Adaptive ecological processes and metabolic independence drive microbial colonization and resilience in the human gut

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Running Title
Drivers of human gut colonization

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Abstract

A detailed understanding of gut microbial ecology is essential to engineer effective microbial therapeutics and to model microbial community assembly and succession in health and disease. However, establishing generalizable insights into the functional determinants of microbial fitness in the human gut has been a formidable challenge. Here we employ fecal microbiota transplantation (FMT) as an in natura experimental model to identify determinants of microbial colonization and resilience. Our long-term sampling strategy and high-resolution multi-omics analyses of FMT donors and recipients reveal adaptive ecological processes as the main driver of microbial colonization outcomes after FMT. We also show that high-fitness donor microbial populations are significantly enriched in metabolic pathways that are responsible for the biosynthesis of nucleotides, essential amino acids, and micronutrients, independent of taxonomy. To determine whether such metabolic competence can explain the microbial ecology of human disease states, we analyzed genomes reconstructed from healthy humans and humans with inflammatory bowel disease (IBD). Our data reveal that such traits are also significantly enriched in microbial genomes recovered from IBD patients, linking presence of superior metabolic competence in bacteria to their expansion in IBD. Overall, these findings suggest that the transfer of gut microbes from a healthy donor to a disrupted recipient environment initiates an environmental filter that selects for populations that can self-sustain. Such ecological processes that select for self-sustenance under stress offer a model to explain why common yet typically rare members of healthy gut environments can become dominant in inflammatory conditions without any need for them to be causally associated with, or contribute to, such disease states.
Introduction

The human gut microbiome is associated with a wide range of diseases and disorders (Almeida et al. 2020; Durack and Lynch 2019; Lynch and Pedersen 2016). However, mechanistic underpinnings of these associations have been difficult to resolve in part due to the diversity of human lifestyles (David et al. 2014) and the limited utility of model systems to make robust causal inferences for microbially mediated human diseases (Walter et al. 2020).

Inflammatory bowel disease (IBD), a group of increasingly common intestinal disorders that cause inflammation of the gastrointestinal tract (Baumgart and Carding 2007), has been a model to study human diseases associated with the gut microbiota (Schirmer et al. 2019). The pathogenesis of IBD is attributed in part to the gut microbiome (Plichta et al. 2019), yet the microbial ecology of IBD-associated dysbiosis remains a puzzle. Despite marked changes in gut microbial community composition in IBD (Ott et al. 2004; Sokol and Seksik 2010; Joossens et al. 2011), the microbiota associated with the disease lacks traditional pathogens (Chow, Tang, and Mazmanian 2011), and microbes that are found in IBD typically also occur in healthy individuals (Clooney et al. 2021), which complicates the search for robust functional or taxonomic markers of health and disease states (Lloyd-Price et al. 2019). One of the hallmarks of IBD is reduced microbial diversity during episodes of inflammation, when the gut environment is often dominated by microbes that typically occur in lower abundances prior to inflammation (Vineis et al. 2016). The sudden increase in the relative abundance of microbes that are common to healthy individuals suggests that the harsh conditions of IBD likely act as an ecological filter that prevents the persistence of low-fitness populations. Yet, in the absence of a complete understanding of the functional drivers of microbial colonization in this habitat, critical insights into the metabolic requirements of survival in IBD remains elusive.

Understanding the determinants of microbial colonization has been one of the fundamental aims of gut microbial ecology (Costello et al. 2012; Messer et al. 2017). To overcome the difficulties of conducting well-controlled studies with humans, researchers have studied the determinants of microbial colonization of the gut in model systems, such
as germ-free mice conventionalized with individual taxa (S. M. Lee et al. 2013) or a consortium of human microbial isolates (Feng et al. 2020). Despite their utility for hypothesis testing, simpler models do not capture the complex ecological interactions fostered by natural systems and thus the insights they yield do not always translate to human gut microbial ecology (Ley et al. 2006; Finucane et al. 2014). Between the extremes of well-controlled but simple mouse models and complex yet uncontrolled human populations, there exists a middleground that provides a window into the microbial ecology of complex human systems through a controlled perturbation: human fecal microbiota transplantation (FMT), the transfer of stool from a donor into a recipient’s gastrointestinal tract (Eiseman et al. 1958).

FMT complements laboratory models of environmental perturbation by colliding two distinct microbial ecosystems, and thus offers a powerful framework to study fundamental questions of microbial ecology, including the determinants of microbial succession and resilience (Schmidt, Raes, and Bork 2018). Here we use FMT as an in natura experimental model to investigate the ecological and functional determinants of successful microbial colonization of the human gut at the level of individual populations. Our findings suggest that adaptive ecological forces are key drivers of colonization outcomes after FMT, reveal taxonomy-independent metabolic determinants of fitness in the human gut, and demonstrate that similar ecological principles determine resilience of microbes in stressful and inflammatory conditions.

Results and Discussion

Our study includes 109 gut metagenomes (Supplementary Table 1) from two healthy FMT donors (A and B) and 10 FMT recipients (five recipients per donor) who had multiply recurrent Clostridium difficile infection (CDI) and received vancomycin for a minimum of 10 days to attain resolution of diarrheal illness prior to FMT. On the last day of vancomycin treatment, a baseline fecal sample was collected from each recipient, and their bowel contents were evacuated immediately prior to FMT. Recipients did not take any antibiotics on the day of transplant, or during the post-FMT sampling period (Supplementary Figure 1). We also collected 24 Donor A samples over a period of 636 days and 15 Donor B
samples over a period of 532 days to establish an understanding of the long-term microbial population dynamics within each donor microbiota. We also collected 5 to 9 samples from each recipient up to 336 days post-FMT. Deep sequencing of donor and recipient metagenomes using Illumina paired-end (2x150) technology resulted in a total of 7.7 billion sequences with an average of 71 million reads per metagenome (Figure 1, Supplementary Table 1, Supplementary Table 2). We employed genome-resolved metagenomics, pangenomics, and microbial population genetics for an in-depth characterization of donor and recipient gut microbiota using these data, and we leveraged publicly available gut metagenomes to benchmark our observations.

Many but not all donor microbes colonized recipients and persisted long-term

We first characterized the taxonomic composition of each donor and recipient sample by aligning metagenomic short reads to reference genomes in the NCBI’s RefSeq database (Supplementary Table 2). The phylum-level microbial community composition of both donors reflected those observed in healthy individuals in North America (Human Microbiome Project Consortium 2012): a large representation of Firmicutes and Bacteroidetes, and other taxa with relatively lower relative abundances, including Actinobacteria, Verrucomicrobia, and Proteobacteria (Figure 1, Supplementary Table 2). In contrast, the vast majority of the recipient pre-FMT samples were dominated by Proteobacteria, a phylum that typically undergoes a drastic expansion in individuals treated with vancomycin (Isaac et al. 2017). After the FMT, we observed a dramatic shift in recipient taxonomic profiles (Supplementary Table 2, Supplementary Figure 2), a widely documented hallmark of this procedure (Khoruts et al. 2010; Grehan et al. 2010; Shahinas et al. 2012). Nearly all recipient samples post-FMT were dominated by Bacteroidetes and Firmicutes as well as Actinobacteria and Verrucomicrobia in lower abundances, resembling qualitatively, but not quantitatively, the taxonomic profiles of their donors (Supplementary Table 2). For example, even though the median relative abundance of Bacteroidetes populations were 5% and 17% in donors A and B, their relative abundance in recipients post-FMT increased to 33% and 45%, respectively
(Figure 1, Supplementary Table 2). A single genus, *Bacteroides*, made up 76% and 82% of the Bacteroidetes populations in the recipients of Donor A and B, respectively (Supplementary Table 2). The success of the donor *Bacteroides* populations in recipients upon FMT is not surprising given the ubiquity of this genus across human populations throughout the globe (Wexler and Goodman 2017) and the ability of its members to survive substantial levels of stress (Swidsinski et al. 2005; Vineis et al. 2016). This result suggests that FMT outcomes in our dataset are unlikely random, and the study design and resulting dataset offers a framework to study ecological principles of the human gut microbiome.

Next, we assembled short metagenomic reads into contiguous segments of DNA (contigs). Co-assemblies of 24 Donor A and 15 Donor B metagenomes independently resulted in 53,891 and 54,311 contigs that were longer than 2,500 nucleotides, and described 0.70 and 0.79 million genes occurring in 179 and 248 genomes, as estimated by the mode of the frequency of bacterial single-copy core genes (Supplementary Table 2). One way to characterize how well a given assembly describes the DNA content of a given metagenome is to calculate the percentage of reads it recruits from the metagenome through read mapping. Donor contigs recruited on average 80.8% of metagenomic reads from donor metagenomes. In contrast, they recruited 43.4% of reads on average from pre-FMT recipient metagenomes. This number increased to 80.2% for recipient metagenomes post-FMT (Figure 1), and the donor contigs continued to represent 76.8% of the recipient metagenomes on average even after a year post-FMT (Supplementary Table 2). These read recruitment results suggest that members of the donor microbiota successfully established in recipient guts upon FMT and largely persisted until the end of the sampling period.

Compared to metagenomic short reads, assembled contigs provide a larger genetic context to study microbial metagenomes. However, a sole focus on contigs may yield misleading results (Kowarsky et al. 2017) that can be ameliorated by reconstructing microbial genomes from metagenomic assemblies (Chen et al. 2020). We reconstructed genomes from co-assembled donor metagenomes by grouping contigs into putative bins based on sequence composition and differential coverage signal as previously described.
We retained bins that were at least 70% complete and had no more than 10% redundancy as predicted by bacterial single-copy core genes (Bowers et al. 2017; Chen et al. 2020) and manually refined them to improve their quality following previously described approaches (Delmont et al. 2018; Shaiber et al. 2020). Our binning resulted in a final list of 128 metagenome-assembled genomes (MAGs) for Donor A and 183 MAGs for Donor B that included members of Firmicutes (n=265), Bacteroidetes (n=20), Actinobacteria (n=14), Proteobacteria (n=7), Verrucomicrobia (n=2), Cyanobacteria (n=2), and Patescibacteria (n=1) (Supplementary Table 3). The taxonomy of donor-derived genomes largely reflected the taxonomic composition of donor metagenomes as predicted by short reads (Figure 1, Supplementary Table 2, Supplementary Table 3). While only 20 genomes (mostly of Bacteroides and Alistipes) explained the entirety of the Bacteroidetes group, we recovered 265 MAGs that represented lower abundance but diverse populations of Firmicutes (Figure 1, Supplementary Table 2, Supplementary Table 3). We found no difference between the delivery method of FMT for the recipients of donor A, where, on average 45% and 43% of donor genomes emerged in recipients who received donor stool through colonoscopy (n=3) versus pills (n=2), respectively. However, there was an increase in the efficiency of pills for donor B, where on average 25% and 54% of donor genomes emerged in recipients who received donor stool through colonoscopy (n=2) versus pill (n=3) (Supplementary Figure 3).

Reconstructing genomes gave us access to microbial populations in metagenomes through metagenomic read recruitment strategies and enabled us to characterize (1) population-level microbial colonization dynamics before and after FMT using donor and recipient metagenomes and (2) the distribution of each donor population across geographically distributed humans using 1,984 publicly available human gut metagenomes (Supplementary Table 4). As expected, we detected each donor population in at least one donor metagenome (see Methods for ‘detection’ criteria). Yet, only 16% of Donor A populations were detected in every Donor A sample, and only 44% of Donor B MAGs were detected in every Donor B sample (Figure 1, Supplementary Table 3), in agreement with the previously documented dynamism of gut microbial community composition over time (David et al. 2014). A marked increase in the detection of donor
populations in recipients after FMT echoed the general pattern of transfer suggested by the short-read taxonomy (Figure 1): while only 38% of Donor A and 54% of Donor B populations were detected in at least one recipient pre-FMT, these percentages increased to 96% and 96% post-FMT (Supplementary Table 3). Not every donor population colonized each recipient, but colonization events did not appear to be random: while some donor populations colonized all recipients, others colonized none (Figure 1), providing us with an opportunity to resolve colonization events and quantify colonization success for each donor population in our dataset.
Resolving colonization events accurately is a challenging task as multiple factors may influence the ability to determine colonization outcomes unambiguously. These factors include (1) the inability to detect low-abundance populations, (2) inaccurate characterization of transient populations observed immediately after FMT as successful colonization events, (3) the reliance on relative abundance of populations to define colonization events when abundance estimates from stool do not always reflect the abundance of organisms in the GI tract (Yasuda et al. 2015; Sheth et al. 2019), and (4) the failure to distinguish between colonization by a donor population or emergence of a pre-FMT recipient population after FMT (where a low-abundance recipient population that is closely related to one or more donor populations becomes abundant after FMT and is mistaken as a bona fide colonization event). To mitigate these factors, we have (1) employed deep-sequencing of our metagenomes which averaged 71 million reads per sample, (2) implemented a longitudinal sampling strategy, that spanned 376 days on average, to observe donor populations in our recipients long after the FMT, (3) leveraged a ‘detection’ metric to define colonization events by presence/absence of populations rather than abundance, and (4) employed microbial population genetics to identify and resolve origins of subpopulations. We also developed an analytical approach (Supplementary Figure 4) to determine whether a given donor population has colonized a given recipient based on the detection of donor subpopulations in the transplant sample, in the recipient pre-FMT, and in the recipient post-FMT (see Materials and Methods,
To determine colonization outcomes, we analyzed 640 genome/recipient pairs for Donor A (128 donor genomes in 5 recipients) and identified 99 successful colonization events, 38 failed colonization events, and 503 ambiguous colonization events (Supplementary Table 6). For Donor B, we analyzed 915 genome/recipient pairs (183 donor genomes in 5 recipients) and identified 106 successful colonization events, 109 failed colonization events, and 700 ambiguous colonization events (Supplementary Table 6). Our stringent criteria (see Materials and Methods, Supplementary Figure 4) classified the vast majority of all genome/recipient pairs as ambiguous colonization events. Nevertheless, due to the relatively large number of donor MAGs and FMT recipients in our study, we were left with 352 MAG/recipient pairs with unambiguous phenotypes for downstream analyses.

Adaptive ecological forces are the primary drivers of microbial colonization

The ability of a microbial population to colonize and persist in a complex ecosystem is influenced by both neutral and adaptive forces (Maignien et al. 2014). Although which of these is the major driver of successful colonization of the human gut remains unclear (Smillie et al. 2018). In the context of FMT, previous studies have suggested neutral processes to determine colonization success based on the abundance of a microbial population in a donor stool sample (Smillie et al. 2018; Podlesny and Florian Fricke 2020). Indeed, ecological drift may have a significant role in a system dominated by neutral processes, where low-abundance donor populations in the transplant would be less likely to be observed in recipients. In contrast, if the system is dominated by adaptive forces, colonization success would be a function of the population fitness in the recipient environment, rather than its abundance in the transplant.

To investigate the impact of neutral versus adaptive processes on colonization in our dataset we first asked whether the prevalence of a donor population in healthy human gut metagenomes, which we define here as a measure of its fitness, was associated with the detection of the same population in donor or recipient metagenomes. Within both FMT cohorts, the mean detection of each population in recipients post-FMT had a stronger
association with population fitness than mean detection in donor samples (Figure 2a).

The fitness of donor A populations explained 4.2% of the variation in mean detection of those populations in donor samples ($R^2=0.042$, $p=0.021$) and 19% of variation in mean detection in recipient post-FMT samples ($R^2=0.19$, $p=2.7e-07$), an increase of approximately 4.5-fold (Figure 2a). Similarly, Donor B population fitness explained 7.3% of the variation in mean detection in donor samples ($R^2=0.073$, $p=2.1e-04$), and 36% of the variation in mean detection in recipient post-FMT samples ($R^2=0.36$, $p=4.5e-19$), an increase of approximately 5-fold (Figure 2a). This suggests that fitness is a better predictor of colonization outcome than it is of the detection of a population in the donor, suggesting that adaptive forces are likely at play. But detecting a donor population in a recipient post-FMT metagenome through metagenomic read recruitment does not prove colonization, since donor genomes can recruit reads from recipient populations that are closely related (i.e., strain variants) and that were low abundance prior to FMT. Single-nucleotide variants in read recruitment results, however, can reveal such cases (Denef 2019) and quantify their dynamics (Quince et al. 2017). Thus, we developed an improved model that took into consideration the presence and absence of distinct subpopulations in our data and their origins (Supplementary Figure 4). We then used this model to test if colonization success was correlated with population fitness or population dose, which we define here as the relative abundance of a given population in the transplanted donor stool sample. For Donor A populations, colonization outcome was significantly correlated with both dose (Wald test, AUC=0.73, $p=7.7e-05$) and fitness (Wald test, AUC=0.76, $p=6.3e-06$) (Figure 2b,c). But combining both measures as predictive variables did not substantially improve the performance of our colonization model (AUC=0.82) (Figure 2c).

This was likely due to the small, but significant, correlation between dose and fitness in Donor A MAGs ($R^2=0.053$, $p=0.0070$) (Figure 2d). When the fitness of a microbial population is reflected in its relative abundance, the effect of fitness on colonization outcome may be masked by an apparent dose effect. In contrast to Donor A, the fitness of Donor B populations and their relative abundance in Donor B samples were not correlated ($R^2=0.0012$, $p=0.61$) (Figure 2d), providing us with an ideal case to analyze these two factors independently. Indeed, there was no correlation between dose of a microbial population in Donor B transplant samples and colonization outcome in
recipients post-FMT (Wald test, AUC=0.56, p=0.09). Instead, we found a significant
correlation between the fitness of each population and the colonization outcome (Wald
test, AUC=0.70, p=9.0e-07) (Figure 2c).

Taken together, our findings suggest that fitness of a microbial population as measured
by its prevalence across global gut metagenomes can predict its colonization success
better than its abundance in the donor stool sample, giving credence to the role of
adaptive rather than neutral ecological processes in colonization. This finding contrasts
with previous studies which suggested that the abundance of a given population in the
donor sample was an important determinant of colonization (Smillie et al. 2018; Podlesny
and Florian Fricke 2020). However, these analyses included many recipient samples
collected less than one week after FMT and it is likely that their observations were
influenced by the presence of transient populations. Indeed, samples collected
immediately after FMT are more likely to inflate the number of colonization events,
whereas longitudinal sampling over a longer time course can distinguish transient
populations from those that successfully colonized the recipients. We cannot definitively
test this hypothesis as we sampled most of our recipients a week after FMT. Still, on
average 12% of the donor populations detected in our recipients a week after FMT were
no longer detected after a month (Figure 1, Supplementary Table 3). Overall, our stringent
criteria to determine colonization outcome and the extended post-FMT sampling period
likely enabled us to study the long-term engraftment of successful and potentially low-
abundance colonizers, instead of high-abundance transient populations that may be
dominant directly after FMT.

Figure 2. Relationships between dose, prevalence, and colonization outcome. Left: Donor A. Right: Donor B. a)
Linear regression models of mean detection of each MAG in either donor or recipient post-FMT samples as a function
of prevalence. b) Colonization outcome of MAG/recipient pairs as a function of MAG dose or MAG prevalence.
Significance calculated by Wald test. c) Receiver operator curves (ROCs) for logistic regression models of colonization.
d) Linear regression models of dose as a function of prevalence.
Accurately distinguishing the role of dose versus fitness in colonization success is further compounded by the fact that microbial populations that are prevalent across human populations may also tend to be more abundant. This is well illustrated by Donor A. Fortunately, the abundant populations in Donor B did not reflect prevalent microbes in healthy adult guts, which demonstrated the importance of fitness as a determinant of colonization success compared to dose without the confounding effect of a correlation between fitness and dose. Thus, it is a theoretical possibility that colonization success is purely driven by adaptive forces and is not influenced by dose, at all. However, while our data assign a larger role to adaptive forces with confidence, a more accurate determination of the proportional influence of adaptive versus neutral processes in colonization requires a much larger dataset.

Colonizer and resilient microbes are enriched in metabolic pathways for the biosynthesis of essential organic compounds

Fitness in a specific environment is conferred to an organism by a combination of functional traits. In the human gut, such traits drive microbial community succession and structure as a response to changing host diet and lifestyle (Koenig et al. 2011; Rothschild et al. 2018). Behind successful colonization and resilience after perturbation are likely similar functional traits that promote fitness. Building on our observation that suggests a primary role of adaptive ecological processes in colonization outcome, we next sought to identify genetic determinants of colonization. For this, we leveraged our access to donor microbial population genomes and global metagenomes to investigate whether a functional enrichment analysis could reveal predictors of success independent of taxonomy.

To generate metabolic insights into colonization success we divided our donor populations into ‘high-fitness’ and ‘low-fitness’ groups by considering both their prevalence in FMT recipients and prevalence across global gut metagenomes (Materials and Methods). The ‘high-fitness’ group included the microbial populations that colonized or persisted in all FMT recipients and were the most prevalent in gut metagenomes from Canada. We assumed that they represented a set of highly fit microbial populations as
they were able to colonize human gut environments systematically, (2) they persisted in these environments long-term regardless of the host genetics or lifestyle, and (3) they were prevalent in gut metagenomes outside of our study. In comparison, the ‘low-fitness’ group comprised microbial populations that failed to colonize or persist in at least three FMT recipients. These populations were nevertheless viable gut microbes as not only our long-term sampling of the donors systematically identified them but also, they sporadically colonized some FMT recipients. Yet, unlike those in the high-fitness group, the distribution patterns of low-fitness populations were sparse, not only within our cohort, but also within publicly available metagenomes. In fact, low-fitness genomes were on average less prevalent than high-fitness genomes in each of the 17 different countries we queried (Supplementary Table 3). Overall, we conservatively categorized 20 populations in each group for downstream analyses (Supplementary Table 7). All populations in the low-fitness group resolved to Firmicutes. The high-fitness group was also dominated by Firmicutes (15 of 20) but it also included five Bacteroidetes and one Actinobacteria (Supplementary Table 7). Genome completion estimates did not differ between high and low-fitness groups (Wilcoxon rank sum test, p=0.42) and averaged to 91% and 93%, respectively. However, genome sizes between the two groups differed dramatically (p=2.9e-06), where high-fitness group genomes averaged to 2.8 Mbp while low-fitness group genomes averaged to 1.6 Mbp. These results suggest that the length difference between genomes in high and low-fitness groups is likely to have biological relevance. Indeed, we found a very high correspondence between the lengths of our MAGs and their best matching reference genomes in the GTDB (r=0.88, p=5e-14) (Supplementary Table 7).

Our metabolic enrichment analysis revealed 33 KEGG pathway modules, each containing genes that form a functional unit in a metabolic pathway. Every module that was enriched differentially between these two groups were enriched in the high-fitness group. The lack of any enriched modules in the low-fitness group is in line with the reduction in genome lengths in the low-fitness group and further suggests that the reduction is associated with the absence of metabolic modules. Of all enriched modules, 79% were modules related to biosynthesis, which indicates an overrepresentation of biosynthetic capabilities in the high-fitness group as KEGG modules for biosynthesis only make up 55% of all KEGG
modules (Figure 3, Supplementary Table 7). Of the 33 enriched modules, 48.5% were associated with amino acid metabolism, 21.2% with vitamin and cofactor metabolism, 18.2% with carbohydrate metabolism, 6% with lipid metabolism and 3% with energy metabolism (Supplementary Table 7). Metabolic modules that were enriched in the high-fitness group included the biosynthesis of seven of the nine essential amino acids, indicating the importance of metabolic competency to synthesize high-demand compounds as a factor increasing fitness in colonizing new gut environments (Supplementary Table 7). This is further supported by the enrichment of biosynthesis pathways for the essential cofactor vitamin B12 (cobalamin), which occurred in 67.5% of the high-fitness populations and only 12.5% of the low-fitness group (Supplementary Table 7). Vitamin B12 is structurally highly complex and costly to produce, requiring expression of more than 30 genes that are exclusively encoded by bacteria and archaea (Martens et al. 2002). Thus, the competitive advantages conferred by metabolic autonomy appear to outweigh the additional costs. In addition to the biosynthesis of tetrahydrofolate, riboflavin, and cobalamin, the high-fitness group had a larger representation of biosynthetic modules for vitamins including biotin, pantothenate, folate, and thiamine (Supplementary Table 7), micronutrients that are equally important in bacterial and human metabolism and are shown to play important roles in mediating host-microbe interactions (Biesalski 2016). Interestingly, enriched metabolic modules in our analysis partially overlap with those that Feng et al. identified as the determinants of microbial fitness using metatranscriptomics and a germ-free mouse model conventionalized with microbial isolates of human origin (Feng et al. 2020).

Even though enriched metabolic modules occurred mostly in high-fitness populations, we did find some of these modules in the low-fitness group as well (Supplementary Table 7), but their distribution was not uniform as they primarily occurred only in a subset of genomes that resolved to Firmicutes (Figure 3). We then sought to identify whether the levels of completion of these modules that occurred in both groups were identical. For this, we matched six low-fitness genomes that encoded modules enriched in high-fitness group genomes to six high-fitness genomes from the same phylum (marked as HF and LF subgroups in Figure 3). Bacterial single-copy core genes estimated that genomes in both subgroups were highly complete with a slight increase in average completion of low-
fitness genomes (93.7%) compared to high-fitness genomes (90.1%). Despite the higher estimated genome completion for low-fitness populations, estimated metabolic module completion values were significantly lower in the low-fitness group (Wilcoxon rank sum test with continuity correction, V=958, p=5e-09) (Figure 3, Supplementary Table 7). This indicates that even when modules that are associated with high-fitness were detected in low-fitness genomes, they were systematically missing genes and were less complete than the same modules in high-fitness genomes.

**Figure 3.** Distribution of metabolic modules across low and high-fitness genomes. Each data point in this heat map shows the level of completion of a given metabolic module (rows) in a given genome (columns). The box-plot on the right-side compares a subset of low-fitness (LF) and high-fitness (HF) genomes, where each data point is the level of completion of a given metabolic module in a genome and shows a statistically significant difference between the overall completion of metabolic modules between these subgroups (Wilcoxon rank sum test, p=5.4e-09).
While gut microbial ecosystems of healthy individuals include both low- and high-fitness microbes, IBD primarily selects for high-fitness populations.

Our results so far show that while the healthy donor environment could support both high-fitness and low-fitness populations (Figure 1, Supplementary Table 3), challenging microbes to colonize a new environment or withstand massive ecosystem perturbation during FMT selects for high-fitness populations (Figure 3, Supplementary Table 7), suggesting that metabolic competence is a more critical determinant of fitness during stress than during homeostasis. Based on these observations, it is conceivable to hypothesize that (1) a healthy gut environment will support a range of microbial populations with a wide spectrum of metabolic competency, and (2) a gut environment under stress will select for high metabolic competency in microbial populations.

To test these hypotheses, we compared genomes reconstructed from a cohort of healthy individuals (Pasolli et al. 2019) to genomes reconstructed from a cohort of pouchitis patients (Vineis et al. 2016), a form of IBD with similar pathology to ulcerative colitis (De Preter et al. 2009). We conservatively excluded genomes that were less than 70% complete from the Pasolli et al. dataset to avoid underestimating pathway completion in genomes from healthy people. In contrast, we included any genomes from the Vineis et al. dataset if they were more than 50% complete. The number of genomes per individual and their level of completeness were similar between IBD and healthy individuals: we compared 44 genomes from 4 IBD patients with an average completion of 89.2% and 264 genomes from 22 healthy individuals with an average completion of 90.4% (Supplementary Table 8). Intriguingly, similar to the length differences between genomes of high-fitness and low-fitness populations (2.8 Mbp versus 1.6 Mbp on average), microbial populations associated with IBD patients had larger genomes compared to healthy people and averaged to 3.3 Mbp versus 2.6 Mbp, respectively (Supplementary Table 8). This suggests that despite the slightly higher completion of microbial genomes from the healthy cohort, these genomes tended to be smaller in size than the genomes reconstructed from IBD patients.
Next, we asked whether the completion of those metabolic modules associated with colonization success and resilience differed between the genomes reconstructed from healthy and IBD individuals. The level of completion of the 33 metabolic modules were almost identical between high-fitness genomes and genomes from IBD patients (Wilcoxon rank sum test, p=0.5), but they were significantly less complete in microbial genomes from healthy individuals (Wilcoxon rank sum test, p=5.39e-08) (Figure 4, Supplementary Table 8). Metabolic modules with the largest differences in completion between genomes from healthy and IBD individuals included biosynthesis of cobalamin, arginine, ornithine, tryptophan, isoleucine as well as the Shikimate pathway (Figure 4, Supplementary Table 8), a seven step metabolic route bacteria use for the biosynthesis of aromatic amino acids (phenylalanine, tyrosine, and tryptophan) (Herrmann and Weaver 1999).

Our findings show that the same set of key metabolic modules that distinguish high-fitness and low-fitness populations in FMT were also differentially associated with populations that occurred in healthy individuals compared to IBD patients. In particular, while healthy individuals seem to harbor microbes with a broad range of metabolic competency, IBD individuals appear to be enriched with organisms with metabolic autonomy. It is conceivable that a stable gut microbial ecosystem is more likely to support low-fitness populations through metabolic cross-feeding, where vitamins, amino acids, and nucleotides are exchanged between microbes (D’Souza et al. 2018). In contrast, host-mediated environmental stress in IBD likely disrupts such interactions and creates an ecological filter that selects for metabolic competence, which subsequently leads to loss of diversity and the dominance of organisms with large genomes that are not necessarily abundant in states of homeostasis.

**Figure 4.** Distribution of metabolic modules in genomes reconstructed from healthy individuals and individuals with IBD. The top panel shows the metabolic module completion values for (1) high- and (2) low-fitness donor genomes identified in this study (blue and orange), (3) genomes from healthy individuals (green), and (4) genomes from individuals with IBD. Next to group averages, shown the distribution of metabolic modules for each individual. Each dot in a given box-plot represents one of 33 metabolic modules that were enriched in high-fitness FMT donor populations and the y-axis indicates its estimated completion. In the bottom panel the completion values for 10 of the 33 pathways shown as ridge-line plots.
These observations have implications for the defining features of healthy gut environments from an ecological point of view. Defining the ‘healthy gut microbiome’ has been a major goal of human gut microbiome research (Bäckhed et al. 2012), and remains elusive (Eisenstein 2020). Despite comprehensive investigations that considered core microbial taxa (Arumugam et al. 2011; Lloyd-Price, Abu-Ali, and Huttenhower 2016) or guilds of microbes that represent coherent functional groups (Wu et al. 2021), the search for ‘biomarkers’ of healthy microbiomes is ongoing (McBurney et al. 2019). Given our data we hypothesize that one of the defining features of a healthy gut environment is its ability to support a diverse community of microbes with a broad spectrum of metabolic competence, where both low-fitness and high-fitness populations live in a coherent ecosystem. Conversely, an enrichment of metabolically competent high-fitness populations would likely indicate the presence of environmental stress. Our analyses demonstrate that this is a quantifiable feature of microbial communities through genome-
resolved metagenomic surveys, although testing this hypothesis rigorously requires a larger number of genomes reconstructed from a larger number of individuals diagnosed with a broader set of IBDs. Our analyses have other limitations. For instance, metabolic insights in our study have been limited to genomic potential and have considered only well-known metabolic pathways, which, given the extent of the unknown coding space in microbial genomes (Vanni et al. 2020), are likely far from complete. As a result, the disproportional enrichment of biosynthetic modules in high-fitness genomes indicates that the ability to synthesize essential biological compounds is necessary but likely insufficient to survive environmental stress in the gut. Nevertheless, the finding that the same metabolic modules that promote colonization success after FMT are also the hallmarks of fitness in IBD suggests the presence of ecological principles that are shared between these systems and warrants deeper investigation.

Subtle differences in key functions distinguish populations of the same genus with differential colonization success

While adaptive processes that favor metabolic independence explain the determinants of colonization and resilience for distantly related taxa, metabolic features that promote high-fitness at this broad level may not explain differences in fitness between more closely related taxa, such as distinct species within a single genus, which are likely to have similar metabolic capabilities (Martiny, Treseder, and Pusch 2013) due to unifying ecological traits in higher ranks of taxonomy (Philippot et al. 2010). We finally investigated whether we could identify determinants of fitness across metabolically similar populations with different levels of success in colonizing unrelated individuals.

Members of the genus *Bifidobacterium* have long been used as probiotics (Gomes and Malcata 1999) and are prevalent occupants of the healthy human gut microbiota (Arboleya et al. 2016). In our dataset, *Bifidobacterium* was the second most abundant genus (14.1%) after *Bacteroides* (15.8%) in Donor A, from whom we reconstructed three MAGs over 98% completion that resolved to three distinct species in this genus: *B. longum*, *B. adolescentis subsp. adolescentis*, and *B. animalis subsp. lactis* (Supplementary Table 3). While each of these *Bifidobacterium* populations occurred in
Donor A metagenomes in a relatively stable fashion, they showed vastly different colonization efficiency upon FMT (Figure 5), enabling us to investigate determinants of colonization among closely related taxa.

In contrast to the *B. longum* and *B. adolescentis subsp. adolescentis* (henceforth *B. adolescentis*) populations that colonized most recipients, *B. animalis subsp. lactis* (henceforth *B. lactis*) did not seem to have colonized any of our recipients (Supplementary Table 3). Overall, we were able to detect *B. longum*, *B. adolescentis*, and *B. lactis* populations in 83%, 75%, and 4% of all post-FMT recipient metagenomes, respectively (Figure 5). Most strikingly, patterns of colonization that emerged from the analysis of FMT recipients reflected those seen in publicly available gut metagenomes from Canada, where *B. longum*, *B. adolescentis*, and *B. lactis* populations occurred in 74%, 39%, and 13% of the population, demonstrating a positive relationship (Pearson’s correlation of 0.9, n.s.) between the colonization efficiency upon FMT and the fitness of these populations. Furthermore, the gut metagenomes from 17 countries confirmed the substantially reduced fitness of *B. lactis* globally (Supplementary Table 9). Interestingly, the *B. lactis* MAG we reconstructed from Donor A was virtually identical (with over 99.99% sequence identity over 99.82% alignment, Supplementary Table 9) to most *B. lactis* strains that are widely used as probiotics (Jungersen et al. 2014), revealing a disagreement between the preferences of commercial microbial therapeutics and human gut microbial ecology.

*Figure 5. Characteristics of three Bifidobacterium species.* Top panel shows the distribution of Donor A MAGs that represent three distinct Bifidobacterium populations across donor and recipient metagenomes before and after FMT. The last two columns in this panel show the prevalence of these populations in post-FMT metagenomes, and publicly available gut metagenomes from Canada. The panel below displays the distribution of KEGG orthologs across the three Bifidobacterium MAGs along with 31 high-quality isolate genomes from the NCBI. Each item shown in concentric circles represents a single function assigned by the database of KEGG Orthologs, and each layer is a distinct genome. The intensity of color indicates the presence of a given function in a given genome. The most outer circle marks groups of functions that are enriched in various groups of Bifidobacterium genomes as well as those functions that are not enriched in any group as they are either in all genomes, or only a very small number of them.
To identify factors that may explain the differences in colonization success between *B. longum*, *B. adolescentis*, and *B. lactis*, we created a collection of *Bifidobacterium* genomes that, in addition to the three metagenome-assembled genomes we reconstructed, included 31 complete genomes obtained from the NCBI (within-group and across-groups average gANI of 98.9% and 77.3%, respectively) (Supplementary Table 9). All three groups of *Bifidobacterium* genomes encoded the majority of the metabolic pathways associated with the high-fitness group (63% ± 5%). However, missing pathways were not uniformly distributed across three: *B. lactis* lacked the largest fraction of these pathways (42%) compared to the more prevalent *B. adolescentis* (36%) and *B. longum* (33%) (Supplementary Table 9). *B. longum* and *B. adolescentis* carried the complete tetrahydrofolate (vitamin B9) biosynthesis pathway in agreement with previous metabolic descriptions of *Bifidobacterium* (D’Aimmo et al. 2012; Sugahara et al. 2015) which
qualifies this group as attractive probiotics (Strozzi and Mogna 2008; Pompei et al. 2007), but this pathway was absent in *B. lactis* genomes. We also found that *B. longum* and *B. adolescentis* genomes encoded histidine and nicotinamide adenine dinucleotide (NAD) biosynthesis which *B. lactis* lacked (Supplementary Table 9). Finally, the average genome lengths of *B. longum* (2.31 Mbp) and *B. adolescentis* (2.18 Mbp) were longer than the average genome length of *B. lactis* (1.94 Mbp), which reflects the pattern we observed previously where high-fitness populations tended to have larger genomes. In summary, even though all *Bifidobacterium* genomes in our pangenome had a higher metabolic overlap with one another compared to high-fitness and low-fitness genomes we have previously studied, the reduced fitness of *B. lactis* compared to *B. longum* and *B. adolescentis* could still be explained by the absence of a small number of metabolic competencies associated with the high-fitness group genomes.

Next, we focused on the enrichment of individual functions across the three groups of genomes using gene annotations from KOfam profiles (Aramaki et al. 2020) from the Kyoto Encyclopedia of Genes and Genomes (KEGG) (M. Kanehisa and Goto 2000) and Clusters of Orthologous Groups (COGs) from the NCBI (Galperin et al. 2021). Of all 954 unique KOfams found in our *Bifidobacterium* pangenome, 272 functions were not common to all genomes but statistically enriched in either one or two groups. Our analysis of these accessory functions showed that *B. longum* encoded 150 (55.5%), *B. adolescentis* encoded 115 (42.3%), and *B. lactis* encoded 95 (34.9%) of all accessory functions that were statistically enriched (Figure 5, Supplementary Table 9). The same analysis with 1,286 unique COGs confirmed these observations: of all 353 COGs enriched in any group, *B. longum* encoded 212 (60.1%), *B. adolescentis* encoded 172 (48.7%), and *B. lactis* encoded 118 (33.4%) (Supplementary Table 9). Overall, these results reveal a striking correlation between the number of accessory functions associated with *B. longum*, *B. adolescentis*, and *B. lactis*, and echo the absence of metabolic pathways in *B. lactis* even at the level of accessory gene functions, explaining their differential ability to colonize new individuals and distribution in global human gut metagenomes.
We finally investigated the contents of the differentially occurring accessory functions to speculate on whether they could be related to differences in fitness. For instance, in contrast to all *B. longum* and *B. adolescentis* in the *Bifidobacterium* pangenome, none of the *B. lactis* genomes encoded a phosphoenolpyruvate phosphotransferase (PEP-PTS) system specific for the uptake of β-glucoside (Supplementary Table 9). As the genus *Bifidobacterium* is characterized by a large array of genes associated with carbohydrate uptake and metabolism (Ventura et al. 2009; Schell et al. 2002; Kleerebezem and Vaughan 2009), *B. lactis* represents a notable exception with a lower number of genes associated with carbohydrate metabolism, fewer genes encoding carbohydrate-specific ABC transporters, and the absence of phosphoenolpyruvate-phosphotransferase (PEP-PTS) systems (Barrangou et al. 2009). The absence of any other PEP-PTS system in *B. longum* and *B. adolescentis* may indicate the catabolic niche occupied by these populations in the human gut that is shaped by their extensive capacity for uptake and metabolism of plant derived glycosides (Chien, Huang, and Chou 2006; Schell et al. 2002). Additional functions that exclusively occurred in *B. adolescentis* and *B. longum* genomes included two multidrug resistance pumps of the ‘multidrug and toxin extrusion’ (MATE) type, three transporters of the major facilitator superfamily (MFS) involved in bile acid tolerance and macrolide efflux, two bile acid:natrium ion symporters, and one proton/chloride ion antiporter conferring acid tolerance (Supplementary Table 9). The drug defense mechanisms may act to protect these populations during periods of inflammation and drug administration, but may also be beneficial with regard to the common ingestion of antibiotics through various food products (Kirchhelle 2018). These results show that in the microbial fitness landscape of the human gut, where the determinants of success across distantly related taxa are primarily defined by the presence or absence of a large number of metabolic pathways, there exists smaller niches equally accessible to closely related organisms with similar metabolic potential, among which success can be speculated by subtle differences in key functions.
Conclusions

Our study points to adaptive ecological processes as primary determinants of both long-term colonization after FMT and microbial fitness in the human gut environment through metabolic competency as conferred by biosynthesis of nucleotides, amino acids, and essential micronutrients. Even when we found these metabolic modules in low-fitness populations, they were systematically less complete compared to their high-fitness counterparts.

Our findings suggest that in a healthy gut environment high- and low-fitness populations co-occur in harmony, with their differential fitness indiscernible by taxonomy or relative abundance. However, transfer to a new gut environment through FMT, or host-mediated stress through IBD, initiates an ecological filter that selects for high-fitness populations that can self-sustain. This model offers a null hypothesis to explain how low-abundance members of healthy gut environments can come to dominate the gut microbiota under stressful conditions, while not being causally associated with disease states. If the association between particular microbial taxa and disease is solely driven by their superior metabolic competence, microbial therapies that aim to treat complex diseases by adding microbes associated with healthy individuals will be unlikely to compete with the adaptive processes that regulate complex gut microbial ecosystems.
Materials and Methods

Sample collection and storage. We used a subset of individuals who participated in a randomized clinical trial (Kao et al. 2017) and conducted a longitudinal FMT study of two human cohorts (DA and DB), each consisting of one FMT donor and 5 FMT recipients of that donor’s stool. All recipients received vancomycin for a minimum of 10 days pre-FMT at a dose of 125 mg four times daily. Three DA and two DB recipients received FMT via pill, and two DA and three DB recipients received FMT via colonoscopy. All recipients had recurrent C. difficile infection before FMT, and two DA recipients and 1 DB recipient were also diagnosed with ulcerative colitis (UC). 24 stool samples were collected from the DA donor over a period of 636 days, and 15 stool samples were collected from the DB donor over a period of 532 days. Between 5 and 9 stool samples were collected from each recipient over periods of 187 to 404 days, with at least one sample collected pre-FMT and 2 samples collected post-FMT. This gave us a total of 109 stool samples from all donors and recipients. Samples were stored at -80°C. (Supplementary Figure 1, Supplementary Table 1)

Metagenomic short-read sequencing. We extracted the genomic DNA from frozen samples according to the centrifugation protocol outlined in MoBio PowerSoil kit with the following modifications: cell lysis was performed using a GenoGrinder to physically lyse the samples in the MoBio Bead Plates and Solution (5–10 min). After final precipitation, the DNA samples were resuspended in TE buffer and stored at ~20 °C until further analysis. Sample DNA concentrations were determined by PicoGreen assay. DNA was sheared to ~400 bp using the Covaris S2 acoustic platform and libraries were constructed using the Nugen Ovation Ultralow kit. The products were visualized on an Agilent Tapestation 4200 and size-selected using BluePippin (Sage Biosciences). The final library pool was quantified with the Kapa Biosystems qPCR protocol and sequenced on the Illumina NextSeq500 in a 2 × 150 paired-end sequencing run using dedicated read indexing.

‘Omics workflows. Whenever applicable, we automated and scaled our ‘omics analyses using the bioinformatics workflows implemented by the program `anvi-run-workflow`
Anvi’o workflows implement numerous steps of bioinformatics tasks including short-read quality filtering, assembly, gene calling, functional annotation, hidden Markov model search, metagenomic read-recruitment, metagenomic binning, pangenomics, and phylogenomics. Workflows use Snakemake (Köster and Rahmann 2012) and a tutorial is available at the URL http://merenlab.org/anvio-workflows/. The following sections detail these steps.

**Taxonomic composition of metagenomes based on short reads.** We used Kraken2 v2.0.8-beta (Wood, Lu, and Langmead 2019) with the NCBI’s RefSeq bacterial, archaeal, viral and viral neighbours genome databases to calculate the taxonomic composition within short-read metagenomes.

**Assembly of metagenomic short and long reads.** To minimize the impact of random sequencing errors in our downstream analyses, we used the program `iu-filter-quality-minoche` to process short metagenomic reads, which is implemented in illumina-utils v2.11 (Eren et al. 2013) and removes low-quality reads according to the criteria outlined by Minoche et al. (Minoche, Dohm, and Himmelbauer 2011). IDBA_UD v1.1.2 (Peng et al. 2012) assembled quality-filtered short reads into longer contiguous sequences (contigs), although we needed to recompile IDBA_UD with a modified header file so it could process 150bp paired-end reads. For the assembly of long-reads and identification of circular contigs we used Flye v2.6 (Kolmogorov et al. 2019) with the `--meta` argument followed by Pilon v1.23 (Walker et al. 2014) correction using the high-quality short-reads.

**Processing of contigs.** We use the following strategies to process both sequences we obtained from our assemblies and those we obtained from reference genomes. Briefly, we used (1) `anvi-gen-contigs-database` on contigs to compute k-mer frequencies and identify open reading frames (ORFs) using Prodigal v2.6.3 (Hyatt et al. 2010), (2) `anvi-run-hmms` to identify sets of bacterial (Campbell et al. 2013) and archaeal (Rinke et al. 2013) single-copy core genes using HMMER v3.2.1 (Eddy 2011), (3) `anvi-run-ncbi-cogs` to annotate ORFs with functions from the NCBI’s Clusters of Orthologous Groups (COGs) (Tatusov et al. 2003), and (4) `anvi-run-kegg-kofams` to annotate ORFs with functions from the KO fam HMM database of KEGG orthologs (KOs) (Aramaki et al. 2020; M. Kanehisa and Goto 2000). To predict the approximate number of genomes in
metagenomic assemblies we used the program `anvi-display-contigs-stats`, which calculates the mode of the frequency of single-copy core genes as described previously (Delmont and Eren 2016).

**Metagenomic read recruitment, reconstructing genomes from metagenomes, determination of genome taxonomy and ANI.** We recruited metagenomic short reads to contigs using Bowtie2 v2.3.5 (Langmead and Salzberg 2012) and converted resulting SAM files to BAM files using samtools v1.9 (Li et al. 2009). We profiled the resulting BAM files using the program `anvi-profile` with the flag `--min-contig-length` set to 2500 to eliminate shorter sequences to minimize noise. Once we have read recruitment results from each metagenome is profiled to store contig coverages into single anvi'o profile databases, `anvi-merge` combined all profiles into an anvi'o merged profile for downstream visualization, binning, and statistical analyses. We then used `anvi-cluster-contigs` to group contigs into 100 initial bins using CONCOCT v1.1.0 (Alneberg et al. 2014), `anvi-refine` to manually curate initial bins with conflation error based on tetranucleotide frequency and differential coverage signal across all samples, and `anvi-summary` to report final summary statistics for each gene, contig, and bin. We used the program `anvi-rename-bins` to identify bins that were more than 70% complete and less than 10% redundant, and store them in a new collection as metagenome-assembled genomes (MAG), discarding lower quality bins from downstream analyses. GTBD-tk v0.3.2 (Chaumeil et al. 2019) assigned taxonomy to each of our MAG using GTDB r89 (Parks et al. 2018), but to assign species- and subspecies-level taxonomy for `DA_MAG_00057`, `DA_MAG_00011`, `DA_MAG_00052` and `DA_MAG_00018`, we used `anvi-get-sequences-for-hmm-hits` to recover DNA sequences for bacterial single-copy core genes that encode ribosomal proteins, and searched them in the NCBI's nucleotide collection (nt) database using BLAST (Altschul et al. 1990). Finally, the program `anvi-compute-genome-similarity` calculated pairwise genomic average nucleotide identity (gANI) of our genomes using PyANI v0.2.9 (Pritchard et al. 2016).

**Criteria for MAG detection in metagenomes.** Using mean coverage to assess the occurrence of populations in a given sample based on metagenomic read recruitment can yield misleading insights since this strategy cannot accurately distinguish reference
sequences that represent very low-abundance environmental populations from those sequences that do not represent an environmental population in a sample yet still recruit reads from non-target populations due to the presence of conserved genomic regions. Thus we relied upon the ‘detection’ metric, which is a measure of the proportion of the nucleotides in a given sequence that are covered by at least one short read, and considered a population was detected in a metagenome if ani’o reported a detection value of at least 0.25 for its genome (whether it was a metagenome-assembled or isolate genome). Values of detection in metagenomic read recruitment results often follow a bimodal distribution for populations that are present and absent (see Supplementary Figure 2 in (Utter et al. 2020)), thus 0.25 is an appropriate cutoff to eliminate false-positive signal in read recruitment results for populations that are absent.

Identification of MAGs that represent multiple subpopulations. To identify subpopulations of MAGs in metagenomes, we used the ani’o command `anvi-gen-variability-profile` with the `--quince-mode` flag which exported single-nucleotide variant (SNV) information for all MAGs after read recruitment. We then used DESMAN v2.1.1 (Quince et al. 2017) to analyze SNVs to determine the number and distribution of subpopulations represented by a single genome. To account for non-specific mapping that can inflate the number of estimated subpopulations, we removed any subpopulation that made up less than 1% of the entire population explained by a single MAG. To account for noise due to low-coverage, we only investigated subpopulation for MAGs for which the mean non-outlier coverage of single-copy core genes was at least 10X.

Criteria for colonization of a recipient by a MAG. We developed a method to determine whether or not a MAG successfully colonized a recipient, and applied this method to each MAG and each recipient within a cohort. In order to confidently assign colonization or non-colonization phenotypes to each MAG/recipient pair, we required that the MAG be detected in the donor sample used for transplant into the recipient. If these criteria were met, we then determined whether the MAG was detected in any post-FMT recipient sample taken more than 7 days after transplant. If not, the MAG/recipient pair was considered a non-colonization event. If the MAG was detected in the recipient greater than7 days post-FMT, we used subpopulation information to determine if any
subpopulation present in the donor and absent in the recipient pre-FMT was detected in the recipient more than 7 days post-FMT. If this was the case, we considered this to represent a colonization event. See Supplementary Figure 4 for a complete outline of all possible cases.

Determination of dose and fitness for MAGs. We defined population dose as the second and third quartile mean coverage of a population in the transplanted stool sample. We defined fitness as the prevalence of a population in 23 healthy adult gut metagenomes (see Materials and Methods: Criteria for MAG detection in metagenomes) from Canada, the same country in which the FMTs were performed.

Regression analysis. To examine the association between dose and/or prevalence with colonization outcome, we built binomial logistic regression models using the R stats `glm` function. We used the R stats `predict` function and the R pROC `roc` function to evaluate our models by creating receiver operating characteristic (ROC) curves and calculating the area under the ROC curve (AUC). To determine the correlation between dose and prevalence, we performed linear regression using the R stats `lm` function. We used the R tidyverse package, including ggplot2, to visualize boxplots, scatterplots, and ROC curves.

Pangenomic analysis and gANI. We used anvi’o to compute and visualize pangenomes of MAGs and reference genomes. We stored all processed MAG and reference genome contigs (see Contig processing methods section) in an anvi’o database using the command `anvi-gen-genomes-storage`. To create the pangenomes, we then passed that database to the command `anvi-pan-genome` which used NCBI’s BLAST (Altschul et al. 1990) to quantify gene similarity within and between genomes and the Markov Cluster algorithm (MCL) (Enright, Van Dongen, and Ouzounis 2002) to cluster groups of similar genes. We set the `anvi-pan-genome` `--min-occurrence` flag to 2 to remove gene clusters only present in one genome (singletons), and visualized pangenomes using `anvi-display-pan`.

Phylogenomic tree construction. To concatenate and align amino acid sequences of 46 single-copy core (Campbell et al. 2013) ribosomal proteins that were present in all of
our *Bifidobacterium* MAGs and reference genomes, we ran the anvi’o command `anvi-get-sequences-for-hmm-hits` with the `--return-best-hit`, `--get-aa-sequence` and `--concatenate` flags, and the `--align-with` flag set to `muscle` to use MUSCLE v3.8.1551 (Edgar 2004) for alignment. We then ran `anvi-gen-phylogenomic-tree` with default parameters to compute a phylogenomic tree using FastTree 2.1 (Price, Dehal, and Arkin 2010).

**Analysis of metabolic modules and enrichment.** We calculated the level of completeness for a given KEGG module (Minoru Kanehisa et al. 2014, 2017) in our genomes using the program `anvi-estimate-metabolism`, which leveraged previous annotation of genes with KEGG orthologs (KOs) (see the section ‘Processing of contigs’). Then, the program `anvi-compute-functional-enrichment` determined whether a given metabolic module was enriched in based on the output from `anvi-estimate-metabolism`. The URL [https://merenlab.org/m/anvi-estimate-metabolism](https://merenlab.org/m/anvi-estimate-metabolism) serves a tutorial for this program which details the modes of usage and output file formats. The statistical approach for enrichment analysis is defined elsewhere (Shaiber et al. 2020), but briefly it computes enrichment scores for functions (or metabolic modules) within groups by fitting a binomial generalized linear model (GLM) to the occurrence of each function or complete metabolic module in each group, and then computing a Rao test statistic, uncorrected p-values, and corrected q-values. We considered any function or metabolic module with a q-value less than 0.05 to be ‘enriched’ in its associated group if it was also at least 75% complete and in at least 50% of the group members. To display the distribution of individual KEGG orthologs across genomes and order them based on their enrichment scores and group affiliations we used the program `anvi-display-functions`.

**Determination of high-fitness and low-fitness MAGs for metabolic enrichment analysis.** We classified MAGs as high-fitness if, in all 5 recipients, they were detected in the donor sample used for transplantation as well as the recipient more than 7 days post-FMT. We classified low-fitness MAGs as those that, in at least 3 recipients, were detected in the donor sample used for FMT but were not detected in the recipient at least 7 days post-FMT. We reduced the number of high-fitness MAGs to be the same as the number...
of low-fitness MAGs for metabolic enrichment analysis by selecting only the high-fitness MAGs which were the most prevalent in the Canadian gut metagenomes.

**Ordination plots.** We used the R vegan v2.4-2 package `metaMDS` function to perform nonmetric multidimensional scaling (NMDS) with Horn-Morisita dissimilarity distance to compare taxonomic composition between donor, recipient, and global metagenomes. We visualized ordination plots using R ggplot2.

**Data Availability**

Raw sequencing data for donor and recipient metagenomes are stored under the NCBI BioProject PRJNA701961 (see Supplementary Table 1 for accession numbers). Supplementary tables are accessible via doi: [10.6084/m9.figshare.14138405](https://doi.org/10.6084/m9.figshare.14138405).

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**Author Contributions**

AME, TL, BJ conceived the study. JZD, MS, TL recruited patients, performed transplantation experiments, and collected samples. ARW, JF, AME performed primary data analyses. IV developed research tools. KL, STML, HGM performed sample processing and sequencing. FT, AS, EF, CQ, MKY, AY contributed to data analyses and interpretation. DTR, BJ, TL, and AME directed research. ARW, JF, AME wrote the paper with critical input from all authors.
Competing Interests

Authors declare no competing financial interests.

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Supplementary Figures

Supplementary Figure 1. Timeline of stool samples collected from FMT study. Each circle represents a stool sample collected from either an FMT donor or FMT recipient. The thicker, red vertical line at day 0 represents the FMT event for each recipient. FMT method (pill or colonoscopy) and FMT recipient health and disease state (C. diff - chronic recurrent Clostridium difficile infection, UC - ulcerative colitis) are indicated on the right.
Supplementary Figure 2. Nonmetric multidimensional scaling (NMDS) ordination of the taxonomic composition of donor, recipient, and Canadian gut metagenomes at the genus level based on Morisita-Horn dissimilarity. Samples from the same participant are joined by lines with the earliest time point labeled. CAN: Canadian gut metagenomes, DA: donor A, DB: donor B, POST: recipients post-FMT, PRE: recipients pre-FMT.

Supplementary Figure 3. Nonmetric multidimensional scaling (NMDS) ordination of the taxonomic composition of the donor and recipient metagenomes at genus level based on Morisita-Horn dissimilarity. Samples from the same participant are joined by lines with the earliest time point labeled. DA_POST: donor A recipients post-FMT, DA_PRE: donor A recipients pre-FMT, DB_POST: donor B recipients post-FMT, DB_PRE: donor B recipients pre-FMT, DA: donor A, DB: donor B.
Supplementary Figure 4. A flowchart outlining our method to assign successful colonization, failed colonization, or undetermined colonization phenotypes to donor-derived populations in the recipients of that donor's stool.
Supplementary Tables

**Supplementary Table 1: Description of FMT study and stool samples collected.**
a) Description of FMT donor stool samples and SRA accession numbers. b) Description of FMT recipient samples and SRA accession numbers. c) Description of transplantation events.

**Supplementary Table 2: Description of FMT metagenomes and co-assemblies.**
a) Metagenome SRA accession numbers and number of metagenomic short-reads sequenced and mapped to co-assemblies and MAGs. b) Phylum level taxonomic composition of metagenomes. c) Genus level taxonomic composition of metagenomes. d) Summary statistics for contigs from metagenome co-assemblies.

**Supplementary Table 3: Description of MAGs.**
a) Summary statistics and taxonomic assignments for MAGs. b) Detection of Donor A and Donor B MAGs in FMT metagenomes, respectively. c) Detection of Donor A and Donor B MAGs in global gut metagenomes, respectively. d) Detection summary statistics of Donor A and Donor B MAGs in global gut metagenomes, respectively.

**Supplementary Table 4: Accession numbers of gut metagenomes from 17 countries.

**Supplementary Table 5: MAG subpopulation information.**
a) and b) Number of Donor A and Donor B MAG subpopulations detected in FMT metagenomes, respectively. c) and d) Subpopulation composition of Donor A and Donor B MAGs in FMT metagenomes, respectively.

**Supplementary Table 6: MAG/recipient pair colonization outcomes and MAG mean coverage in the 2nd and 3rd quartiles in stool samples used for transplantation.

**Supplementary Table 7: Description of high vs. low-fitness populations.**
a) Taxonomic assignments and genome size estimates for high- and low-fitness populations. b) KEGG module completeness information for high- and low-fitness populations. c) Raw KEGG module enrichment information for high- and low-fitness populations. d) KEGG module enrichment and categorical information for the 33 modules enriched in high-fitness populations. e) and f) Completeness information for the 33 modules enriched in high-fitness populations in all high- and low-fitness populations.

**Supplementary Table 8: a) List of genomes from healthy individuals and individuals with IBD. b) Module completion values across genomes.

**Supplementary Table 9: Bifidobacteria functional analysis.**
a) Accession numbers for *Bifidobacteria* reference genomes. b) Summary statistics for *Bifidobacteria* MAGs and reference genomes. c) Prevalence of *Bifidobacteria* MAGs in global gut metagenomes. d) gANI percent identity between *Bifidobacteria* genomes. e) gANI percent alignment coverage between *Bifidobacteria* genomes. f) KOfams enriched in different *Bifidobacteria* species. g) KOfam presence and absence in *Bifidobacteria* genomes. h) COG functions enriched in different *Bifidobacteria* species. i) COG function presence and absence in *Bifidobacteria* genomes. j) KEGG modules enriched in different *Bifidobacteria* species. k) KEGG module completeness in *Bifidobacteria* genomes.