# The Microbiota and Individual Community Members in Colorectal Cancer: Is There a Common Theme?

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## **Abstract**

Background. An increasing body of literature suggests that there is a role for the microbiota in colorectal cancer. Important drivers within this axis have ranged from individual microbes to the whole community. A recent meta-analysis investigated whether consistent biomarkers could be identified across studies. This report expands on this previous research and tests the hypothesis that the bacterial community is an important driver of disease from control to adenoma to carcinoma. To test this hypothesis we examined both feces (total individuals = 1737) and tissue (total samples = 492) across 14 different studies.

Results. Fecal samples had a significant decrease from control to adenoma to carcinoma for both Shannon diversity and evenness (P-value < 0.05) after correcting for study effect and variable region sequenced. Lower Shannon diversity and evenness in fecal samples resulted in a significant increase in relative risk for carcinoma (P-value < 0.05) while only evenness for adenoma (P-value < 0.05) resulted in a slightly increased relative Previously associated colorectal cancer genera (Fusobacterium, Parvimonas, Peptostreptococcus, or Porphyromonas) followed a similar pattern with a significantly 16 increased relative risk by their presence for carcinoma (P-value < 0.05) but not adenoma 17 (P-value > 0.05) with the exception of *Porphyromonas* (P-value < 0.05). Using the whole 18 community versus only CRC associated genera to build a prediction model resulted in a higher classification AUC for both adenoma and carcinoma for fecal and tissue samples. For the studies that were analyzed, most were adequately powered for large effect size differences which may be more amenable for carcinoma than for adenoma microbiota research.

Conclusions. This data provides support for the importance of the bacteral community
 to both adenoma and carcinoma genesis. The evidence that has been collated within

- 26 this study is much stronger for carcinoma and this may be in part due to the low power to
- <sup>27</sup> detect more subtle changes in the majority of studies that have been performed to date.

# 28 Keywords

<sup>29</sup> microbiota; colorectal cancer; polyps; adenoma; meta-analysis.

# 30 Background

Colorectal cancer (CRC) is a growing world wide health problem [1] in which the microbiota
has been purported to play an active role in disease pathogenesis [2]. Numerous
studies have shown the importance of both individual microbes [3–7] and the overall
community [8–10] in polyp formation in mouse models. There has also been numerous
case/control studies investigating the microbiota in both adenoma and carcinoma.
Recently, a meta-analysis was published investigating whether specific biomarkers could
be consistently identified using multiple data sets [11]. Many of the studies along with the
current meta-analysis focus on identifying biomarkers or individual microbes but do not
critically investigate the community role in the disease.

Using both fecal (total individuals = 1737) and tissue samples (total samples = 492) totalling over 2229 total samples across 14 studies [12–25] we expand both the breadth and scope of the previous meta-analysis; to investigate whether the bacterial community is an important risk factor for both adenoma and carcinoma. To accomplish this we first assessed whether the diversity changes across disease (control to adenoma to carcinoma) and if it results in an increased relative risk for adenoma or carcinoma. Next, we assessed how typical CRC associated genera (*Fusobacterium*, *Parvimonas*, *Peptostreptococcus*, or *Porphyromonas*) affect the relative risk of adenoma or carcinoma. Third, using Random Forest models we analyzed whether the full community or only the CRC associated genera resulted in better model classification area under the curve (AUC). Finally, we examined at what effect size and sample n the studies that were used were adequately powered for.

Our analysis found a continuous decrease in Shannon diversity from control to adenoma to carcinoma and a significantly increased relative risk for carcinoma with this lower diversity. Using only the CRC associated genera this relative risk was higher than what was observed for Shannon diversity. However, by using the full community instead of only

- the CRC associated genera the AUC of the classification models increased. Although
- we analyze a data set with a large number of total individuals each individual study was
- 57 underpowered for effect size differences of 10% or below between the case and control
- 58 group.

## **Results**

Diversity is Lower in Those with Carcinoma and Increases Relative Risk: Using power transformed and Z-score normalized alpha diversity metrics, both evenness and 61 Shannon diversity in feces and not tissue were lower in those with carcinoma [Figure 1]. Using linear mixed-effect models to control for study and variable region there was a 63 significant decrease from control to adenoma to carcinoma for both evenness (P-value 64 = 0.025) and Shannon diversity (P-value = 0.043). This effect was not observed in 65 tissue when additionally controlling for whether the sample came from the same individual (P-value > 0.05). For fecal samples a decrease in Shannon diversity and evenness 67 resulted in a significantly increased relative risk for carcinoma (P-value = 0.01 and P-value = 0.0011, respectively) [Figure 2]. Although these values were significant, the effect size 69 was relatively small for both metrics (Shannon RR = 1.31 and evenness RR = 1.34) [Figure 2]. Only a decrease in evenness had an increased relative risk for adenoma (P-value = 0.032) [Figure 2A & S1] but this effect size was even smaller than what was observed for carcinoma (RR = 1.16).

Using the Bray-Curtis distance metric, there was a significant difference across studies in
the bacterial community of fecal samples between carcinoma and controls but not between
adenoma and controls [Table S1 & S2]. Many studies with unmatched tissue samples had
a significant difference for both adenoma and carcinoma versus controls [Table S3 & S4]
while studies with matched tissue samples had no differences [Table S3 & S4].

Genera Previously Associated with Carcinoma Predominately Increases Relative
Risk for Carcinoma but not Adenoma: The majority of CRC associated genera for
both feces and tissue had a significantly increased relative risk for carcinoma but not for
adenoma [Figure 3]. In fecal samples the relative risk due to CRC associated genera
was greater than either evenness or Shannon diversity [Figure 2 & 3]. Additionally, the

- relative risk of carcinoma continuously increased as individuals tested positive for more
  CRC associated genera [Figure 3B & 3D]. The relative risk effect size was greater for stool
  (RR range = 1.61 2.74) then for tissue (RR range = 1.21 1.81). This decrease may be
  explained by the fact that the tissue analysis include matched samples.
- Two measures in stool for adenoma were significant when investigating these CRC associated genera. The first was *Porphyromonas* (P-value = 0.023) and the second was being positive for three CRC associated genera (P-value = 0.022) [Figure 3A]. For tissue three measures for adenoma were significant; being positive for one CRC associated genera (P-value = 0.032), being positive for two CRC associated genera (P-value = 0.008), and being positive for four CRC associated genera (P-value = 0.039) [Figure 3C].

## <sup>94</sup> Using the Whole Community Increases Model AUC over CRC Associated Genera:

For both fecal and tissue samples (matched and unmatched) the AUC decreases when only OTUs from the CRC associated genera are used [Figure 4 & 5]. This decrease is observed in both adenoma and carcinoma groups [Figure 4 & 5]. The genus models generally had similar trends as observed for the OTU based models with the full genera models performing better then the CRC associated genera models [Figure S2-S3]. Both genus models perform similarily in their ability to be able to predict lesion (adenoma or carcinoma) with carcinoma having a higher AUC then adenoma [Figure S4-S5]. Matched tissue samples for those with carcinoma had an AUC that was more similar to the adenoma models [Figure S4A, S5B, & S6] then carcinoma models [Figure S4B & S5A].

### Majority of Studies are Underpowered for Detecting Small Effect Size Differences:

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When assessing the power of each study at different effect sizes the majority of studies for both adenoma and carcinoma have an 80% power to detect a 30% difference [Figure 6A & B]. No single study that was analyzed had the standard 80% power to detect an effect size difference that was eqaul to or below 10% [Figure 6A & B]. In order to achieve adequate power for small effect sizes it would be necessary to recruit over 1000 individuals for each

arm of the study [Figure 6C].

## 11 Discussion

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Our study identifies clear diversity changes both at the community level and within individual genera that are present in individuals with carcinoma versus those without the disease. 113 Although there was a step wise decrease in diversity from control to adenoma to carcinoma; 114 this did not translate into large effect sizes for the relative risk of either of these two 115 conditions. Even though CRC associated genera increases the relative risk of carcinoma 116 it does not consistently increase the relative risk of adenoma. This information suggests 117 that these specific genera are important in carcinoma genesis but may not be the primary 118 members of the microbial community that contribute to the formation of an adenoma. 119 Additionally, our data shows that by using the whole community our models perform better 120 then when they only use the CRC associated genera. CRC associated genera are clearly 121 important to carcinoma but the context or community in which these microbes are a part 122 of can drastically increase the ability of models to make predictions. This data supports 123 the concept that small localized changes within the community may be occuring that are 124 important in the disease progression of colorectal cancer and that they may not directly involve CRC associated genera. 126

The driver-passenger model of the microbial role in CRC, as summarized by Flynn [2], can be supported with this data for carcinoma but not necessarily for adenoma. The drasitically increased relative risk of disease, when considering the CRC associated genera, is highly supportive of this type of process, especially in the context of increasing relative risk with more CRC associated genera positivity. It is also possible that in a driver-passenger scenario that simply having the driver present or only identifying the passenger is a good enough proxy that the event is occuring. This would account for the observation that there is no constant additive effect on relative risk for increasing positivity. Additionally, the initial establishment of the driver within the system is also dependent on the community that is present and this is supported by the observation that when adding the community context

to our models along with the CRC associated genera the model AUC increases.

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Our carcinoma observations fit the driver-passenger model and support this concept 138 within the framework of the transition from adenoma to carcinoma. In contrast, with 139 the present data we can only suggest that the adenoma observations might fit with this 140 model but the changes that occur at this timepoint are small and possibly focal to the adenoma. The stepwise decrease in diversity suggests that the adenoma community 142 is not normal but this change is subtle. Although there may be localized changes that do depend on the driver-passenger model, supported by an increased relative risk for one, two, and four positive CRC associated genera in tissue [Figure 3C], there may be 145 other processes involved that ultimately exacerbates the condition from a subtle localized change to a global community one. The poor performance of the Random Forest models for classifying adenoma based only on the microbiota would suggest that this is the case. It is possible to hypothesize that at early stages of the diease, how the host interacts to these subtle changes could be the catlyst that ultimately leads to this larger global dysfunctional 150 community.

Although there are still questions that need to be answered for the microbiota and carcinoma, a clearer framework is beginning to develop as to how this occurs. The 153 role of the microbiota in adenoma is still not clear and part of the reason this may be is 154 because many studies are not powered effectively to observe the small changes reported here. It is realistic to suspect that many changes in carcinoma could easily result in effect sizes that are 30% or more between the case and control. Most of the studies analyzed have sufficient power to detect these type of changes. In contrast, our data suggests that the adenoma effect size is relatively small. None of the studies analyzed were properly 159 powered to detect a 10% or lower change between case and controls and this may well be 160 the range in which differences consistently occur in adenoma. Future studies investigating 161 adenoma and the microbiota need to take these factors into consideration if we are to work out the role of the microbiota in adenoma formation.

## 64 Conclusion

By aggregating together a large collection of studies from both feces and tissue we are able to provide information in support of the importance of the bacterial community in both adenoma and carcinoma. We are also able to provide support for the driver-passenger model in the context of carcinoma. However, within the context of adenoma it is less clear that this relationship exists. These observations highlight the importance of power and sample number considerations when considering investigations into the microbiota and adenoma due to the subtle changes in the community.

#### 72 Methods

Obtaining Data Sets: Studies used for this meta-analysis were identified through the review articles written by Keku, et al. and Vogtmann, et al. [26,27]. Additional studies not mentioned in the reviews were obtained from the authors knowledge of the literature. All 175 studies were included that used tissue or feces as their sample source for 16S rRNA gene 176 sequencing analysis. Studies using either 454 or Illumina sequencing technology were 177 included. Only data sets that had sequences available for analysis were included. Some 178 studies did not have publically available sequences or did not have meta data in which 179 the authors were able to share. After these filtering steps the following studies remained: 180 Ahn [21], Baxter [24], Brim [17], Burns [22], Chen [14], Dejea [19], Flemer [13], Geng [25], 181 Hale [12], Kostic [28], Lu [16], Sanapareddy [20], Wang [15], Weir [18], and Zeller [23]. 182 The Zackular [29] study was not included becasue the 90 individuals analyzed within the 183 study are contained within the larger Baxter study. The Kostic study was not used since 184 after sequence processing all the case samples did not have more than 100 sequences remaining. This left a total of 14 studies in which complete analysis could be completed.

Data Set Breakdown: In total there were 7 studies with only fecal samples (Ahn, Baxter,
Brim, Hale, Wang, Weir, and Zeller), 5 studies with only tissue samples (Burns, Dejea,
Geng, Lu, Sanapareddy), and 2 studies with both fecal and tissue samples (Chen and
Flemer). The total number of individuals that were analyzed after sequence processing for
feces was 1737 [Table 1]. The total number of matched and unmatched tissue samples
that were analyzed after sequence processing was 492 [Table 2].

Sequence Processing: For the majority of studies raw sequences were downloaded from the SRA (ftp://ftp-trace.ncbi.nih.gov/sra/sra-instant/reads/ByStudy/sra/SRP/) and metadata was obtained from the following website: http://www.ncbi.nlm.nih.gov/Traces/study/ by searching the respective accession number of the study. Of the studies that did not have

sequences and meta data on the SRA one study had the data stored on DBGap [21] and for four studies the data was obtained directly from the authors [12,13,18,20]. Each study was processed using the mothur (v1.39.3) software program [30]. Where possible quality filtering utilized the default methods used in mothur for either 454 or Illumina based 200 sequencing. If it was not possible to use these defaults the author stated quality cut-offs 201 were used instead. Chimeras were identifed and removed using the VSEARCH [31] 202 program and de novo OTU clustering at 97% similarity using the OptiClust algorithm [32] 203 was utilized. 204

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Statistical Analysis: All statistical analysis after sequence processing utilized the R software package (v3.4.2). For the alpha diversity analysis values were power transformed using the rcompanion (v1.10.1) package and then Z-score normalized using the car (v2.1.5) package. Testing for alpha diversity differences utilized linear mixed-effect models created using the lme4 (v1.1.14) package to correct for study and variable region effects in feces and study, variable region, and individual effects in tissue. Relative Risk was analyzed using both the epiR (v0.9.87) and metafor (v2.0.0) packages. Relative risk significance testing utilized the chi-squred test. Beta-diversity differences utilized a Bray-Curtis distance matrix and PERMANOVA executed with the vegan (v2.4.4) package. Random Forest models were built using both the caret (v6.0.77) and randomForest (v4.6.12) packages. Random Forest testing of the obtained AUC versus a random model AUC utilized T-tests. Power analysis and estimations were made using the pwr (v1.2.1) and statmod (v1.4.30) packages. All figures were created using both ggplot2 (v2.2.1) and gridExtra (v2.3) packages.

Study Analysis Overview: Alpha diversity was first assessed for differences between controls, adenoma, and carcinoma. We analyzed the data using linear mixed-effect models 219 and relative risk. Beta-diversity was then assessed for each inidividual study. Next, four specific CRC-associated genera (Fusobacterium, Parvimonas, Peptostreptococcus, and Porphyromonas) were assessed for differences in relative risk. We then built Random

Forest models based on all genera or the select CRC-associated genera. The models were trained on one study then tested on the remaining studies for every study. The data was split between feces and tissue samples. Within the tissue groups the data was further divided between matched and unmatched tissue samples. Where applicable for each 226 study predictions for adenoma and carcinoma were tested. This same approach was then 227 applied at the OTU level with the exception that instead of testing on the other studies 228 a 10-fold cross validation was utilized and 100 different models were created based on 229 random 80/20 splitting of the data to generate a range of expected AUCs. For OTU based 230 models the CRC associated genera included all OTUs that had a taxonomic classification 231 to Fusobacterium, Parvimonas, Peptostreptococcus, or Porphyromonas. The power of 232 each study was assessed for an effect size ranging from 1% to 30%. An estimated sample 233 n for these effect sizes was also generated based on 80% power. 234

Reproducible Methods: The code and analysis can be found here https://github.com/
SchlossLab/Sze\_CRCMetaAnalysis\_Microbiome\_2017. Unless mentioned otherwise the
accession number for the raw sequences for the studies used in this analysis can be found
directly in the respective batch file, on the GitHub repository or in the original manuscript.

## **Declarations**

#### 240 Ethics approval and consent to participate

Ethics approval and informed consent for each of the studies used is mentioned in the respective manuscript used in this meta-analysis.

#### 243 Consent for publication

Not applicable.

#### 245 Availability of data and material

A detailed and reporducible description of how the data were processed and analyzed for each study can be found at https://github.com/SchlossLab/Sze\_CRCMetaAnalysis\_
Microbiome\_2017. Raw sequences can be downloaded from the SRA in most cases and can be found in the respective studies batch file in the GitHub repo or within the original publication. When sequences were not publicly available contacting the corresponding author for raw sequences needs to be undertaken.

#### 252 Competing Interests

All authors declare that they do not have any relevant competing interests to report.

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#### 257 Authors' contributions

All authors helped to design and conceptualize the study. MAS identified and analyzed the data. MAS and PDS interpreted the data. MAS wrote the first draft of the manuscript and both he and PDS reviewed and revised updated versions. All authors approved the final manuscript.

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Table 1: Studies with Stool Samples Included in the Analysis

Study	Data Stored	16S Region	Controls	Adenoma	Carcinoma
Ahn	DBGap	V3-4	148	0	62
Baxter	SRA	V4	172	198	120
Brim	SRA	V1-3	6	6	0
Flemer	Author	V3-4	37	0	43
Hale	Author	V3-5	473	214	17
Wang	SRA	V3	56	0	46
Weir	Author	V4	4	0	7
Zeller	SRA	V4	50	37	41

Table 2: Studies with Tissue Samples Included in the Analysis

Study	Data Stored	16S Region	Controls	Adenoma	Carcinoma
Burns	SRA	V5-6	18	0	16
Chen	SRA	V1-V3	9	0	9
Dejea	SRA	V3-5	31	0	32
Flemer	Author	V3-4	103	37	94
Geng	SRA	V1-2	16	0	16
Lu	SRA	V3-4	20	20	0
Sanapareddy	Author	V1-2	38	0	33

- Figure 1: Alpha Diversity Differences between Control, Adenoma, and Carcinoma
  Across Sampling Site. A) Alpha diversity metric differences by group in stool samples. B)
  Alpha diversity metric differences by group in unmatched tissue samples. C) Alpha diversity
  metric differences by group in matched tissue samples. The dashed line represents a
  Z-score of 0 or no difference from the median.
- Figure 2: Relative Risk for Adenoma or Carcinoma based on Alpha Diversity

  Metrics in Stool. A) Alpha metric relative risk for adenoma. B) Alpha metric relative risk

  for carcinoma. Colors represent the different variable regions used within the respective

  study.
- Figure 3: Colorectal Cancer Associated Genera Relative Risk for Adenoma and
  Carcinoma in Stool and Tissue. A) Adenoma relative risk in stool. B) Carinoma relative
  risk in stool. C) Adenoma relative risk in tissue. D) Carcinoma relative risk in tissue. For all
  panels the relative risk was also compared to whether one, two, three, or four of the CRC
  associated genera were present.
- Figure 4: OTU Random Forest Model of Stool Across Studies. A) Adenoma random forest model between the full community and CRC associated genera OTUs only. B)
  Carcinoma random forest model between the full community and CRC associated genera
  OTUs only. The dotted line represents an AUC of 0.5 and the lines represent the range in which the AUC for the 100 different 80/20 runs fell between.
- Figure 5: OTU Random Forest Model of Tissue Across Studies. A) Adenoma random forest model between the full community and CRC associated genera OTUs only. B)
  Carcinoma random forest model between the full community and CRC associated genera
  OTUs only. The dotted line represents an AUC of 0.5 and the lines represent the range in
  which the AUC for the 100 different 80/20 runs fell between.
  - Figure 6: Power and Effect Size Analysis of Studies Included. A) Power based on

effect size for studies with Adenoma individuals. B) Power based on effect size for studies with Carcinoma individuals. C) The estimated sample number needed for each arm of each study to detect aneffect size of 1-30%. The dotted red lines in A) and B) represent the generally used power of 0.8.

- Figure S1: Relative Risk for Adenoma or Carcinoma based on Alpha Diversity

  Metrics in Tissue. A) Alpha metric relative risk for adenoma. B) Alpha metric relative risk

  for carcinoma. Colors represent the different variable regions used within the respective

  study.
- Figure S2: Random Forest Genus Model AUC for each Stool Study. A) AUC of
  Adenoma models using all genera or CRC associated genera only. B) AUC of Carcinoma
  models using all genera or CRC associated genera only. The black line represents the
  median within each group.
- Figure S3: Random Forest Genus Model AUC for each Tissue Study. A) AUC of
  Adenoma models using all genera or CRC associated genera only divided between
  matched and unmatched tissue. B) AUC of Carcinoma models using all genera or CRC
  associated genera only. The black line represents the median within each group divided
  between matched and unmatched tissue.
- Figure S4: Random Forest Prediction Success Using Genera for each Stool Study.

  A) AUC for prediction in Adenoma using all genera or CRC associated genera only. B)

  AUC for prediction in Carcinoma using all genera or CRC associated genera only. The

  dotted line represents an AUC of 0.5. The x-axis is the data set in which the model was

  initially trained on.
- Figure S5: Random Forest Prediction Success of Carcinoma Using Genera for each
  Tissue Study. A) AUC for prediction in unmatched tissue for all genera or CRC associated
  genera only. B) AUC for prediction in matched tissue using all genera or CRC associated
  genera only. The dotted line represents an AUC of 0.5. The x-axis is the data set in which
  the model was initially trained on.
- Figure S6: Random Forest Prediction Success of Adenoma Using Genera for each
  Tissue Study.