Differences in the fecal Microbiome Before and After Colorectal Cancer Treatment

Running Title: Human Microbiome and Colorectal Cancer

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Abstract

2 Background: Colorectal cancer (CRC) continues to be a worldwide health problem with

early detection being used as a key component in mitigating deaths due to the disease.

Previous research suggests a link between fecal bacterial microbiome and CRC. The

overall objective of our study was to investigate the changes in the bacterial microbiome

after surgery in patients with lesion (i.e. adenoma or carcinoma). Specifically, we wanted to

 $_{ au}$ identify what within the community was different within those undergoing surgical removal

of lesion. We also wanted to investigate the use of the bacterial microbiome and Fecal

Immunoglobulin Test (FIT) to build models which classify individuals as having a lesion or

whether samples could be classified as before or after surgery.

Results: Adenoma individual's bacterial microbiome were more similar to their pre-surgery sample then those with carcinoma (P-value = 0.00198) and this was also reflected in FIT (P-value = 2.15e-05). There was no significant difference in any individual OTU between samples before and after surgery (P-value > 0.125). A model with a total of 37 variables was able to classify lesion (AUC = 0.847 - 0.791) while the model to classify samples as before and after surgery had 33 variables (AUC = 0.79 - 0.651). Both models had a significant decrease in the positive probability of a lesion or being a before surgery sample (P-value = 1.91e-11 and 6.72e-12). In total there were 14 OTUs that were common to both models and the majority of these classified to commensal bacteria with the largest representation belonging to Bacteroides, Blautia, Streptococcus, and Clostridiales.

Conclusions: Our data suggests that treatment not only significantly reduces the probability of having a colonic lesion within our models but also causes detectable changes in the bacterial microbiome. Further surveillance of these individuals will enable us to determine whether models such as the one we present here can also be used to predict recurrence of colorectal cancer.

26 Keywords

²⁷ bacterial microbiome; colorectal cancer; polyps; FIT; detection; risk factors

28 Background

Colorectal cancer (CRC) continues to be a leading cause of cancer related deaths and is the second most common cancer death among men aged 40-79 years of age [1,2].

Over the last few years death due to the disease has seen a significant decrease, thanks mainly to improvements in screening [1]. However, despite this improvement there are still approximately 50,000 deaths from the disease a year [2]. It is estimated that around 5-10% of all CRCs can be explained by autosomal dominant inheritance [3]. The vast majority of CRCs are not inherited and the exact etiology of the disease has not been well worked out [2]. Although many risk factors have been identified [2] and non-invasive screening techniques have started to be put into consistent use [4,5] there has been an additional increase in the incidence of CRC in the younger population.

This increased incidence of CRC in the younger population is concerning since having
either an adenoma or carcinoma increases ones risk for future adenomas or carcinomas
[6–8]. This increased risk can also carry with it an increased chance of mortality due to
this recurrence [9,10]. Therefore, there has been a great amount of interest in early risk
stratification tools [11,12] that can help identify those that may be at most susceptibility
to reccurence. Concurrently, there has also been a lot of interest in new areas that could
have a role in disease pathogenesis, such as the gut bacterial microbiome.

There has been promising work on the bacterial microbiome and it's ability to be able to complement existing screening methods such as Fecal Immunoglobulin Test (FIT) or act alone as a screening tool [13,14]. There has also been research into how this microbiome could be altered directly on tumor tissue itself [15]. A few studies have now even shown how this microbiome [16] or specific members within it [17] could be directly involved with the pathogenesis of CRC. These studies have helped to provide a tantilzing link between the bacterial microbiome and CRC. However, at this present time there remains limited

- information on the bacterial microbiome before and after successful surgery for removal of the adenoma or carcinoma and whether it changes at all.
- In this study we investigated what happened to the bacterial microbiome before and after surgery for indivdiuals with either adenoma and carcinoma. Our anlaysis includes both alpha and beta diversity analysis along with investigation of individual operational taxonomic units (OTUs). We also utilized Random Forest models and observed how these models as well as specific OTUs within this model performed on before (initial) and after (follow up) surgery samples. We used these models to look for similar important OTUs to identify the crucial OTUs for not only classifying initial and follow up samples but also lesion or normal.

Results

Bacterial Community and FIT Changes before and after Surgery Based on thetayo distance metrics, comparing the initial to the follow up samples, there was a significant difference between the adenoma and carcinoma groups (P-value = 0.00198) [Figure 1a]. There was also a significant difference in change in FIT between initial and follow up 67 samples (P-value = 2.15e-05) [Figure 1b]. The whole community structure before and after 68 surgery were visualized on NMDS graphs for both adenoma [Figure1c] (PERMANOVA = 69 0.997) and carcinoma [Figure 1d] (PERMANOVA = 0.002). When all initial and follow up 70 samples were compared to each other there was no significant overall difference between 71 them (PERMANOVA = 0.085). There was no significant difference between initial and follow up samples for observed OTUs, Shannon diversity, or evenness after correction for multiple comparisons [Table S1]. There was also no significant difference between initial and follow up samples for any single OTU [Figure S1].

Differences in Adenoma and Carcinoma of Previously Associated Cancer Bacteria We next examined whether there were differences in previously well described carcinoma associated OTUs. These included the OTUs that aligned with *Porphyromonas* 78 asaccharolytica (Otu000153), Fusobacterium nucleatum (Otu000226), Parvimonas 79 micra (Otu000460), and Peptostreptococcus stomatis (Otu000653). First, the carcinoma 80 samples showed a significant difference between initial and follow up samples for 81 Peptostreptococcus stomatis (P-value = 0.0183) and Porphyromonas asaccharolytica (P-value = 0.0154) whereas there was no signficant differences in any of these OTUs in 83 the adenoma samples [Table S4]. Second, when these OTUs were present, there was 84 a clear magnitude difference based on whether they were from adenoma or carcinoma 85 [Figure 2]. However, only a small percentage of those with adenoma or carcinoma were positive for any of these OTUs.

Full and Reduced Model Results Since differences were observed between initial and follow up samples and only a small number of individuals were positive for previously associated CRC bacteria; we next investigated if we could create a model that could adequately classify and adjust lesion probability based on the bacterial community and FIT.

The lesion model had an AUC range of 0.723 to 0.795 versus the initial follow up model which had an AUC range of 0.451 to 0.67 after 100 20 repeated 10-fold cross validations. Interestingly, identification of the most important variables and reducing the models to only these factors increased the AUC in the lesion model (0.791 - 0.847) and initial and follow up model (0.651 - 0.79).

The test set AUC range for the full and reduced lesion model were similar to that reported for the training set AUC ranges and the ROC curve ranges overlap each other [Figure 3a]. The ROC curve for the final lesion model used was within the range of both the full and reduced lesion model [Figure 3a]. Interestingly, the test set AUC range for the initial and follow up performed much better then the training set AUCs. Both the full and reduced initial and follow up models overlapped with each other [Figure 3b] there was a marked decrease in the ROC curve for the final model used.

Most Important Variables to the Models The reduced models were built based on the most important variables to the respective full model. For the lesion model there were a total of 37 variables [Figure S2] whereas for the initial and follow up model there were a total of 33 variables [Figure S3]. For both models FIT resulted in the largest decrease in MDA [Figure S2a & S3a].

Positive Probability Prediction after Surgical Removal of Adenoma or Carcinoma
Regardless of model used there was a significant decrease in the positive probability of
either the sample being lesion or an initial sample on follow up [Figure 4 & S4] (full lesion
P-value = 1.11e-11, reduced lesion P-value = 1.91e-11, initial and follow up P-value =
6.71e-12, and reduced initial and follow up P-value = 6.72e-12).

For the full and reduced lesion model there was a significant difference in the classification for the lesion model between predicted and actual (P-value = 4.19e-10 and 6.98e-10, respectively) but not for the initial follow up model (P-value = 1.00 and 1.00). However, the lesion model correctly kept the one individual who still had a carcinoma on follow up above the cut off point [Figure 4a & S4a] for a positive call while the initial and follow up models did not [Figure 4b & S4b].

Common OTUs to both Lesion and Initial and Follow Up Models There were a total of
14 OTUs that were common to both models. Of these OTUs the most common taxonomic
121 identifications were to Blautia, Bacteroides, Streptococcus, and Clostridiales. The majority
123 of these OTUs had taxonomic identification to bacteria typically thought of as commensal
124 [Table S2].

Treatment and Time Differences There was no difference in the amount of change in positive probability for either the full or reduced lesion model for either chemotherapy (P-value = 0.821 and 0.821) or radiation therapy (P-value = 0.69 and 0.981). Although the initial follow up model was similar there was a significant decrease in positive probability for those treated with chemotherapy (P-value = 7.04e-04 and 5.07e-03). Time elapsed between the collection of the follow up sample from the initial sample, did not have a significant difference between adenoma and carcinoma (uncorrected P-value = 0.784).

32 Discussion

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There was no difference in alpha diversity metrics between the initial and follow up samples [Table S1]. Yet based on thetayc there was a significant difference between inital and 134 follow up samples [Figure 1a] with carcinoma samples being more dissimilar to their 135 initial samples then adenoma samples. The change in FIT was also different between 136 initial and follow up for those with adenoma or carcinoma [Figure 1b]. There was also a 137 significant difference in the overall bacterial community structure for those in the carcinoma 138 group [Figure 1d] before and after surgery but not for the adenoma group [Figure 1c]. 139 Investigation of the OTU relative abundance before and after surgery found no single OTU 140 to be significantly different [Figure S1].

Interestingly, when only previously associated cancer bacteria were investigated only 2
out of the 4 had a significant decrease in relative abundance between initial and follow
up for the carcinoma group and 0 out of the 4 were significant for the adenoma group.
This data suggests that the changes that may be important in the bacterial microbiome
after surgery are not necessarily any one specific bacterium, a clear depression of CRC
related bacteria, nor any addition of new or depressed bacterium but rather that a shift of
the bacterial community as a whole occurs with the existing members of this microbiome.

We next created a model that incorporated FIT and the bacterial microbiome to either be able to classify lesions (adenoma or carcinoma) or initial and follow up samples in order to confirm that the community was what was responding or changing due to surgery. We found that the OTUs that made up the most important variables to the model overwhelmingly belonged to comensal bacteria. With only the lesion model having a single OTU from previously associated cancer bacteria (*Porphyromonas asaccharolytica*). Using only these important OTUs and FIT both models (lesion and initial and follow up) had a significantly decreased positive probability of either lesion or being an initial sample on follow up

[Figure 4 & S4]. Confirming the importance of the changes of comensal bacteria to these classifications was that a total of 14 OTUs were common to both models and the vast majority belonged to regular residents of our gut community.

There was no difference for the majority of models tested for differences in positive probablity based on whether chemotherapy or radiation was received. There was also no difference in the thetayc distance metric based on length of time between initial and follow up sample between adenoma and carcinoma. These results would indicate that the findings described were specific to the surgical intervention and that differences observed between carcinoma and adenoma samples can not be simply attributed to collection time between samples.

This study builds upon previous work from numerous labs that have looked into the bacterial 167 microbiome as a potential screening tool [13,14] by exploring what happens to the bacterial 168 community after surgical removal of a lesion. Based on previous work by Arthur, et al. 169 [18] it may not be surprising to have E.coli as one of the most important OTUs and one 170 that was common to both models. Interestingly, many of the most important OTUs had 171 taxonomic identification for resident gut microbes. This could suggest that the bacterial 172 community is one of the first components that could change during the pathogenesis of 173 disease. These bacterial microbiome changes could be the first step in allowing more inflammatory bacterium to gain a foothold within the colon [19].

Curiously, we observed that the typical CRC associated bacteria were not predictive within our models. One potential explanation is that even with surgery and a shift of the bacterial community these specific bacteria still persisted within the colon. An alternative reason would be that even if they were reduced in relative abundance they were still present. Another potential explanation is that they were not present in enough individuals to be able to classify those with and without disease with a high degree of accuracy. It is possible that our Random Forest models were able to get the same information from measures such as

FIT or other OTUs. Finally, it is also possible that all three of these potential explanations could have played a role. Regardless, our observations would suggest that an individual's 184 resident bacteria have a large role to play in disease initiation and could change in a way 185 that allows predictive models to lower the positive probability of a lesion after surgery 186 [Figure 4]. It should be stated that our study does not argue against the importance of 187 these CRC associated bacteria in the pathogenesis of disease but rather that the models 188 do not utilize these specific bacteria for classification purposes (lesion or initial and follow 189 up). In fact, it is possible that these CRC associated bacteria are important in the transition 190 from adenoma to carcinoma and would be one explanation as to why in our data we not 191 only see high initial relative abundances, in certain individuals, but also large decreases in 192 relative abundance in those with carcinoma but not in those with adenoma after surgery 193 [Figure 2]. 194

One limitation of our study is that we do not know whether individuals who were still classified as positive by the lesion model eventually had a subsequent CRC diagnosis. 196 This information would help to strengthen the case for our Random Forest based model 197 to have kept a number of individuals above the cutoff threshold even though at follow up 198 they were diagnosed as no longer having a lesion. Another limitation is that we do not 199 know if adding modern tests such as the fecal DNA test [20] could help improve our overall 200 AUC. Another limitation is that this study drew heavily from those with caucasian ancestry. 201 The results may not be immediately representative of those with either Asian or African 202 ancestry. Finally, although our training and test set are relatively large we still run the 203 risk of overfitting or having a model that may not be immediately extrapolateable to other 204 populations. We've done our best to safeguard against this by not only running 10-fold 205 cross validation but also having over 100 different 80/20 splits to try and mimic the type of 206 variation that might be expected to occur. 207

Another interesting outcome was that Within figure 3 the initial and follow up model showed

better test AUC results then the training set AUC. This may have occured because the training AUC that was determined from 20 repeated 10 fold cross validation removed samples at random and did not take into account that they were matched samples. Another potential reason is that the model itself may be overfit since the total number of samples 212 was not that large. However, the lesion model did not suffer from these discrepencies and 213 similar conclusions can be drawn solely from this model. Regardless, further independent 214 studies will need to be carried out to verify our findings since not only are we dealing with 215 feces, which could be very different than the communities present on the actual tissue, but 216 also are dealing with correlations that may not be representative of the true pathogensis of 217 disease.

Despite these limitations we think that these findings significantly add to the existing scientific knowledge on CRC and the bacterial microbiome. The ability for machine learning algorithms to take bacterial microbiome data and successfully lower positive probability after either adenoma or carcinoma removal provides evidence that there are specific signatures, mostly attributable to commensal organisms, associated with these lesions. It also shows that these algorithms can not only successfully react to successful treatment regimens but also may be able to one day stratify CRC disease risk with a high level of accuracy.

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27 Methods

Study Design and Patient Sampling The sampling and design of the study was similar to that reported in Baxter, et al [13]. In brief, study exclusion involved those who had already undergone surgery, radiation, or chemotherapy, had colorectal cancer before a 230 baseline fecal sample could be obtained, had IBD, a known hereditary non-polyposis 231 colorectal cancer, or Familial adenomatous polyposis. Samples used to build the models 232 for prediction were collected either prior to a colonoscopy or between 1 - 2 weeks after. The 233 bacterial microbiome has been shown to nomralize within this time period [21]. Our follow 234 up data set had a total of 67 individuals that not only had a sample as described previoulsy 235 but also a follow up sample between 188 - 546 days after surgery and treatment had been 236 completed. This study was approved by the University of Michigan Institutional Review 237 Board. All study participants provided informed consent and the study itself conformed to 238 the guidelines set out by the Helsinki Declaration. 239

FIT and 16S rRNA Gene Sequencing FIT was analyzed as previously published using
both OC FIT-CHEK and OC-Auto Micro 80 automated system (Polymedco Inc.) [22]. 16S
rRNA gene sequencing was completed as previously described by Kozich, et al. [23]. In
brief, 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 three sequencing runs with control,
adenoma, and carcinoma evenly represented on each run. Each group was randomly
assigned to avoid biases based on sample collection location.

Sequence Processing The mothur software package (v1.37.5) was used to process the 16S rRNA gene sequences. This process has been previously described [23]. The general processing workflow using mothur was as follows: Paired-end reads were first merged into contigs, quality filtered, aligned to the SILVA database, screened for chimeras,

classified with a naive Bayesian classifier using the Ribosomal Database Project (RDP), and clustered into Operational Taxonomic Units (OTUs) using a 97% similarity cutoff with an average neighbor clustering algorithm. The numer of sequences for each sample was rarified to 10521 in an attempt to minimize uneven sampling.

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Lesion Model Creation The Random Forest [24] algorithm was used to create the model used for prediction of lesion (adenoma or carcinoma) with the main testing and training of the model completed on a data set of 490 individuals. This model was then applied to our follow up data set of 67 individuals. The model included data on FIT and the bacterial microbiome. Non-binary data was checked for near zero variance. Data columns that had near zero variance were removed. This pre-processing was performed with the R package caret (v6.0.73). Optimization of the mtry hyperparameter involved taking the samples and making 100 80/20 (train/test) splits in the data where control and lesion were equally represented in the 80 and 20 split, respectively. This 80% portion was then split again into an 80/20 split, and run through 20 repeated 10-fold cross validations to optimzie the model's AUC (Area Under the Curve of the Receiver Operator Characteristic). This resulting model was then tested on the 20% of the data that was originally held out from this overall process. Once the ideal mtry was found the entire 490 sample set was used to create the final Random Forest model on which testing on the 67-person cohort was completed. The default cutoff of 0.5 was used as the threshold to classify individuals as positive or negative for lesion. The hyperparameter, mtry, defines the number of variables to investigate at each split before a new division of the data is created.

Initial Follow Up Model Creation We also investigated whether a model could be created that could identify before and after surgery samples. The main difference was that only the 67-person cohort follow up data set was used at all stages of model building and classification. Other than this difference the creation of this model and optimization of the mtry hyperparameter was completed using the same procedure that was used to create

278 the lesion model.

Selection of Important OTUs In order to assess which variables were most important to all the models we counted the number of times a variable was present in the top 10% of mean decrease in accuracy (MDA) for each different 80/20 split model and then filtered this list to variables that were only present more than 50% of the time. This final collated list of variables was what was considered the most important for the lesion or initial follow up models.

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

Comparisons between bacterial community structure utilized PERMANOVA [25] in the

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

differences between initial and follow up samples utilized a paired wilcoxson ranked sum

test. Where multiple comparison testing was needed a Benjamini-Hochberg (BH) correction

was applied [26] and a corrected P-value of less than 0.05 was considered significant.

Unless otherwise stated the P-values reported are those of that were BH corrected.

Analysis Overview Initial and follow up samples were analyzed for differences in alpha and beta diversity. Next, differences in FIT between initial and follow ups for either 293 adenoma or carcinoma were investigated. From here, all OTUs that were used in the 294 lesion model were then analyzed using a paired wilcoxson test. We then investigated 295 the relative abundance of specific previously CRC associated bacteria, specifically, 296 OTUs that taxonomically classified to Fusobacterium nucleatum, Parvimonas micra, 297 Peptostreptococcus assacharolytica, and Porphyromonas stomatis. We wanted to test if 298 there were any differences based on whether the individual had an adenoma or carcinoma. 299 From here the lesion model was then tested for accuracy in prediction and whether it 300 reduced the positive probability of lesion after surgery. The most important OTUs for this 301 were used to build a reduced model and it was assessed for similarity to the original model. 302 We then used the initial follow up model to assess whether it could classify samples better 303

then the lesion model. The most important OTUs were then identified from this model and used to create a reduced feature initial follow up model. This reduced feature model, as was done with the lesion model, was compared to the full model for loss of accuracy. Finally, a list of common OTUs were found for the two different models used.

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).

Figure 1: General Differences between the Adenoma or Carcinoma Group. A) A significant difference was found between the adenoma and carcinoma group for thetayc (P-value = 0.00198). B) A significant difference was found between the adenoma and carcinoma group for change in FIT (P-value = 2.15e-05). C) NMDS of the initial and follow up samples for the Adenoma group. D) NMDS of the initial and follow up samples for the Carcinoma group. For C) and D) the teal represents initial samples and the pink represents follow up samples.

Figure 2: Previously Associated CRC Bacteria in Initial and Follow up Samples. A)

Carcinoma initial and follow up samples. There was a significant difference in initial and

follow up sample for the OTUs classfied as *Peptostreptococcus stomatis* (P-value = 0.0183)

and *Porphyromonas asaccharolytica* (P-value = 0.0154). B) Adenoma initial and follow up

samples. There were no significant differences between initial and follow up.

Figure 3: Graph of the Receiver Operating Characteristic Curve for lesion and Initial and Follow up Models. The shaded areas represents the range of values of a 100 different 80/20 splits of the test set data using either all variables (grey) or reduced variable (red) models. The blue line represents the reduced variable model using 100% of the data set.

A) Lesion model. B) Initial/Follow up model

Figure 4: Breakdown by Carcinoma and Adenoma of Prediction Results for Lesion 329 and Initial Follow Up Reduced Variable Models A) Lesion positive probability adjustment 330 of those with carcinoma from initial to follow up sample B) Initial follow up positive probability 331 adjustment of those with carcinoma from initial to follow up sample C) Lesion positive 332 probability adjustment of those with adenoma as well as those with SRN and the probability 333 adjustment from initial to follow up sample. D) Initial follow up positive probability adjustment 334 of those with adenoma as well as those with SRN and the probability adjustment from initial 335 to follow up sample. The dotted line represents the threshold used to make the decision of whether a sample was positive or not.

Figure S1: Distribution of P-values from Paired Wilcoxson Analysis of OTUs in Initial versus Follow Up

Figure S2: Summary of Important Variables in the Lesion Model A) MDA of the most important variables in the lesion model. The black point represents the median and the different colors are the different runs up to 100. B) The total number of appearances of each variable in the 100 different lesion models. The cutoff of 50% was used to assess importance.

Figure S3: Summary of Important Variables in Initial Follow Up Model A) MDA of the most important variables in the lesion model. The black point represents the median and the different colors are the different runs up to 100. B) The total number of appearances of each variable in the 100 different lesion models. The cutoff of 50% was used to assess importance.

Figure S4: Breakdown by Carcinoma and Adenoma of Prediction Results for Lesion
and Initial and Follow Up Full Variable Models A) Lesion positive probability adjustment
of those with carcinoma from initial to follow up sample B) Initial follow up positive probability
adjustment of those with carcinoma from initial to follow up sample C) Lesion positive
probability adjustment of those with adenoma as well as those with SRN and the probability
adjustment from initial to follow up sample. D) Initial follow up positive probability adjustment
of those with adenoma as well as those with SRN and the probability adjustment from initial
to follow up sample. The dotted line represents the threshold used to make the decision of
whether a sample was positive or not.

59 Figure S5: Thetayc Graphed Against Time of Follow up Sample from Initial

Declarations

- 361 Ethics approval and consent to participate
- 362 Consent for publication
- 363 Availability of data and material
- 364 Competing Interests
- All authors declare that they do not have any relevent competing interests to report.

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370 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.

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