Telomere length and genetics are independent colorectal tumour risk factors in an evaluation of biomarkers in normal bowel

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Background: Colorectal cancer (CRC) screening might be improved by using a measure of prior risk to modulate screening intensity or the faecal immunochemical test threshold. Intermediate molecular biomarkers could aid risk prediction by capturing both known and unknown risk factors.

Methods: We sampled normal bowel mucosa from the proximal colon, distal colon and rectum of 317 individuals undergoing colonoscopy. We defined cases as having a personal history of colorectal polyp(s)/cancer, and controls as having no history of colorectal neoplasia. Molecular analyses were performed for: telomere length (TL); global methylation; and the expression of genes in molecular pathways associated with colorectal tumourigenesis. We also calculated a polygenic risk score (PRS) based on CRC susceptibility polymorphisms.

Results: Bowel TL was significantly longer in cases than controls, but was not associated with blood TL. PRS was significantly and independently higher in cases. Hypermethylation showed a suggestive association with case:control status. No gene or pathway was differentially expressed between cases and controls. Gene expression often varied considerably between bowel locations.

Conclusions: PRS and bowel TL (but not blood TL) may be clinically-useful predictors of CRC risk. Sample collection to assess these biomarkers is feasible in clinical practice, especially where population screening uses flexible sigmoidoscopy or colonoscopy.

Colorectal carcinoma (CRC) is one of the most common forms of malignancy (Ferlay et al, 2013). It is considered a complex disease, with both inherited and environmental factors involved in predisposition (Houlston and Peto, 2004). Genome-wide association studies (GWAS) have so far identified over 30 common polymorphisms (SNPs) associated with CRC and adenoma risk in the general population (Broderick et al, 2007; Tomlinson et al, 2007, 2008, 2011; Houlston et al, 2008, 2010; Tenesa et al, 2008; Dunlop et al, 2012; Whiffin et al, 2014).

Screening for benign polyps and their removal by colonoscopy can reduce CRC risk (Zauber et al, 2012). Many European countries now offer population screening programmes, based primarily on faecal occult blood tests (FOBT) (Robertson and Imperiale, 2015). Despite the effectiveness of these programmes,
the burden of screening is likely to increase considerably and CRC will continue to be a major killer. It would therefore be useful to improve screening by targeting it to those at highest risk. Individualised risk prediction is generally infeasible outside the Mendelian CRC syndromes (Dunlop et al., 2013). It remains possible, however, that risk prediction is still clinically useful when applied to population-level screening because of the following reasons: (i) the unknown and/or unmeasurable risks are distributed among large numbers of individuals and (ii) population stratification by prior risk is an improvement on the current screening that is targeted solely by age.

Another potentially useful measure of disease risk is the intermediate molecular phenotype, since this can hypothetically capture both known and unknown factors that act through a common mechanism. For example, there are at least 7 CRC SNPs with putative effects on bone morphogenetic protein (BMP) signalling, and a measure of BMP pathway activity might, therefore, capture the effects on BMP signalling of the following: (i) these SNPs, (ii) other unknown SNPs, (iii) rare variants in the intermediate phenotypes with potential effects on CRC risk include telomere length (TL) (von Zglinicki, 2002) and DNA methylation (Ehrlich, 2002). These phenotypes have usually been assessed in a more convenient sample such as peripheral blood, rather than in the bowel itself. While many biomarkers have been postulated to predict CRC risk (Ishihara et al., 2008), most of these are not established risk factors and there exist relatively few previous studies of intermediate molecular markers (Alexander et al., 2017).

The polygenic risk score (PRS) – a quantitative score of susceptibility caused by common genetic variants (Dudbridge, 2013) – is another measure of CRC risk that may be clinically useful (Machiela et al., 2011). The predictive value of the CRC PRS has been assessed in previous studies (Dunlop et al., 2013; Frampton et al., 2016). However, these have only considered carcinoma risk and none included polyps or performed a prospective assessment.

In this study, we have searched for associations between a set of candidate intermediate molecular markers and case/control status in individuals undergoing colonoscopic screening for colorectal tumours. We have also assessed a PRS based on the major CRC predisposition SNPs. Our results have potential importance and clinical applicability for risk prediction models that can be used to modulate population screening for CRC.

### MATERIALS AND METHODS

#### Patients and samples

Colorectal biopsies were obtained from 317 individuals of white UK origin undergoing colonoscopy in Oxford. Indications for colonoscopy included: family history of CRC (N = 15); symptoms compatible with CRC (N = 164); follow-up for previous polyps/CRC (N = 97); FOBT-positive UK National Bowel Cancer Screening patients (N = 91), and rectal prolapse or diverticulitis (N = 16). ~3 mm3 biopsies of normal bowel epithelium were taken from up to three sites (rectum, distal colon and proximal colon), together with 10 ml peripheral blood. Personal and family histories of bowel disease and the basic demographic data were recorded. Participants were placed into one of the following three categories: (1) polyp cases—individuals with personal history of adenomatous and/or serrated polyps; (2) CRC cases—individuals with personal history of carcinoma; and (3) controls—individuals with no polyps or CRC in the present or past. Individuals not fulfilling these criteria or with other pathologies, such as inflammatory bowel disease, were excluded (Table 1). Ethical approval was obtained from the Oxfordshire Research Ethics Committee A (REC 10/H0604/72). All patients provided informed consent, according to the tenets of the Declaration of Helsinki.

#### DNA/RNA extraction

Each biopsy was submerged in RNALater (Life Technologies). DNA and RNA were extracted with the AllPrep DNA–RNA Mini kit (Qiagen, Manchester, UK). Where multiple biopsies had been taken from the same patient, each individual biopsy was analysed, except that for assays in which a single DNA-based measure per patient was required, where an aliquot of DNA from each available location was pooled in equal proportions. For RNA samples, an aliquot was DNase-treated and reverse transcribed into cDNA. We estimated the typical ratio of epithelium–to-mesenchyme in biopsies to be 1:1.

#### Telomere length assays

We assessed telomere length (TL) in blood and bowel DNA in 262 patients using a modification of Cawthon’s method (Cawthon, 2009; Jones et al., 2012). DNA samples were run in duplicate on a CFX96 system (Bio-Rad, Watford, UK). A curve was established relating the quantity of a standard DNA to ΔCt for the telomere repeat and a single-copy gene (gamma haemoglobin). This provided the expected Ct value (EIC) for the telomere repeat. Corrected Ct (CorrΔCt) was calculated by subtracting ECI from observed (OIC), providing a relative measure of telomere length. CorrΔCt was always 0 and the higher the CorrΔCt, the longer the telomere.

#### Genomic DNA methylation assay

Global DNA methylation analyses were performed using reverse-HPLC (Quinlivan and Gregory, 2008) by measuring total 5-methylcytosine (5mC) content in 136 patients. Briefly, 2500 ng genomic DNA were RNase-treated in NEB2.1 buffer and incubated for 1 h at 37°C, purified with ethanol and lyophilised. The sample was re-suspended in 50 μl RNase-free water and hydrolysed overnight in a solution containing 45 mM NaCl, 9 mM MgCl2, 9 mM Tris pH 7.9, ≥250 U ml−1 Benzonase (Sigma), 50 μM U ml−1 Phosphodiesterase I, ≥20 U ml−1 Alkaline phosphatase, 46.8 ng ml−1 EHNase hydrochloride, 8.64 μM deoxoamine. Protein components were removed with Amicon centrifugal filter units (3 kDa cutoff, Millipore, Watford, UK) and resolved with an Agilent UHPLC 1290 (Stockport, UK) instrument fitted with Eclipse Plus C18 RRHD 1.8 μm, 2.1 × 150 mm column. Buffer A was 100 mM ammonium acetate, pH 6.5; buffer B was 40% acetonitrile, and the flow rate 0.4 ml min−1. The gradient was between 1.8 and 100

| Table 1. Numbers of patients included in the study by case |
| --- |
| **N** | **Male/ female** | **Mean age (SD), range** | **Clinical phenotype (N)** |
| **Phase 1** | **Controls** | 47 | 24/23 | 54 (17–3), 22–80 | A (46), CRC (8) |
| **Cases** | 54 | 33/21 | 67 (8.2), 42–84 | A (46), CRC (8) |
| **Phase 2** | **Controls** | 59 | 27/32 | 52 (16.4), 21–83 | A (79), S (37), CRC (41) |
| **Cases** | 157 | 79/78 | 65 (9.4), 30–89 | A (79), S (37), CRC (41) |

**Abbreviations:** A = adenoma; CRC = colorectal carcinoma; S = serrated polyp; SD = standard deviation. Numbers of patients included in the study by case:control status, sex, age, and tumour histology.
of 40% acetonitrile. Methylation was also assayed in blood and bowel DNA separately.

**Molecular pathway activity assessment.** Using real-time, quantitative PCR (qPCR), eight molecular pathways were investigated, which were either related to CRC development or reflecting cellular processes linked to tumourigenesis (Supplementary Table 1). To assess these, we identified 33 genes as pathway end points. Gene expression assays were conducted on an ABI1990HT real-time instrument (Applied Biosystems, Foster City, CA, USA), using TaqMan (ThermoScientific, Waltham, MA, USA) molecular probes. As per the recommendations in the current qPCR guidelines, we optimised the normalisation procedure by choosing the best-suited set of housekeeping genes. Given that a literature search did not yield did not yield any unambiguous results, we tested 11 endogenous genes (Supplementary Table 1). The optimal set for normalisation was identified using qBase+ . Normalisation was then carried out, using a ΔΔCT method for relative quantification.

**SNP genotyping and polygenic risk scores.** KASPar genotyping technology (LCGenomics) was used to generate a 25 SNP genotype profile of each of 310 patients (Supplementary Table 2). PRSs were calculated for the 293 patients with ≥22 successfully genotyped SNPs according to:

\[
\log(S_{\beta Ni}) = \frac{\beta_1 + \ldots + \beta_{25} \times N}{1}
\]

where \(\beta\) is the Ln(odds ratio) for each risk allele and \(N\) is the number of risk alleles. The missing data points were substituted by the population mean.

**Statistical analyses.** These were performed with Stata-v.11 (StataCorp LP, TX, USA) and R (http://www.r-project.org/). For gene expression analysis, individual genes were assessed for associations with case:control status at nominal \(P=0.05\). Pathway analysis revealed \(P=0.05\) thus was independent of family history of colorectal neoplasia \((P=0.201)\). Within-patient analysis showed no significant variation in bowel TL by location (Supplementary Figure 1), and this remained true when comparing within-patient TL in biopsies from locations close to the polyp/CRC, compared with those further away \((P=0.436\) for proximal colon, \(P=0.591\) for distal colon, \(P=0.055\) for rectum).

Of note, bowel TL was associated with neither casecontrol status \((P=0.989\), OR = 1.00, 95% CI = 0.88–1.14) nor bowel TL \((P=0.195\), OR = 1.06 95% CI = 0.97–1.15; Supplementary Figure 2), although both bowel and blood TL were negatively correlated with age \((P=0.000\), \(r^2 = -0.03\); and \(P=0.000\), \(r^2 = -0.04\), respectively).

**Methylation.** Global methylation (5mC) levels, as assessed from pooled DNA derived from all three colorectal regions, increased with age as expected \((P=4 \times 10^{-6}\), OR = 1.09; 95% CI = 1.05–1.14). We observed a suggestive, but non-significant, increase in global methylation levels in cases compared with controls \((P_{meta} = 0.070\); OR = 1.58; 95% CI = 0.96–2.64; Supplementary Figure 3). Blood 5mC was assayed in a subset of 31 samples, but there was no good evidence of a correlation with bowel 5mC \((P=0.139\); OR = 0.94; 95% CI = 0.86–1.02; \(r^2 = -0.27\); Supplementary Figure 4). For TL, we did not find any significant differences between patients with present or past polyps \((P=0.44\), with or without family history \((P=0.117\), polyps or CRC \((P=0.259\) or present vs previous CRC \((P=0.117\), and methylation levels were independent of family history \((P=0.304\).

Analyses by location showed significantly lower methylation in the proximal colon compared to the distal colon or the rectum (Supplementary Figure 5). Analyses using only the location-specific measurements of methylation did not yield any significant associations (data not shown).

**Gene expression analysis.** We assayed 273 samples from 100 patients for 33 selected genes from 8 pathways (Supplementary Table 1). There was an effect of location on gene expression, with 18 out of 33 genes differentially expressed \((q\leq0.05\) between the proximal and distal colon, 21 out of 33 between the proximal colon and rectum, and 16 out of 33 between the distal colon and rectum (Supplementary Figure 6); some genes showed monotonically decreased \((\text{ID1, ID3, OLFM4, PLK1})\) or increased \((\text{AXIN2, CCND1, CDK6, E2F1, EPHB2, ID2, KCNHA1, LRIG1, MCM2, PCBP4, SCNN1D, SCNN1G, TFDP1})\) levels along the proximal colon-distal colon-rectum axis (Supplementary Figure 7). Correlations between the expression of gene pairs are depicted in Supplementary Figure 8. Casecontrol status was not significantly associated with the expression of any individual gene at a nominal \(P\leq0.05\) (Supplementary Table 3). Pathway analysis revealed nominally higher expression of colonocyte differentiation markers in cases \((P=0.010\), \(q = 0.080\); Supplementary Table 4).

We then examined bowel location-specific gene expression in cases who had one or more tumours in that location vs controls with no colorectal tumour anywhere. Cases with distal colon tumour(s) had significantly higher SCNN1A expression in that location than controls \((P=0.004\); \(q = 0.032\); OR = 4.09; 95% CI = 1.56–10.76; Supplementary Figure 9). This gene showed no evidence of association with status in the other locations and no additional significant associations were found for other genes. With regards to pathway expression by location, colonocyte differentiation and Wnt levels were higher in cases in the proximal colon \((q = 0.040\) for both). Stem cell markers and ion channels showed varying expression levels depending on the bowel segment (Supplementary Table 4).

**Assessment of SNP genotypes, bowel TL and status.** In logistic regression analysis conditioning on age, sex and family history,
PRS was significantly higher in cases than controls ($P = 0.031$; OR = 51.1; 95% CI = 1.45–1803; per quartile OR = 1.22; Supplementary Figure 10). PRS was not significantly correlated with bowel TL, methylation or, interestingly, family history ($P = 0.134$; $P = 0.114$ and $P = 0.640$, respectively, full data not shown).

Two of the three telomerase SNPs were significantly associated with bowel TL (for rs10936599, $P = 0.161$; for rs2736100, $P = 0.009$; for rs2735940, $P = 0.001$; non-parametric trend test; Supplementary Figure 11) and also independently with case:control status. No other SNP was associated with case:control status (details not shown).

In reverse stepwise logistic regression analysis, PRS ($P = 0.046$; OR = 63.5; 95% CI = 1.08–3740) and bowel TL ($P = 0.026$; OR = 1.74; 95% CI = 1.07–2.82) were significantly and independently of greater magnitude in cases than controls.

**DISCUSSION**

In this study, we examined whether intermediate molecular phenotypes derived from the bowel mucosa were associated with case:control status in a series of individuals undergoing colonoscopic screening for colorectal neoplasia. Such molecular markers potentially predict CRC risk and could be used to modulate population screening. Our principal finding was that cases with colorectal polyps or CRC had longer bowel telomeres, independent of other risk factors. There is a controversy about the effects of telomere length on the risk of many cancers, including CRC (Quin et al, 2014; Segui et al, 2014), but almost all studies have measured telomere length in peripheral blood. Our finding of no correlation between telomere length in the colorectum and blood may help to explain the inconsistent results of previous studies. Despite the assay’s speed and simplicity, telomere measurements from the mixed population of cells in peripheral blood may be inappropriate for predicting cancer risk.

SNPs rs2736100 and rs2735940 near TERT and rs10936599 near TERC have been associated with CRC risk (Houlston et al, 2010; Kinnersley et al, 2012; Yang et al 2015), but associations with normal bowel TL length have not been explored. rs2735940 and rs2736100 were significantly associated with bowel TL and each was also independently associated with case:control status, consistent with previous reports (Hofer et al, 2012).

It has been posited that hypermethylation, resulting from ageing or environmental causes, is a cause of increased cancer risk (Ehrlich, 2002). However, most studies have investigated methylation levels in circulating blood cells. In our study, we found borderline significant differences in global methylation levels between cases and controls. A true association might have been missed due to the fact that our global methylation measure was too general and did not sufficiently capture finer differential methylation patterns in CpG islands and other regions. In addition, the proximal colon had significantly less methylation than the distal colon and rectum, potentially influencing tumourigenic pathways in the different regions of the large bowel (Bufill, 1990).

Gene expression analysis throughout the colorectum detected no significant associations with case:control status for individual genes, and only a nominal association for molecular pathways in the case of colonocyte differentiation markers. Given the non-uniform gene expression patterns observed throughout the bowel for many of the genes analysed (LaPointe et al, 2008), we performed association analyses in each location independently and found that expression of SCNN1A in the distal colon was different between cases and controls. SCNN1A is a potential marker of differentiation and encodes a membrane calcium channel that plays an essential role in maintaining the osmotic gradient in the intestinal epithelium. We were a little surprised, given the importance of variation in the BMP pathway for CRC risk, that we found no evidence of differential activity in this pathway between cases and controls. Many explanations are possible, including limited assay sensitivity, sub-optimal choice of target genes, and spatial and temporal heterogeneity in gene expression.

Our study’s conclusions should be interpreted in the light of limitations and potential confounding factors. While the association between status and bowel TL could have been caused by the presence of tumour(s) rather than *vice versa*, we found no evidence of this, as there were no significant differences between TL in cases with incident CRC or polyps and previous tumours. Moreover, it seems unlikely that polyps could have major effects on sampled normal mucosa that can be more than a metre away in the large bowel, and indeed we did not observe significant intra-patient differences when comparing TL in biopsies close to and distant from a tumour. Second, the fact that both gene expression and global methylation vary throughout the large bowel suggests that location-specific risk models may be more powerful. Our exploratory analyses (details not shown) have demonstrated that because patients frequently have polyp(s) in only one bowel location, location-specific risk models are required with larger sample sets in order to provide adequate statistical power. Third, we caution that, although we deliberately used simple biopsies without the separation of epithelium and mesenchyme, because these represent the sample type most likely to be available in a clinical setting, our measurements were derived from a mixed cellular population, potentially affecting all assays except the PRS. Fourth, the performance of the PRS is likely to be improved in the future by taking into account recently discovered predisposition loci (Zeng et al, 2016, amongst others) and polymorphisms specific to the risks of adenomas, serrated polyps and CRC.
In summary, we have shown the potential value of using intermediate molecular markers, and the independent value of the PRS, for predicting colorectal tumour risk. Our data also show the virtue of using biomarkers derived from the colorectum rather than blood. Bowel cancer screening programmes are increasingly using colonoscopy or flexible sigmoidoscopy as a primary modality and rectal biopsies could be obtained safely from participants at limited cost and subjected to molecular analyses that could then be incorporated into algorithms for subsequent screening. While additional studies are required, we suggest that stratification by prior risk could offer a way to improve the cost-effectiveness of population-level large bowel cancer screening populations, whether limited cost and subjected to molecular analyses that could then be incorporated into algorithms for subsequent screening by FIT.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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