Diet Dominates Host Genotype in Shaping the Murine Gut Microbiota

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SUMMARY

Mammals exhibit marked interindividual variations in their gut microbiota, but it remains unclear if this is primarily driven by host genetics or by extrinsic factors like dietary intake. To address this, we examined the effect of dietary perturbations on the gut microbiota of five inbred mouse strains, mice deficient for genes relevant to host-microbial interactions (MyD88−/−, NOD2−/−, ob/ob, and Rag1−/−), and >200 outbred mice. In each experiment, consumption of a high-fat, high-sugar diet reproducibly altered the gut microbiota despite differences in host genotype. The gut microbiota exhibited a linear dose response to dietary perturbations, taking an average of 3.5 days for each diet-responsive bacterial group to reach a new steady state. Repeated dietary shifts demonstrated that most changes to the gut microbiota are reversible, while also uncovering bacteria whose abundance depends on prior consumption. These results emphasize the dominant role that diet plays in shaping interindividual variations in host-associated microbial communities.

INTRODUCTION

Although humans and other mammals exhibit many shared features of their resident gut microbial communities (Muegge et al., 2011), each individual harbors an idiosyncratic mixture of microbial strains and species (Faith et al., 2013). In healthy adults, the component members of the gut microbiota can be stable for years (Faith et al., 2013), whereas the relative abundance of each member (community structure) is highly dynamic (David et al., 2014). The underlying causes and consequences of these interindividual and temporal variations remain poorly characterized.

Studies in animal models have led to the proposal that the gut microbiota might be considered a complex polygenic trait shaped by both environmental and host genetic factors (Benson et al., 2010). However, it remains unclear if host genotype or environment (e.g., diet) plays a more dominant role in shaping microbial ecology. Associations between genetic loci and the abundance of bacterial taxa have been described in mice fed a controlled diet (Benson et al., 2010; McKnite et al., 2012) and in humans (Li et al., 2012; Smeeckens et al., 2014), the genetic distance between mouse strains was recently linked to the overall structure of the gut microbiota (Hildebrand et al., 2013), and numerous differences have been shown between transgenic animals and matched controls (Couturier-Maillard et al., 2013; Hashimoto et al., 2012; Spor et al., 2011).

Conversely, time series analyses of inbred mice have shown that the consumption of a high-fat, high-sugar diet dramatically alters the gut microbiota in a single day (Turnbaugh et al., 2009b; Zhang et al., 2012). Endpoint analyses of multiple inbred mouse strains support the importance of current dietary intake (Parks et al., 2013), as does a recent study, which demonstrates that the microbial responses to the fucosylation of host glycans are diet dependent (Kashyap et al., 2013). Finally, comparisons of human twins at various ages, from infants to adults, have failed to detect significantly more similar microbial communities in monozygotic versus dizygotic pairs, suggesting that environmental factors predominate over host genetics in shaping microbial ecology (Turnbaugh et al., 2009a; Yatsunenko et al., 2012).

Here, we systematically test the relative impacts of dietary intake and host genetics on the gut microbiota, through the combined analysis of five inbred mouse strains, four transgenic lines, and a recently developed outbred mouse resource, the Diversity Outbred (DO) population (Churchill et al., 2012; Svenson et al., 2012). The DO population was derived from partially inbred lines of the Collaborative Cross (Threadgill and Churchill, 2012) that were outbred using a randomized breeding scheme with avoidance of sibling matings to obtain high genetic diversity, heterozygosity, and fine recombination block structure (Svenson et al., 2012). These traits make the DO population ideal for investigating the relative contributions of host and environmental factors in shaping complex traits.

We selected two diets that reflect distinctive macronutrient profiles, are widely used to study diet-induced obesity, and represent modern human dietary regimes: a low-fat, high-plant polysaccharide diet (LFPP: 22.2% kcal protein, 16.0%...
fat, 61.7% carbohydrate) and a high-fat, high-sugar diet (HFHS: 14.8% kcal protein, 40.6% carbohydrate, 44.6% fat). Consecutive dietary shifts, longitudinal sampling, dietary mixtures, and computational modeling of microbial dynamics showed that interindividual and interstrain differences are rapidly and consistently reset by dietary perturbations. Our results also provide evidence that members of the gut microbiota undergo hysteresis in response to diet: their abundance is shaped by both current dietary intake and past exposures. Together, these findings have important implications for understanding host-microbial interactions during the consumption of a wide range of dietary regimes and suggest that attempts to understand the genetic underpinnings of the human gut microbiota will require a careful consideration of our past and present dietary habits.

RESULTS

Consistent Microbial Responses to a High-Fat, High-Sugar Diet in Inbred Mice

16S rRNA gene sequencing was used to profile the fecal microbiota of 73 mice representing 5 distinct inbred founder strains used in the Collaborative Cross: 129S1/SvImJ, A/J, C57BL/6J, NOD/LtJ, and NZO/HILtJ (n = 7–20 mice genotype; 3–9 cages genotype). For each genotype, we fed mice either a low-fat, high-plant-polysaccharide diet (LFPP) or a high-fat, high-sugar diet (HFHS) for ≥15 weeks (Figure S1A and Table S1A available online). 16S rRNA gene sequences were clustered into operational taxonomic units (OTUs) using a reference tree-guided approach, prior to unsupervised ordination using Bray-Curtis dissimilarity-based principal coordinates analysis (PCoA).

Consumption of the HFHS diet consistently modified the gut microbiota of all five strains such that all strains formed two distinct clusters defined by each diet (Figure 1A; p value < 0.001, permutational multivariate analysis of variance [PERMANOVA] of Bray-Curtis distances). This shift was evident at the phylum level: the HFHS diet significantly increased the relative abundance of the Firmicutes (21.39% ± 1.29% LFPP versus 52.92% ± 4.45% HFHS), increased the Verrucomicrobia (5.90% ± 1.20% LFPP versus 26.15% ± 3.98% HFHS; see Supplemental Results), and decreased the Bacteroidetes (71.02% ± 1.84% LFPP versus 16.35% ± 1.48% HFHS) (all comparisons p value < 0.0001, Wilcoxon rank-sum test; Figure 1B). These trends were also detectable at finer taxonomic levels, ranging from class to genus (Table S2A).

When considering each diet independently, we detected significant clustering by host genotype (p value < 0.001, PERMANOVA of Bray-Curtis distances). We also detected multiple taxa with significantly altered relative abundance between inbred strains; however, only three bacterial genera were significantly enriched in one mouse line relative to the other four (Table S3A). De novo clustering of our 16S RNA gene sequences into OTUs confirmed these trends, while also identifying a member of the Erysipelotrichaceae family (OTU13400) that was consistently enriched in C57BL/6J mice regardless of diet (Table S3B). Interestingly, both the magnitude and direction of change for multiple bacterial genera were different between genotypes (Figure 1C; Table S2A), suggesting that the impact of the HFHS diet may depend on the broader host or microbial community context.

To test if these correlations were driven by host genotype or reflected a recent shared history of microbial exposures, we sampled multiple cages from each inbred line, each representing a unique set of littermates cohoused since birth and split according to sex at 3 weeks of age. Principal coordinates analysis confirmed that many of the differences between genotypes were consistently found in all cages (Figure 2A). Correspondingly, we identified bacterial genera that were significantly enriched in 129S1/SvImJ, C57BL/6J, and NZO/HILtJ mice from multiple cages (Figures 2B–2D). However, we also detected one genus that was only significantly altered in one cage (Figure 2C), and we did not detect any genera (or higher-level taxa) that were consistently enriched in the same host genotype on both diets (Table S3). Thus, our results imply that the effects of dietary intake overshadow any preexisting differences between strains due to host genotype. Furthermore, we cannot fully exclude the possibility that some of the observed associations with host genotype may actually be the result of a longer-term shared history of environmental exposures.

Next, we sought to test if diet is capable of reproducibly shaping the gut microbiota in the context of more dramatic perturbations to host genotype and phenotype. Male C57BL/6J wild-type mice were fed a LFPP diet until they reached 7 weeks of age, at which point they were switched to the HFHS diet for 1 week. Fecal samples from multiple days prior to and after the diet shift were used for 16S rRNA gene sequencing (Figure S1B; Table S1B). The same procedure was performed for animals homozygous for mutations in four genes that have been previously shown to impact the gut microbiota (n = 5 mice genotype): MyD88−/− (Wen et al., 2008); NOD2−/− (Couturier-Maillard et al., 2013); ob/ob (Ley et al., 2005); and Rag1−/− (Scholz et al., 2014).}

The HFHS diet consistently modified the gut microbiota of wild-type animals and all four transgenic lines (Figure 3A), with significant clustering by diet across the entire data set (p value < 0.001, PERMANOVA of Bray-Curtis distances). Within each diet, there was significant clustering according to host genotype (p value < 0.001, PERMANOVA on the final time point for each diet). Analysis of microbial community structure over time revealed that all genotypes responded within 2 days (Figure 3B); however, MyD88−/− animals exhibited a blunted response to the HFHS diet (p value < 0.05, Kruskal-Wallis test with Dunn’s correction for multiple comparisons using data from the final time point). In order to account for differences in sample number between groups, we reanalyzed the data for each transgenic mouse strain and the wild-type control (n = 15 samples/group; the final time point on the HFHS diet was excluded to allow for matched comparisons on both diets). These analyses again identified diet as the primary factor driving the observed variations in microbial community structure, with genotype providing secondary clustering (Figures 3C–3F). Pairwise statistical analyses of the final time point on each diet confirmed these trends, with diet explaining more of the observed variation than host genotype (R² ranges: 0.08–0.23 [genotype] and 0.35–0.48 [diet]; ADONIS test). Diet also consistently altered community membership as indicated by the unweighted UniFrac metric (data not shown).

The HFHS diet significantly altered bacteria at taxonomic levels ranging from phylum to genus (Table S2A), confirming
the overall trend of increased Firmicutes and decreased Bacteroidetes. Although not all diet-associated groups reached significance in both the transgenic and inbred strain experiments, the direction of change was consistent in all cases where a significant association was found in both data sets. The identified diet-dependent bacterial taxa were also generally consistent in their direction of change in each mouse strain following consumption of the HFHS diet; in all cases where there was a disagreement, it was due to a single genotype (Table S2 A).

One explanation for the marked impact of the HFHS diet across multiple inbred strains, including transgenic animals, is that this represents a relatively strong dietary shift: 16.0% to 40.6% fat accompanied by a shift from plant polysaccharides to more readily digestible carbohydrates. Ecological theory predicts that the gut microbiota may be able to tolerate more subtle dietary interventions (Costello et al., 2012). To test if the magnitude of microbial response is directly proportional to the degree of dietary perturbation, we fed adult male C57BL/6J wild-type mice repelleted mixtures of the LFPP and HFHS diet (0, 1, 10, 25, 50, 75, and 100% HFHS by weight; n = 4–5 mice/diet). Fecal samples were collected prior to and 7 days after each dietary intervention and analyzed by 16S rRNA gene sequencing (Figure S1 C; Table S1 C). Dietary HFHS content influenced host adiposity over the 7 days of gradient feeding. Despite reducing food intake as the proportion of the HFHS diet increased (Figure 4 A), mice consumed more calories overall on HFHS-rich diets (Figure 4 B). Correspondingly, we observed HFHS-dependent increases in epididymal fat pad mass, whether measured
on an absolute basis ($R^2 = 0.272$; $F = 11.93$; $p$ value $= 0.002$) or as a fraction of body mass (Figure 4 C).

Including an increased proportion of Firmicutes (Figure 4 E) and of microbial taxa previously associated with the HFHS diet, early as HFHS content increased (Figure 4 D). We also observed coordinate (PC1, 37% of variance). PC1 values decreased linearly ($p$ value $< 0.01$; PERMANOVA of Bray-Curtis distances), with revealed significant clustering of the gut microbiota by diet group between the LFPP and HFHS diets (Table S2A). Together, these linear associations were also detectable at the genus level (Table S2B).

To more systematically vary host genetics in mice with a common combined experiment, was either below the limit of detection or present at low abundance in all samples from this experiment, regardless of diet (mean % of 16S sequences: $0.0013 \pm 0.0001$). These linear associations were also detectable at the genus level (Table S2B), confirming many of the previously observed differences between the LFPP and HFHS diets (Table S2A). Together, these data suggest that the gut microbiota responds to diet in a dose-dependent manner with even subtle perturbations detectably changing microbial community structure.

**A Rapid and Sustained Microbial Response to the High-Fat, High-Sugar Diet in Outbred Mice**

To more systematically vary host genetics in mice with a common history of environmental exposures, we obtained 60 Diversity Outbred (DO) mice (30 females, 30 males). All mice were from generation 6 of outbreeding, each representing a unique combination of eight founder alleles from the five strains surveyed previously, in addition to CAST/EiJ, PWK/PhJ, and WSB/EiJ (Churchill et al., 2012; Svenson et al., 2012). Genome-wide analyses of our animals confirmed that we successfully recaptured nearly all of the genetic diversity in the DO population ($F_{ST} = 0.05$ for the 60 animals in the DO time series experiment; $F_{ST} = 0.03$ for the expanded set of 203 animals). Consecutive dietary shifts were employed to assess the speed, reproducibility, and reversibility of the microbial responses to dietary perturbations. Individually housed mice were fed a LFPP diet from weaning until 7 weeks of age, at which time they were fed a HFHS diet for 1 week, returned to the LFPP diet for 1 week, and then maintained on the HFHS diet for 2 months. Fecal pellets were analyzed from the 60 mice at 18 time points, spanning 14 weeks ($n = 977$ samples; Figure S1D; Table S1D).

Consistent with our analysis of inbred strains and transgenic mice, the HFHS diet rapidly altered microbial community membership (Figure S2A) and structure (Figures 5A and 5B) despite substantial variations in host genotype (Svenson et al., 2012). The diet shifts also reset interindividual variability in community structure (i.e., $\beta$-diversity patterns). Mantel tests were performed for all pairwise comparisons of Bray-Curtis distance matrices representing differences in gut community structure between animals at each time point. Although within each diet $\beta$-diversity was stable over time (median $p$ value $< 0.001$), microbial community structure was rapidly altered by the HFHS diet such that the differences between mice could not be accurately predicted using samples collected on different diets (median $p$ value $> 0.1$; Figure S2B).

As with the inbred mice, consumption of the HFHS diet by outbred mice led to a significant increase in the Firmicutes phylum (39.10% $\pm$ 1.17% LFPP versus 64.65% $\pm$ 0.73% HFHS) and a decrease in the Bacteroidetes (57.81% $\pm$ 1.25%
A Bray-Curtis PC2 (12% variance)

B Bray-Curtis PC1 (25% variance)

C Bray-Curtis PC2 (23% variance)

D Bray-Curtis PC1 (34% variance)

E Bray-Curtis PC2 (17% variance)

F Bray-Curtis PC1 (38% variance)

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**Microbial Responses to Sequential Diet Shifts**

Given the rapid and reproducible microbial response to HFHS intake, we next sought to test whether this phenomenon could be attributed fully to current dietary intake or whether there were lingering effects of past dietary history. Analysis of the consecutive dietary shifts in outbred mice suggested that both community membership and structure were markedly altered after 3–7 days of HFHS diet consumption in naive mice, whereas the gut microbiota of mice previously exposed to the HFHS diet took 1–2 weeks to respond (Figure 5B). To more rigorously address this question, we collected daily fecal samples from inbred C57BL/6J mice (n = 15) oscillating between LFPP and HFHS diets every 3 days. All animals were maintained on a LFPP diet in individual cages prior to the beginning of the experiment at 7 weeks of age. In total, we analyzed four groups, representing two sets of oscillating mice staggered by 3 days, and two control groups maintained on either a continuous LFPP or a continuous HFHS diet. All animals were switched to the HFHS diet for the final week. Daily fecal samples were subjected to 16S rRNA gene sequencing, representing 536 total samples across the 38 day experiment (Figure S1E; Table S1F).

Mouse physiology and chow consumption were rapidly altered by diet. Body weight consistently increased during consumption of the HFHS diet, while decreasing on the LFPP diet (Figure S5A). Similarly, control mice continuously fed a HFHS diet had an elevated weight relative to LFPP controls (Figure S5B). On average, each 3 day diet period resulted in 3.40% ± 0.54% weight loss on the LFPP diet and 5.95% ± 0.56% weight gain on the HFHS diet (Figure S5C). Weight gain during the 5 week experiment was significantly higher in control mice continuously fed the HFHS diet, as compared to the mice on oscillating diets or LFPP-fed controls (Figure S5D). The mice on oscillating dietary regimens displayed excess caloric intake on the HFHS diet and diminished intake on the LFPP diet during the first day following each shift (Figures S5E and S5F).

Analysis of microbial community membership (Figure S6A) and structure (Figures 6A and 6B) over time revealed that the gut microbiota was rapidly and consistently reshaped by both diets. Control mice continuously fed the LFPP or HFHS diets were clearly distinguishable (Figure 6B). The observed changes to

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**Figure 3. Microbial Responses to the High-Fat, High-Sugar Diet in Transgenic Mice**

(A) Microbial community structure is primarily determined by diet (see PC1; F = 60.8, p value < 0.001, PERMANOVA on Bray-Curtis distances). Secondary clustering is by host genotype (see PC2; F = 17.9, p value < 0.001). Bray-Curtis dissimilarity-based principal coordinates analysis (PCoA) was performed on 16S rRNA gene sequencing data; the first two coordinates are shown (representing 37% of the total variance). Values are mean ± SEM (n = 15–20 samples/group). The asterisk represents significant differences at the final time point relative to wild-type controls (p value < 0.05, Kruskal-Wallis test with Dunn’s correction for multiple comparisons).

(B–F) Bray-Curtis-based PCoA of the fecal microbiota of animals on a LFPP (white filled symbols) or HFHS (black filled symbols) diet (n = 15 samples/group). Wild-type controls are included in all panels, indicated by white triangles (LFPP diet) and inverted black triangles (HFHS diet). Transgenic mice include: (C) MyD88.Delete−/− LFPP (white diamonds) and HFHS (black leftward triangles); (D) NOD2−/− LFPP (white circles) and HFHS (black circles); (E) ob/ob LFPP (white pentagons) and HFHS (black squares); and (F) Rag1−/− LFPP (white squares) and HFHS (black rightward triangles). See also Figure S1 and Tables S1 and S2.
overall microbial community structure corresponded to rapid shifts in the relative abundance of the two major bacterial orders in the distal gut, the Bacteroidales (phylum: Bacteroidetes) and Clostridiales (phylum: Firmicutes) (Figures 6C and 6D). These high-level changes occurred consistently during each successive diet shift and were detectable after only a single day. These trends were consistent with our previous experiments and significant at multiple taxonomic levels (Table S2A).

Figure 4. Microbial Responses Are Proportional to the Degree of Dietary Perturbation

(A–C) Physiological responses of mice to diets with differing HFHS contents: (A) food intake decreases as dietary HFHS content increases; nevertheless both (B) caloric intake and (C) body fat increase on HFHS-rich diets.

(D) Dose-dependent relationship between dietary HFHS content and the first principal coordinate from a Bray-Curtis dissimilarity-based PCoA of microbial community composition.

(E and F) Dose-dependent relationships between dietary HFHS content and the two most abundant diet-associated bacterial phyla: (E) Firmicutes increase with HFHS content and (F) Bacteroidetes decrease with HFHS content.

Within each panel, the upper graph (colored circles) represents data collected during gradient feeding, whereas the lower graph (gray squares) represents data collected during the baseline week, when mice had been assigned to a diet group but had not yet initiated gradient feeding. R^2 and p values reflect linear regression (n = 4–5 animals/group). See also Figure S1 and Tables S1 and S2.
Figure 5. A Rapid and Reproducible Microbial Response to the High-Fat, High-Sugar Diet in Outbred Mice
(A) Bray-Curtis-based PCoA of the fecal microbiota of animals on a LFPP (blue) or HFHS (red) diet. The first two principal coordinates are shown (representing 27% of the total variance), which clearly separate the 977 fecal samples by diet. (B) Analysis of the microbial response to the HFHS diet over time, using the first principal coordinate from the Bray-Curtis-based PCoA. Points are labeled based on the current diet: LFPP (blue) or HFHS (red). Samples were collected weekly, with daily sampling during the first week of the HFHS diet (indicated by the number of days post shift, dps). On average, 52 mice were sampled at each time point. Values are mean ± SEM. (C) Time map of consistently responsive species-level bacterial operational taxonomic units (OTUs) in outbred mice. The selected OTUs were present, responsive, and had consistent temporal patterns in ≥50% of mice. Each row represents a consensus temporal signature (aggregated model estimates across mice) for an OTU, ordered by agglomerative clustering of signatures. Blue indicates relative abundances below the mean abundance for the entire signature, and red indicates relative abundances above the mean. Values represent model estimates, in units of log transformed and standardized relative abundances. The taxonomic assignment for each OTU is indicated on the right of the heatmap: Bacteroidales (black), Clostridiales (orange), and Lactobacillales (pink). (D) Relaxation time constant distributions on the second HFHS diet regimen, for OTUs belonging to the bacterial orders Clostridiales and Bacteroidales (data from taxonomic assignment for each OTU is indicated on the right of the heatmap: Bacteroidales (black), Clostridiales (orange), and Lactobacillales (pink). Normalized abundance

To analyze the impact of successive dietary perturbations on the time-dependent responses of species-level bacterial OTUs, we modified our MC-TIMME algorithm to model temporal signatures using simple linear models with constant levels for each dietary regimen (LFPP and HFHS) and linear increases or decreases with subsequent dietary switches (oscillation number) (Figure 5A). Our model merged data from the staggered (counter-oscillatory) experiments to produce consensus signatures for each OTU (see Supplemental Experimental Procedures). One hundred and twenty-five OTUs were consistently responsive (present in >50% of mice, with significant changes on the first or last dietary shifts) and exhibited patterns of change in response to diet that were consistent in >50% of mice. Thirty-two OTUs exhibited dependence of their levels over time on the serial dietary switches (Figures 7A and S7A–S7E; Table S4B), whereas 93 had a consistent difference in abundance on the LFPP versus HFHS diets but displayed no detectable change in abundance after each sequential shift (Figures S6C, S7F, and S7G; Table S4C).

We performed three analyses to assess whether these trends were primarily dependent on the dietary oscillations, rather than being dependent on temporal drift caused by other host or environmental factors. First, we found no bias in the temporal consistency of the behavior of the detected OTUs between the two groups of mice subjected to staggered dietary oscillations (p value = 0.94, paired t test corrected for differing group sizes). Second, we tested if the 32 OTUs exhibiting dependence on the serial dietary shifts were also altered over time in control mice maintained on a constant LFPP or HFHS diet. Of the 32 OTUs, only 8 demonstrated any significant change in abundance.
over time on either the LFPP or HFHS diet (Table S4D; \( q \) value < 0.05, F test and slope \( R^2 > 0.1 \)).

Finally, we tested if the observed hysteresis patterns of species-level OTUs were accompanied by a change in the abundance of groups of functionally coherent bacterial genes. 16S rRNA gene sequencing data were used to predict the abundance of enzyme-level orthologous groups, which were then filtered and clustered according to their temporal dynamics with MC-TIMME (see Experimental Procedures). We identified 47 clusters of orthologous groups (containing on average 68 orthologous groups each) with a consistent difference in abundance on the LFPP versus HFHS diets (Figure 7B). Thirty-seven of these clusters exhibited dependence of their levels over time on the serial dietary switches (hysteresis, using the same criteria as for OTUs). The clusters that consistently increased on the HFHS diet were significantly enriched for orthologous groups from pathways for the metabolism of sucrose, the dominant carbohydrate in the HFHS diet, including a phosphotransferase system for sucrose import (K02808/K02809) and a key enzyme for sucrose catabolism (levansucrase, K00692) (Table S4E). Of note, on the HFHS diet we also found a steady increase in the abundance of orthologous groups for urea metabolism (found within the arginine metabolism pathway), including urease (K01428-30), allophanate hydrolase (K01457), and urea carboxylase (K01941). We also observed significant enrichments for orthologous groups in the sucrose and arginine metabolism pathways in our outbred mouse time series experiment (data not shown).

Taken together, these results suggest that gut microbial community structure and metabolic activity are, at least in part, determined by prior dietary history (i.e., oscillation number) and not simply by current dietary intake. Our use of a relatively simple linear model provides a conservative estimate of these effects, which could conceivably include changes in equilibration time.
with serial dietary shifts or nonlinear dependencies on oscillation number that our model did not capture.

**DISCUSSION**

A recent endpoint analysis of 52 matched inbred strains of mice fed a comparable LFPP or HFHS diet reported a significant association between host genotype and microbial community structure (Parks et al., 2013). However, our current results, based on both endpoint and extensive time series analyses, emphasize that the microbial response to the consumption of the HFHS diet consistently overshadows preexisting genetic associations. We found that (1) the gut microbia of inbred, transgenic, and outbred mice were consistently shaped by diet despite widely divergent genotypes, (2) interstrain and interindividual differences in community structure were reset after dietary perturbation, and (3) the changes in microbial community structure were rapid, reproducible, and reversible. These results suggest that environmental factors, namely host diet, play a dominant role in shaping gut microbial ecology.

The robust linkage between dietary intake and gut microbial community structure is exemplified by our analysis of C57BL/6J mice alternating between the LFPP and HFHS diets every 3 days. Remarkably, each diet shift rapidly and reproducibly reshaped the gut microbiota, detectable after a single day. Additional work is necessary to identify the mechanisms responsible for the observed changes to the gut microbiota during the consumption of the HFHS diet. Although total caloric intake has been associated with changes to the human gut microbiota (Jumpertz et al., 2011), studies in both mice and humans suggest that these effects are minimal relative to that of dietary composition (David et al., 2014; Ravussin et al., 2012). But what components of the LFPP and HFHS diets matter most? The answer will likely require systematic comparisons of defined mixtures that span the nutritional space encompassed by these two diets (Faith et al., 2011). Whether changes in the microbial community

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**Figure 7. Identification of Bacterial Species and Genes Dependent on Prior Dietary Intake**

(A) Relative abundance of species-level OTUs that were consistently present, responsive to diet, had consistent temporal patterns, and exhibited dependence of levels on serial dietary changes (see Experimental Procedures for thresholds used). Each row represents a temporal signature for an OTU (model estimate from combined data from the staggered dietary oscillation groups). Blue indicates relative abundances below the mean abundance for the entire signature, and red indicates relative abundances above the mean. Values represent model estimates, in units of log transformed and standardized relative abundances. The top 37 clusters (above the white line) exhibited dependence of their levels over time on the serial dietary switches (hysteresis).

(B) Bacterial gene content (KEGG orthologous groups) was inferred using an ancestral state reconstruction method (Langille et al., 2013). The MC-TIMME algorithm identified 47 clusters of orthologous groups (mean of 68 orthologous groups per cluster) showing consistent differences in abundance on the LFPP versus HFHS diets. PTS, phosphotransferase system. Each row in the time map represents a consensus temporal signature for the indicated cluster. Blue indicates relative abundances below the mean abundance for the entire signature, and red indicates relative abundances above the mean. Values represent model estimates, in units of log transformed and standardized relative abundances. The top 37 clusters (above the white line) exhibited dependence of their levels over time on the serial dietary switches (hysteresis).

See also Figures S1, S4, S6, and S7 and Tables S1 and S4.
arise directly from the altered nutrient environment in the gastrointestinal tract or indirectly due to effects of altered diets on host physiology that are consistent across genotypes, remains a fascinating area for future inquiry.

These studies also provide a basis from which to consider the many functional consequences of a gut microbiota altered by diet. Rapid restructuring of the gut microbiota may contribute to the beneficial effects of time-restricted feeding (Hatori et al., 2012) and could broadly alter the microbial contributions toward metabolism and immunity. It will be important to determine if similar shifts occur in the context of humans undergoing analogous “yo-yo” diets reflecting alternating periods of increased and decreased caloric intake (Atkinson et al., 1994). More broadly, it is interesting to consider whether or not the observed microbial plasticity may be a selective trait that helped our ancestors maintain energy balance given a volatile diet that was dependent on season and foraging success (Hawkes et al., 1991). Diet-induced changes in the gut microbiota could also alter key metabolic activities, as shown recently for the interaction between dietary protein and the bacterial inactivation of cardiac drugs (Haiser et al., 2013).

Our data also support a role for host genetics and/or other environmental factors in mediating variations in community structure within each dietary regime. Controlling for diet, microbial communities clustered by genotype among inbred strains and transgenic mice. We identified 12 outbred mice with divergent community structure (2 that resembled LFPP during consumption of the HFHS diet, and 10 with the opposite response; Figure S3B). Our computational modeling also highlighted species-level bacterial OTUs that behaved inconsistently between mice. The range of relative abundances found within each diet group was substantial, especially for the Akkermansia genus (phylum: Verrucomicrobia), which dominated the gut microbiota of some mice and was nearly undetectable in many others (Figures S3C and SSD and Supplemental Results). Additional studies are warranted to determine what other host and/or environmental factors may drive variations in microbial community structure within and between animal facilities (Ivanov et al., 2009).

These results also demonstrate that during dietary perturbations the gut microbiota can exhibit hysteresis: the dependence of a complex system not only on its current environment but also on past exposures. These effects have been described previously in a wide range of fields, including ecology (Scheffer et al., 2001), immunology (Das et al., 2009), developmental biology (Sha et al., 2003), economics (Blanchard and Summers, 1986), and physics (Jiles and Atherton, 1986). There is also preliminary evidence for hysteresis in the human gut microbiota following repeated exposure to broad-spectrum antibiotics (Gerber et al., 2012). Additional work is needed to test the underlying source of this effect, whether it is due to a dynamic lag between the dietary input and the change in bacterial abundance (i.e., rate-dependent hysteresis) or to another as-of-yet unknown mechanism that allows the gut microbiota to remember its past dietary exposures (e.g., through bacterial interaction networks) (Levy and Borenstein, 2013).

Finally, this study provides a cautionary note for ongoing efforts to link host genetics to the composition and function of the human gut microbiota. Perhaps more so than any other complex trait, the gut microbiota is shaped by a wide range of environmental factors, including diet (David et al., 2014), antibiotics and other drugs (Maurice et al., 2013), and geography (Yatsunenko et al., 2012). It will be difficult to identify host factors that are reproducibly associated with alterations to the gut microbiota without adequately controlling for these environmental factors. We anticipate that studies of large cohorts of outbred mice, fed a wide range of diets, coupled to experiments with inbred, transgenic, and gnotobiotic animal models will be an important strategy to elucidate the complex interrelationships between diet, host genetics, and the gut microbiota.

**EXPERIMENTAL PROCEDURES**

**Animal Husbandry**

Animal procedures were approved by the Jackson Laboratory and Harvard University. The LFPP diet (LabDiet; SK52) contained 16.0% kcal from fat (primarily soybean oil) and 61.7% from carbohydrates (primarily whole grains). The HFHS diet (Harlan Laboratories; TD.08811) contained 44.6% kcal from fat (primarily milk-derived saturated fat) and 40.6% from carbohydrates (sucrose, maltodextrin, and corn starch). Mice in the dietary gradient experiment were fed custom diets representing defined mixtures of TD.08811 and TD.96338, a Harlan product that is the nutritional equivalent of LabDiet SK52.

**16S rRNA Gene Sequencing and Analysis**

DNA was extracted using the PowerSoil bacterial DNA extraction kit (MoBio) and PCR amplified using barcoded universal bacterial primers targeting variable region 4 of the 16S rRNA gene. Barcoded amplicons from multiple samples were pooled and sequenced using the Illumina HiSeq platform. 16S rRNA gene sequences were analyzed on the Harvard Odyssey cluster using the QIIME (Quantitative Insights Into Microbial Ecology) software package (Caporaso et al., 2010) along with custom Perl scripts. LefSe was used to identify taxa associated with each experimental group (Segata et al., 2011).

**Data Preprocessing for Dynamical Modeling**

An operational taxonomic unit (OTU) was defined as present in a given mouse if there were ≥10 sequence counts at ≥5 time points. For the outbred mouse time series, we defined each OTU as responsive in a given mouse if its relative abundance on the first LFPP diet interval was significantly different from that on the final HFHS diet interval (q value < 0.05, Wilcoxon rank-sum test) (Bennamini and Hochberg, 1995). For the dietary oscillation experiment, we defined each OTU as responsive in a given mouse if its relative abundance on all LFPP time points was significantly different compared to its relative abundance on all HFHS time points (q value < 0.05, Wilcoxon rank-sum test).

**Modeling of Dynamics**

We customized the MC-TIMME model (Gerber et al., 2012) to capture dynamics of the outbred mouse time series and dietary oscillation experiments. This model uses nonparametric Bayesian methods to infer prototypical patterns of change in OTU relative abundances over time (signatures), and groups of OTUs with similar temporal behavior (signature groups). For the outbred mouse experiments, we used exponential relaxation process models to capture the dynamics of dietary shifts. For the dietary oscillation experiments we adjusted the MC-TIMME model to assess the dependence of changes in the microbiota on repeated dietary shifts. See Supplemental Experimental Procedures for justifications of our assumptions and a detailed model validation.

**Analysis of Gene and Pathway Abundances**

PICRUSt version 0.9.1 (Langille et al., 2013) was used to impute the abundance of KEGG orthologous groups (KOs). We retained KOs with >100 predicted counts in ≥5 time points in >50% of mice and a significant response to diet in >50% of mice (Wilcoxon rank-sum test, q value < 0.05). Postfiltering, we clustered the remaining KOs based on their temporal abundance patterns using the MC-TIMME algorithm. We then used the hypergeometric test to evaluate pathway-level enrichment.
Statistical Analysis
Statistical analyses were performed using GraphPad Prism. Data are expressed as mean ± SEM, and significance was set at a two-tailed p value < 0.05, unless otherwise indicated.

ACCESSION NUMBERS
Sequencing reads are in MG-RAST (Meyer et al., 2008) under the accession number 11172.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures, seven figures, and four tables and can be found with this article online at http://dx.doi.org/10.1016/j.chom.2014.11.010.

AUTHOR CONTRIBUTIONS
R.N.C., G.K.G., J.M.L., and P.J.T. designed the experiments. R.N.C., J.M.L., and L.S. did the animal husbandry. R.N.C., J.M.L., and P.J.T. did the microbial sequencing and analysis. G.K.G. constructed and analyzed the computational model. D.M.G. performed host genotypic analyses. R.N.C., G.K.G., and P.J.T. prepared the manuscript with comments from all of the other authors.

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