Revisiting Short-Chain Fatty Acids and the Microbiota in Colorectal Tumors

| Running title: SCFAs and colorectal tumors |
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Abstract

- 2 Background. Colorectal cancer (CRC) is a growing health concern with a large percentage of
- the risk of developing disease being due to environmental factors. Certain bacterial community
- 4 members have been associated to increase CRC while other taxa have been associated with normal
- 5 colons. Some of the taxa associated with normal colons can use fiber to produce short-chain
- 6 fatty acids (SCFAs) that can inhibit tumor growth in model systems. However, the data supporting
- the importance of SCFAs in human CRC is less certain. Here, we test the hypothesis that SCFA
- 8 concentration changes in individuals with colorectal tumors.
- Methods. We analyzed a cross-sectional (n=490) and pre- and post-treatment (n=67) group for
- their concentrations of acetate, butyrate, and propionate. Analysis also included tumor classification
- models using Random Forest, imputed gene relative abundance with PICRUSt, and metagenomic
- sequencing on a subset (n=85) of the total cross-sectional group.
- 13 **Results.** No difference in SCFA concentration was found between normal individuals and patients
- with adenoma or carcinoma (P-value > 0.15). There was no difference in classification models
- 15 with or without SCFAs in their ability to predict adenoma or carcinoma (P-value > 0.05). Using
- metagenomic seguencing, there was also no difference in genes involved with SCFA synthesis
- $_{17}$ (P-value > 0.70).
- ¹⁸ Conclusions. Although our data does not support the hypothesis that SCFAs decrease in
- individuals that have colorectal tumors, there may be specific scenarios where SCFAs may still be
- beneficial for treatment of CRC. Alternatively, there may be other metabolites that have not been
- thoroughly investigated that are more important to the development of human CRC.

2 Introduction

Colorectal cancer (CRC) is currently the third leading cancer-related cause of death within the US (1, 2). Although there is a genetic component to the disease, the envrionment has been attributed as an important and large risk factor for CRC (3). This has lead to the investigation of how the microbiota may exacerbate or cause tumorigensis (4–6) and whether the bacterial community is altered (7, 8). Multiple reports in case/control studies on the microbiota and CRC have identified taxa commonly associated with the normal colons to be decreased in patients with carcinoma tumors (9–11). Many of these taxa actively produce short-chain fatty acids (SCFAs) from fiber that are a part of our general diet (12). The most extensively studied of these SCFAs are acetate, butyrate, and propionate (13). Overall, CRC is a growing health problem and the microbiota through SCFAs have shown promise as a potential treatment option.

Specific SCFAs, like butyrate, have shown positive results for CRC treatment within model systems
(14). Butyrate has been shown to inhibit cancer cell growth in *in vitro* systems (15). Additionally,
supplementation with food sources that bacteria use to create these SCFAs may also be able to
confer beneficial effects. For example, fiber supplementation in mouse models of CRC caused an
overall reduction in tumor burden while also increasing SCFA concentrations (16). Although these
model systems provide important preliminary evidence towards the ability of SCFAs to reduce and
treat tumors, the studies reporting benefit in humans has been less convincing.

There is a lack of evidence on the benefit of increasing SCFA concentrations to protect against CRC in human populations. The initial case/control studies that investigated SCFA concentrations in CRC found that patients with carcinomas had lower concentrations of acetate, butyrate, and propionate versus either patients with adenomas or individuals with normal colons (17). Although this would argue that increasing SCFA concentrations could be protective against tumorigenesis, fiber supplementation in randomized controlled trials have consistently failed to protect against tumor recurrence (18). These findings would argue against the utility of treatments that aim to use SCFAs to reduce or protect against tumorigenesis. Given the lack of clear evidence in human studies of the benefit of SCFAs in CRC, there is a need for more investigation into this area.

Our current study will fill some of the current gaps in the literature that relate to the study of SCFAs and CRC in human populations. Specifically, it will attempt to replicate previous case/control findings on SCFA concentrations in individuals with and without tumors. Additionally, by increasing the adenoma tumor cohort it we will identify if the observation of a continuous gradual reduction in SCFA concentrations occurs as tumor severity increases. It also will build upon these observations and assess the utility of using SCFAs and Operational Taxonomic Units (OTUs) as a potential risk stratification tool of colorectal tumors (adenoma or carcinoma). This study will provide important information on the replicability of previous findings in humans by extensively studying how SCFAs could be associated with colorectal tumors.

To accomplish this task we will directly measure the concentration of acetate, butyrate, and propionate within fecal samples for two different groups. The first group had a sample obtained at a single cross sectional point in time while the second group had samples obtained before (pre-) and after (post-)treatment for colorectal tumors. Additionally, we also assessed whether adding SCFA concentrations to OTU data improves the classification of adenoma or carcinoma using the Random Forest algorithm (19), used PICRUSt (20) and metagenomic sequencing to assess the genes involved in SCFA synthesis, and analyzed how well 16S rRNA gene sequencing can predict SCFA concentrations. This investigation provides additional information as to whether the taxa that are decreased in carcionma use SCFAs to help protect against colon tumorigenesis.

7 Results

Decreased SCFA concentrations are not associated with tumors. We used high-performance liquid chromatography (HPLC) to measure acetate, butyrate, and propionate concentrations of frozen fecal samples from 490 individuals at a cross-sectional point in time. There was no difference between individuals with normal colons (n=172) and patients with either an adenoma (n=198) or carcinoma (n=120) for any of the SCFAs measured after multiple comparison correction (P-value > 0.15) [Figure 1A - 1C]. We next measured the concentration of SCFAs in 67 patients with an adenoma (n=41) or carcinoma (n=26) in which we had pre- and post-treatment fecal samples. Although there was a general trend for increasing acetate, butyrate, and propionate concentrations after treatment for tumors, there was no significant difference pre- and post-treatment for patients with adenoma (P-value > 0.20) or carcinoma (P-value > 0.80) [Figure 1D - 1F]. Even though there was no change in SCFA concentrations between individuals with normal colons and those with tumors, this information could still be important to help classify disease.

Random Forest models with SCFA concentrations do not classify tumors better. Using the
Random Forest algorithm we built models with OTUs only or OTUs and SCFA concentrations to
classify normal versus adenoma and normal versus carcinoma fecal samples. For adenoma and
carcinoma models, there was no difference between the median AUC of models with or without
SCFA concentrations (P-value > 0.05) [Figure 2]. Although including SCFA concentrations did
not add extra information for classification of colorectal tumors using Random Forest models, it
is possible that the genes for enzymes involved in SCFA synthesis may vary based on the type
of colorectal tumor; where a smaller number of microbes are responsible for the observed SCFA
concentrations in patients with carcinoma.

Changes in genes for enzymes involved in SCFA synthesis are not associated with tumors.

Using a list of specific genes that are important for the synthesis of SCFAs [Table S1], we looked for differences in gene abundance based on individuals having normal colons, adenomas, or carcinomas. First, using imputed gene relative abundance based on 16S rRNA gene sequencing we found no difference in any of the genes inolved with acetate, butyrate, or propionate synthesis (P-value > 0.90) [Table S2]. This similarity between groups is highlighted by visualizing genes

important in butyrate synthesis [Figure 3A]. Using a Wilcoxson rank-sum test, there also was no difference in imputed gene relative abundance between pre- and post-treatment samples for any genes involved with SCFA synthesis (P-value > 0.70) [Table S3]. Next, we took a subset of these 490 fecal samples (n=85) and used metagenomic sequencing to confirm these results. Like the imputed gene results, metagenomic analysis found that there was no difference in any of the genes involved in SCFA synthesis between individuals with normal colons (n=29) and patients 100 with adenoma (n=28) or carcinoma (n=28) (P-value > 0.70) [Table S4]. This lack of difference 101 is highlighted when we visualize the results for butyrate kinase [Figure 3B]. These observations 102 provide evidence that the gene content does not change due to colorectal tumors. Although these 103 results suggest that SCFA concentrations do not change due to colorectal tumor, there may be 104 some errors present within our results that may account for the lack of associations. 105

Expected taxa are associated with higher SCFA concentrations regardless of tumor status.

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Using OTU data we built Random Forest models to classify higher than median and lower than median SCFA concentrations. Overall, OTU data had a reasonable ability to classify high and low SCFA concentrations [Figure S1A]. However, these models tended to be overfit, suggesting that rarer taxa may be important for this classification [Figure S1A]. The most important OTUs to these models (assessed with mean decrease in accuracy (MDA)) were to taxa that are normally associated with SCFA production [Figure S1B]. These results highlight that there are no immediately apparent errors in our data because SCFA concentrations are associated with taxa known to produce acetate, butyrate, and propionate. Additionally, OTUs associated with these taxa are the most important to models that can classify high and low SCFA concentrations. Overall, our results are robust and do not support the hypothesis that differences in SCFA concentrations are associated with colorectal cancer.

Discussion

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The observations from this study do not support the hypothesis that SCFA concentrations are lower 119 in individuals with tumors. Whether we directly measured the SCFA concentration or investigated 120 genes associated with their production, no difference could be identified [Figure 1 & 3]. Further, 121 no benefit could be found in using the concentrations to help classify individuals with and without 122 tumors [Figure 2]. These observations are in stark contrast to some of the previous literature. 123

Much of the previous research on SCFA benefit to human CRC has been illustrated in model systems (14). Many SCFAs are produced through the breakdown of fiber (12) and a recent study in 125 mice found that fiber supplementation increased SCFA concentrations and decreased tumor burden 126 (16). Additionally, SCFAs such as butyrate can inhibit tumor growth in in vitro experiments (15). 127 Yet, observations in humans has been mixed. Previous case/control studies found associations 128 with lower SCFA concentrations in individuals with carcinoma tumors (17). However, individual 129 randomized controlled trials and a recent meta-analysis on fiber supplementation to prevent 130 tumor recurrence has found no benefit (18, 21). Our results align with what has been reported 131 in randomized-controlled trials, that SCFAs do not provide general protection against colorectal 132 tumors. It is possible though that there are specific instances where SCFAs may be beneficial. 133

One limitation of ours and others research into the effect of SCFAs in CRC has been that all tumors 134 are treated as the same type. However, there are known differences in the types of mutations that 135 occur (22) and treating all tumors as equal may actually hide any benefit that could be found in 136 specific subsets of individuals. Similar to the idea of specific immunotherapy for specific tumors (23), SCFAs may have beneficial effects for specific types of colorectal tumors. Future research will 138 need to test if this is a valid hypothesis. Although there are limitations with the current research 139 on SCFAs and colorectal tumors, evidence is building that these specific metabolites may not be protective. Yet, taxa that are associated with SCFA production are consistently higher in normal colons than those with carcinomas (7, 8, 24). 142

The potential protection against colorectal cancers may not be from SCFAs even though taxa associated with their production are higher in individuals with normal colons versus patients with carcinomas (7, 8, 24). Protection could be via a different pathway and by extension other metabolites that have not been extensively studied. Alternatively, protection may not occur via a metabolite but instead through niche exclusion of mouth-associated microbes (e.g. *Fusobacterium*, *Porphyromonas, Parvimonas, Peptostreptococcus* (9, 10)). The idea of niche exclusion is similar to how the community protects against *Clostridium difficile* infection (25) with chronic inflammation replacing the role of antibiotics. Although we did not find lower concentrations of SCFAs associated with colorectal tumors, we think that there are many exciting new avenues to explore because of these results

Conclusions

Our observations found no difference in SCFA concentration, their utility as a classification tool, and for genes for enzymes involved in SCFA synthesis between individuals with normal colons and patients with either adenoma or carcinoma tumors. Although these results are different than other reports in the literature, they do align with the randomized controlled trials that have tested fiber use in preventing colorectal tumor recurrence. Overall, these results suggest that the SCFAs produced by resident microbes do not protect against tumor. By focusing on other types of metabolites, the identification of more promising therapeutic options for use in treating colorectal cancer may be found.

Materials and Methods

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Study design and sampling. The overall protocol has been described in detail previously (26, 163 27). In brief, this study used fecal samples obtained at either a single cross-sectional time point 164 (n=490) or from before (pre-) and after (post-) treatment of a patient's tumor (n=67). For patients undergoing treatment for their tumor the length of time between their initial and follow up sample 166 ranged from 188 - 546 days. Our use of treatment has been previously defined as encompassing 167 removal of a tumor with or without chemotherapy and radiation (26). Diagnosis of tumor was made by colonoscopic examination and histopathological review of biopsies obtained (26, 27). The 169 University of Michigan Institutional Review Board approved the study and informed consent was 170 obtained from all participants in accordance to the guidelines set out by the Helsinki Decleration. 171

Measuring specific SCFAs. Our protocol for the measurement of acetate, butyrate, and propionate followed a previously published protocol that used a High-Performance Liquid Chromatography 173 (HPLC) machine (28). The following changes to this protocol included the use of frozen fecal 174 samples suspended in 1ml of PBS instead of fecal suspensions in DNA Genotek OmniGut tubes, and the use of the acutal weight of fecal samples instead of the average weight for SCFA 176 concentration normalizations. These methodological changes did not affect the overall median concentrations of these SCFAs between the two studies (see Table 1 (28) and Figure 1 here).

16s rRNA gene sequencing. The workflow and processing have been previously described (26, 179 29, 30). The major differences from these previous reports include: the use of version 1.39.5 of the 180 mothur software package and clustering Operational Taxonomic Units (OTUs) at 97% similarity 181 using the OptClust algorithm (31). 182

Generating imputed metagenomes. The use of PICRUSt version 1.1.2 with the recommended 183 standard operating protocol (20) was used. Briefly, the mothur shared file and metadata was converted into a biom formated table using the biom convert function, the subsequent biom 185 file was processed with the 'normalize by copy number.py' function, and subsequent imputed 186 metagenomes created using the 'predict metagenomes.py' function.

Obtaining Operational Protein Families from metagenomes. A subset of the cross-sectional

group (n=490) containing a total of 85 individuals (normal n=29, adenoma n=28, and carcinoma n=28) was shotgun sequenced on an Illumina HiSeq using 125 bp paired end reads and a previously described method (32). Briefly, the sequences were quality filtered and sequences aligning to the human genome were removed prior to contig assembly with MEGAHIT (33). Open Reading Frames (ORFs) were identified using Prodigal (34), counts generated using Diamond (35), subsequent clustering into Operational Protein Families (OPFs) used mmseq2 (36), and OPF alignment used the KEGG database (37).

Pulling genes involved with SCFA synthesis. Specific genes located near the end of the pathways involved in the synthesis of acetate, butyrate, and propionate were analyzed for any differences between individuals with normal colons and those with tumors. These genes were based on pathways from KEGG as well as previous research (37, 38) and a list can be found in the supplemental material [Table S1].

Random Forest models. The model was first trained on 80% of the data and then tested on the
held out 20% (80/20 split) using the Random Forest algorithm for classification models (19). This
was repeated on 100 different 80/20 splits of the data to generate a reasonable range for the AUC of
the model. The reported AUCs, unless otherwise specified, are for the test sets. The classification
models were built to group normal versus adenoma, normal versus carcinoma, and high versus low
SCFA concentrations.

Statistical analysis workflow. All analysis was performed using the statistical language R (39). 207 Generally, a Kruskal-Walis rank sum test with a Dunn's post-hoc test was used to assess differences 208 between the groups used. Where appropriate Benjamini-Hochberg was used to correct for multiple comparisons (40). First, we assessed differences in SCFA concentrations measured by HPLC 210 between individuals with normal colons and patients with tumors (adenoma or carcinoma). We 211 then analyzed whether SCFA concentrations changed in patients with an adenoma or carcinoma pre- versus post-treatment. Next, we assessed whether OTUs alone or OTUs and SCFAs were 213 better able to classify individuals with and without tumor using Random Forest models. Next, the 214 imputed gene counts of important mediators of SCFA synthesis was tested. Additionally, the counts 215 generated for OPFs that matched important genes involved with SCFA creation were analyzed.

- 217 Finally, models to classify high or low SCFA concentration based on the median of each SCFA and
- ²¹⁸ 16S rRNA gene sequencing data was created using the Random Forest algorithm.

Acknowledgements

The authors thank the Great Lakes-New England Early Detection Research Network for providing the fecal samples that were used in this study. We would also like to thank Kwi Kim and Thomas M Schmidt for their help in running the short-chian fatty acid analysis on the High-Performace Liquid Chromatography machine at the University of Michigan. Salary support for Marc A. Sze came from the Canadian Institute of Health Research and NIH grant UL1TR002240. Salary support for Patrick D. Schloss came from NIH grants P30DK034933 and 1R01CA215574.

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Figure 1. Using HPLC no change in SCFA measurements was observed between normal, adenoma, and carcinoma individuals. Acetate concentrations in fecal samples of individuals with normal colons, adenomas, and carcinomas (A). Butyrate concentrations in fecal samples of individuals with normal colons, adenomas, and carcinomas (B). Propionate concentrations in fecal samples of individuals with normal colons, adenomas, and carcinomas (C). The black links indicate the median SCFA concentration. Acetate concentrations in fecal samples before and after treatment for adenoma (yellow) and carcinoma (red) (D). Butyrate concentrations in fecal samples before and after treatment for adenoma (yellow) and carcinoma (red) (E). Propionate concentrations in fecal samples before and after treatment for adenoma (yellow) and carcinoma (red) (F). The black dots and lines represent the median change in SCFA concentration.

Figure 2. SCFAs do not improve OTU-based Random Forest models. Difference between the area under the curve of 100 different 80/20 split OTU-based normal versus adenoma 10-fold CV models with and without SCFAs (A). Difference between the area under the curve of 100 different 80/20 OTU-based normal versus carcinoma 10-fold CV models with and without SCFAs (B). The black linke represents the median AUC. The dotted line highlights an AUC of 0.5.

Figure 3. No change in butyrate producing genes identified between normal, adenoma, and carcinoma individuals. Imputed gene relative abundance of important butyrate pathway genes using PICRUSt (A). Counts per million (corrected for size and number of contigs in an OPF) for the Butyrate Kinase gene (B). The other genes from the PICRUSt analysis did not align to any of the OPFs in the metagenome analysis.

Figure S1. OTU-based Random Forest models of SCFA concentrations. Classification
Random Forest train and tests of 100 different 80/20 OTU-based models with 10-fold CV based
on higher or lower than the medain SCFA concentration (A). The top 10 OTUs based on mean
decrease in accuracy (MDA) for each model, colored by their lowest taxonomic identification (B).
Regression Random Forest train and tests of 100 different 80/20 OTU-based models with 10-fold
CV based on correlation to actual SCFA concentration (C). The top 10 OTUs based on mean
decrease in accuracy (MDA) for each model, colored by their lowest taxonomic identification (D).