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**Urinary volatile organic compounds and faecal microbiome profiles in colorectal cancer**

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Running title: Urinary VOCs and faecal microbiomes in CRC

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Author contributions:

MM: Recruited subjects, analysed samples, data analysis, manuscript writing
AM: data analysis
HH: Analysed samples
RS: data analysis
CC: data analysis
RA: data analysis, manuscript writing
CN: project co-ordinator, manuscript writing

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Key words: Colorectal cancer, genetics, environment, VOCs, microbiome

Abbreviations:

ANOSIM: Analysis of Similarity
AUC: Area Under Curve
BCSP: Bowel cancer Screening Programme
CRC: colorectal cancer
FAIMS: Field Asymmetric Ion Mobility Spectrometry
FIT: Faecal Immunochemical Testing
FOBT: faecal occult blood test
LC-FAIMS-MS: Liquid Chromatography- Field Asymmetric Ion Mobility Spectrometry- Mass Spectrometer
MDT: Multi-Disciplinary Team
OTU: operational taxonomic units

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Abstract:

Background

Volatile organic compounds are potential biomarkers for diagnosing colorectal cancer (CRC). We characterised urinary VOCs from CRC patients, their spouses/co-habitors (spouses) and first-degree relatives (relatives) to determine any differences. Correlation with stool-derived microbiomes was also undertaken.

Methods

Urine from 56 CRC patients, 45 spouses and 37 relatives were assayed using Liquid Chromatography: Field Asymmetric Ion Mobility Spectrometry (FAIMS): Mass Spectrometer technology. Analysis was performed using 5-fold cross-validation and a Random Forrest classifier. Faecal microbiome 16s RNA was sequenced using Illumina Miseq protocols and analysed using UPARSE and QIIME pipelines.

VOC and microbiome profiles were also compared before, and after, cancer treatment.

Results

Urinary VOC profiles of CRC patients were indistinguishable from either spouses or relatives. When spouses and relatives were grouped together to form a larger non-cancer control group (n=82), their VOC profiles became distinguishable from CRC patients (n=56) with 69% sensitivity and specificity, area under curve 0.72 (p<0.001).

Microbiome analysis identified >1300 operational taxonomic units (OTUs) across all groups. Analysis of Similarity (ANOSIM) R value was 0.067 (p=<0.001), with significantly different bacterial abundances in 82 OTUs (6.2%) by Kruskal-Wallis testing.

CRC patients’ VOC or stool microbiome profiles were unchanged after treatment.
Conclusion

Although CRC patients’ urinary VOC profiles cannot be differentiated from spouses or relatives they can be differentiated from a larger non-cancer control group. Comparison of the groups’ microbiomes confirmed differences in bacterial species abundance. The current FAIMS-based assay can detect a unique, but modest, signal in CRC patients’ urinary VOCs, which remains unaltered after treatment.

What does this add?

This study adds to the growing evidence of the utility of urinary VOC testing as a non-invasive biomarker for the distinction of patients with colorectal cancer from healthy controls. Here we use crude genetic and environmental controls, whereas previous studies had utilised unconnected controls.

Introduction

There is a worldwide increase in both the incidence and mortality from Colorectal Cancer (CRC) (1). In the United Kingdom (UK), CRC is the fourth commonest cancer and second commonest cause of cancer-related death, with 15,900 deaths in 2014 (2,3). CRC is associated with genetic and environmental factors, including age, sex, family and past medical history, including colorectal polyps, smoking and diet (3, 4, 5, 6, 7, 8, 9, and 10).

Current non-invasive CRC screening tests have had their utility questioned. The UK Bowel cancer Screening Programme (BCSP) utilizes guaiac-based faecal occult blood testing (gFOBT), with a positive predictive value (PPV) of ~10% (11). gFOBT will be superseded by Faecal Immunochemical Testing (FIT). FIT is simpler to perform, both for patients and laboratory staff, with a higher PPV of 41% (12). FIT shows a sensitivity of 66–88%, depending on haemoglobin cut-off values (50–200 ng/ml), and a specificity of 87–96% (13 – 15). Screening based on faecal samples had a reported ~50% uptake rate after the first 1 million cases of the BCSP, although there was variation across the country (11). Uptake in France was reported as 34.3% after the first 2 years (16). Whilst Finland has a 71% uptake (17). This variable uptake reduces the overall efficacy of this strategy. Given this mixed performance, a more effective point-of-care CRC screening test involving non-stool samples is required.

An approach for non-invasive detection of cancers is volatile organic compounds (VOCs) analysis. The detection of VOC patterns by both invasive and non-invasive methods, and their utility as disease-specific gas phase biomarkers, has developed in recent years (18).
CRC has been distinguished from other cancers and healthy controls using exhaled VOC analysis in several studies (19-21). This has been replicated in urinary VOC analysis using Mass Spectrometry (MS) technology (22, 23), and subsequently Electronic Nose and Field Asymmetric Ion Mobility Spectrometry (FAIMS) technology (24, 25). Faecal VOCs have also been studied and shown promise, however, as the uptake of stool-based screening programmes is ~50%, this may well prove of limited clinical benefit long term (26). No studies have identified any consistently unique VOCs, suggesting the overall pattern or "smell print" allows differentiation, rather than individual chemicals. The presence of intestinally derived VOCs in urine is believed to occur via VOC migration into the blood stream and subsequent renal filtration allowing their presence, and detection, in urine (27).

There have been limited studies that have demonstrated a role for the detection of pre-cancerous polyps (15, 28). Indeed one study showed urinary VOCs could distinguish CRC from controls and CRC from advanced adenomas, but poor distinction of adenoma from control (28). The current stool-based assays have lower sensitivity and specificities for pre-malignant lesions (29).

VOCs themselves are a diverse group of carbon-based chemicals which are products of bodily metabolic processes. Pathological states are believed to alter VOC profiles, resulting in disease specific patterns, thus allowing their use as biomarkers (18). Within the context of CRC, this could include the interaction of malignant colonocytes altered metabolism and ingested dietary factor fermentation by the subject’s intestinal microbiome (30). The true origin of VOCs remains uncertain.

Recent years has seen expanding interest into the role of gut micro-organisms in health and disease. Studies of the microbiome in CRC patients have demonstrated greatly reduced microbial diversity, termed dysbiosis (31, 32). Specific bacteria identified as over-represented in CRC include; Streptococcus bovis, Helicobacter pylori, Bacteroides fragilis, Clostridium septicum, and Escherichia coli strains (31, 33, and 34). Conversely, butyrate-producing bacteria, including Roseburia and Fecalibacterium are reduced (33, 35). It remains unclear whether these changes are a cause, or consequence, of CRC.

This study aimed to characterise urinary VOC and stool microbiome profiles of CRC patients, to ascertain whether these profiles could be distinguished from those of controls whom shared environment exposures (spouses/co-habitors) and genetic factors (first degree relatives). Previous studies have used healthy controls with no links to cancer patients (19-25). The objective was to determine whether VOCs and stool microbiomes remained distinguishable from control groups when the controls shared potential risk factors for CRC, and to provide evidence for the potential utility of urinary VOCs as a non-invasive CRC screening test.
Methods

Ethics:
Scientific and ethical approval was granted by the University Hospitals Coventry and Warwickshire (UHCW) Research and Development Office, and Solihull Ethics committee, ref: 13/WM/0136. Written, informed consent was obtained from all study participants.

Patient recruitment:
Patients were recruited between September 2015 and December 2016 from the Lower GI Multi-Disciplinary Team (MDT). Inclusion criteria were adult patients with confirmed CRC, with no family history to suggest hereditary CRC conditions, and relatives and spouses/co-habitors who had consented. Exclusion criteria were patients/relatives/spouses that had concurrent malignancy, non-malignant gastrointestinal conditions and urological/renal conditions requiring secondary hospital care. Patients with polyps showing advanced neoplasia were excluded.

Relatives and spouses were recruited after consent had been confirmed by the CRC patients. Relatives living with the CRC patients were excluded, as were spouses/partners not co-habiting with the CRC patients. Post-operative samples were collected 3 and 6 months after initial surgery. CRC patients who were 24-36 months post initial treatment and in endoscopic and radiological disease-free surveillance provided urine for analysis. All recruited subjects provided data regarding smoking, alcohol consumption, current medications and a dietary questionnaire. No patient had received bowel preparation or antibiotics within the 4 weeks prior to sample collection.

Sample Collection and storage:
Urine and stool samples were collected and stored at -80°C within 72 hours of voiding/elimination. Urinary VOCs have been shown to be stable at room temperature during this period (unpublished data). They were defrosted in a laboratory fridge at 3°C overnight, prior to analysis.

Urinary VOC and statistical analysis
Urine samples were analysed on the LC-FAIMS-MS equipment (Owlstone, UK) which is a bespoke device combining mass spectrometry and FAIMS technology, in series with an LC column to allow greater separation of chemicals and more sensitive assays of VOC concentrations. The LC column was pre-conditioned using a pooled sample of all 3 groups and samples applied to the column from a chilled autosampler (4°C). The column was
washed with an increasing chromatographic gradient of acetonitrile (5-90%). The eluted fractions from the column were aerosolized by an electrospray and then passed into the FAIMS and time of flight mass spectrometer (TOF-MS). FAIMS and MS settings can be found in supplementary file 1.

Due to the hybrid instrument set-up acquired chromatograms were pseudo MS/MS files with each collision detected corresponding to one of 10 FAIMS compensation fields (CF). This meant it was not possible to extract features directly from raw LC-FAIMS-MS data. To resolve this, acquired data was split into individual chromatograms corresponding to FAIMS CF settings using a custom Python script and the data subjected to feature extraction using XCMS package in R. The output data contained a list of features, or peaks, for the sample. The peaks were force aligned, allowing direct comparison across samples and improved the signal-to-noise ratio. All features were then normalized to have zero mean and unit variance.

For subgroup analysis, the relevant sample subset was extracted, and the different groups defined. 5-fold cross-validation was then used to assess classification accuracy across these groups, using a Random Forest multi-class classifier. This analysis generated outputs of one-vs-all Receiver Operator Curves (ROC) i.e. comparing a single group vs all other groups, for example CRC vs relatives and spouses, relatives vs CRC and spouses etc. Other results generated included the Area-Under-Curve (AUC) statistic, sensitivity/specificity values, and a p-value, comparing the results to that expected by random chance (AUC=0.5), using a Wilcoxon rank-sum test. All analyses were carried out using R programming language.

*Faecal 16S microbiome sequencing and statistical analysis*

200mg of each stool underwent DNA extraction using a QIAamp Fast DNA stool mini-kit (QIAGEN). The 16S V3-V4 region was then amplified using V3-V4 specific primers and polymerase chain reaction (PCR). The DNA was purified using AMPure magnetic beads (Beckman Coulter) and further PCR performed using unique forward and reverse primers to allow identification of individual subject’s sequences. The amplified products were purified using AMPure magnetic beads, pooled and diluted to 4nM concentration before sequencing on an Illumina Miseq platform (Illumina) using a Miseq V3 2 x 300bp paired end protocol.

The raw sequence data was merged and quality controlled using UPARSE software. Contiguous sequences were assigned to operational taxonomic units (OTUs) then clustered and filtered to a default 97% identity level before being linked to corresponding samples and assigned taxonomic classification. QIIME (Qualitative Insights Into Microbial Ecology) software was used to perform analysis using Analysis of Similarity (ANOSIM) and Kruskal-Wallis testing.
Results

Urinary VOC analysis

Seventy-two CRC patients 56 spouses/co-habitors (spouses) and 61 first degree relatives (relatives) and were approached. Urine samples were returned by 56 CRC patients, 45 spouses and 37 relatives.

There was no significant difference in the mean ages between the CRC patients and the spouses, although there was a significant difference between the CRC and spouse groups and the relative group, as a result of children of the CRC patients being included. There were more males in the cancer group and as expected this was reversed in the spouses group. The proportion of males and females in the relatives group was equal. The demographics are shown in table 1.

Urinary VOC analysis was performed using a 5-fold cross-validation, the Random Forest multi-class classifier and Wilcoxon rank-sum test. An analysis in which only CRC patients with a paired with spouse and relative (n=35 in each group) showed no significant differences in their VOC profiles. A larger analysis was performed using all recruited subjects (CRC n=56, Spouses n=45, relatives n=37). This analysis also could not distinguish CRC patients from either their spouses or first degree relatives. There was also no difference when the spouses were compared with the relatives. However when the spouses and relatives were grouped together (n=82) and compared to CRC patients (n=56) the technology was able to detect a difference with a sensitivity of 69% (95% confidence intervals: 54% – 81%), a specificity of 69% (57% – 79%) and Area under Curve (AUC) of 0.71 (0.62 – 0.8). The Bonferroni corrected p value was <0.001. The positive predictive value was 60%. The ROC curve is shown in figure 1.

Analysis of CRC subjects by stage of CRC, site of CRC and pathway for detection (2 week wait, BCSP etc) did not reveal any statistically significant differences.

Post-treatment urine sample analysis:

Of the original 56 CRC patients who provided urine samples before treatment, 23 and 9 returned samples at 3 and 6 months respectively.

Thirty patients who were 24-36 months post CRC resection, and in disease free surveillance, also provided urine samples. Demographics can be found in table 2.

An analysis was again performed using a 5-fold cross-validation and Random Forrest classifier. Comparison of the CRC patients VOC profiles with samples taken at 3 months, 6 months, combined 3 and 6 months, and 24-36 months after treatment revealed no difference.
Faecal 16S microbiome analysis

Seventy-two CRC patients 61 first degree relatives and 56 spouses were approached. Stool samples were returned by 44 CRC patients, 34 first degree relatives and 39 spouses.

The mean ages of the CRC patients, spouses and relatives and spouses were 65.3 years (SD 11.2), 59.9 years (SD 12.4) and 51.2 years (SD 12.6) respectively. Group demographics can be found in table 3.

The 16S V3-V4 region was sequenced using an Illumina Miseq platform. Once the raw sequence data had been merged, quality controlled and filtered to exclude low quality reads, with counts <5000, 5 samples were excluded. This left sample sizes of 41 for CRC, 33 relatives and 38 spouses.

There were 1346 OTUs identified across all samples. Relative abundance plots of the CRC samples (n=41) and the combined non-cancer control group (relatives and spouses: n=71) are in figure 2. The non-parametric analysis of similarity (ANOSIM) test was run using QIIME software. This returned an R value of 0.067 (p=<0.001), indicating CRC samples have a very similar microbiome profile to non-cancer controls. Principal component analysis of the overall diversity of CRC patients and non-cancer controls showed no statistically significant differences in microbiome composition (See figure 3).

The non-parametric Kruskal-Wallis test was performed to identify any significant differences between the CRC and non-cancer controls. It showed 82/1346 OTUs (6.2%) were significantly different between CRC and non-cancer control group, i.e. >93% were not significantly different. Of the 82 OTUs, 46 showed increased abundances and 36 decreased abundances in CRC samples relative to the control group, with 64/82 identified as clostridiales sp.

Comparison of the relative and spouse control sub-groups using Kruskal-Wallis testing identified 567 OTUs, with 25 (4.4%), showing statistically significant differences between the two groups.

Analysis of CRC subjects by stage of CRC, site of CRC and pathway for detection (2 week wait, BCSP etc) did not reveal any statistically significant differences.

Post-treatment stool sample analysis:

Of the 44 CRC patients, 15 and 14 returned additional samples at 3 and 6 months respectively. The demographics can be found in table 4.

Microbiome profiles of the CRC patients compared to the profiles at 3 months, 6 months and pooled 3 and 6 months after treatment revealed no significant differences.
The rarefaction curve plots of observed OTUs, plotted against average number of sample reads shows an increase in the number of observed OTUs in post-operative samples, though not to statistically significant levels. See Figure 4.

Discussion

An individual’s urinary VOC profile is a sum of the different volatile compounds when exposed to assay methodology. The origin and role of VOCs in human metabolism remains unclear, but likely represents the complex interaction of malignant colonocytes, dietary factors and the individual’s microbiome (30). While the specificity of each compound is unknown the VOC profile can give a unique identifier to diseased individuals.

The strength of the VOC signal detectable in urine is variable given the heterogeneity of the earlier pilot studies, which showed diversity of assay methodology, and the comparisons were between cancer and non-cancer controls. The number of subjects in groups studied in these pilot studies was about 20-30 subjects. In this study we aimed to have at least 20 in each group. A non-cancer control group included the spouses/co-habitors of CRC patients who shared their environment and diet and first degree relatives of CRC patients. The CRC patients shared a gene pool with their first degree relatives but did not have a hereditary CRC syndrome.

We found no difference in VOC profiles between the CRC patients and either their spouses or first degree relatives, either in a smaller analysis, with only paired CRC/spouse/relative samples or when all collected samples were included. Our interpretation of this result is that the cancer patients and the two non-cancer control groups shared enough aspects of the make-up of their VOC signals to mask any cancer-specific signal. When the non-cancer controls were grouped and then compared with the CRC patients a significant difference in the VOC signal was identified (AUC, sensitivity and specificity of 0.71, 0.69 and 0.69 respectively). Increasing the number of non-cancer controls raised the power of the study and allowed a modest cancer-specific signal to be detected. There is no validated database to allow identification of individual VOCs and so this was not possible.

These results though modest, conform to previous studies, which have shown distinct patterns of urinary VOCs in CRC patients compared to disease-free controls, (22-25). The PPV was 60%, which compares favourably to the ~8-10% demonstrated for FOBT and FIT (36-39), however, the specificity compares less favourably to FOBT ~98% (39). The relatively poor specificity represents a major obstacle for urinary VOCs entering clinical use for CRC screening. Another study by our group, comparing urinary VOC profiling to FIT testing has shown that FIT proved superior to VOCs in isolation but that a combination of FIT and VOCs raised sensitivity and specificity to 0.97 and 0.72 respectively (40).
There was no difference in urinary VOC profiles of the first-degree relatives and spouse/co-habitor groups. This suggests a heterogeneous VOC composition among non-cancer individuals, and neither environmental nor genetic factors appear to significantly affect it.

The spouses/ co-habitors were all consuming the same food as the CRC patients and none of the main groups were consuming special diets (vegan, vegetarian, gluten free). This minimises the potential for VOC profile being confounded by dietary factors. None of the relatives group was consuming special diets. It is possible that each sub-group of the non-cancer control groups were underpowered to detect VOC profile suggesting that any group specific VOC signal is relatively weak, or that any variations have occurred by chance.

A potential weakness in the study design is the absence of endoscopic screening of the first-degree relatives, who are at increased risk of CRC, compared to the co-habitors group. Currently the risk of CRC in asymptomatic first-degree relatives of patients with sporadic CRC is not high enough to attract endoscopic screening in the UK.

This is the first study to include healthy control subjects that are connected to CRC subjects, by a shared environment, or shared gene pool. All previous studies had used unrelated healthy controls (19-25). Despite the relatedness of the non-cancer controls in this study, they could still be distinguished from the CRC group, with similar sensitivities, specificities and AUCs to those reported previously.

There was no significant difference in VOC signals between the CRC subjects when sub-analyses of stage of CRC, site of CRC and pathway for detection (2 week wait, BCSP etc) were performed. This is most likely due to under powering as a result of diminishing sample sizes.

There was no significant difference in VOC’s and microbiome between pre-treatment and post-treatment CRC samples. This suggests that either the urinary VOC profile detected in CRC patients does not originate from the cancerous tissue or confirms the weakness of the signal. Either scenario would make VOC analysis unsuitable for CRC surveillance using the current FAIMs technology. Altomare et al found that exhaled breath analysis, 2 years after curative surgery, using GC-MS, distinguished pre- from post-operative samples with a sensitivity of 100%, specificity of 97%, accuracy of 98.8% and AUC 1 (41). The sample size of 32 was similar to our cohort, so may indicate GC-MS is better suited than LC-FAIMS-MS. Ma et al had previously demonstrated a significant reduction in 2 urinary VOCs post-operatively, using GC-MS, although they did not suggest it's use for CRC surveillance (22).

The microbial 16s profiling of CRC samples and healthy non-cancer controls (relatives and spouses) was performed using Illumina 16s sequencing and bioinformatics analysis. It revealed no significant difference for >93% of identified OTUs between CRC patients and the non-cancer control group. Of 1346 identified OTUs, only 82 (6.2%) were significantly different between CRC patients and non-cancer controls, with 64 identified as clostridiales.
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Conclusions

This study demonstrates that CRC patients have a unique urinary VOC profile different from non-cancer controls. This cancer-specific signal is modest, requiring the pooling on non-cancer controls, and becomes detectable at n/> 50, however, this adds to growing evidence that urinary VOCs may have a potential future role in distinguishing CRC patients from non-cancer controls, whether they are independent controls as in previous studies, or related controls as in this study. Current technology is improving and should in future be able to detect cancer-specific signals in urine with greater specificity and sensitivity. Comparison with the stool-based screening tests will continue but the poor uptake of stool testing will hand an advantage to any urine-based screening tests developed.

References

<table>
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<td>37</td>
<td>45</td>
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</tr>
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<td>50.0 (14.1)*</td>
<td>60.7 (12.1)</td>
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<td>17:20</td>
<td>15:30</td>
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<td>Number of cigarettes</td>
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<td>smoked per day (SD)</td>
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<tr>
<td>Alcohol units per week (SD)</td>
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<td>6.7 (10.1)</td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>A</td>
<td>8 (14.2%)</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>17 (30.4%)</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>C1</td>
<td>20 (35.7%)</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>C2</td>
<td>9 (16.1%)</td>
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<td>Right</td>
<td>24 (42.8%)</td>
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<td>Left</td>
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<td>Rectal</td>
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<tr>
<td>Referral Route (%)</td>
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<td>2 Week Wait pathway</td>
<td>31 (55.4%)</td>
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<tr>
<td>Other</td>
<td>7 (12.5%)</td>
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<td>-</td>
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</table>

Table 1. Demographic data of CRC patients, their first degree relatives and spouses who provided urine samples for VOC analysis.
Table 2. Demographic data of pre-operative, post-operative and 24-36 month disease free surveillance CRC patients who provided urine samples for VOC analysis.
<table>
<thead>
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<td>*&lt;0.01</td>
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<td>A</td>
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<td>15 (34.1%)</td>
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<td>C1</td>
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<td>C2</td>
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<td>Site (%)</td>
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<td>Rectal</td>
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<td>2 Week Wait pathway</td>
<td>21 (50%)</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>6 (9.5%)</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Demographic data of CRC patients, their first degree relatives and spouses who provided stool samples for 16S microbiome analysis.

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<table>
<thead>
<tr>
<th>Group</th>
<th>Pre-operative CRC</th>
<th>3 months post-operative</th>
<th>6 months post-operative</th>
<th>P values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Samples</td>
<td>44</td>
<td>15</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td>Mean age (SD)</td>
<td>65.3 (11.2)</td>
<td>63.2 (12.1)</td>
<td>66.9 (11.4)</td>
<td>0.69</td>
</tr>
<tr>
<td>Sex (M:F)</td>
<td>25:19</td>
<td>8:7</td>
<td>7:7</td>
<td>0.90</td>
</tr>
<tr>
<td>Number of cigarettes smoked per day (SD)</td>
<td>0.73 (2.6)</td>
<td>0.67 (2.5)</td>
<td>0.7 (2.6)</td>
<td>1</td>
</tr>
<tr>
<td>Alcohol units per week (SD)</td>
<td>7.8 (11.6)</td>
<td>6.1 (11.2)</td>
<td>7.2 (12.5)</td>
<td>0.91</td>
</tr>
<tr>
<td>Mean BMI (SD)</td>
<td>27.4 (4.3)</td>
<td>28.5 (7.7)</td>
<td>26.8 (4.6)</td>
<td>0.61</td>
</tr>
<tr>
<td>Dukes stage (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>9 (21.4%)</td>
<td>3 (20.0%)</td>
<td>3 (21.4%)</td>
<td>0.99</td>
</tr>
<tr>
<td>B</td>
<td>15 (35.7%)</td>
<td>5 (33.3%)</td>
<td>5 (35.7%)</td>
<td></td>
</tr>
<tr>
<td>C1</td>
<td>13 (30.1%)</td>
<td>5 (33.3%)</td>
<td>6 (42.9%)</td>
<td></td>
</tr>
<tr>
<td>C2</td>
<td>5 (11.9%)</td>
<td>2 (13.3%)</td>
<td>0 (0%)</td>
<td></td>
</tr>
<tr>
<td>Site (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right</td>
<td>18 (42.9%)</td>
<td>9 (60%)</td>
<td>9 (64.3%)</td>
<td>0.45</td>
</tr>
<tr>
<td>Left</td>
<td>13 (31.0%)</td>
<td>4 (26.7%)</td>
<td>4 (28.6%)</td>
<td></td>
</tr>
<tr>
<td>Rectal</td>
<td>11 (26.2%)</td>
<td>2 (13.3%)</td>
<td>1 (7.1%)</td>
<td></td>
</tr>
</tbody>
</table>

Table 4. Demographic data of pre-operative, post-operative and 24-36 month disease free surveillance CRC patients who provided stool samples for 16S microbiome analysis.
Figure 1. ROC plot of sensitivity and specificity for Random Forrest analysis of urinary VOC data.
Figure 2. Relative abundance plots of pre-treatment CRC patients (left) and all healthy controls: relatives and spouses (right).
Figure 3. Principle component analysis of the overall diversity of pre-treatment CRC (red) and healthy controls (blue).
Figure 4. Rarefaction curve of observed number of OTUs against sequences per sample for pre-treatment samples (blue) and 6 month post-treatment samples (red).