



Cohort Profile

Cohort Profile: The Colon Cancer Family Registry Cohort (CCFRC)

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Why was the cohort set up?

Colorectal cancer has long been one of the most frequently diagnosed cancers in the world, with an estimated 1.4 million new cases diagnosed each year (9.8% of worldwide cancer diagnoses) and the cause of 694 000 deaths (8.5% of all worldwide cancer deaths) in 2012.¹ In 1996, as a commitment to reduce morbidity and mortality from this disease, the National Cancer Institute (NCI) of the U.S. National Institutes of Health invited investigators to apply

for funding to establish a ‘Cooperative Family Registry for Colorectal Cancer Studies’ (RFA: CA-96-011). The main NIH stated aims were: to collect pedigree information, epidemiological data and related biological specimens from participants with and without colorectal cancer and with and without a family history of the disease, as a resource for interdisciplinary studies on the aetiology of colorectal cancer; and to identify a population at high risk of colorectal cancer that could benefit from preventive strategies.

This cohort profile provides an update of the Colon Cancer Family Registry, described in detail in Newcomb *et al.*²

The basic premise of this initiative is that family-based designs across the spectrum of risk, in which cases, controls and their relatives are all recruited into a single research infrastructure, would enable the study of genetic aetiology, gene penetrance, gene-gene interaction and interaction with lifestyle factors. Thus, in 1997, the Colon Cancer Family Registry was established with funding support from the NCI. For Phase I (1998–2002), 5 years of funding was awarded to six Colon Cancer Family Registry sites:

- Cancer Care Ontario (Toronto, ON, Canada);
- Fred Hutchinson Cancer Research Center (Seattle, WA, USA);
- Mayo Clinic (Rochester, MN, USA);
- University of Hawaii (Honolulu, Hawaii, USA);
- University of Southern California Consortium (comprising the Universities of Southern California, Minnesota, North Carolina, Colorado and Arizona, Dartmouth University and the Cleveland Clinic Foundation, USA);
- University of Queensland (Brisbane, QLD, Australia).

The Colon Cancer Family Registry received funding renewals for Phase II (2003–07) and Phase III (2008–12) with the addition of:

- University of Melbourne (Melbourne, VIC, Australia) substituting for the University of Queensland;
- Memorial University (Newfoundland, Canada) as a collaborative site within the Cancer Care Ontario site.

In 2004–11, the ethnic/racial minority component of the Colon Cancer Family Registry was expanded through the recruitment of additional African American and Japanese American families with a separate NCI grant that included the University of Hawaii, the University of Southern California, the University of North Carolina, the Fred Hutchinson Cancer Research Center and the Cancer Prevention Institute of California.

Phase IV (2013–18) of the Colon Cancer Family Registry was funded by the NCI as a Cancer Epidemiology Cohort, and consequently renamed the Colon Cancer Family Registry Cohort (CCFRC). This phase saw the addition of:

- Stanford University (CA, USA) as the administering site for the Colon Cancer Family Registry, and
- Mayo Clinic (Scottsdale, AZ, USA) as the administering site for the Mayo Clinic CCFRC site.

Who is in the cohort?

Recruitment sampling schemes and inclusion and exclusion criteria varied by CCFRC site and funding phase. Details

of the recruitment methods at each CCFRC site have been published previously.² Briefly, recruitment protocols fall broadly into two main categories: population-based and clinic-based. Population-based probands were either people with a diagnosis of recently diagnosed colorectal cancer (case-probands) identified from cancer registries, or people without a prior diagnosis of colorectal cancer (control-probands) randomly sampled from the general population living in the relevant recruitment area using Medicare and Driver's License files, telephone subscribers lists or electoral rolls, who were frequency-matched for age to the case-probands. Clinic-based probands were people with or without colorectal cancer who were attendees at a family cancer clinic or genetics clinic. Cases with known familial adenomatous polyposis were excluded. Once recruited, probands were asked for permission to contact their relatives for recruitment. The CCFRC recruited 42 489 participants—from 15 049 families—who completed a baseline questionnaire between 1998 and 2012 (Table 1). Recruitment numbers within clinic-based families was, on average, twice that for population-based families (5.3 vs. 2.6 relatives per family, respectively). The majority of participants self-reported as Caucasian/White followed by Asian ethnicities and African American/Black (Table 2).

How often have they been followed up?

We have used both active and passive follow-up methods to update the cohort, where active follow-up includes direct contact with participants and passive follow-up includes indirect methods; details as follows.

Active follow-up

Approximately every 4–5 years after completing their baseline questionnaire, all participants of population-based case-families (but not control-families) and clinic-based families were asked, either by telephone interview or by self-completed questionnaire (mailed or online), for updates on their personal and family history of cancer as well as history of surgery, cancer screening and some risk factors.

Of the 37 436 participants who completed baseline questionnaires and were approached for follow-up, 27 918 completed the first follow-up questionnaire [response proportion (or response 'rate' of those alive) 83%], 3549 died before being approached for the first follow-up and 5969 could not be contacted or refused follow-up. Of the 27 918 participants who had completed the first follow-up, 18 958 completed their second follow-up questionnaire (response rate 87%), 1934 had died, 2824 were either uncontactable or refused, and 4202 are still in process. Of the 18 958 participants who had completed the second follow-up,

Table 1. Number of families and participants of the Colon Cancer Family Registry Cohort by sex and colorectal cancer (CRC) status at baseline recruitment

| | Males N (%) | Females N (%) | Total N (%) |
|--|----------------|------------------|----------------|
| Population-based families ^a | | | 13,190 |
| Probands with CRC (case-probands) | 4321 (29.2) | 4419 (24.6) | 8740 (26.7) |
| Relatives with CRC | 332 (22) | 372 (21) | 704 (21) |
| Relatives without CRC | 7769 (52.5) | 10516 (58.5) | 18285 (55.8) |
| Probands without CRC (control-probands) | 2071 (14.0) | 2205 (12.3) | 4276 (13.0) |
| Relatives of control-probands ^b | 310 (21) | 467 (26) | 777 (24) |
| Total population-based individuals | 14803 | 17979 | 32782 |
| Clinic-based families ^c | | | 1859 |
| Probands and relatives with CRC | 1139 (26.1) | 1108 (20.7) | 2247 (23.1) |
| Probands and relatives without CRC | 3221 (73.9) | 4239 (79.3) | 7460 (76.9) |
| Total clinic-based individuals | 4360 | 5347 | 9707 |
| Total families | | | 15049 |
| Total participants | 19163 | 23326 | 42489 |

^aProbands recruited from a population-based source.

^bOnly the University of Melbourne recruited relatives of control-probands.

^cProbands recruited from a family cancer clinic source.

Table 2. Distribution of participants by race

| Race ^a | Proportion of participants (%) |
|------------------------|--------------------------------|
| Native American | 0.9 |
| Asian | 5.5 |
| Pacific Islander | 0.3 |
| African American/Black | 4.8 |
| Caucasian/White | 86.0 |
| More than one race | 1.0 |
| Not reported | 1.5 |

^aSelf-reported by questionnaire.

8371 had completed their third follow-up questionnaire (response rate 95%), 1536 had died, 368 were either uncontactable or refused and 8683 are still in process (Figure 1).

The total number of person-years of follow-up by participants who completed a follow-up questionnaire is 276 762 person-years. As this is a family study, the vital status and cancer diagnoses of participants were also ascertained, even if they did not participate in the follow-up themselves, based on interviews of any relatives who were also participants. Including the reports by relatives, the total number of person-years of follow-up of all participants who completed a baseline questionnaire was 338 970 person-years, an average of 9.1 years per participant. These comprise approximately: 49 000 person-years for those recruited within 2 years after colorectal cancer diagnosis (thus relevant for studies of colorectal cancer survival and risk of metachronous cancer); 39 000 person-years for relatives with CRC and probands recruited more than 2 years after colorectal cancer diagnosis (thus relevant for studies of survivors of

colorectal cancer); and 251 000 person-years for those with no previous diagnosis of colorectal cancer (thus relevant for studies of colorectal cancer risk and aetiology)—Table 3.

Passive follow-up

One or more of the following passive follow-up activities have been conducted at each site of the CCFRC: data linkage with local and national death files, population-based cancer registries and electoral rolls; annual newsletters; reviews by genetic counsellors; and other mailings to participants. Passive follow-up was regularly conducted on all participants—at intervals that varied by site, type of follow-up activity and cost—to obtain information on new cancers, vital status and cause of death, and to update contact information.

Incident cancers and deaths during follow-up

During active and passive follow-up, all new reports of colorectal polyps and all cancers were recorded. Attempts were made to verify cancers using medical records, cancer registry data and confirmatory reports from relatives. To date, 824 (2.2%) participants have been diagnosed with an incident colorectal cancer since baseline; of those, 170 were diagnosed before the age of 50 years (Table 3); and 3582 (9.5%) participants have been diagnosed with an incident non-colorectal cancer since baseline. The total 4164 incident non-colorectal cancers were as follows: 772 skin, 568 breast, 599 prostate, 97 gastric, 52 small bowel, 103 hepatobiliary, 102 pancreas, 147 renal, 40 ureteric,

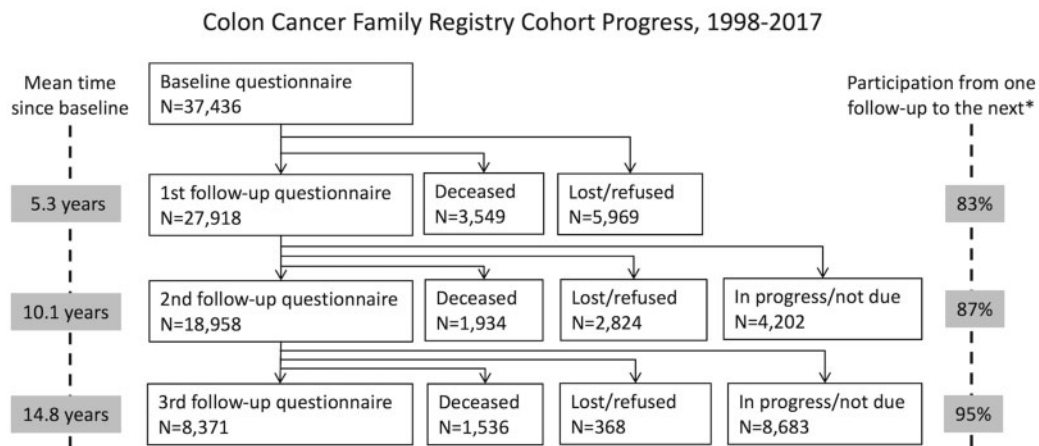


Figure 1. Progress of follow-up of participants of the Colon Cancer Family Registry Cohort (as of June 2017). Participation is defined as the percentage of those who were alive at contact attempt who completed the questionnaire.

Table 3. Numbers of incident colorectal cancer diagnosis and deaths occurring in study participants (except controls) of the Colon Cancer Family Registry since baseline recruitment by different cohort types, as of June 2017

| | Number of participants | Number of incident colorectal cancer at any age (%) | Number of incident colorectal cancer under age 50 years (%) | Number of deaths (%) | Average follow-up (years) ^d |
|---|------------------------|---|---|----------------------|--|
| Colorectal cancer within 2 years preceding recruitment ^a | 6765 | 144 (2.1) | 31 (0.5) | 2694 (39.8) | 7.5 |
| Colorectal cancer >2 years preceding recruitment ^b | 4623 | 202 (4.4) | 25 (0.5) | 1411 (30.5) | 8.8 |
| No history of colorectal cancer prior to recruitment ^c | 26048 | 478 (1.8) | 114 (0.4) | 2914 (11.2) | 10.0 |
| Total | 37436 | 824 (2.2) | 170 (0.5) | 7019 (18.7) | 9.4 |

^aCohort useful for studies of colorectal cancer survival and risk of metachronous cancer.

^bCohort useful for studies of survivors of colorectal cancer.

^cCohort useful for studies of colorectal cancer risk and aetiology.

^dBased on follow-up interview or report from participating relative.

150 urinary bladder, 76 brain, 355 lung, 27 bone, 219 blood, 163 endometrial, 73 ovarian and 35 cervical cancers, and 586 in other organs. A total of 7019 (19%) participants (including those with and without colorectal cancer at baseline) are known to have died since baseline (Figure 1).

What has been measured?

At the baseline recruitment, CCFRC participants were asked to complete a detailed family history of cancer, a risk factor questionnaire, permission to access medical records pertaining to any colorectal cancer diagnoses, permission to access colorectal cancer tumours and, depending on the degree of relationship to the proband, to provide a blood (or buccal wash) sample—Table 4.

Baseline risk factor questionnaires

All participants (proband and their participating relatives) were asked to complete the same detailed baseline risk

factor survey using standardized questionnaires via personal or telephone interviews or mailed questionnaires. Items included demography, lifestyle factors, screening, medication and family history.² Four CCFRC sites also asked participants to complete a self-administered food frequency dietary questionnaire. Three CCFRC sites (University of Hawaii, Cancer Care Ontario and University of Southern California consortium) used the questionnaire developed by the Multiethnic Cohort study in Hawaii and California.³ The University of Melbourne used the questionnaire developed by the Melbourne Collaborative Cohort Study.⁴

Follow-up risk factor questionnaires

At each follow-up, participants were asked for the following events that might have occurred since the previous contact: cancer diagnoses; bowel and gynaecological surgery; screening for colorectal cancer; polyps; and cancer diagnoses and deaths in relatives. Some CCFRC sites opted to

Table 4. Resources available from the Colon Cancer Family Registry Cohort, as of June 2017

| | | | Males | Females | Total | |
|---|---|--------------------------------|------------------------------|--------------------|--------------------|-------------|
| | | | N (%) ^f | N (%) ^f | N (%) ^f | |
| Population-based case families ^a | Probands | Baseline questionnaire | 4321 (22.5) | 4419 (18.9) | 8740 (20.6) | |
| | | Food frequency questionnaire | 2096 (22.7) | 2409 (20.5) | 4505 (21.5) | |
| | | Blood/buccal samples | 3759 (27.3) | 3886 (22.8) | 7645 (24.8) | |
| | | Polyp material | 14 (7.9) | 8 (4.0) | 22 (5.8) | |
| | | Cancer material | 3561 (73.6) | 3447 (70.2) | 7008 (71.9) | |
| | | Diagnosis and treatment | 1563 (84.2) | 1526 (85.7) | 3089 (85.0) | |
| | Relatives ^b | Baseline questionnaire | 7740 (40.4) | 10328 (44.3) | 18068 (42.5) | |
| | | Food frequency questionnaire | 3323 (36.0) | 4700 (40.0) | 8023 (38.3) | |
| | | Blood/buccal samples | 4751 (34.4) | 6731 (39.5) | 11482 (37.2) | |
| | | Polyp material | 7 (3.9) | 8 (4.0) | 15 (3.9) | |
| | | Cancer material | 265 (5.5) | 325 (6.6) | 590 (6.1) | |
| | | Diagnosis and treatment | 40 (2.2) | 44 (2.5) | 84 (2.3) | |
| | Spouse controls ^c | Baseline questionnaire | 361 (1.9) | 560 (2.4) | 921 (2.2) | |
| | | Food frequency questionnaire | 135 (1.5) | 197 (1.7) | 332 (1.6) | |
| | | Blood/buccal samples | 149 (1.1) | 225 (1.3) | 374 (1.2) | |
| | Population-based control families ^d | Probands | Baseline questionnaire | 2071 (10.8) | 2205 (9.5) | 4276 (10.1) |
| | | | Food frequency questionnaire | 1142 (12.4) | 1023 (8.7) | 2165 (10.3) |
| | | | Blood/buccal samples | 1399 (10.1) | 1497 (8.8) | 2896 (9.4) |
| Relatives ^b | | Baseline epi data | 310 (1.6) | 467 (2.0) | 777 (1.8) | |
| | | Food frequency questionnaire | 260 (2.8) | 383 (3.3) | 643 (3.1) | |
| | | Blood/buccal samples | 6 (0.0) | 5 (0.0) | 11 (0.0) | |
| Clinic-based Colon Cancer families ^e | | Probands with CRC ^a | Baseline questionnaire | 699 (3.6) | 644 (2.8) | 1343 (3.2) |
| | | | Food frequency questionnaire | 247 (2.7) | 270 (2.3) | 517 (2.5) |
| | | | Blood/buccal samples | 645 (4.7) | 625 (3.7) | 1270 (4.1) |
| | Polyp material | | 24 (13.5) | 29 (14.4) | 53 (13.9) | |
| | Cancer material | | 561 (11.6) | 526 (10.7) | 1087 (11.2) | |
| | Diagnosis and treatment | | 239 (12.9) | 204 (11.5) | 443 (12.2) | |
| | Probands no CRC ^c | Baseline questionnaire | 137 (0.7) | 304 (1.3) | 441 (1.0) | |
| | | Food frequency questionnaire | 62 (0.7) | 164 (1.4) | 226 (1.1) | |
| | | Blood/buccal samples | 101 (0.7) | 250 (1.5) | 351 (1.1) | |
| | | Polyp material | 13 (7.3) | 27 (13.4) | 40 (10.5) | |
| | | Cancer material | 20 (0.4) | 67 (1.4) | 87 (0.9) | |
| | | Diagnosis and treatment | 0 (0.0) | 0 (0.0) | 0 (0.0) | |
| | Relatives ^b | Baseline questionnaire | 3524 (18.4) | 4399 (18.9) | 7923 (18.6) | |
| | | Food frequency questionnaire | 1953 (21.2) | 2596 (22.1) | 4549 (21.7) | |
| | | Blood/buccal samples | 2983 (21.6) | 3814 (22.4) | 6797 (22.0) | |
| | | Polyp material | 120 (67.4) | 130 (64.4) | 250 (65.8) | |
| | | Cancer material | 429 (8.9) | 542 (11.0) | 971 (10.0) | |
| | | Diagnosis and treatment | 14 (0.8) | 6 (0.3) | 20 (0.6) | |
| Total | All population- and clinic-based probands and relatives | Baseline questionnaire | 19163 | 23326 | 42489 | |
| | | Food frequency questionnaire | 9218 | 11742 | 20960 | |
| | | Blood/buccal samples | 13793 | 17033 | 30826 | |
| | | Polyp material | 178 | 202 | 380 | |
| | | Cancer material | 4836 | 4907 | 9743 | |
| | | Diagnosis and treatment | 1856 | 1780 | 3636 | |

^aProband had a history of colorectal cancer (CRC) at baseline interview.

^bAffected or unaffected with colorectal cancer at baseline interview.

^cSpouse of proband, had no history of colorectal cancer at baseline interview.

^dProband had no history of colorectal cancer at baseline interview.

^eProband was recruited from a family cancer clinic.

^f% of total items obtained. For example 20.6% of all baseline questionnaires completed by probands of population-based case families.

include additional questions pertaining to colorectal cancer risk factors. All baseline and follow-up questionnaires used by each CCFRC site can be accessed at: [http://www.coloncfr.org/questionnaires].

Family history

One or more participants from each family was asked to provide their family history of cancer by answering a standard set of questions for each of their relatives (irrespective of cancer history) including: sex and date of birth; cancer sites (except non-melanoma skin cancer), and ages or dates at diagnoses (for those with a cancer history); vital status and, if deceased, date of death. All CCFRC sites recorded detailed family history information for each first- and second-degree relative, and some sites expanded to third-degree relatives, depending on site-specific protocols (detail in Newcomb *et al.*²). Attempts were made to verify the anatomical site, extent of disease, age at diagnosis and pathology of tumours. Sources of verification used included pathology reports, medical and surgical records, cancer registry information and death certificates.

Blood/mouthwash samples

Participants were asked to provide a blood or mouthwash sample. Of those who agreed, 93% provided a blood sample (Table 4).⁵ DNA was extracted from blood and mouthwash samples under CCFRC quality-control protocols, to maximize target DNA concentration and fragment size. To provide an unlimited supply of DNA and RNA for probands and selected relatives, lymphoblastoid cell lines of case-probands were immortalized using Epstein-Barr virus.⁶

Tumours and pathology

Paraffin-embedded colorectal cancer tumours, as well as diagnostic pathology reports, were obtained from treating facilities with the consent of the participant or the next-of-kin if the participant was deceased. In addition, some sites also obtained polyps and non-colorectal tumours, especially cancers commonly identified as part of Lynch syndrome. Multiple sections were cut from each tumour and normal-tissue block, stained with haematoxylin and eosin (H&E) and reviewed by pathologists. For each colorectal cancer, a pathology review was completed (either by examination of the H&E slides or extraction of relevant data from available pathology reports) to obtain the following standardized set of tumour features: grade, histological type, stage (depth of infiltration in large bowel wall and spread to regional lymph nodes), lymphovascular invasion

and perineural invasion. Sections were stored for future research at each CCFRC site. Two sites (Ontario and University of Southern California consortium) have made tumour microarrays (TMAs) from colorectal cancers (*n* 1278).

Virtual tissue repository

CCFRC has created a digitized library of pathology slides (electronic representations of traditional glass slides). A total of 4510 H&E stained slides of histological sections of colorectal tumours from the probands were scanned using either the NanoZoomer Digital Pathology scanner (Hamamatsu Corp.) or the Aperio ScanScope digital slide scanner. Each image is stored as a series of 752 x 480 pixel jpeg image tiles that are reconstructed with relevant software. Typical size of these images is between 200mb and 1.5gb per slide. All images were archived on five image servers: one for short-term storage and four for long-term storage.

Clinical records

Clinical treatment and outcome records were requested from 3830 case-probands and for 111 relatives with an incident colorectal cancer diagnosed since baseline, and have been abstracted into standardized items for analysis.

Molecular characterization of tumours

Probands' colorectal cancers were characterized for DNA mismatch repair (MMR) deficiency by polymerase chain reaction (PCR)-based microsatellite instability (MSI) tests and/or by immunohistochemistry (IHC) for the four DNA MMR proteins.⁷ Colorectal cancer tumour DNA was tested for the *BRAF* V600E somatic point mutation and somatic mutations in codons 12 and 13 of *KRAS*.^{8,9} Tumours were also tested for methylation of the *MLH1* gene promoter (an epigenetic phenotype used to indicate that MMR deficiency is more likely to have been caused by a somatic epigenetic event in *MLH1* than by a germline mutation in *MLH1*).¹⁰ Characterization of the CpG Island Methylator Phenotype (CIMP) was also performed by assessing quantitative methylation across five gene promoters (*CACNA1G*, *IGF2*, *NEUROG1*, *RUNX3* and *SOCS1*)¹¹—Table 5.

Molecular characterization of germline DNA

Screening for germline mutations in *MLH1*, *MSH2*, *MSH6*, *PMS2* and *EPCAM* was performed for all population-based probands who had a colorectal tumour displaying an MSI-High phenotype or a loss of expression

Table 5. Molecular characterizations of participants (probands and relatives) at the Colon Cancer Family Registry Cohort, as of June 2017

| Molecular test | Number of participants tested | Number of colorectal cancer tumours tested | Test results |
|------------------------------|----------------------------------|--|--|
| Tumour | | | |
| DNA MSI | 5147 | 5305 | 1065 high 665 low 3575 stable |
| IHC for MMR proteins | 8036 | 8338 | 1671 loss 6667 present |
| CIMP ^a | 3855 | 3888 | 479 positive 3409 negative |
| <i>MLH1</i> methylation | 3041 | 3412 | 465 methylated 2947 normal |
| <i>BRAF</i> V600E mutation | 7080 | 7322 | 679 positive 6643 negative |
| <i>KRAS</i> mutation | 4014 | 4154 | 1299 positive 2855 negative |
| Blood | | | |
| <i>MMR</i> gene ^b | 2895 probands 4106 relatives | | 710 mutations 1408 mutations |
| <i>MUTYH</i> ^b | 10649 probands 3571 relatives | | 47 biallelic 195 monoallelic 12 biallelic 197 monoallelic |

CIMP, CpG island methylator phenotype.

^aTumours were classified as CIMP-positive if more than three of five genes gave percentage of methylated reference value ≥ 10 .

^bSequencing (probands) or predictive testing (relatives).

of one or more of the MMR proteins expression by IHC, and for the youngest-onset colorectal cancer case participant from each clinic-based family, regardless of MSI or MMR-protein expression status. All case-probands were genotyped for 12 previously identified *MUTYH* variants: c.536A>G p.(Tyr179Cys), c.1187G>A p.(Gly396Asp), c.312C>A p.(Tyr104Ter), c.821G>A p.(Arg274Gln), c.1438G>T p.(Glu480Ter), c.1171C>T p.(Gln391Ter), c.1147delC p.(Ala385ProfsTer23), c.933 + 3A>C p.(Gly264TrpfsX7), c.1437_1439delGGA p.(Glu480del), c.721C>T, p.(Arg241Trp), c.1227_1228dup p.(Glu410 GlyfsX43) and c.1187-2A>G p.(Leu397CysfsX89) (detail in Cleary *et al.*¹²). Where available, blood samples from the relatives of probands with a pathogenic mutation were tested for the specific mutation (MMR gene or *MUTYH*) identified in the proband (predictive testing). Of the CCFRC participants: 2118 were identified as carrying a mutation in one of the MMR genes (761 in *MLH1*, 976 in *MSH2*, 243 in *MSH6*, 109 in *PMS2* and 29 in *EPCAM*); and 451 were identified as carrying either a monoallelic ($n = 392$) or biallelic mutation ($n = 59$) in *MUTYH*. Since baseline, these mutation carriers have contributed a total of 18 514 MMR-mutation person-years and 3732

MUTYH-mutation person-years. In addition, targeted sequencing was conducted of 36 known or putative colorectal cancer susceptibility genes (including the MMR genes) for 1231 cases including cases with familial colorectal cancer type X;¹³ early-onset (age < 50 years at colorectal cancer diagnosis), or suspected Lynch syndrome.

CCFRC has genome-wide single nucleotide polymorphism (SNP) genotyping data for 10 716 participants (6732 cases and 2435 controls) by various platforms (including OncoArray), all imputed to the 1000 Genomes Project.¹⁴ We excluded known carriers of germline mutations MMR genes and *MUTYH*.

What has it found? Key findings and publications

The CCFRC resource has been used for more than 400 original peer-reviewed publications—see [http://coloncfr.org/publications]. Here, we highlight a few findings that illustrate the power of this cohort to understand genetic and environmental risk factors for colorectal cancer.

Lynch syndrome (cancer caused by inherited mutations in DNA MMR genes *MLH1*, *MSH2*, *MSH6* and *PMS2* or

EPCAM) has been a major research focus of the CCFRC. These studies have included analyses of the identification of Lynch syndrome^{15–18} and associated rare variant classification,^{19–21} age-specific risk of cancer (penetrance),^{22–25} effect of extent of colon resection on the risk of metachronous cancer,²⁶ prevalence of Lynch syndrome in colorectal cancer cases²⁷ and in the general population,²⁸ pathology of Lynch syndrome tumours,²⁹ acceptability and impact of genetic testing,^{30–33} and modifiers of penetrance for Lynch syndrome-associated cancers.^{34–42} A prime example is the prospective cohort analysis of MMR gene mutation carriers to estimate cancer risks, i.e. penetrance of Lynch syndrome.²² A total of 446 MMR gene mutation carriers with an average age of 40 years and who had no cancer diagnosis were followed up for 5–10 years—the largest prospective study ever of Lynch syndrome in which participants completed a risk factor questionnaire. The 5- and 10-year risks of several cancer types were calculated, with the highest risk observed for colorectal cancer (8% 10-year risk 20 times population risk), endometrial cancer (10% 10-year risk 30 times population risk) and ovarian cancer (3% 10-year risk 19 times population risk). A novel finding of this study was that these mutation carriers also appear to have an increased risk of breast cancer. A total of 1029 of their relatives who were not mutation carriers were also assessed for cancer risk. They were found to be at the same risk as the general population and therefore can be screened as someone at average risk, despite often having a strong family history of cancer.

The CCFRC also makes major contributions to the search for new genetic risk factors, primarily due to its large sample size and because of its family-based design. This allows for the collection of DNA samples from case-probands as well as their relatives, and the verification of reports of cancer by relatives rather than relying on self-report only. This research includes the investigation of SNP associations,^{43–50} genome panel testing⁵⁵ and statistical modelling of risk.^{28,51,52} Many of these studies stemmed from the research within the CCFRC on syndromic classification of familial colorectal cancer, primarily the work leading to the description of ‘familial colorectal cancer type X’, the phenotype of MMR-proficient colorectal cancer cases who fulfilled the Amsterdam Criteria for hereditary non-polyposis colon cancer,¹³ which has become an accepted category of familial colorectal cancer.⁵³

What are the main strengths and weaknesses?

The unique strength of the CCFRC is its prospective, observational design, with familial enrichment and molecular

characterization. Participants have deliberately been over-sampled for familial risk, and therefore the CCFRC differs from the usual cancer research cohort in novel ways that allow inferences not otherwise possible.⁵⁴ This facilitates a deeper and broader research agenda that covers aetiological factors (both genetic and environmental), molecular characterization, behavioural issues and clinical research relevant to people at increased familial risk.

Participants can be categorized on their underlying familial risk profile based on their genotype, family history and risk factor data, which allows the effects of environmental risk factors to be investigated for varying levels of putative genetic risk: i.e. for studies of gene-environment interactions. The availability of genotype data allows prospective studies of the risk-modifying effects of genetic and non-genetic factors, and the effectiveness of targeted screening/surveillance by genetic sub-groups. The CCFRC can be, and has been, used for a range of gene discovery research including classic linkage studies, genome-wide association studies and whole-exome and whole-genome studies.⁵⁵ Furthermore, because a large proportion of CCFRC participants were diagnosed with colorectal cancer just before recruitment and have risk factor data as well as blood samples, powerful studies of prognostic factors can be undertaken. The CCFRC also facilitates novel behavioural, psychosocial and health utilisation research for clinical translation.

From a practical perspective, conducting family studies can be challenging because of the often complex nature of familial relationships, as well as the additional layers of protocol that need to be incorporated to protect privacy within families (for example, procedures to ensure that sensitive information is not inadvertently passed to other family members). We have demonstrated that these issues, however, can be managed through carefully designed study protocols and training. We strongly believe that the benefits of a family cohort far outweigh its limitations, and that more epidemiologists should consider this design when conducting aetiological research focused on environmental risk factors across the risk spectrum.

Can I use the data? Where can I find out more?

From its inception, the CCFRC has functioned under the principle that it is a resource for research on the aetiology, risk and prognosis of colorectal cancer for all researchers, including those not affiliated with CCFRC. To this end, CCFRC welcomes collaborative applications to access and analyse both electronic data (questionnaire, genotypes, medical records, family history etc.) and biospecimens (DNA, blood, serum, and tumour specimens). Of the total

294 approved applications to use CCFRC resources, 157 (53%) have come from external investigators.

The CCFRC provides internal and external researchers fair and equitable access to this unique resource. Collaborating investigators have established numerous funded projects. For information on how to collaborate and access data for the CCFRC, including cohort data described here, please see [<http://coloncfr.org/>].

Profile in a nutshell

- The Colon Cancer Family Registry Cohort (CCFRC) was established for the purpose of research on the genetic and environmental aetiology of colorectal cancer.
- The 42 489 study participants from 15 049 families were recruited between 1998 and 2012 in the USA, Canada, Australia and New Zealand. They include: recently diagnosed colorectal cancer cases from population-based cancer registries; controls from population-based sources; patients from family cancer clinics with a strong family history of colorectal cancer; and their relatives.
- Every 4–5 years after baseline, all population-based case-families and clinic-based families were followed up and re-surveyed. The total follow-up of 37 436 participants covers approximately 339 000 person-years (mean follow-up 9.1 years). Since baseline, 824 (2.2%) participants were diagnosed with a colorectal cancer and 3582 (9.5%) were diagnosed with a non-colorectal cancer.
- At baseline, all participants completed the same risk factor questionnaire for a detailed personal and family history of cancer, and a wide range of risk factors. At each follow-up, participants were asked for updates on their personal and family history of cancer, screening, surgery, death and some risk factors. Blood samples and tumour specimens have been collected and used for extensive genetic and molecular characterization including Lynch syndrome.
- CCFRC resources are available for collaborative research [<http://www.coloncfr.org/>].

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