- The fecal microbiome as a tool for monitoring and
- predicting response outcomes in Ustekinumab-treated,
- anti-TNF-alpha refractory Crohn's Disease patients.
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- 6 Running title: The fecal microbiome as a tool for monitoring and predicting response outcomes in
- ⁷ Ustekinumab-treated, anti-TNF-alpha refractory Crohn's Disease patients.

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- $_{13}$ α Currently at ...

4 Abstract

Abstract: The 16S rRNA gene from patient stool samples was sequenced using the Illumina MiSeq platform. The resulting sequences were curated and assigned to taxonomic groups using the mothur software package to determine the bacterial communities and relative abundance of bacterial species present in these patients. The relative abundance among the fecal microbiota, patient demographic data, and clinical metadata were used as input to a random forest machine-learning algorithm to predict disease severity and response to treatment with UST.

Fecal microbial diversity at baseline significantly correlates with markers for disease severity, such as Crohn's Disease Activity Index (CDAI), stool frequency, and disease duration. Additionally, the overall community structure of the microbiome was significantly different based on stool frequency, CRP, fecal lactoferrin, fecal calprotectin, corticosteroid use, disease duration, and tissue involvement. Baseline fecal microbiome community structures and species diversity were significantly different among responders and non-responders to UST treatment. Faecalibacterium, among other taxa, was significantly more abundant in responders/remitters. Additionally, the microbiome of clinical responders changed over time, in contrast to nonresponsive subjects. Using AUC-RF, differences in the baseline microbiome and clinical metadata were able to predict response to UST, especially remission, with some AUCs approaching 0.85.

Importance: Crohn's disease (CD) is a global health issue characterized by patches of ulceration and inflammation along the gastrointestinal tract. Individuals with CD have reduced microbial diversity in their guts, compared to healthy individuals. It remains unclear if this reduced diversity is a result or cause of pathogenesis. We investigated the relationship between the fecal microbiome and clinical phenotypes in subjects with moderate to severe CD treated with Ustekinumab (UST) in a Phase 2b study to determine whether the fecal microbiome at baseline is predictive of disease severity and therapeutic response, as well as if the fecal microbiota changes due to therapy.

The ability to predict and monitor response to treatment using the microbiome will likely provide another clinical tool in treating CD patients. Additionally, the observed baseline differences in fecal microbiota and changes due to therapeutic response will allow further investigation into the microbes important in CD pathogenesis as well as establishing and maintaining CD remission. Finally, beneficial microbes

- associated with response to treatment could be developed as probiotics to increase the likelihood of
- response while undergoing treatment.
- 44 Keywords: Crohn's Disease, fecal microbiome, biologics, prediction

15 Introduction

Crohn's disease (CD), an incurable inflammatory bowel disease (IBD), is a global health issue with 46 increasing incidence. CD affects approximately 3 million people worldwide, causing large economic and 47 healthcare utilization impacts on society (1-3). CD is characterized by patches of ulceration and inflammation affecting the entire bowel wall along the gastrointestinal tract, most commonly in the ileum and 49 colon. Individuals with CD experience frequent diarrhea, abdominal pain, fatigue, and weight loss resulting in significant health care costs, lower quality of life, and economic impacts due to loss of productivity (2, 4, 5). Current treatments for CD include antibiotics, anti-inflammatory drugs, immunomodulators, 52 surgery, and biologic agents targeting tumor necrosis factor alpha (TNF- α), such as Infliximab (Remi-53 cade). Within 10 years of diagnosis, approximately half of individuals with CD will require surgery and the majority will experience escalating immunosuppressive treatment (6). Currently, individuals with CD are treated based on disease location and risk of complications using escalating immunosuppressive treatment and/or surgery with the goal of achieving and sustaining remission (5, 7). Faster induction of remission following diagnosis reduces the risk of irreversible intestinal damage and disability (7-9) (7-9). Anti-TNF- α therapy in combination with thiopurines has emerged as the preferred treatment for 59 CD, but up to half of individuals with CD fail to respond or lose response to anti-TNF- α therapy (6, 7). Ustekinumab (UST), a monoclonal antibody directed against the shared p40 subunit of IL-12 and IL-23, has been proposed as an alternative therapy for these patients (10). While clinical trials have demonstrated that UST is a viable option for the treatment of CD (7, 10-12) (7, 10-12), some patients 63 within these trials were non-responsive to UST, which we hypothesized was due to differences in the patients' gut microbiota at the initiation of treatment. The precise etiology of CD remains unknown, but host genetics, environmental exposure, and the gut microbiome appear involved (1, 13). Genome-wide association studies of individuals with CD identified 67 several susceptibility genes including NOD2, a receptor involved in bacterial killing and innate immunity. 68 Defects in NOD2 function affects microbial sensing, the regulation of IL-23 driven Th17 responses, and indirect modulation of the gut microbiome (5, 14). The gut microbiome has also been shown to play a 70 key role in inflammation, immunity, and IBD (15). Individuals with CD have reduced microbial diversity 71 in their guts, compared to healthy individuals, with a lower relative abundance of Firmicutes and an

increased relative abundance of Enterobacteraciae and Bacteroides, at the phylum level (14, 16-19) .

Additionally, previous studies have shown that the gut microbiome correlates with disease severity in new-onset, pediatric CD patients (19, 20).

The microbiome has shown promise as a predictive tool for a variety of disease. Colon cancer has been show to etc etc prediction...other success stories leading into the Schloss lab's success predicting cancer? -> can we do for CD It remains to be determined, however, whether the microbiome can predict response to therapy in CD (14). Additionally, the effect of biologic treatment on the gut microbiome is not well understood (14). If the fecal microbiome can be used as a theraprognostic tool to non-invasively determine and monitor disease severity as well as predict response to specific treatment modalities, then more targeted treatment could result in reduced adverse effects of less effective therapies and faster achievement of remission.

Our lab was approached to analyze the gut microbiomes of individuals who participated in a placebocontrolled Phase II clinical trial to determine the safety and efficacy of UST in treating CD (10).

Using stool samples taken prior to, and following, the start of the study, 16S rRNA gene sequence
data from these patients' stool allowed us to determine associations between clinical metadata, disease
severity, and the fecal microbiome, as well as whether clinical responders have a microbiome that is
distinct from non-responders at baseline. We also show that the microbiome changed in subjects who
responded to treatment. Our study demonstrates that the fecal microbiome is associated with baseline
clinical metadata and that these associations and differences are useful in predicting disease severity and
treatment outcome.

3 Results

Characteristics of Study Population Using 16S rRNA gene sequencing, we studied the fecal microbiota in a subset of TNF- α refractory CD patients who took park in the CERTIFI clinical trial (10). Demographic and baseline disease characteristics of this subset are summarized in Table 1. Patients with a history of moderate to severe CD were randomly assigned to a treatment group in the induction phase of the study (Figure 1A). At Week 8 patients were re-randomized into maintenance therapy groups. Both patients and clinicians were blinded to their induction and maintenance treatment groups. Subjects provided stool sample at screening (week 0), week 4, week 6, and week 22 post induction (Figure 1B). Therapeutic response was evaluated based on the change in CDAI at week 6 (primary endpoint) and week 22 (secondary endpoint) post induction. For this study, "response" was defined as a decrease in CDAI of 30% from baseline and "remission" was defined as a CDAI below 150.

Comparison of microbiome at screening based on clinical variables *Question: disease severity and*microboime at screening?

Following sequence curation using the mothur software package, we obtained between 1 and 130,074 sequences per sample (median 13786) (21). Parallel sequencing of a mock community revealed an error rate of 0.017%. To limit effects of uneven sampling, we rarefied the dataset to 3,000 sequences per sample. Samples from subjects that completed the clinical trial and had complete clinical metadata were included in our analysis. Of these samples, 306 were provided prior to treatment as well as 258 provided at week 4, 289 at week 6, and 205 at week 22 post-treatment, for a total of 1058 samples.

To determine whether there are any significant associations between microbial diversity and clinical 112 variables of interest, we compared the microbiome with clinical data at week 0 (Supplemental Table 113 1). We observed a weak, but significant correlation between CDAI and species diversity, with higher 114 CDAI correlating to lower diversity (rho =-0.161, p = 0.00483). The overall community structure was 115 not different based on CDAI. When looking at CDAI subscores, we observed a weak, but significant 116 association between species diversity and the frequency of loose stools per week (rho =-0.193, p = 117 0.000693). The overall community structure was also significantly different based on weekly loose 118 stool frequency (p=0.012). No significant association was observed between CRP, fecal calprotectin, 119 or fecal lactoferrin and species diversity, following multiple comparison correction. However, the overall 120 community structure was significantly different based on CRP (p = 0.022), fecal calprotectin (p =121 (0.002), and fecal lactoferrin (p = 0.001). No significant differences in the microbiome were observed for 122 BMI, weight, or sex. Overall community structure was different based on age (p = 0.019). The overall 123 community structure was also different based on the tissue affected (p = 0.001). Species diversity 124 (p = 0.000196) and the overall community structure (p = 0.004) were significantly different based on 125 corticosteroid use. The community structure was significantly different (p =0.001) based on disease 126 duration and a weak, but significant correlation was seen between species diversity and disease duration 127 (rho =-0.225, p = 0.0000713), with lower diversity corresponding to longer disease. 128

Comparison of clinical responders and non-responders *Q*: Are responder different from non respon-

We hypothesized that there are associations between the microbiome at baseline and response to treat-131 ment. To test this, we compared the week 0 microbiomes of subjects based on treatment group and 132 response status at week 6 and week 22. Only week 6 remitters induced with UST were significantly 133 different from non-remitters in terms of species diversity (p = 0.020), having higher diversity based 134 on inverse simpson (put inv simp medians here?). Baseline community structures were significantly 135 different for each outcome status (response/remission) and treatment group at each clinical endpoint 136 (week 6 response p = 0.012, week 6 remission p = 0.017, week 22 response p = 0.012, week 22 137 remission p = 0.012), as seen in Table 2. No phyla were significantly different by treatment and 138 response, however Fusobacteria was less frequently observed in week 6 remitters than non-remitters 139 treated with UST (median relative abundance, respectively). Two taxa, Bacteroides (OTU0019) (p 140 = trtd.REMISSwk6.Otu19p) and Faecalibacterium (OTU0007) (p = trtd.REMISSwk6.Otu7p) were significantly more abundant in UST-induced, week 6 remitters compared to non-remitters induced with 142 UST (Figure ?). No individual taxa were significantly different among subjects receiving placebo for 143 induction, regardless of response/remission status at week 6. 144

The microbiome by treatment and response over time Does microbiome change following treatemnt 145 Having oberved differences in the microbiomes of subjects who responded to treatment compared to those who did not, we hypothesized that treatment may affect the microbioime. The effects of biologic 147 treatment of IBD on the microbiome are not yet well described, but are hypothesized to be indirect. 148 We tested whether treatment with UST affects the microbiome using subjects who provided samples at weeks 0, 4, and 6. This allowed for us to analyze 156 treated subjects and 48 placebo subjects at each 150 time point. Using the adonis function in the vegan R package (22), we performed a PERMANOVA 151 stratified on each subject, as a proxy for a repeated measures ANOVA, to determine if the community 152 structure of microbiome changed over time. We included induction treatment group, response at each 153 clinical endpoint, and sample week as parameters. 154

We found that treatment only affects the microbiome over time in subjects who responded to UST at week 22. No significant difference was seen in community structure or species diversity based on sample date when looking at all treatment groups and week 6 response status, but there was a significant

interaction between week 22 response and sample date 0.001. There was also a significant interaction and between week 22 response, induction group, and sample date 0.044 (Supplemental Table 2). This led us to further examining the microbial community structures in week 22 responders and nonresponders over time by induction treatment. No significant difference was observed in Week 22 non-responders over time, regardless of treatment. In week 22 responders, we saw a significant change in community structure over time in both placebo (p = 1) and UST induction groups (p = 1).

Since we observed significant changes in the community structure of week 22 repsonders, we also 164 hypothesized that treatment may also affect species diversity. We tested this by performing a freidman 165 test comparing species diversity at each sample date within each induction treatment group based on 166 their week 22 response status. As seen in Figure 4, we saw no significant difference in species diversity over time in subjects who did not responde at week 22, regardless of induction treatment. However, in 168 UST treated-week 22 responders species diversity increased significantly from week 0 to week 4 (0.0022) 169 and remained higher than baseline at week 6. This change was not observed in subjects induced with 170 placebo who responded at week 22, unlike the community structure analysis. We hypothesize that this 171 reflects decreased inflammation in the subjects who responded to treatment. 172

The microbiome following treatment reflects disease status Does microbiome relfect disease status

at wk 6

A paper recently published by Tedjo et al. demonstrated a link between the microbiome and disease 175 severity, where specific microbes were associated with remission capared to active CD (23). We hy-176 pothesized that the microbiome could be used to monitor response to therapy in a similar manner. We 177 used AUC-RF in order to determine if the fecal microbiome at Week 6 could be used to determine if a 178 study participant responded to therapy or was in remission at Week 6. As seen in Figure 4, using the 179 microbiome alone we achieved an AUC of 0.708 for response with a sensitivity of 0.769 and a specificity 180 of 0.606. For remission we had an AUC of 0.866 with a sensitivity of 0.833 and specificity of 0.832. We 181 were better able to distinguish remitters from non-remitters than responders from non-responders. 182

The top microbes that were indicitive of disease status in this model included blanks at enriched in remitters and blanks enriched in nonremitters.

Prediction of response based on the microbiome at screening can differences at baseline predict

later response to treatment Given the observed differences in the fecal microbiome at baseline and week 6 in responders/remitters compared to nonresponders/nonremitters, we hypothesized that the fecal 187 microbiome could predict response to therapy. To test this hypothesis, we used the AUCRF package in 188 R to develop a random forest classification model to differentiate responders from non-responders, as 189 well as remitters from non-remitters, based on the relative abundance of fecal microbiome community 190 members, clinical metadata, and the combination of microbiome and clinical data (24, 25). We ran 191 these models for response and remission at Week 6 and 22 of the study. The optimal models for 192 response and remission at the primary endpoint (Week 6) are shown in Figure ?. Using only clinical 193 metadata to predict response the model did not perform well, with an AUC of 0.693, a specificity of 194 0.76, and a sensitivity of 0.598. Using only microbiome data, the model predicted response with an 195 AUC of 0.737 with a specificity of 0.807 and a sensitivity of 0.585. When combining clinical metadata 196 with the microbiome, the model predicted response with an AUC of 0.745, a specificity of 0.727, and a 197 sensitivity of 0.744. With respect to Week 6 remission, using solely clinical metadata we achieved AUC 198 of 0.616a with a specificity of 0.801 and a sensitivity of 0.452. Using only fecal microbiome data we 199 achieved an AUC of 0.838 with a specificity of 0.766 and a sensitivity of 0.806. When combining clinical 200 metadata with the microbiome AUC of 0.844 with a specificity of 0.831 and a sensitivity of 0.774. 201

Across all weeks and responses, prediction with clinical metadata alone did not perform as well as models using the fecal microbiome at screening. Also, combining microbiome data with clinical metadata did not consistently improve prediction compared to microbiome data alone. Additionally we found several OTUs occurred frequently across models including *Faecalibacterium*, among other taxa that were more abundant in responders/remitters. Their abundances can be seen in figure 4.

207 Discussion

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With this study we sought to gain a more detailed understanding of if and how biologic treatment affects
the microbiome, to determine whether the microbiome can be used to identify patients who will respond
to therapy, and to gain a better understanding of the interaction between the human gut microbiome
and CD pathogenesis in adult patients. We found the fecal microbiome to be useful in uncovering
associations between the microbiome and aspects of CD severity metrics and treatment outcomes. We
also demonstrated that the microbiome of treated responders changed over time, though it is not yet

possible to determine any direct effects of treatment on the microbiome. Finally, we were able to show that the microbiome could be useful in predicting response to therapy, especially clinical remission, compared to clinical metadata alone in our unique patient cohort.

We observed several associations between the microbiome and clinical variables that could play a role 217 in how CD is monitored and treated in the future. Given that serum CRP, fecal calprotectin, and fecal 218 lactoferrin are used as biomarkers to measure intestinal inflammation and CD severity, the observation 219 that the microbial community structure is different among patients based on these markers supports the hypothesis that the microbiome could function as a biomarker for measuring disease activity in 221 patients, especially in concert with these established inflammatory biomarkers (23, 26, 27). Higher 222 CDAI was associated with lower microbial diversity. This appears to be consistent with other studies 223 on the microbiome in individuals with CD compared to healthy individuals and studies looking at active 224 disease compared to remission (19, 23, 28). However, these differences may have been driven by weekly 225 stool frequency, one component of the CDAI, where higher stool frequency is also negatively associated 226 with microbial diversity. This finding is consistent with the association between loose stools and lower 227 diversity (29). We also observed differences in the microbial community structure based on disease 228 localization. These results are consistent with a study by Naftali et al finding distinct microbiotas 229 for ileal versus colonic CD using mucosal tissue (30). Our study also found that corticosteroid use 230 impacts the composition of the human fecal microbiome. This supports data seen in the mouse model 231 where corticosteroid injections altered the fecal mouse microbiome (31). As corticosteroid use appears 232 to impact diversity, corticosteroid therapy may be useful when trying to positively impact microbial 233 diversity during biologic therapy and thereby increase the possibility of response to CD therapies. We 234 also observed that longer disease duration is associated with a reduction in fecal microbial diversity. 235 This decreased diversity may be due to the long duration of inflammatory conditions in the gut. This 236 observation and the increased likelyhood of remission and mucosal healing in individuals treated with 237 biologics earlier in the course of their disease is an argument for earlier biologic intervention (32-34). 238 Hypothetically, earlier biologic intervention could 'preserve' a more diverse microbiome that promotes 239 remission and reduces the likelihood of relapse. However, the cost of biologics for patients is hindrance to early biologic intervention. Using aptamers in place of monoclonal antibodies may reduce this cost and 241 make earlier intervention possible. Aptamers are short strands of DNA or RNA capable of specifically 242

binding small molecules, proteins, and whole cells. Anti-TNF aptamers have been published that could potentially be used to test this in the mouse model (35).

An important question for the microbiome and IBD is whether or not the microbiome is affected by 245 treatment with biologics. This study attempted to answer that question by looking at the microbiome of 246 our CD subjects across multiple time points during treatment. While we were unable to see direct effects 247 of the drug on the fecal microbiome, we observed that the microbiome of clinical responders changed 248 over time, in contrast to nonresponsive subjects. This was observed for responsive patients regardless of induction treatment, leading us to think we are observing the effects of change in disease activity 250 and health, leading to lower inflammation, rather than any effects from treatment. This interpretation 251 is consistent with studies using the microbiome to distinguish between remission and active CD (23). 252 We did however observe a significant difference in community structure based on treatment and cannot 253 eliminate the possibility of a direct effect on the microbiome in treated responders, however the change 254 in community structure observe in responders treated with placebo supports the hypothesis that the 255 change in community structure reflects a change in inflammation. 256

Another important question in for the importance of the microbiome in IBD is whether response to 257 therapy can be predicted with the microbiome. We attempted to address this by developing a random-258 forest model that used relative microbial abundance data and/or clinical metadata for input. We 259 found we were better able to predict remission status compared to response status. Response may be less predictable due to the "floating target" nature of a relative decrease (>30% decrease) in CDAI 261 compared to the hard threshold for remission (CDAI<150). We were also better able to distinguish 262 remission/non-remission than response/non-response, using samples provided 6 weeks after treatment 263 induction. This is consistent with other studies, again suggesting the microbiome could be useful as a 264 biomarker in detecting remission versus active disease (23). 265

While using the presented model may not be useful clinically to predict response to therapy at this time,
it is useful for hypothesis generation about the biology of CD as it relates to the microbiome. Some of
the frequently occurring factors in our predictive models have already been linked to CD pathogenesis.
As far as clinical biomarkers, fecal lactoferrin and fecal calprotectin occurred in the majority of models
where clinical metadata was combined with the microbiome, supporting their importance as biomarkers
for CD activity, especially in relation to the fecal microbiome (26, 27). Faecalibacterium was the most

frequently occurring OTU in our models. It is associated with health, comprising up to 5% of the relative 272 abundance in healthy individuals (source) and has been shown to be low in CD patients (14, 17, 30, 36). 273 Remission was much more likely in individuals who had measurable Faecalibacterium present at baseline. 274 This supports the hypothesis that Faecalibacterium impacts CD pathogenesis. Escherichia/Shigella also 275 occurred frequently in our models. This OTU is associated with inflammation and has been shown to 276 negatively impact CD pathogenesis (36). Fusobacterium also appeared in our predictive models and 277 is associated with CD and CRC, something CD patients are more likely to develop than individuals 278 without IBD (36). Many other taxa observed in our analysis had low abundance, but in many cases 279 these taxa are related and may serve similar ecologic and metabolic roles in teh gut envronment. We 280 hypothesize that these microbe may have genes that perform similar metabolic functions that could 281 be revealed by perfroming metagenomics on the week 0 stool samples in future studies, especially in 282 subjects who achieved remission. These observations and the positive/negative associations of these 283 microbes and CD also allow us to hypothesize on ways to alter the microbiome to increase the likelihood 284 therapeutic response. Prior to the initiation of therapy, patients could get a fecal microbiome analysis. 285 The community data could then be used to direct the patient to undergo a round of antibiotics to 286 target and reduce the levels of Escherichia in the patient's gut. Alternatively, the microbes found to 287 be positively associated with response could be formulated into a daily probiotic patients could take 288 while receiving therapy with the goal of increasing the likelihood of remission and mucosal healing. 289 Additionally, altering the weighting or binning of important factors in the model could make prediction 290 of response or remission more reliable. This could eventually allow for pre-screening of patients with 291 stool samples to predict successful treatment or better direct treatment. If the fecal microbiome can 292 be used as a theraprognostic tool to non-invasively predict response to specific treatment modalities 293 or inform treatment, then more personalized treatment could result in faster achievement of remission, 294 thereby increasing patients' quality of life and reducing economic and healthcare impacts. 295

296 Methods

297 Study Design and Sample Collection

Janssen Research and Development conducted a phase II clinical study of approximately 500 patients to 298 assess the safety and efficacy of UST for treating anti-TNF- α refractory CD patients (10). Participants 299 provided a stool sample prior to the initiation of the study and were then divided into 4 groups of 300 125 individuals receiving placebo or 1, 3, or 6 mg/kg doses of UST by IV. Additional stool samples 301 were provided at week 4. At week 6 an additional stool sample was collected, patients were scored for 302 their response to UST based on CD Activity Index (CDAI), and divided into groups receiving either 303 subcutaneous injection of UST or placebo at weeks 8 and 16 as maintenance therapy. Finally, at 22 weeks patients provided an additional stool sample and were then scored using CDAI for their response 305 to therapy. Frozen fecal samples were shipped to the University of Michigan and stored at -80°C prior 306 to DNA extraction

308 DNA extraction and 16S rRNA gene sequencing

Microbial genomic DNA was extracted using the PowerSoil-htp 96 Well Soil DNA Isolation Kit (MoBio 309 Laboratories) using an EPMotion 5075 pipetting system, as previously described (24, 37). The V4 region 310 of the 16S rRNA gene from each sample was amplified and sequenced using the Illumina MiSeq Personal 311 Sequencing platform as described elsewhere (27). Sequences were curated as described previously using 312 the mothur software package (38). Briefly, we reduced sequencing and PCR errors, aligned the resulting 313 sequences to the SILVA 16S rRNA sequence database, and removed any chimeric sequences flagged 314 by UCHIME (39). Sequences were clustered into operational taxonomic units (OTU), as previously 315 described (40). Briefly, OTUs were clustered at a 97% similarity cutoff and the relative abundance 316 was calculated for OTUs in each sample. All sequences were classified using a naive Bayesian classifier 317 trained against the RDP training set (version 11) and OTUs were assigned a classification based on 318 which taxonomy had the majority consensus of sequences within a given OTU (41). All fastq files and 319 the MIMARKS spreadsheet with de-identified clinical metadata are available at TBD.

Gut microbiome biomarker discovery analysis

Mothur as well as the R software package were used for our data analysis. Alpha diversity metrics 322 (e.g. Shannon, Inverse Simpson) were calculated for each sample in the dataset, and compared using non-323 parametric statistical tests (i.e Kruskal-Wallace and Wilcox Test) (42, 43). Beta diversity was calculated 324 the distance between samples using the theta YC metric, which takes into account the types of bacteria 325 and their abundance to calculate the differences between the communities (44). These distance matrices 326 were visualized by generating non-metric dimensional scaling (NMDS) plots of the distances. Overlap 327 between sets of communities was assessed using the non-parametric analysis of molecular variance 328 (AMOVA) and homogeneity of variance (HOMOVA) tests (22, 45). Differentially abundant OTUs 329 were selected using the biomarker discovery algorithm, LEfSe [linear discriminant analysis (LDA) effect 330 size for each pairwise comparison of clinical groups (46). In short, This method uses the Wilcox non-331 parametric test to identify OTUs where there is a P-value less than 0.05 and then applies a LDA step to 332 identify the effect sizes that are the most meaningful (i.e. greater than 2.0). We also used the relative abundance of each OTU across the samples and clinical metadata as input to the AUC-Random forest 334 package available to identify phylotypes/clinical variables that would allow us to distinguish between 335 various treatment and response groups (47).

Statistical analysis

Tables

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Table 1: Summary of clinical metadata of chort at baseline

Clinical Variable	Treated	Placebo	Total
	n = 232	n = 74	n = 306
Age (years)	38 ± 13	40 ± 14	39 ± 13
Sex (% Male)	36.6	43.2	38.2
Race (% Caucasian)	91.8	93.2	92.2
Corticosteroid Use (%)	40.1	52.7	43.1
BMI (kg/m^2)	26 ± 6.7	25 ± 4.9	25 ± 6.3
Disease Duration (years)	12 ± 8.4	13 ± 10	12 ± 8.8
CDAI	330 ± 62	310 ± 69	320 ± 64
Bowel Stricture (%)	12.5	10.8	12.1
Tissue Involvement (%) Colon/Ileocolic/Ileal	28.9/51.3/19.8	24.3/39.2/36.5	27.8/48.4/23.9

341 Supplemental_Table 1: Diversity differences based on clinical metadata of chort at baseline

Clinical Variable	Correlation	Species Diveristy (p-value)	Community Structure (p-value)
CDAI	rho = -0.2	0.014	0.324
Loose Stool Frequency (per week)	rho = -0.2	0.003	0.024
C-Reactive Protein (mg/L serum)	rho = 0.06	0.394	0.033
Fecal Calprotectin (µg/g)	rho = 0.08	0.254	0.006
Fecal Lactoferrin (µg/g)	rho = 0.1	0.07	0.004
ВМІ	rho = 0.07	0.299	0.277
Weight (kg)	rho = 0.07	0.299	0.112
Age (years)	rho = -0.05	0.472	0.033
Sex (F/M)	-	0.539	0.277
Corticosteroid Use (Y/N)	-	0.001	0.01
Disease Duration (years)	rho = -0.2	0.001	0.004
Tissue Involvement	-	0.19	0.004

Table 2: Diversity differenced bases on Response/Remission in treated subjects.

Clinical Variable	Species Diversity (p-value)	Community Structure (p-value)		
Week 6 Response (No, Yes)	0.440	0.012		
Week 6 Remission (No, Yes)	0.020	0.017		
Week 22 Response (No, Yes)	0.900	0.012		
Week 22 Remission (No, Yes)	0.440	0.012		

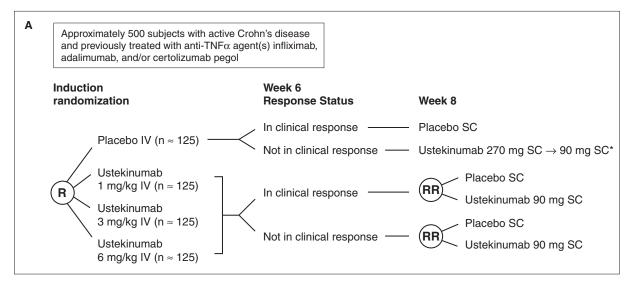
345 Supplemental Table 2: adonis

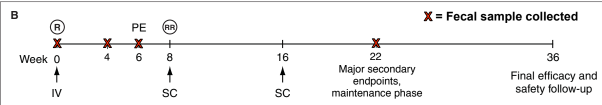
	Df	SumsOfSqs	MeanSqs	F.Model	R2	Pr(>F)
visit	2	0.52	0.26	0.66	0.0022	0.089
TRTGR	1	0.64	0.64	1.6	0.0026	0.0025
RelRSPwk22	1	1.1	1.1	2.7	0.0045	0.0025
visit:TRTGR	2	0.31	0.16	0.4	0.0013	0.87
visit:ReIRSPwk22	2	0.8	0.4	1	0.0033	0.001
TRTGR:ReIRSPwk22	1	0.74	0.74	1.9	0.0031	0.0025
visit:TRTGR:ReIRSPwk22	2	0.59	0.3	0.75	0.0025	0.044
Residuals	600	240	0.4	NA	0.98	NA
Total	610	240	NA	NA	1	NA

347 Figures

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Figure 1: Experimental design.



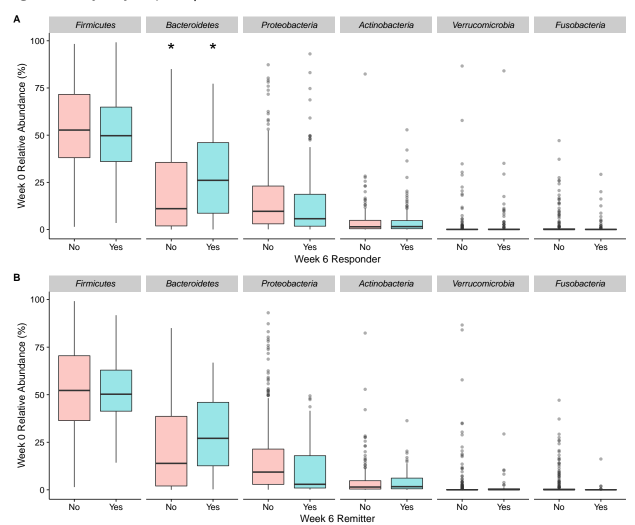


IV = Intravenous; SC = Subcutaneous; \uparrow = Study agent administration

PE = Primary Endpoint; R = Randomization; RR = Rerandomization only for subjects receiving ustekinumab induction therapy

^{*} Subjects receiving placebo at Week 0 who are not in clinical response at Week 6 will receive ustekinumab 270 mg SC and 90 mg SC at Weeks 8 and 16, respectively.

Figure 2: Phyla by response/remisison week 6 and OTU abundance week 6 remission



SF 1: Phyla by response/remission week 22

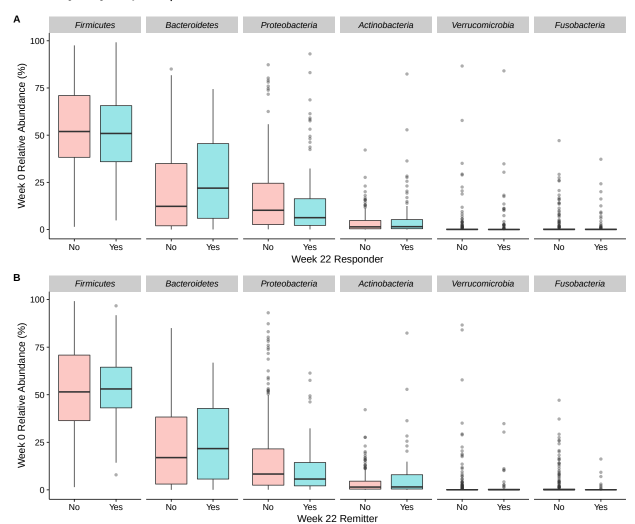


Figure 3: OTUS at screening

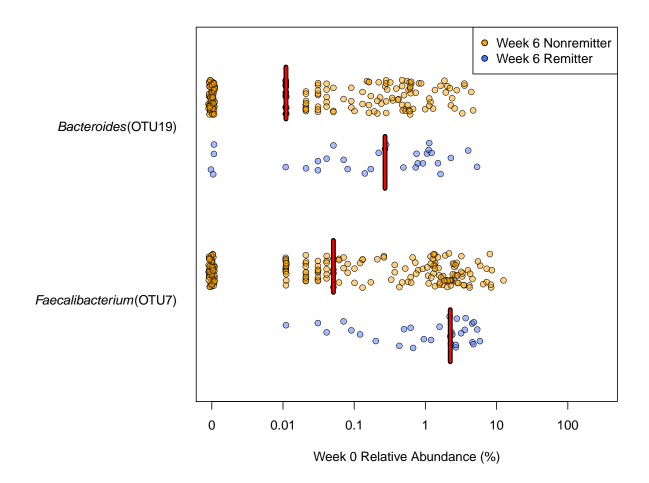


Figure 4: Change in alpha diversity over time

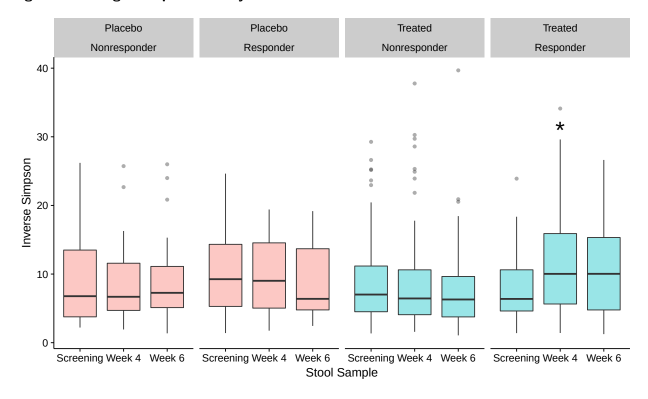


Figure 5: Determine week 6 status by week 6 stool and impt OTUs and abunds

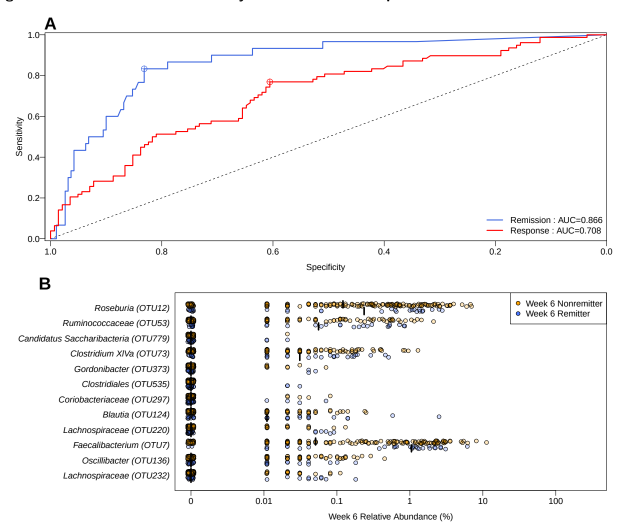
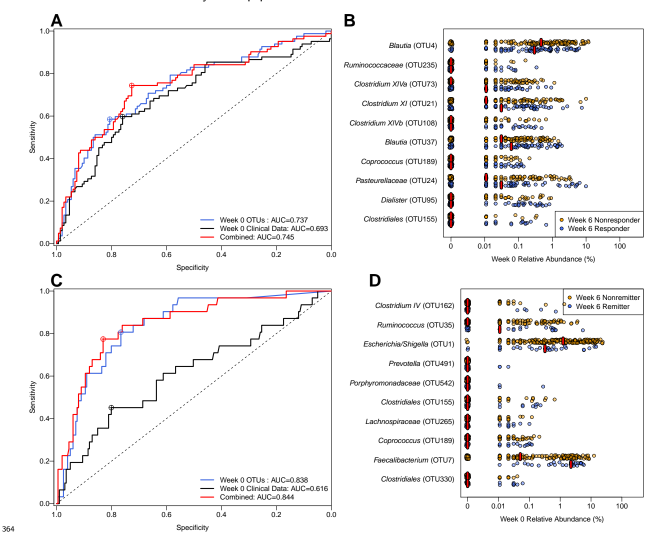
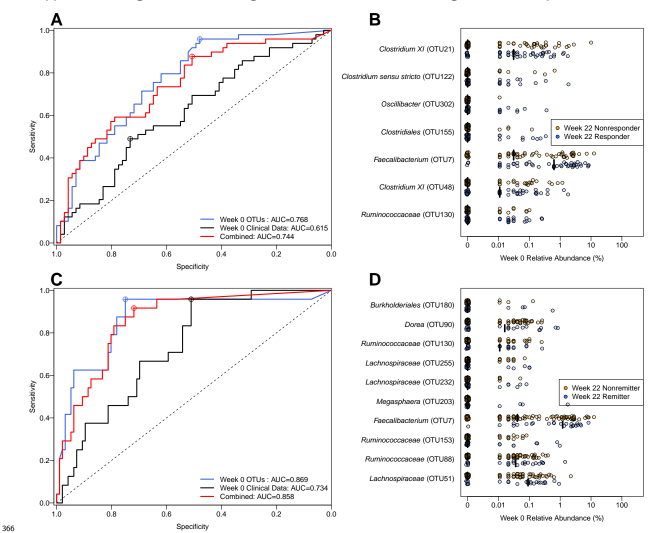


Figure 6: Prediction of RESPONSE/REMISSION in treated subjects using all clinical metadata,
baseline microbiome alone, and combined A. Response ROCs B. Response Model Performance
vs. reality C. Top predictive taxa and abundance based on response D. REMISSION ROCs E. REMISSION
Model Performance vs. reality F. Top predictive taxa and abundance based on remission



Supplemental Figure 2: Predicting Week 22 disease status using Week 0 samples



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