# The effect of treatment on the microbiota of patients diagnosed with colonic lesions

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#### Abstract

Background. Colorectal cancer (CRC) is a worldwide health problem. Despite growing evidence that members of the gut microbiota can drive tumorigenesis, little is known about what happens to the microbiota after treatment for an adenoma or carcinoma. This study tested the hypothesis that treatment for adenoma or carcinoma alters the abundance of bacterial populations associated with disease to those associated with a normal colon. We tested this hypothesis by sequencing the 16S rRNA genes in the feces of 67 individuals before and after treatment for adenoma (N=22), advanced adenoma (N=19), and carcinoma (N=26).

There were large changes to the bacterial communities associated with treatment across the three groups. The communities from patients with carcinomas changed significantly more than those with adenoma following treatment (P-value < 0.001). There was no significant change in the microbiota between patients with adenoma and advanced adenoma, or between patients with advanced adenoma and carcinoma (P-value > 0.05). Although treatment was associated with intrapersonal changes, the change in the abundance of individual OTUs to treatment was not consistent within 16 diagnosis groups (P-value > 0.05). Because the distribution of OTUs across patients and 17 diagnosis groups was irregular, we used the Random Forest machine learning algorithm 18 to identify groups of OTUs that allowed us to successfully distinguish between pre and post-treatment samples for each of the diagnosis groups. Although the three models successfully differentiated between the pre and post-treatment samples, there was little 21 overlap between the OTUs that were indicative of treatment. Next, we used a larger cohort that contained individuals with normal colons and those with adenomas, advanced adenomas, and carcinomas to determine whether individuals who underwent treatment were more likely to have OTUs associated with normal colons. We again built Random Forest models and measured the change in the positive probability of having one of the three diagnoses. Only patients who had carcinomas experienced a significant decrease in positive probability of having a lesion after treatment (P-value < 0.05), indicating that the microbial milieu of the colon more closely resembled that of a normal colon. Finally, we tested whether the type of treatment impacted the microbiota of those diagnosed with carcinomas and were unable to detect any significant differences in characteristics of these communities between individuals treated with surgery alone and those treated with chemotherapy or chemotherapy and radiation (P-value > 0.05).

Conclusions. Further exploration of the relationship between diagnosis, treatment, and
 the impact on the microbiota will yield improvements in disease management.

#### 36 Keywords

microbiota; colorectal cancer; polyps; treatment; risk factor.

# Background

tumorigenesis [14].

Colorectal cancer (CRC) is the third most common cause of cancer deaths in the United States [1,2]. Disease mortality has significantly decreased, predominately due to improvements in screening [2]. Despite these improvements, there are still approximately 50,000 CRC-related deaths per year in the United States [1]. Current estimates indicate that 20-30% of those who undergo treatment will experience recurrence and 35% of all patients will die within five years [3–5]. Identification of methods to assess patients' risk of recurrence is of great importance to reduce mortality and healthcare costs.

There is growing evidence that the gut microbiota is involved in the progression of CRC.

Mouse-based studies have identified populations of *Bacteroides fragilis*, *Escherichia coli*,

and *Fusobacterium nucleatum* that alter disease progression [6–10]. Furthermore, studies

that shift the structure of the microbiota through the use of antibiotics or inoculation of

germ free mice with human feces have shown that varying community compositions can

result in varied tumor burden [11–13]. Collectively, these studies support the hypothesis

that the microbiota can alter the amount of inflammation in the colon and with it the rate of

Building upon this evidence, several human studies have identified unique signatures of colonic lesions [15–20]. One line of research has identified community-level differences between those bacteria that are found on and adjacent to colonic lesions and have supported a role for *Bacteroides fragilis*, *Escherichia coli*, and *Fusobacterium nucleatum* in tumorigenesis [21–23]. Others have proposed feces-based biomarkers that could be used to diagnose the presence of colonic adenomas and carcinomas [24–26]. These studies have associated *Fusobacterium nucleatum* and other oral pathogens with colonic lesions (adenoma, advanced adenoma, and carcinoma). They have also noted that the loss of bacteria generally thought to produce short chain fatty acids, which can suppress

inflammation, is associated with colonic lesions. This suggests that gut bacteria have a role in tumorigenesis with potential as useful biomarkers for aiding in the early detection of disease [21–26].

Despite advances in understanding the role between the gut microbiota and colonic tumorigenesis, we still do not understand how treatments including resection, chemotherapy, and/or radiation affect the composition of the gut microbiota. If the microbial community drives tumorigenesis then one would hypothesize that treatment to remove a lesion would affect the microbiota and risk of recurrence. To test this hypothesis, we addressed two related questions: Does treatment affect the colonic microbiota in a predictable manner? If so, does the treatment alter the community to more closely resemble that of individuals with normal colons?

We answered these questions by sequencing the V4 region of 16S rRNA genes amplified from fecal samples of individuals with adenoma, advanced adenoma, and carcinomas pre and post-treatment. We used classical community analysis to compare the alpha and beta-diversity of communities pre and post treatment. Next, we generated Random Forest models to identify bacterial populations that were indicative of treatment for each diagnosis group. Finally, we measured the predictive probabilities to assess whether treatment yielded bacterial communities similar to those individuals with normal colons. We found that treatment alters the composition of the gut microbiota and that, for those with carinomas, the gut microbiota shifted more towards that of a normal colon after treatment. In the individuals with carcinomas, no difference was found by the type of treatment (surgery alone versus surgery with chemotherapy). Understanding how the community responds to these treatments could be a valuable tool for identifying biomarkers to quantify the risk of recurrence and the likelihood of survival.

#### Results

Treatment alters the bacterial community structure of patients diagnosed with colonic lesions. Within our 67-person cohort we tested whether the microbiota of patients with adenoma (N=22), advanced adenoma (N=19), or carcinoma (N=26) had any broad differences between pre and post-treatment samples [Table 1]. The structure of the 91 microbial communities of the pre and post-treatment samples differed, as measured by the 92  $\theta_{YC}$  beta diversity metric [Figure 1A]. We found that the communities obtained pre and post-treament among the patients with carcinomas changed significantly more than those 94 patients with adenoma (P-value = 5.4e-05). There were no significant differences in the 95 amount of change observed between the patients with adenoma and advanced adenoma or between the patients with advanced adenoma and carcinoma (P-value > 0.05). Next, 97 we tested whether there was a consistent direction in the change in the community 98 structure between the pre and post-treatment samples for each of the diagnosis groups [Figure 1B-D]. We only observed a consistent shift in community structure for the patients 100 with carcinoma when using a PERMANOVA test (adenoma P-value =0.999, advanced 101 adenoma P-value =0.945, and carcinoma P-value =0.005). Finally, we measured the 102 number of observed OTUs, Shannon evenness, and Shannon diversity in the pre and post-treatment samples and did not observe a significant change for any of the diagnosis groups (P-value > 0.05) [Table S1].

The effects of treatment are not consistent across treatment groups. We used two approaches to identify those bacterial populations that change between the two samples for each diagnosis group. First, we sought to identify individual OTUs that could account for the change in overall community structure. However, using a paired Wilcoxon test we were unable to identify any OTUs that were significantly different in the pre and post-treatment groups (P-value NA 0.05). It is likely that high inter-individual variation and the irregular distribution of OTUs across individuals limited the statistical power of the test. To overcome

these problems we developed Random Forest models to identify collections of OTUs that would allow us to differentiate between pre and post-treatment samples from each of the diagnosis groups. Because of the relatively small number of subjects in each group, we restricted our models to only incorporate 10 OTUs to limit the likelihood that the models 116 would overfit the data. Despite this restriction, the models performed well (adenoma AUC 117 range = 0.69 - 0.92, advanced adenoma AUC range = 0.80 - 1.00, carcinoma AUC range 118 = 0.82 - 0.98). Interestingly, the 10 OTUs that were used for each model had little overlap 119 with each other [Figure 2]. These results support the earlier community-wide analysis 120 where we observed that the treatment had an impact on the overall community structure; 121 however, the effect of treatment was not consistent across patients and diagnosis groups. 122

Post-treatment samples from patients with carcinoma more closely resemble those of a normal colon. Next, we determined whether treatment changed the microbiota in a 124 way that the post-treatment communities resembled that of patients with normal colons. To test this, we used an expanded cohort of 423 individuals that were diagnosed under 126 the same protocol as having normal colons or colons with adenoma, advanced adenoma, 127 or carcinoma [Table 2]. We then constructed Random Forest models to classify the study 128 samples, with the 3 diagnosis groups (adenoma, advanced adenoma, or carcinoma), or 129 having a normal colon. The models performed well (adenoma AUC range =0.62 - 0.72, 130 advanced adenoma AUC range = 0.68 - 0.77, carcinoma AUC range = 0.84 - 0.90; Figure 131 S1). The OTUs that were incorporated into the adenoma and advanced adenoma models 132 largely overlapped and those OTUs that were used to classify the carcinoma samples were 133 largely distinct from those of the other two models [Figure 3A]. Among the OTUs that were 134 shared across the three models were those populations generally considered beneficial to 135 their host (e.g. Faecalibacterium, Lachnospiraceae, Bacteroides, Dorea, Anaerostipes, and 136 Roseburia) [Figures 3B]. Although many of these OTUs were also included in the model differentiating between patients with normal colons and those with carcinoma, this model also included OTUs affiliated with populations that have previously been associated with

carcinoma (Fusobacterium, Porphyromonas, Parvimonas) [24-26] [Figure S2] with some individuals showing are marked decrease in relative abundance [Figure S3]. Finally, we 141 applied these three models to the pre and post-treatment samples for each diagnosis group 142 and quantified the change in the positive probability of the model. A decrease in the positive 143 probability would indicate that the microbiota more closely resembled that of a patient 144 with a normal colon. There was no significant change in the positive probability for the 145 adenoma or advanced adenoma groups [Figure 4]. The positive probability for the pre and 146 post-treatment samples from patients diagnosed with carcinoma significantly decreased 147 with treatment, suggesting a shift toward a normal microbiota for most individuals. Only, 6 148 of the 26 patients (23.08%) who were diagnosed with a carcinoma had a higher positive 149 probability after treatment; one of those was re-diagnosed with carcinoma on the follow up 150 visit. These results indicate that, although there were changes in the microbiota associated 151 with treatment, those experienced by patients with carcinoma after treatment yielded gut 152 bacterial communities of greater similarity to that of a normal colon. 153

#### Difficult to identify effects of specific treatments on the change in the microbiota.

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The type of treatment that the patients received varied across diagnosis groups. Those 155 with adenomas and advanced adenomas received surgical resection (adenoma, N=4; 156 advanced adenoma, N=4) or polyp removal during colonoscopy (adenoma, N=18; advanced adenoma, N=15) and those with carcinomas received surgical resection (N=12), 158 surgical resection with chemotherapy (N=9), and surgical resection with chemotherapy 159 and radiation (N=5). We focused on the patients with carcinoma and pooled those patients 160 that received chemotherapy with those that received chemotherapy and radiation to improve our statistical power. We did not observe a significant difference in the effect 162 of these treatments on the number of observed OTUs, Shannon diversity, or Shannon 163 evenness (P-value > 0.05). Furthermore, there was not a significant difference in the effect of the treatments on the amount of change in the community structure (P-value = 0.298). Finally, the change in the positive probability was not significantly different between the

two treatment groups (P-value = 0.999). Due to the relatively small number of samples in
each treatment group, it was difficult to make a definitive statement regarding the specific
type of treatment on the amount of change in the structure of the microbiota.

### Discussion

Our study focused on comparing the microbiota of patients diagnosed with adenoma, advanced adenoma, and carcinoma before and after treatment. For all three groups of patients, we observed changes in their microbiota. After treatment, the microbiota of 173 patients with carcinoma changed significantly more than the other groups. This change 174 resulted in communities that more closely resembled those of patients with a normal colon. 175 This may suggest that treatment for carcinoma is not only successful for removing the 176 carcinoma but also at reducing the associated bacterial communities. Understanding 177 the effect of treatment on the microbiota of those diagnosed with carcinomas may have 178 important implications for reducing disease recurrence. It is intriguing that it may be 179 possible to use microbiome-based biomarkers to not only predict the presence of lesions 180 but to assess the risk of recurrence. 181

Patients diagnosed with adenoma and advanced adenoma, however, did not experience a shift towards a community structure that resembled those with normal colons. This may 183 be due to the fundamental differences between the features of adenomas and advanced 184 adenomas and carcinoma. Specifically, carcinomas may create an inflammatory milieu that 185 would impact the structure of the community and removal of that stimulus would alter said 186 structure. In addition, it is possible that the difference between the microbiota of patients 187 with adenoma and advanced adenoma and those with normal colons is subtle. This is 188 supported by the reduced ability of our models to correctly classify patients with adenomas 189 and advanced adenomas relative to those diagnosed with carcinomas [Figure S1]. Given 190 the irregular distribution of microbiota across patients in the different diagnosis groups, it is 191 possible that we lacked the statistical power to adequately characterize the change in the 192 communities following treatment. 193

There was a subset of patients (6 of the 26 with carcinomas) who demonstrated an elevated

probability of carcinoma after treatment. This may reflect an elevated risk of recurrence. The 23.08% prevalence of increased carcinoma probability from our study is within the expected rate of recurrence (20-30% [3,4]). We hypothesized that these individuals may 197 have had more severe tumors; however, the tumor severity of these 6 individuals (3 with 198 Stage II and 3 with Stage III) was similar to the distribution observed among the other 20 199 patients. We also hypothesized that we may have sampled these patients later than the 200 rest and their communities may have reverted to a carcinoma-associated state; however, 201 there was not a statistically significant difference in the length of time between sample 202 collection among those whose probabilities increased (382  $\pm$  116 days) or decreased (341 203  $\pm$  98 days) (Wilcoxon Test; P-value = 0.56). Finally, it is possible that these patients may 204 not have responded to treatment as well as the other 20 patients diagnosed with carcinoma 205 and so the microbiota may not have been impacted the same way. Again, further studies 206 looking at the role of the microbiota in recurrence are needed to understand the dynamics 207 following treatment. 208

Our final hypothesis was that the specific type of treatment altered the structure of 209 the microbiome. The treatment to remove adenomas and advanced adenomas was 210 either polyp removal or surgical resection whereas it was surgical resection alone or 211 in combination with chemotherapy or with chemotherapy and radiation for individuals 212 with carcinoma. Because chemotherapy and radiation target rapidly growing cells, these 213 treatments would be more likely to cause a turnover of the colonic epithelium driving 214 a more significant change in the structure of the microbiota. Although, we were able 215 to test for an effect across these specific types of treatment, the number of patients in 216 each treatment group was relatively small. Finally, those undergoing surgery would have 217 received antibiotics and this may be a potential confounder. However, our pre-treatment 218 stool samples were obtained before the surgery and the post-treatment samples were obtained long after any effects due to antibiotic administration on the microbiome would be 220 expected to occur (351  $\pm$  102 days).

This study expands upon existing research that has established a role for the microbiota in tumorigenesis and that demonstrated the utility of microbiome-based biomarkers to predict the presence of colonic lesions. The most exciting future direction from the current study is the possibility that markers within the microbiota could be used to evaluate the effect of treatment and predict recurrence for those diagnosed with carcinoma. If such an approach is effective, it might be possible to target the microbiota as part of adjuvant therapy. Our data provides additional evidence on the importance of the microbiota in tumorigenesis by addressing the recovery of the microbiota after treatment and opens interesting avenues of research into how these changes may affect recurrence.

## 31 Methods

Study Design and Patient Sampling. Sampling and design have been previously reported in Baxter, et al [24]. Briefly, study exclusion involved those who had already 233 undergone surgery, radiation, or chemotherapy, had colorectal cancer before a baseline 234 fecal sample could be obtained, had IBD, a known hereditary non-polyposis colorectal 235 cancer, or familial adenomatous polyposis. Samples used to build the models for 236 prediction were collected either prior to a colonoscopy or between one and two weeks 237 after initial colonoscopy. The bacterial community has been shown to normalize back to 238 a pre-colonoscopy community within this time period [27]. Our study cohort consisted 239 of 67 individuals with an initial sample as described and a follow up sample obtained 240 between 188 - 546 days after treatment of lesion [Table 1]. Patients were diagnosed by 241 colonoscopic examination and histopathological review of any biopsies taken. Patients 242 were classified as having advanced adenoma if they had an adenoma greater than 1 243 cm, more than three adenomas of any size, or an adenoma with villous histology. This study was approved by the University of Michigan Institutional Review Board. All study 245 participants provided informed consent and the study itself conformed to the guidelines set 246 out by the Helsinki Declaration.

16S rRNA Gene Sequencing. Sequencing was completed as described by Kozich, et al. [28]. DNA extraction used the 96-well Soil DNA isolation kit (MO BIO Laboratories) and an epMotion 5075 automated pipetting system (Eppendorf). The V4 variable region was amplified and the resulting product was split between four sequencing runs with normal, adenoma, and carcinoma evenly represented on each run. Each group was randomly assigned to avoid biases based on sample collection location. The pre and post-treatment samples were sequenced on the same run.

Sequence Processing. The mothur software package (v1.37.5) was used to process

the 16S rRNA gene sequences and has been previously described [28]. The general workflow using mothur included merging paired-end reads into contigs, filtering for low quality contigs, aligning to the SILVA database [29], screening for chimeras using UCHIME [30], classifying with a naive Bayesian classifier using the Ribosomal Database Project (RDP)[31], and clustered into Operational Taxonomic Units (OTUs) using a 97% similarity cutoff with an average neighbor clustering algorithm [32]. The number of sequences for each sample was rarefied to 10523 to minimize the impacts of uneven sampling.

**Model Building.** The Random Forest [33] algorithm was used to create the three models 263 used. The adenoma model classified normal versus adenoma, the advanced adenoma model classified normal versus advanced adenoma, and the carcinoma model classified normal versus carcinoma. The total number of individuals in the treatment models was 67 individuals. There were a total of 22 individuals in the treatment adenoma model, 267 19 individuals in the treatment advanced adenoma model, and 26 individuals in the treatment carcinoma model [Table 1]. The total number of individuals in the normal versus 269 diagnosis models was 423 individuals [Table 2]. There were a total of 239 individuals in 270 the adenoma model, 262 individuals in the advanced adenoma model, and 266 individuals 271 in the carcinoma model [Table 2]. Each model was then applied to our 67-person cohort 272 [Table 1] which assesed the prediction of pre-treatment adenoma (adenoma n = 22 and 273 disease free n = 0) versus post-treatment adenoma (adenoma n = 0 and disease free n 274 = 22), pre-treatment advanced adenoma (advanced adenoma n = 19 and disease free n 275 = 0) versus post-treatment advanced adenoma (advanced adenoma n = 0 and disease 276 free n = 19), pre-treatment carcinoma (carinoma n = 26 and disease free n = 0) versus 277 post-treatment carcinoma (carcinoma n = 1 and disease free n = 25). 278

The model included only OTU data obtained from 16S rRNA sequencing. Non-binary data was checked for near zero variance and OTUs that had near zero variance were removed.

This pre-processing was performed with the R package caret (v6.0.73). Optimization of

the mtry hyper-parameter involved making 100 different 80/20 (train/test) splits of the data where normal and adenoma, normal and advanced adenoma, or normal and carcinoma were represented in the same proportion within both the whole data set and the 80/20 split. For each of the different splits, 20 repeated 10-fold cross validation was performed on the 80% component to optimize the mtry hyper-parameter by maximizing the AUC (Area Under the Curve of the Receiver Operator Characteristic). The resulting model was then tested on the hold out data obtained from the 20% component. All three models had an optimized mtry of 2. The hyper-parameter, mtry, defines the number of variables to investigate at each split before a new division of the data was created with the Random Forest model. 

Assessment of the most important OTUs to the model involved counting the number of times an OTU was present in the top 10% of mean decrease in accuracy (MDA) for each of the 100 different splits run. This was then followed with filtering of this list to variables that were only present in more than 50% of these 100 runs. The final collated list of variables was then run through the mtry optimization again. Once the ideal mtry was found the entire sample set specific to normal versus adenoma, normal versus advanced adenoma, or normal versus carcinoma was used to create the final Random Forest model on which classifications on the 67-person cohort was completed. For all three models the final optimized mtry was 2. The only difference other than the sample set used between treatment models and normal versus diagnosis models was that only the top 10 OTUs were used to build each respective treatment model by diagnosis group to help avoid model overfitting.

Statistical Analysis. The R software package (v3.3.2) was used for all statistical analysis.

Comparisons between bacterial community structure utilized PERMANOVA [34] in the

vegan package (v2.4.1). Comparisons between probabilities as well as overall OTU

differences between pre and post-treatment samples utilized a paired Wilcoxon ranked

sum test. Where multiple comparison testing was appropriate, a Benjamini-Hochberg (BH)

- correction was applied [35] and a corrected P-value of less than 0.05 was considered significant. Unless otherwise stated the P-values reported are those that were BH corrected.
  Model rank importance was determined by obtaining the median MDA from the 100, 20 repeated 10-fold cross validation and then ranking from largest to smallest MDA.
- Reproducible Methods. A detailed and reproducible description of how the data were processed and analyzed can be found at https://github.com/SchlossLab/Sze\_followUps\_ 2017. Raw sequences have been deposited into the NCBI Sequence Read Archive (SRP062005 and SRP096978) and the necessary metadata can be found at https://www. ncbi.nlm.nih.gov/Traces/study/ and searching the respective SRA study accession.

- Figure 1: General Differences between Adenoma, Advanced Adenoma, and
  Carcinoma Groups After Treatment. A) Thetayc distance from pre versus post sample
  within each individual. A significant difference was found between the adenoma and
  carcinoma group for thetayc (P-value = 5.36e-05). Solid black points represent the median
  value for each diagnosis group. B) NMDS of the pre and post-treatment samples for
  the adenoma group. C) NMDS of the pre and post-treatment samples for the advanced
  adenoma group. D) NMDS of the pre and post-treatment samples for the carcinoma group.
- Figure 2: The 10 OTUs used to classify treatment for Adenoma, Advanced
  Adenoma, and Carcinoma. A) Adenoma OTUs. B) Advanced Adenoma OTUs. C)
  Carcinoma OTUs.
- Figure 3: Common OTUs to All Models. A) Venn diagram showing the OTU overlap between each model. B) For each common OTU the lowest taxonomic identification and importance rank for each model run is shown.
- Figure 4: Treatment Response Based on Models Built for Adenoma, Advanced
  Adenoma, or Carcinoma. A) Positive probability change from initial to follow up sample in
  those with adenoma. B) Positive probability change from initial to follow up sample in those
  with advanced adenoma. C) Positive probability change from initial to follow up sample in
  those with carcinoma.

Table 1: Demographic Data of pre and Post Treatment Cohort

	Adenoma	Advanced Adenoma	Carcinoma
n	22	19	26
Age (Mean ± SD)	61.68 ± 7.2	63.11 ± 10.9	61.65 ± 12.9
Sex (%F)	36.36	36.84	42.31
BMI (Mean ± SD)	26.86 ± 3.9	25.80 ± 4.7	28.63 ± 7.2
Caucasian (%)	95.45	84.21	96.15
Post-Treatment Sample (Mean ± SD)	255.41 ± 42	250.16 ± 41	350.85 ± 102

Table 2: Demographic Data of Training Cohort

	Normal	Adenoma	Advanced Adenoma	Carcinoma
n	172	67	90	94
Age (Mean ± SD)	54.29 ± 9.9	63.01 ± 13.1	64.07 ± 11.3	64.37 ± 12.9
Sex (%F)	64.53	46.27	37.78	43.62
BMI (Mean ± SD)	26.96 ± 5.3	25.68 ± 4.8	26.66 ± 4.9	29.27 ± 6.7
Caucasian (%)	87.79	92.54	92.22	94.68

Figure S1: ROC Curves of the Adenoma, Advanced Adenoma, and Carcinoma **Models.** A) Adenoma ROC curve: The light green shaded areas represent the range of values of a 100 different 80/20 splits of the test set data and the dark green line represents the model using 100% of the data set and what was used for subsequent classification. B) Advanced Adenoma ROC curve: The light yellow shaded areas represent the range of 341 values of a 100 different 80/20 splits of the test set data and the dark yellow line represents 342 the model using 100% of the data set and what was used for subsequent classification. C) 343 Carcinoma ROC curve: The light red shaded areas represent the range of values of a 100 344 different 80/20 splits of the test set data and the dark red line represents the model using 345 100% of the data set and what was used for subsequent classification. 346

Figure S2: Summary of Important OTUs for the Adenoma, Advanced Adenoma, and
Carcinoma Models. A) MDA of the most important variables in the adenoma model. The
dark green point represents the mean and the lighter green points are the value of each
of the 100 different runs. B) Summary of Important Variables in the advanced adenoma
model. MDA of the most important variables in the SRN model. The dark yellow point
represents the mean and the lighter yellow points are the value of each of the 100 different
runs. C) MDA of the most important variables in the carcinoma model. The dark red point
represents the mean and the lighter red points are the value of each of the 100 different
runs.

Figure S3: Pre and Post Treatment Relative Abundance of CRC associated OTUs within the Carcinoma Model.

# **Declarations**

#### **Ethics approval and consent to participate**

The University of Michigan Institutional Review Board approved this study, and all subjects provided informed consent. This study conformed to the guidelines of the Helsinki Declaration.

#### **Consent for publication**

Not applicable.

#### Availability of data and material

A detailed and reproducible description of how the data were processed and analyzed can be found at https://github.com/SchlossLab/Sze\_followUps\_2017. Raw sequences have been deposited into the NCBI Sequence Read Archive (SRP062005 and SRP096978) and the necessary metadata can be found at https://www.ncbi.nlm.nih.gov/Traces/study/ and searching the respective SRA study accession.

#### Competing Interests

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

#### Funding

This study was supported by funding from the National Institutes of Health to P. Schloss (R01GM099514, P30DK034933) and to the Early Detection Research Network (U01CA86400).

#### Authors' contributions

All authors were involved in the conception and design of the study. MAS analyzed the data. NTB processed samples and analyzed the data. All authors interpreted the data. MAS and PDS wrote the manuscript. All authors reviewed and revised the manuscript. All authors read and approved the final manuscript.

## 82 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 Amanda Elmore for reviewing and correcting code error and providing feedback on manuscript drafts. We would also like to thank Nicholas Lesniak for providing feedback on manuscript drafts.

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