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

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- 2 Background.
- 3 Results.
- 4 Conclusions.
- 5 Keywords
- 6 microbiota; colorectal cancer; polyps; treatment; risk factor.

7 Background

- 8 Results
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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. [1,2]. All studies were included that used tissue or feces as their sample source for 16S rRNA gene sequencing analysis. Studies using either 454 or Illumina sequencing technology were included. Only data sets that had the raw sequences available for analysis were included. Some studies did not have publically available raw sequences or did not have meta data in which the authors were able to share. After this filtering step the following studies remained: Ahn [3], Baxter [4], Brim [5], Burns [6], Chen [7], Dejea [8], Flemer [9], Geng [10], Hale [11], Kostic [12], Lu [13], Sanapareddy [14], Wang [15], Weir [16], and Zeller [17]. The Zackular [18] study was not included becasue the 90 individuals analyzed within the study are contained within the larger Baxter study. The Kostic study was not used since after sequence processing all the case samples did not have more than 100 sequences remaining. This left a total of 13 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 initially run through the sequence processing for
the fecal samples was 1899 and for the tissue samples was 462.

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
[3] and four studies the data was obtained directly from the authors [9,11,14,16]. Each

study was processed using the mothur (v1.39.3) software program [19]. Where possible quality filtering utilized the default methods used in mothur for either 454 or Illumina based sequencing. If it was not possible to use these defaults the author stated quality cut-offs were used instead. Chimeras were identifed and removed using the VSEARCH [20] program and *de novo* OTU clustering at 97% similarity using the OptiClust algorithm [21] was utilized.

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 both T-tests and ANOVA. Linear mixed-effect models were created using the lme4 (v1.1.14) package to correct for both study and variable region effect in the diversity measures when analyzing colorectal cancer groups. 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) 55 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 57 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 and adenoma versus cancer and controls versus adenoma. We analyzed the data using standard T-tests, ANOVA, linear mixed-effect models, and relative risk. 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. Both prediction for adenoma
and carcinoma were tested. This same approach was then applied at the OTU level with
the exception that instead of testing on the other studies a 10-fold cross validation was
utilized and 100 different models were created based on random 80/20 splitting of the data
to generate a range of expected AUCs. The power of each study was assessed for and
effect size ranging from 1% to 30%. An estimated sample n for these effect sizes was also
generated based on 80% power.

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.

79 Declarations

80	Ethics approval and consent to participate
81	Need to fill in.
82	Consent for publication
83	Not applicable.
84	Availability of data and material
85	Need to fill in.
86	Competing Interests
87	All authors declare that they do not have any relevant competing interests to report.
88	Funding
89	Need to fill in.
90	Authors' contributions
91	Need to fill in.

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Table 1:

Table 2:

- Figure 1:
- Figure 2:
- Figure 3:
- Figure 4:

- 161 Figure S1:
- Figure S2:
- Figure S3: