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Fecal Viral Community Responses to High-Fat Diet in Mice

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ABSTRACT Alterations in diet can have significant impact on the host, with high-fat diet (HFD) leading to obesity, diabetes, and inflammation of the gut. Although membership and abundances in gut bacterial communities are strongly influenced by diet, substantially less is known about how viral communities respond to dietary changes. Examining fecal contents of mice as the mice were transitioned from normal chow to HFD, we found significant changes in the relative abundances and the diversity in the gut of bacteria and their viruses. Alpha diversity of the bacterial community was significantly diminished in response to the diet change but did not change significantly in the viral community. However, the diet shift significantly impacted the beta diversity in both the bacterial and viral communities. There was a significant shift away from the relatively abundant Siphoviridae accompanied by increases in bacteriophages from the Microviridae family. The proportion of identified bacteriophage structural genes significantly decreased after the transition to HFD, with a conserved loss of integrase genes in all four experimental groups. In total, this study provides evidence for substantial changes in the intestinal virome disproportionate to bacterial changes, and with alterations in putative viral lifestyles related to chromosomal integration as a result of shift to HFD.

IMPORTANCE Prior studies have shown that high-fat diet (HFD) can have profound effects on the gastrointestinal (GI) tract microbiome and also demonstrate that bacteria in the GI tract can affect metabolism and lean/obese phenotypes. We investigated whether the composition of viral communities that also inhabit the GI tract are affected by shifts from normal to HFD. We found significant and reproducible shifts in the content of GI tract viromes after the transition to HFD. The differences observed in virome community membership and their associated gene content suggest that these altered viral communities are populated by viruses that are more virulent toward their host bacteria. Because HFD also are associated with significant shifts in GI tract bacterial communities, we believe that the shifts in the viral community may serve to drive the changes that occur in associated bacterial communities.

KEYWORDS virome, gut, microbiome, 16S rRNA, antibiotic perturbations, antibiotics, metabolism, high-fat diet

Obesity is a problem worldwide, and much of the epidemic in the Western world is related to dietary excess. The consumption of diets high in fat content can be directly correlated with obesity (1–3) and its associated conditions, including diabetes (4), coronary artery disease, obstructive sleep apnea, and fatty liver (5–7). More recent work has demonstrated that the vast communities of microbes inhabiting human body surfaces (8), collectively known as the microbiome, also are affected by alterations in

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diet; these communities characteristically change in response to high-fat diets (HFD) (9–11). The organisms forming the bacterial biota in the human gastrointestinal (GI) tract have been the primary microbes studied in response to diet changes, and their responses to HFD suggest that they may play central roles in the metabolic changes that result in both obesity and diabetes (12–15).

Our body surfaces also are inhabited by robust communities of viruses (16–20), many of which are bacteriophages, whose role and responses to shifts in diet have not been well characterized (16, 21). The relationships of bacteriophages and their prokaryotic hosts may be antagonistic (killing their hosts) or mutualistic (integrating into host genomes and potentially providing beneficial gene functions) (22). Because these viral communities have the capacity to alter the resident bacterial biota, they could play important roles in shaping cellular microbiomes.

The study of the human virome lags considerably behind that of the bacteriome. Particular viruses may persist in the human gut virome, with characteristics reflecting the diet of the human host (16). This relationship may primarily represent changes in gut bacteria and with their associated phages (16). In mice, antibiotic administration increases the reservoir for putative antibiotic resistance genes in the virome (23). Gut bacteria also respond to bacteriophage-mediated perturbations (24). There is an approximate 20:1 ratio of virus particles to bacterial cells in the gut mucosa, which reflects the relationship of most of these phages with their host bacteria but may also reflect the ability of some bacteriophages to bind to mucosal layers in the gut (25). Binding of phages to mucosal layers serves as one type of de facto immune system, protecting the host from susceptible invading bacteria (25). Phages bound to mucosal layers may have broader host ranges than can be observed using plaque assays (26), reflecting their putatively broad immune functions based on their parasitism. As with humans (16), mouse gut viromes also respond to diet changes, with temperate phages that are associated with phylum Bacteroidetes becoming more prominent in mice receiving HFD (21).

The gut epithelial barrier may become impaired in mice receiving HFD (27, 28), due to reduced tight junction protein expression (29), with increased gut permeability, translocation of bacterial lipopolysaccharides into the bloodstream, and increased intestinal inflammation in humans (4, 30) and in mice (31). HFD also have been associated with downregulation of immunoglobulin synthesis and of components of the major histocompatibility complex, which may have further downstream immune effects (32).

Previously, we have studied the effects of exposure to low-dose penicillin during maturation and determined that it has long-term metabolic effects in mice (33). These mice also develop enhanced insulin resistance and liver disease when fed HFD (34). The effects of low-dose penicillin G appear to be microbiota related, as phenotypes associated with the microbes are transferable to germfree mice (35). In this study, we utilized the low-dose penicillin G model from our prior studies to characterize the effects of HFD on gut microbiota over 28 weeks (36). Our goals were to identify changes in the taxonomy of both gut bacterial and viral communities in response to HFD and to characterize the overall virome community composition and structure in response to diet changes.

RESULTS

Experimental design and G+C content differences. We exposed C57BL/6J pregnant female mice to low-dose penicillin G (subtherapeutic antibiotic treatment [STAT]) in their drinking water (Fig. 1), which continued until their pups were 28 days old and had been weaned (33, 35, 36). Control mice and their pups were exposed in parallel to untreated water without antibiotics (CTL). To study the transmissible factors of the microbiota, pups were litter-mixed at day 32 of life and half of the CTL and half of the STAT group were cohoused at a 1:1 ratio to create 4 distinct groups (36). The CTL group \((n = 6)\) consisted of mice that had only been housed with other CTL mice. The STAT group \((n = 6)\) consisted of mice that had been housed only with other STAT mice. The
CTL-COHO group (n = 6) was composed of mice that had never been exposed to penicillin G and then were cohoused with mice that had received penicillin G. The STAT-COHO group (n = 6) consisted of mice that had received penicillin G until day 28 of life and then were cohoused with CTL mice. All mice remained with these cagemates for the duration of the study. Fecal samples were collected and combined from the same six females per group at weaning (week 4, sample 1) and 5 days after cohousing (week 5, sample 2). All groups were exposed to normal diets from weeks 4 to 15 and then were transitioned to a 45% high-fat diet (HFD) from weeks 16 to 36. Fecal samples were collected just before (week 15, sample 3) and just after (week 16, sample 4) transition to the HFD. The final sample was collected at week 28 of life (sample 5) after the mice had consumed HFD for 12 weeks (Fig. 1). Feces from each sample (samples 1 to 5) were pooled for each of the four treatment groups (CTL, STAT, CTL-COHO, and STAT-COHO), yielding 20 total samples.

We isolated the viromes from feces of each group of mice, as described previously (17), which involved sequential filtering to remove cellular debris, CsCl density gradient ultracentrifugation, and DNA extraction from intact virions. Resulting DNA was sequenced using semiconductor sequencing (37) for a total of 10,590,368 reads after quality filtering, with a mean length of 216 nucleotides. We sequenced 2.65 million reads per group with 0.53 million reads per time point collected. The average G+C content for all the quality reads was 42.3% overall; however, there were significant differences (P = 0.009, t test) in G+C contents of viromes from mice receiving normal diets (40.8%) compared to those from mice on HFD (44.5%) (see Fig. S1 in the supplemental material). These differences suggested a shift in the composition of the viral community with the shift to HFD.

Changes in alpha diversity after transitioning to HFD. We assembled virome reads from each time point to construct larger viral contigs to improve searchability for sequence similarities. For each sample, we assembled a mean of 3,572 ± 1,497 contigs, with 94.9% ± 1.8% of the total sequence reads assembling into viral contigs (Table S1). We first examined the alpha diversity of both the viral and bacterial communities. The alpha diversity for both the viral and bacterial communities was not significantly
different based on STAT and cohousing status both when mice were fed normal chow compared to an HFD (P > 0.05, analysis of variance [ANOVA]). Additionally, when mice were separated into their respective STAT and cohousing status groups, there was no significant differences in either the bacterial or viral diversity based on diet (P > 0.05, t test [Fig. 2A and B]). When grouping mice regardless of STAT and cohousing status, we found that after the transition from normal chow to HFD, there was a nonsignificant increase in the alpha diversity of the viral communities (Fig. 2C). In contrast, in the bacterial communities, alpha diversity was significantly reduced after the transition to HFD (Fig. 2D), consistent with prior studies (38). We further parsed the alpha diversity by comparing each time point following transition from normal chow to HFD (Fig. 3A). In the short term, we identified a significant increase in alpha diversity in the viromes.
after the transition to HFD ($P = 0.03$, ANOVA); however, this trend in alpha diversity was significantly reduced by week 28 ($P = 0.005$, ANOVA) (Fig. 3A). Together, these data indicate that the community richness of the virome did not passively follow the bacteriome after the dietary transition to HFD.

**Beta diversity in response to high-fat diets.** We next examined the beta diversity among the viral communities across the dietary transition. We found that viral communities clearly differed before and after the transition in diet ($P = 0.001$ and $R^2 = 0.26$, Adonis [Fig. 4A]). The viral communities of mice were also distinct based on the time point from which they were derived ($P = 0.006$ and $R^2 = 0.09$, Adonis). A similar trend in response to HFD also was observed in bacterial communities, with two distinct groups based on diet ($P = 0.001$ and $R^2 = 0.83$, Adonis [Fig. 4B]). However, there was
no significant demarcation due to time ($P = 0.31$, Adonis). For both bacterial and viral communities, there was no significant difference based on STAT and cohousing status.

To determine whether there were significant differences in the numbers of homologous viruses shared between different groups of mice, we used permutation testing (Table 1), as we have described previously (39, 40). There was a significant conservation of viromes in mice receiving normal diets ($P = 0.02$), but none among the mice receiving HFD ($P = 0.32$). We also examined whether time might be a factor in the conservation of virome contents and found that the only significant conservation of virome contents occurred in the mice receiving HFD at week 28 ($P = 0.03$). There were no significant distinctions between the viromes of the mice based on their cohousing status. In total, these results indicate that the beta diversity of both the viral and bacterial communities was affected by the transition to HFD, with lesser effects of age and antibiotic or cohousing exposure status. There was no consistent variation in the viromes to indicate a link between mouse exposure to penicillin G early in life and the fecal virome compositions (Fig. 4 and Table 1).

**Gene content differences in viromes in response to diet.** Using BLASTX to identify virome contigs that had homologies to known viral genes in the NCBI Non-redundant (NR) database, we identified substantial numbers of homologues (Fig. S2). We found that 84% ± 9% of the reads were assembled into contigs with homologies to known viruses, 6% ± 3% were homologous to bacteria, and 10% ± 9% had no known homologies. It is important to note that there were bacteriophages with putative lysogenic lifestyles present in these viromes, which can result in homologies to bacterial genes in virome studies (17). However, there were no significant differences between the proportions of reads in contigs that were homologous to known viruses in the samples based on diet (83% in normal chow and 86% in HFD).

Next, we characterized the viral homologues according to function. The proportion of each of the phage structural genes, aside from the Collar, was significantly lower after the transition from normal chow to HFD ($P < 0.001$, t test [Fig. 5]). In addition, similar to what was observed previously, the proportion of phage structural genes was not significantly different based on STAT and cohousing status ($P > 0.05$, ANOVA [Fig. 5]). We also identified nonsignificant increases in the proportions of lysins ($P = 0.23$, t test) and concomitant, significant decreases in integrases ($P = 0.0002$, t test) associated with the transition to HFD (Fig. 6). Again, we found no significant difference in the proportion of integrases or lysins based on STAT and cohousing status ($P > 0.05$, ANOVA [Fig. 5]).

| Grouping          | % homologous within group$^a$ | % homologous between groups$^a$ | $P$ value$^b$ |
|-------------------|------------------------------|-------------------------------|-------------|
| **By diet**       |                              |                               |             |
| Normal chow       | 74.11 ± 9.54                 | 43.59 ± 16.8                  | 0.02        |
| HFD               | 50.00 ± 18.96                | 42.91 ± 17.24                 | 0.32        |
| **By time**       |                              |                               |             |
| 4 wks (normal chow) | 42.11 ± 10.40               | 32.00 ± 14.16                 | 0.31        |
| 5 wks (normal chow) | 58.07 ± 8.24                | 40.07 ± 18.38                 | 0.18        |
| 15 wks (normal chow) | 53.80 ± 9.56                | 35.39 ± 17.50                 | 0.19        |
| 16 wks (HFD)      | 33.43 ± 10.40                | 26.95 ± 8.44                  | 0.37        |
| 28 wks (HFD)      | 33.80 ± 9.87                 | 17.48 ± 5.38                  | 0.03        |
| **By cohousing**  |                              |                               |             |
| No cohousing      | 57.61 ± 20.11                | 56.28 ± 21.27                 | 0.41        |
| Cohousing         | 50.03 ± 18.72                | 55.93 ± 21.42                 | 0.61        |
| **By penicillin G exposure** | |                           |             |
| No penicillin     | 51.96 ± 25.64                | 54.68 ± 19.27                 | 0.52        |
| Penicillin        | 59.33 ± 16.99                | 54.15 ± 19.34                 | 0.45        |

$^a$Based on the mean of 10,000 iterations. A total of 1,000 random contigs were sampled per iteration.

$^b$$P$ value based on the fraction of times the estimated percent homologous contigs for each group exceeded that between groups. Bold text denote significant $P$ values ($P < 0.05$).
Moreover, we found that regardless of antibiotic exposure, the ratio of integrase to lysin genes was significantly lower at week 28 than at all weeks on the normal diet ($P < 0.05$, ANOVA [Fig. 6E]). These data suggest a shift in the types of viruses, according to lifestyle and host (bacterial) interactions.

**Shifts in virome contents.** We next examined the types of viruses present in the fecal viral communities. Using TBLASTX analysis, we profiled the optimal matches for each virome contig and categorized each by virus family. We weighted the results based on contig read coverage to provide additional representation to those contigs assembled from greater numbers of reads in each fecal virome. For the mice on normal chow, most contigs were homologous to caudoviruses, including the families Siphoviridae, Myoviridae, and Podoviridae (Fig. 7). However, with change to HFD, the relative proportions of caudoviruses decreased, while the proportions of phages from the family Microviridae increased substantially by week 28. There was little variation in the profiles that could be observed regardless of whether the mice had penicillin G exposure (STAT [Fig. 7A]) or not (CTL [Fig. 7B]) or were cohoused with mice that had been exposed to penicillin G (STAT-COHO [Fig. 7C]) or not (CTL-COHO [Fig. 7D]). To search for a more subtle signal, we combined the viromes from all mice on normal diets and all mice on HFD. We observed a significant reduction in the representation of Siphoviridae after the change to HFD and a significant increase in the representation of eukaryotic viruses Phycodnaviridae and Mimiviridae ($P < 0.05$, t test [Fig. 8]). Additionally, when grouping the HFD samples by week, the Siphoviridae decreased significantly at both week 16 and week 28 compared to the normal chow samples, while the proportion of Phycodnaviridae was only significantly higher at week 16 with the HFD and that of Mimiviridae only significantly higher at week 28 compared to that with normal chow ($P < 0.05$, ANOVA [Fig. S3]). Using a Pearson correlation test, we also determined that the relative abundance of Siphoviridae ($R = 0.69$ and $P = 0.0008$, Pearson) correlated positively and those of Microviridae ($R = -0.44$ and $P = 0.049$, Pearson), Mimiviridae ($R = -0.52$ and $P = 0.02$, Pearson), Phycodnaviridae ($R = -0.57$ and $P = 0.009$, Pearson), and Iridoviridae ($R = -0.45$ and $P = 0.047$, Pearson) correlated negatively with the percentage of integrase homologues.

**FIG 5** Assignment of virome contigs with homologues to structural genes in the viromes of mice, according to time point and diet type. The y axis represents the percentage of putative structural genes identified. Shape represents the different treatment groups: STAT, CTL, STAT-COHO, and CTL-COHO.
Characterizing the viral communities by quantifying the relative number of contigs assigned to specific viral families rather than the number of reads belonging to contigs that were homologous to specific virus families, we identified similarly significant trends (Fig. S4 to S6). Specifically, we found a significant decrease in the proportion of contigs assigned as *Siphoviridae* when the mice were transitioned to HFD. While we identified large numbers of reads assigned to the family *Microviridae* (Fig. 7), especially after transition to HFD, only a small proportion of contigs were assigned to this family (Fig. S4 to S6). These results indicate that there were relatively few highly abundant *Microviridae* family members. We also identified a greater proportion of contigs representing eukaryotic viruses, including *Phycodnaviridae*, *Herpesviridae*, and *Poxviridae*, in the mice after the transition to HFD (*p* < 0.05, t test [Fig. S5]). Although none of these viruses were highly represented according to read abundances (Fig. 7 and 8), the profiles of virus families changed significantly after the transition to HFD, regardless of the analytical methodology. We found similar, significant trends when we evaluated the proportion of contigs assembled from reads that were normalized to a minimum sampling depth (Fig. S5B and S6B).

**Changes in bacterial communities.** To determine whether shifts in virome contents followed changes in the compositions of the bacterial community, we characterized the fecal bacterial biota. Sequencing of the V1-V2 segment of 16S rRNA across the 20 samples showed that there were a total of 2,486,606 reads after quality filtering, with a mean length of 340 nucleotides. In total, there were ~497,321 reads per group, with ~124,330 reads per time point. The phylum *Bacteroidetes* dominated the bacterial community when the mice were fed normal chow (weeks 4 to 15), while *Firmicutes* predominated after the mice were begun on HFD (weeks 16 to 28), a shift that was statistically significant (*p* < 0.001) and was apparent regardless of cohousing status.
These data indicate that HFD also had significant and profound impacts on the fecal bacterial community, suggesting that the changes observed in viromes (Fig. 7) and the bacterial communities (Fig. 9) could be interrelated.

DISCUSSION

While it has been well described that changes in diet significantly affect gut bacterial communities (41, 42), substantially less is known about how viral communities respond. Some members of the human gut viral community are highly individual specific, appear remarkably stable over time, and have similar ecologies to one another based on diet.
In mice, luminal and mucosal viral communities differ, reflecting the associated bacterial biota in the GI tract, but HFD consumption resulted in greater numbers of temperate bacteriophages in the mucosa than in the lumen (21). Transitioning mice to HFD to allow direct comparisons of gut virome contents in the same mice before and after consuming HFD allowed us to examine this issue more precisely. The decreases in the proportions of all 3 major caudovirus families (Myoviridae, Siphoviridae, and Podo-viridae) after the transition to HFD, regardless of analytical method (Fig. 8 and Fig. S3 to S6), were consistent. However, in contrast to the previous research on mucosal and luminal viromes of mice fed an HFD (21), we identified significant decreases in the proportion of integrases, suggesting a reduction in the overall numbers of temperate bacteriophages. Indeed, bacteriophages from the family Siphoviridae, which often have lysogenic lifestyles (43, 44), were significantly reduced on HFD. This phage phenotype transition may have consequences for gut health, as a proper balance between lysis and lysogeny is suggested to maintain a healthy microbial community (45). Specifically, improper balance between these lifestyles in the gut is associated with leukemic diseases (46) and inflammatory bowel disease (47).

The significant changes we observed in the viromes were accompanied by shifts in

![Graphs](fig9.png)

**Fig 9** Bacterial taxonomy based on 16S rRNA sequences identified at the phylum level. The y axis represents the percentage of operational taxonomic units (OTUs) assigned to each phylum. The difference in the representation of the phyla Bacteroidetes and Firmicutes in mice on normal diets and HFD was statistically significant ($P < 0.001$) by two-tailed t test for each group.
the bacterial communities in response to HFD. After the transition to HFD, the shift from *Bacteroidetes* to *Firmicutes* (Fig. 9) was accompanied by significant reductions in cauloviruses and increases in *Microviridae*, especially at week 28 (Fig. 7 and Fig. S3). A recent study has found that microviruses are persistent inhabitants of the human gut (48). However, because they have been identified in both *Bacteroidetes* and *Firmicutes* (49), the source of their increased representation in this study could not be determined. We did not identify greater numbers of microviruses in response to the diet change (Fig. S5 and S6), but we did find rather significant changes in their relative abundances (Fig. 7 and 8). It is uncertain whether the virome differences merely reflect changes in the bacterial communities or whether phages actively promoted shifts in the bacterial communities; the observed shifts could reflect the *Firmicute* predominance if these microviruses infect *Firmicutes*, or conversely, they could represent increased predation of the *Bacteroidetes* if *Bacteroidetes* are their hosts.

Early-life STAT has a substantial impact on murine phenotypes after transitioning to HFD (34, 36), which is likely associated with the composition of the gut bacterial community (35). Our data did not indicate that the antibiotics had any significant effects on the composition of the gut virome (Fig. 4). In this case, the change caused by the diet transition may have masked any substantial effects of the STAT on the virome. Moreover, after transitioning to HFD, alpha diversity of the bacterial biota was significantly reduced, as observed previously (50–52), but the viral diversity did not change significantly in the long term (Fig. 2 and 3). We have previously observed a parallel trend in the GI tract of human subjects receiving long-term antibiotics (53), where viral diversity was not significantly affected despite the significant reduction in bacterial diversity. In that study, we observed greater numbers of eukaryotic viruses that likely compensated for the reduction in bacteriophages. In this study, we also observed an increase in the number of eukaryotic viruses following the transition to HFD, suggesting commonalities in the effect of antibiotics and diet on virome community composition (Fig. S4 and S5).

While the association of *Herpesviridae* and *Poxviridae* with human disease is well understood, the potential impact of *Mimiviridae* and *Phycodnaviridae* on health is only now being recognized. The family *Mimiviridae* has been linked to clinical cases of pneumonia, and recent evidence suggests that *Phycodnaviridae* may affect substantially more than just algal species (54–56). For example, the inoculation of a *Phycodnaviridae* virus (ATCV-1) into the intestinal tracts of 9- to 11-week-old mice resulted in a decrease in cognitive performance (56). However, it is important to note that *Mimiviridae* (size, ~0.7 μm) were likely unable to pass through the 0.22-μm filter used to isolate the viral community in this study. Therefore, the *Mimiviridae* homology identified here was likely due to smaller unknown viruses, such as virophage, carrying *Mimiviridae*-like elements (57, 58).

The beta diversity shifts in viromes and the bacterial microbiome were most pronounced between weeks 15 and 16 when the mice were transitioned to HFD (Fig. 4). While a few studies have characterized murine GI tract viromes (21, 24, 59, 60), none have observed shifts in bacterial and viral ecology in the absence of perturbations such as dietary shifts. We do not believe the shift in ecology between weeks 15 and 16 is due to age (61); it more likely represents a direct response to HFD. While there was a clear and significant distinction between the viral communities observed across the specific time points (Fig. 4A), none was observed within the bacterial communities (Fig. 4B). It is possible that these differences reflect methodologic variation (shotgun metagenomics for viruses versus 16S rRNA amplicon sequencing for bacteria).

The virome changes we observed after the transition to HFD included reductions in *Siphoviridae* accompanied by gains in others such as *Microviridae* and several eukaryotic viruses (Fig. 8 and Fig. S3). However, it is important to note that the amplification step (MDA) used in the production of the viromes is known to introduce biases (62, 63). The bacterial community changes included increases in *Firmicutes* with a concomitant decline in *Bacteroidetes* (Fig. 9). The reduction in integrase gene homologues found in the viral community of all four groups of mice (Fig. 6) after the start of the HFD is...
striking and is consistent with a global shift from a more temperate bacteriophage community to a community with more lytic phages. This phenotypic shift in the viral community was unanticipated but quite consistent, and it supports the hypothesis that the viral community plays a role in driving the significant changes observed in the bacterial communities and/or in maintaining these altered bacterial community phenotypes. In any event, exposure to HFD has led to this emergent property of the virome with a loss in abundance of integrase genes, which are markers for lytic versus lysogenic viral lifestyles.

Despite these novel findings, there were some limitations to our study. The long duration of our study limited our ability to reproduce the experiment multiple independent times. Because we observed no significant differences between STAT or cohousing status, these groups (the STAT and STAT-COHO groups and the CTL and CTL-COHO groups) may be considered replicates, with each group demonstrating similar, significant trends. Moreover, multiple confounding factors, such as genetic lineage and animal husbandry practices, may drive interlab variability in all aspects of the microbiome (64). Therefore, although these factors were controlled in this study, future studies should aim to ensure the generalizability and confirm the identities and relative abundances of the viral community members after the HFD transition in other genetic lines.

MATERIALS AND METHODS

Animal experiments. The protocol was approved by the New York University School of Medicine (NYUSoM) Institutional Animal Care and Use Committee (IACUC). The mice used in this experiment were part of a larger cohort described by Schulfer and colleagues (36). Male and female C57BL/6J (stock no. 000664) mice were received at 6 weeks of age from Jackson Laboratories and allowed to adjust to the NYUSoM animal facility for 1 week prior to breeding. After 5 days, breeding pairs were separated and pregnant dams were randomized into control (CTL) or subtherapeutic antibiotic treatment (STAT) groups. Penicillin G (6.8 mg/liter) was added or not (CTL) to dams’ drinking water at ~ day 14 of gestation, as described previously (49). To maintain a fresh supply of penicillin G, water containers were changed twice weekly. Pups were weaned at day of life 28, at which point antibiotics were stopped in the STAT group. All mice had ad libitum access to water and chow (Purina Mills International Diet 5001; 4.07 kcal/g with 13.5% kcal from fat) and were maintained on a 12-h light/dark cycle. After removing dams from pups and stopping antibiotic treatment at day 28 of life, pups were moved to clean cages for 3 to 4 days to remove residual antibiotics from their environment. After this washout phase, mice were randomly assigned to new cages. Half of each STAT or CTL treatment group was placed with other mice of the same treatment, and half were cohoused at a 1:1 ratio of STAT to CTL mice. At week 16, all mice were switched to HFD (4.73 kcal/g with 45% kcal from fat; rodent diet D12451; Research Diets, New Brunswick, NJ). For each treatment group, 6 mice were individually placed into sterile containers and fresh fecal pellets were collected from the container and immediately frozen at ~80 °C until use in this study. Fecal samples were collected at defined time points: week 4, week 5, week 15, week 16, and week 28. Mice continued on the treatment until week 36 as part of a larger cohort described by Schulfer and colleagues (36).

Analysis of viromes. Fecal viromes were prepared by diluting 0.4 g of feces in 4 ml of a sodium chloride, magnesium sulfate buffer (SM buffer) and vortexing for 40 min to separate viral particles, with spinning at 4,000 × g for 10 min to pellet the remaining solid material. The supernatants then were filtered sequentially using 0.45-μm and 0.2-μm filters (VWR) to remove cellular and other debris and then purified on a cesium chloride gradient according to previously described protocols (17). Only the fraction with a density corresponding to most known bacteriophages (65) was retained, further purified on Amicon YM-100 protein purification columns (Millipore, Inc.), treated with DNase I, and subjected to lysis and DNA purification using the Qiagen UltraSens virus kit (Qiagen). Recovered DNA was screened for the presence of contaminating bacterial nucleic acids by quantitative 16S rRNA gene PCR using primers 8F (AGAGTTTGATCCTGGCTCAG) and 357R (CTGCTGCCTYCCGTA) in Power SYBR green PCR master mix (Thermo Fisher Scientific). Viral DNA then was amplified using GenomiPhi Hy MDA amplification (GE Healthcare), fragmented to roughly 200 to 400 bp using a Bioruptor (Diagenode), and utilized as input to create libraries using the Ion Plus fragment library kit according to the manufacturer’s instructions. Libraries then were sequenced using 316 chips on an Ion Torrent personal genome machine. We trimmed sequence reads according to modified Phred scores of 0.5 using CLC Genomics Workbench 9.0, removed any low-complexity reads with ≥8 consecutive homopolymers, and removed any reads with substantial length variation (<150 nucleotides or >300 nucleotides) or ambiguous characters prior to further analysis. Each virome was screened for human nucleic acids using BLASTN analysis (E value < 10−7) against the human reference database available at ftp://ftp.ncbi.nlm.nih.gov/genomes/H_sapiens/. Any reads with significant similarity to human sequences were removed prior to further analysis using Ion Assist (www.thepridelaboratory.org).

Prior to assembly, read statistics (e.g., quantity, GC content) were calculated via the CLC Genomics Workbench 9.0. Sequence reads were then assembled using CLC Genomics Workbench 9.0 based on 98%
identity with a minimum of 50% read overlap, which were more stringent than criteria developed to discriminate between highly related viruses (66). Because the shortest reads were 50 nucleotides, the minimum tolerable overlap was 25 nucleotides, and the average overlap was no less than 100 nucleotides depending on the characteristics of each virome. The consensus sequence for each contig was constructed according to majority rule, and any contigs of <200 nucleotides or with ambiguous characters were removed prior to further analysis.

Virome contigs were annotated using BLASTX against the NCBI NR database with an E value cutoff value of 10^-5. Specific viral sequences were identified using Ion Assist (www.thepridelaboratory.org) by parsing BLASTX results for known viral genes, including replication, structural, transposition, restriction/modification, hypothetical, and other genes previously found in viruses for which the E value was at least 10^-3. Each individual virome contig was annotated using this technique; however, if the best hit for any portion of the contig was to a gene with no known function, lower-level hits were used as long as they had known function and still met the E value cutoff. Virus types were determined by parsing the virus families from the TBLASTX best hits of each viral contig with an E value of <10^-10. Analysis of shared sequence similarities present in each virome was performed by creating custom BLAST databases for each virome, comparing each database with all other viromes using BLASTN analysis (E value < 10^-10), and these compiled data were used to calculate Jaccard distances using Ion Assist (www.thepridelaboratory.org). These distances then were used as input for principal-coordinates analysis using QIIME (67). Alpha diversity was determined using the homologous virus diversity index (based on Shannon Diversity) as we have previously described (68, 69). Data were visualized using the R package ggplot2 ver. 3.1.0 (70).

**Statistical analysis of differences among viromes.** Statistically significant differences in gene categories (integrases and lysins), alpha diversity, relative abundances of viral families, and G+C content between HFD and normal chow groups were determined by two-tailed t tests. Differences in alpha diversity and the relative abundance of viral families with HFD grouped by week and the ratio of integrases and lysins by week were determined by analysis of variance (ANOVA) with post hoc Tukey HSD (honestly significant difference) test. To assess whether viromes had significant overlap within or between groups, we performed a permutation test using Ion Assist (www.thepridelaboratory.org) based on resampling (10,000 iterations). We previously have used this test to identify significant differences between viromes of individuals and different treatment groups (39, 40, 71). Briefly, we simulated the distribution of the fraction of shared virome homologues from 2 different groups within individual mice that were randomly chosen. For each set, we computed the summed fraction of shared homologues using 1,000 random contigs between and within different mice, and from these computed an empirical null distribution of our statistic of interest (the fraction of shared homologues). The simulated statistics within each mouse were referred to the null distribution of intermouse comparisons, and the P value was computed as the fraction of times the simulated statistic for the each exceeded the observed statistic. To perform statistical tests on viral beta diversities, represented in principal-coordinate analysis plots, we performed permutational multivariate analyses of variance (PERMANOVA) in the R Vegan package using the Adonis function with 999 permutations.

**Analysis of 16S rRNA.** Genomic DNA was prepared from the fecal pellets of each subject and time point using the QIAamp DNA Stool MINI kit (Qiagen). We amplified the bacterial 16S rRNA gene V1-V2 hypervariable region using the forward primer 8F (AGAGTTTGATCCTGGCTCAG) fused with the Ion Torrent adaptor A sequence and one of 70 unique 10-bp barcodes, and reverse primer 357R (CTGCTG CCTYCCGTA) fused with the Ion Torrent adaptor P1 from each donor and sample type (72). PCRs were performed using Platinum high-fidelity PCR SuperMix (Invitrogen) with the following cycling parameters: 94°C for 10 min, followed by 30 cycles of 94°C for 30 s, 55°C for 30 s, and 72°C for 30 s and a final elongation step of 72°C for 10 min. Resulting amplicons were purified on a 2% agarose gel stained with SYBR Safe (Invitrogen) using the MinElute PCR purification kit (Qiagen). Amplicons were further purified with Ampure XP beads (Beckman-Coulter), and molar equivalents were determined for each sample by quantifying the amplicons using PicoGreen (Invitrogen) using a plate reader. Samples were pooled into equimolar proportions and sequenced on 316 chips using an Ion Torrent PGM according to manufacturer’s instructions (Life Technologies) (37). Resulting sequence reads were removed from the analysis if they were <180 nucleotides or >500 nucleotides, had any barcode or primer errors, contained any ambiguous characters, or contained any stretch of >8 consecutive homopolymers. Sequences then were trimmed according to any site that had a Phred score of less than 15 (73). Sequences then were assigned to their respective samples based on a 10-nucleotide barcode sequence and were further processed to remove reads that were greater than 3 standard deviations from the mean read lengths in any specimen.

We sequenced a minimum of 10,000 reads from each sample and analyzed the sequence data using Quantitative Insights Into Microbial Ecology (QIIME 1.5) (67). Representative OTUs from each set were chosen at a minimum sequence identity of 97% using the QIIME script pick_otus_through_otu_table, which uses the Greengenes database (74). Principal-coordinate analysis was performed based on beta diversity using weighted UniFrac distances (75) via the QIIME script beta_diversity_through_plots. Alpha diversity using the Shannon diversity index (76) was determined using QIIME. Statistical differences in alpha diversity were determined by two-tailed t tests. For beta diversity, statistical differences were determined with PERMANOVA (999 permutations).

**Availability of data and material.** All sequences are available for download in the NCBI Sequence Read Archive under accession number PRJNA437977. Ion Assist software is available for download at www.thepridelaboratory.org and runs on Windows XP or higher.
SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, PDF file, 0.03 MB.
FIG S2, PDF file, 0.03 MB.
FIG S3, PDF file, 0.03 MB.
FIG S4, PDF file, 0.03 MB.
FIG S5, PDF file, 0.03 MB.
FIG S6, PDF file, 0.04 MB.

TABLE S1, DOCX file, 0.02 MB.

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A.S., M.J.B., and D.T.P. conceived and designed the experiments. A.S., T.M.S.-R., and M.L. performed the experiments. T.M.S.-R., A.S., J.C., J.M.B., M.J.B., and D.T.P. analyzed the data. A.S. and M.J.B. contributed reagents. A.S., M.J.B., J.C., J.M.B., and D.T.P. wrote the manuscript.

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We declare no competing interests.

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