analysis and interpretation of data.

Notes on contributors

Jakob Søgaard Juul is a GP trainee and PhD from the research unit for general practice in Aarhus, Denmark.

Berit Andersen is a professor and head consultant at the department of Public Health Programmes, Randers Regional Hospital, Denmark.

Søren Laurberg is a professor and consultant at the department of Surgery, Aarhus University Hospital, Denmark.

Anders Helles Carlsen is a statistician at the research unit for general practice in Aarhus, Denmark.

Frede Olesen is a professor at the research unit for general practice in Aarhus, Denmark.

Peter Vedsted is a professor at the research unit for general practice and research Centre for Cancer Diagnosis in Primary Care in Aarhus, Denmark.

References
[1] Ferlay J, Soerjomataram I, Dikshit R, et al. Cancer incidence and mortality worldwide: Sources, methods and major patterns in GLOBOCAN 2012. Int J Cancer. 2015;136:E359–E386.
[2] Maringe C, Walters S, Rachet B, et al. Stage at diagnosis and colorectal cancer survival in six high-income countries: A population-based study of patients diagnosed during 2000–2007. Acta Oncol (Madr). 2013;52:919–932.
[3] Danish Health and Medicine Authority. Anbefalinger vedrørende screening for tyk- og endetarmskraeft [Recommendations for colorectal cancer screening]. 2012 [cited 2018 May 1]. Available from: https://www.sst.dk/~media/1327A2433DDDB54C86D031D50FE6D9D6.ashx.
[4] Ahmed S, Leslie A, Thaha MA, et al. Lower gastrointestinal symptoms are not predictive of colorectal neoplasmia in a faecal occult blood screen-positive population. Br J Surg. 2005;92:478–481.
[5] Rajasekhar PT, Ritchie M, Rutter MD, et al. Lower gastrointestinal symptoms are prevalent among individuals colonosced within the Bowel Cancer Screening Programme. Color Dis. 2012;14:e603–e607.
[6] Vaughan-Shaw PG, Cutting J, Borley N, et al. Two-week wait symptoms are prevalent in screened patients with a positive faecal occult blood test but do not predict cancer. Colorectal Dis. 2014;16:40–47.
[7] Hamilton W. Five misconceptions in cancer diagnosis. Br J Gen Pract. 2009;59:441–445.
[8] Woolf A, Dapper EA, de Leeuw JRJ. Colorectal cancer screening participation: a systematic review. Eur J Public Health. 2016;26:158–168.
[9] Frederiksen BL, Jørgensen T, Brasso K, et al. Socioeconomic position and participation in colorectal cancer screening. Br J Cancer. 2010;103:1496–1501.
[10] Egeberg R, Halkjaer J, Rottmann N, et al. Social inequality and incidence of and survival from cancers of the colon and rectum in a population-based study in Denmark, 1994-2003. Eur J Cancer. 2008;44:1978–1988.

[11] Pedersen K, Andersen J, Søndergaard J. General practice and primary health care in Denmark. J Am Board Fam Med. 2012;25:S34–S38.

[12] Andersen JS, Olivarius NDF, Krasnik A. The Danish National Health Service Register. Scand J Public Health. 2011;39:34–37.

[13] Gjerstorff ML. The Danish Cancer Registry. Scand J Public Health. 2011;39:42–45.

[14] International Statistical Classification of Diseases and Related Health Problems 10th Revision (ICD-10)-WHO Version; 2016. Available from: http://apps.who.int/classifications/icd10/browse/2016/en/

[15] Thomsen MK, Njor SH, Rasmussen M, et al. Validity of data in the Danish Colorectal Cancer Screening Database. Clin Epidemiol. 2017;9:105–111.

[16] Lyne E, Sandegaard JL, Rebolj M. The Danish National Patient Register. Scand J Public Health. 2011;39:30–33.

[17] Charlson ME, Pompei P, Ales KL, et al. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. J Chronic Dis. 1987;40:373–383.

[18] Quan H, Sundararajan V, Halfon P, et al. Coding algorithms for defining comorbidities in ICD-9-CM and ICD-10 administrative data. Med Care. 2005;43:1130–1139.

[19] Kildemoes HW, Sørensen HT, Hallas J. The Danish National Prescription Registry. Scand J Public Health. 2011;39:38–41.

[20] Statistics Denmark [Internet]. [cited 2018 June 01]. Available from: http://www.dst.dk/da/

[21] UNESCO Institute for Statistics. International Standard Classification of Education. 2011 [cited 2018 May 1]. Available from: http://uis.unesco.org/sites/default/files/documents/international-standard-classification-of-education-isced-2011-en.pdf.

[22] Bøcker Pedersen C. The Danish Civil Registration System. Scand J Public Health. 2011;39:22–25.

[23] Garnier Christensen K, Fenger-Grøn M, Rud Flarup K, et al. Use of general practice, diagnostic investigations and hospital services before and after cancer diagnosis -a population-based nationwide registry study of 127,000 incident adult cancer patients. BMC Health Serv Res. 2012;12:8.

[24] Libach Hansen P, Hjertholm P, Vedsted P. Increased diagnostic activity in general practice during the year preceding colorectal cancer diagnosis. Int J Cancer. 2015;137:615–624.

[25] Campbell SM, Roland MO. Why do people consult the doctor? Fam Pract. 1996;13:75–83.

[26] Juul JS, Vedsted P. Uncharacteristic colorectal cancer symptoms and their value in general practice. Ugeskr Laeger 2012;174:710–713.

[27] Högberg C, Karling P, Jörgen R, et al. Diagnosing colorectal cancer and inflammatory bowel disease in primary care: the usefulness of tests for faecal haemoglobin faecal calprotectin. Scand J Gastroenterol. 2017;52:69.

[28] Mcdonald PJ, Digby J, Innes C, et al. Low faecal haemoglobin concentration potentially rules out significant colorectal disease. Colorectal Dis. 2013;15:1463–1469.

[29] Mowat C, Digby J, Strachan JA, et al. Faecal haemoglobin and faecal calprotectin as indicators of bowel disease in patients presenting to primary care with bowel symptoms. Gut 2016;65:1463–1469.

[30] Kok L, Elias SG, Witteman BJ, et al. Diagnostic accuracy of point-of-care faecal calprotectin and immunochemical occult blood tests for diagnosis of organic bowel disease in primary care: the cost-effectiveness of a decision rule for abdominal complaints in primary care (CEDAR) study. Clin Chem. 2012;58:989–998.