Community-Wide Response of the Gut Microbiota to Enteropathogenic Citrobacter rodentium Infection Revealed by Deep Sequencing

Christian Hoffmann,1 David A. Hill,2 Nana Minkah,1 Thomas Kirn,2 Amy Troy,2 David Artis,2* and Frederic Bushman1*

University of Pennsylvania School of Medicine, Department of Microbiology, Philadelphia, Pennsylvania 19104,1 and University of Pennsylvania School of Veterinary Medicine, Department of Pathobiology, Philadelphia, Pennsylvania 191042

Received 4 May 2009/Returned for modification 6 May 2009/Accepted 22 July 2009

We investigated the spatial and temporal response of the murine gut microbiome to infection with Citrobacter rodentium, an attaching-and-effacing bacterium that provokes innate and adaptive immune responses, resulting in transient bacterial colitis. Previous studies have suggested that C. rodentium-induced inflammation is associated with an increased abundance of Enterobacteriaceae. We report here a deeper analysis of this model using DNA bar coding and 454 pyrosequencing to characterize 101,894 partial 16S rRNA gene sequences from 85 microbial samples from tissue-adhered and luminal bacteria of the cecum, proximal colon, and distal colon, which allowed us to identify previously unappreciated spatial and kinetic changes in multiple bacterial lineages. The deep sequencing data revealed that C. rodentium was most abundantly associated with the cecal mucosa at day 9 postinfection and then diminished in abundance, providing the first reported use of deep sequencing to track a pathogen in vivo through the course of infection. Notable changes were associated with both the mucosally adhered and luminal microbiota at both day 9 and day 14 postinfection. Alterations in abundance were seen for Proteobacteria, Deferribacteres, Clostridia, and others; however, changes in Enterobacteriaceae could be accounted for by the presence of C. rodentium itself, which is a member of this family. The Lactobacillus group decreased in abundance during infection, which may be important for pathogenesis because members of this lineage modulate the composition of the gut microbiota and are used as probiotics. Thus, deep sequencing provides previously inaccessible information on how Citrobacter infection and clearance reshapes the gut microbial community in space and time.

The human intestine is home to some 100 trillion microorganisms. The density of bacterial cells in the colon has been estimated at $10^{11}$ to $10^{12}$ per ml, which make the colon one of the most densely populated microbial habitats known on Earth (22, 42). The pooled genome size of the intestinal bacteria is estimated to exceed the size of the human nuclear genome by 2 orders of magnitude (22, 43). Mutualistic bacteria allow digestion of otherwise undigestible foods, protect against epithelial injury, regulate fat metabolism, boost intestinal angiogenesis, and promote proper development of the immune system (see reference 43 and references therein). The great majority of these gut bacterial species have not been cultured outside the mammalian host, so most species are little studied. Recent reports have surveyed the intestinal microbiota using DNA sequencing of uncultured communities (9, 21, 29, 34), microarray-based methods (31, 32), and targeted full-genome sequencing (12). Many of these studies used 16S rRNA sequences to query the different bacterial taxa, since 16S sequencing (12). Many of these studies used 16S rRNA sequencing to track a pathogen in vivo through the course of infection. Notable changes were associated with proteobacterial, Deferribacteres, Clostridia, and others; however, changes in Enterobacteriaceae could be accounted for by the presence of C. rodentium itself, which is a member of this family. The Lactobacillus group decreased in abundance during infection, which may be important for pathogenesis because members of this lineage modulate the composition of the gut microbiota and are used as probiotics. Thus, deep sequencing provides previously inaccessible information on how Citrobacter infection and clearance reshapes the gut microbial community in space and time.

(30). These and other studies have revealed that major bacterial phyla in the mammalian gut include Bacteroidetes, Firmicutes, Proteobacteria, Actinobacteria, and Fusobacteria.

Inappropriate innate and adaptive immune responses to the commensal gut bacteria are implicated in the pathogenesis of inflammatory bowel disease (IBD; e.g., see references 6, 11, and 37). The development of IBD in rodent models can be suppressed by raising animals in a germfree environment, where commensal bacteria are absent, but can be induced in IBD-prone animals after colonization. Both animal models of IBD and human IBD patients are characterized by an increase in mucosal adherence and translocation of bacteria, T-cell responses to bacterial antigens, and amelioration with antibiotic treatment.

A few studies have begun to characterize the extensive variation in the human-associated microbiota at different anatomic sites (2, 9, 13, 29). For example, populations of 16S rRNA gene sequences have been isolated from the mouth (33), stomach (2), six major subdivisions of the colon (cecum, ascending colon, transverse colon, descending colon, sigmoid colon, and rectum), and stool (9, 13). A recent comparison among human individuals indicate extreme variation in gastrointestinal (GI) bacterial communities, to the point where it is unclear whether a “core” human microbiome even exists (43). In comparison, laboratory mice show much more similarity between individual animals, due at least in part to the homogeneity of diet and genotype possible with these models.

Relatively few studies have investigated the spatial and temporal effects of immune activation on the gut microbiota, which is critical to understanding the factors leading to disregulation...
during IBD. In one study (27), inflammation was induced in mice by infection with *Citrobacter rodentium* as a model for enteropathogenic *Escherichia coli* infection in humans or by chemical means. Studies using fluorescence in situ hybridization and Sanger sequencing suggested that *Enterobacteriaceae* were increased in abundance in the colonic microbiota with *C. rodentium* infection. An increase in *Enterobacteriaceae* was also suggested to be associated with chemical induction of inflammation by treatment with dextran sodium sulfate or deletion of the regulatory cytokine IL-10 but not with infection with *Campylobacter jejuni*, which does not cause inflammation. In humans, a variety of changes in gut microbial communities have been proposed to accompany GI inflammation, but the generality of observations to date is uncertain, particularly because diet has profound effects on gut microbial composition and has not been controlled in human (21, 39) or macaque (21, 29) studies.

Given that alterations in gut microbial communities are widely expected to be associated with disease states but that the nature of these transitions are mostly unknown, we investigated how infection with *C. rodentium* affects the spatial and temporal dynamics of the full gut microbial community. We used DNA bar coding (3, 4, 14, 15) and pyrosequencing (28) to characterize a total of 85 microbial communities, 48 from infected mice and 37 from uninfected mice, by sequencing 102,398 gene segments from the 16S rRNA gene. Luminal contents and tissue-associated bacteria obtained from the cecum, proximal colon, and distal colon were compared. Samples were compared at day 9 after infection, the time of maximal colonization by treatment with dextran sodium sulfate or deletion of the regulatory cytokine IL-10 but not with infection with *Campylobacter jejuni*, which does not cause inflammation. In the present study, we observed an increase in *Enterobacteriaceae* upon infection, as was previously reported (27), but in contrast to the previous report, the increase was accounted for by colonization with *C. rodentium* itself, which could be quantified in the deep-seqencing data. A statistically rigorous analysis revealed alterations in the *Proteobacteria, Deferribacteres, Clostridia*, and *Lactobacillus* lineages accompanying infection, and their distributions differed in time postinfection (p.i.) and in space within the gut. Thus, this comprehensive study of the progression of a bacterial infection revealed unanticipated and extensive changes in bacterial community structure in space and time.

### MATERIALS AND METHODS

**Animals, bacterial infections, cytokine responses, histological analysis, and sample acquisition.** Twenty C57BL/6 mice in three litters (both males and females) were obtained from Charles River Laboratory. Animals were housed as littersmates from birth and throughout the study. Four-week-old animals were orally gavaged on day 0 with $5 \times 10^6$ *C. rodentium* CFU strain DBS100 (formerly *C. freundii* biotype 4280 [provided by Bruce Vallance, University of British Columbia, Vancouver, British Columbia, Canada]). Mice were fed acidified water and autoclaved LabDiet 5010 mouse chow (LabDiet). Areas of the large intestine areas were defined by anatomical criteria, i.e., the cecum, and the proximal (1-cm portion immediately following the cecum) or distal (the last 1-cm portion of the large intestine immediately adjacent to the anus) colon. The expression of *Ifn*/*Ifng*, *Il-22*, *Il-17a*, and *Il-17f* was analyzed by intestinal RNA isolation using an RNAseq kit (Qiagen, Inc., Valencia, CA), followed by cDNA generation and real-time PCR using SYBR green technology (Applied Biosystems, Foster City, CA) and transcript-specific primers (Qiagen). For real-time PCR analysis of gene expression, RNA was isolated from a section of the mid-colon. The primers used are specific to the cytokine transcripts for gamma interferon (IFN-γ), interleukin-22 (IL-22), IL-17a, and IL-17f and were either purchased from Qiagen (IFN-γ [QT0138821]) and IL-22 ([QT0102823)]) or synthesized (IL-17a, forward [5′-TCCAGAAGGGCCCTGACTA-3′] and reverse [5′-TTCAATGCCTTGAGAGATC-3′]; IL-17f, forward [5′-CCTGAGTACACTGTTGAG-3′] and reverse [5′-AATTGACGAGAAAGAAATG-3′].

To determine the number of *C. rodentium* CFU per gram of stool, four C57BL/6 mice obtained from Charles River Laboratory were orally gavaged on day 0 with $5 \times 10^5$ naldixic acid-resistant *C. rodentium* CFU, and pellets were collected at days 0, 6, 11, 20, and 27. Pellets were homogenized and plated at appropriate dilutions on LB-Nal plates. CFU were enumerated after overnight incubation at 37°C.

Recovery of 16S DNA was analyzed by quantitative TaqMan PCR using universal 16S primers to determine the numbers of 16S DNA copies per microgram of total DNA. Comparison of infected and uninfected animals showed no decrease in abundance upon infection, in contrast to a published report (27). The only statistically significant differences were for the 2 of 12 samples (tissue associated from the distal colon [day 9] and luminal contents from the distal colon [day 9]), where the number of 16S copies actually increased after *Citrobacter rodentium* infection.

For historical analysis, IkkKb/Il-10−/− mice on a mixed B6×129 background of wild-type mice obtained from Charles River Laboratory (cecum and colon samples, respectively) were orally gavaged on day 0 with $5 \times 10^5$ *C. rodentium* CFU and euthanized at day 13 or 14 p.i. Paraaffin-embedded 4% paraformaldehyde-fixed cecal and colon tissue sections were stained with hematoxylin and eosin (H&E). To separate luminal from mucosal attached bacteria, luminal content was removed by sterile squeezing from 1-cm sections of intestine (luminal bacteria). The intestinal samples were then cut open and gently washed by shaking the tissue in sterile phosphate-buffered saline. Whole DNA was isolated from these washed tissue samples (mucosal attached bacteria). For histological analyses, samples were collected from either commercially purchased C57BL/6 mice or B6 mice in which an IkkKb/Il-10−/− allele was inserted. IkkKb/Il-10−/− mice were generated using the 129 embryonic stem cell line and backcrossed at least six times to a C57BL/6 background. In all of our studies with these mice, they were immunologically and pathologically indistinguishable from C57BL/6 mice. Stool samples were obtained at the time of voiding (fresh from the mice). Pellets were frozen and −80°C for up to several weeks. DNA was extracted from tissue frozen at −80°C.

All experiments were performed following the guidelines of the University of Pennsylvania Institutional Animal Care and Use Committee.

**DNA manipulations.** Total DNA from stool samples was extracted from frozen samples by using a QIAamp DNA stool mini kit (Qiagen) according to the manufacturer’s protocol for pathogen detection. Total DNA extracted from tissue samples was obtained by using the DNeasy blood and tissue kit (Qiagen) according to the protocol for DNA extraction of gram-positive bacteria. The 16S rRNA gene fragment used in the present study was obtained as previously described (29). Briefly, each sample was amplified by using a bar-coded primer pair (3, 14, 15) based on the universal bacterial primers BSF8 and BSR357. These primers were chosen on the basis of extensive optimization studies that compared segments of the 16S gene for (i) the ability to recover known clustering among bacterial communities and (ii) support accurate phylogenetic placement. In addition to the eight-nucleotide bar code, the primers used to obtain the 16S rRNA gene fragment were also modified to contain the 454 Life Sciences primers A and B.

A 50-μl PCR was carried out using in the AmpliTaq System (Applied Biosystems). The amplification system used AmpliTaq Buffer I and contained 1.5 mM MgCl₂, 0.05% Triton X-100, 100 μM concentrations of each deoxyribonucleoside triphosphates, 0.4 μM concentrations of each primer, 0.1 μg of bovine serum albumin, and 2.5 U of AmpliTaq DNA polymerase. A total of 100 ng of template DNA was used for stool samples and 500 ng for tissue samples (to account for mouse DNA extracted together with the bacterial DNA). Amplification was carried out using the following cycling parameters: 5 min denaturing at 95°C, followed by 20 cycles (for stool samples) or 25 cycles (for tissue samples) of 30 s at 95°C (denaturing), 30 s at 56°C (annealing), and 30 s at 72°C (elongation), with a final extension at 72°C for 7 min. Four independent PCRs were performed for each sample, along with a no-template negative control.

Each PCR product was gel purified from a 0.8% agarose gel. DNA was isolated by using a QIAquick gel extraction kit (Qiagen). A 100-ng portion of each of the 100 gel-purified DNAs was added to a master pool of DNA, which was evaluated by pyrosequencing. Of the 120 samples available, 98 were successfully amplified and sequenced; 85 of these yielded 100 sequences or more for
RESULTS

C. rodentium infection elicits robust expression of proinflammatory cytokines and transient bacterial colitis. To examine the kinetics of bacterial burden over the course of infection, C57BL/6 mice were infected with nalidixic acid-resistant C. rodentium, stool samples were collected at various times, and CFU were enumerated on LB-Nal plates (Fig. 1A). The bacterial burden peaked by day 11, fell significantly by day 20, and was undetectable by day 27, indicating clearance kinetics similar to those published previously (1). To characterize the inflammatory changes associated with C. rodentium infection, mice were infected with Citrobacter rodentium DBS100 (formerly C. freundii biotype 4280 [provided by Bruce Vallance]) per os or mock infected as controls. Mice were sacrificed on day 13 or 14 p.i. for immunological, pathological, and microbiological analyses.

Consistent with previous studies, quantitative reverse transcription-PCR analysis revealed that mRNA levels of IL-17A, IL-17F, IL-22, and IFN-γ in the gut peaked on day 8 p.i. and subsequently declined by day 11 with resolution of infection (Fig. 1B and data not shown) (44). Histological analysis of cecal tissue obtained from infected mice showed characteristic changes in the brush border by day 13 p.i., including elongation of the crypts, a mononuclear cell infiltrate, and a reactive epithelial layer compared to mock-infected controls (Fig. 1C). Intestinal mucosa in the colon also demonstrated heightened activation of epithelial cells, as well as focal ulcerations, a prominent neutrophilia, and crypt abscesses (Fig. 1D). Periodic acid-Schiff staining showed characteristic goblet cell loss accompanying infection (Fig. 1E). Collectively, these findings indicate that C. rodentium infection elicits a robust proinflammatory cytokine response that peaks by day 8 and is associated with the development of bacterial colitis by day 14.

Deep-sequencing analysis and distribution of C. rodentium sequence reads. To determine whether C. rodentium infection and the anti-C. rodentium immune response influenced the structure or stability of gut microbial communities, we used pyrosequencing of 16S DNA segments. Nine mice were analyzed at 9 days p.i. (five infected and four controls), and eleven mice were analyzed at 14 days p.i. (five infected and six controls). Luminal contents were isolated from the cecum, proximal colon, and distal colon of each mouse. Tissue samples were also harvested from each of these sites for analysis of mucosal attached communities. DNA was extracted from each sample, PCR amplified using 16S primers, and subjected to pyrosequencing. The 16S sequence reads were then aligned by using the NAST and GreenGenes servers (7) and inserted into a common phylogenetic tree generated using ARB (26), allowing phylogenetic placement of each sequence read. Sequence reads for C. rodentium could be distinguished unambiguously in the pyrosequence data, allowing the dissemination of the infection to be monitored in time and space (Table 1). C. rodentium sequences were detected only in the infected samples and not in the naïve controls (P < 0.0001). By far the largest accumulation of C. rodentium sequences was detected in mucosally adhered samples from the cecum at day 9 p.i., where C. rodentium 16S sequences reached 2.79% of all sequences. Smaller numbers of sequences were found in communities adhered to the proximal colon and in the luminal contents of the cecum, proximal colon, and distal colon. By day 14 p.i., C. rodentium constituted a smaller proportion of sequences at all sites, indicating control of the infection, although infection at the cecum still constituted 0.91% of sequences.

Table 1 also summarizes counts of Enterobacteriaceae sequences that do not correspond to C. rodentium (which is itself a member of the Enterobacteriaceae family). No obvious differences were seen between infected and uninfected samples. In fact, the greatest number of sequences detected was in the distal colon of an uninfected control mouse. Thus, we did not observe a generalized bloom of Enterobacteriaceae as previously reported (27); rather, the alterations in Enterobacteriaceae appear to have been due to colonization by C. rodentium itself.

UniFrac analysis discloses changes in global community structure accompanying C. rodentium infection. As a first step in analyzing the global effects of C. rodentium infection we compared microbial communities using UniFrac (23–25), which quantifies the similarities among microbial communities based on phylogenetic distances (Fig. 2). To compare two communities, sequences from both communities are placed on a common phylogenetic tree generated using ARB (26). The fraction of the branch length on the tree unique to each community is then measured. This provides an objective measure of community similarity based on the amount of shared evolutionary history. To compare multiple communities, distances between all pairs of communities are computed to generate a distance matrix, and principal coordinate analysis is used to
plot communities in a scatter plot along orthogonal axes of maximal variance. Such scatter plots can be generated, taking into account the abundance, as well as the presence of each taxa (weighted UniFrac), or using only the presence or absence information (unweighted UniFrac). The two methods thus address different questions: weighted analysis allows differences in proportional representation of community members to be assessed, whereas unweighted analysis discloses changes in community composition. In Fig. 2 we present examples of both weighted and unweighted analysis. The specific bacterial taxa that change significantly in abundance during infection are discussed below (see Table 2).

A variety of factors are known to influence the murine gut microbiota, including gender and caging history. The comparison in Fig. 2A and B eliminates these variables by focusing on males from a single litter in which three uninfected mice could be compared to three infected mice at day 9 p.i.

In the unweighted analysis (presence or absence information only), the first principal coordinate (15% of sample variation) separated the samples by tissue-associated versus luminal communities (Fig. 2A). The second principal coordinate (10% of sample variation) separated samples by C. rodentium infection state. A nonparametric test of the difference in means along the second principal coordinate achieved high-level significance (P < 0.0001), indicating that the C. rodentium infection caused a community-wide change in the microbial taxa present.

For weighted UniFrac (Fig. 2B), which takes into account abundance information, the first principal coordinate explained a relatively high proportion of the variation in the sample (67%). Again, this first principal coordinate separated the communities by sample type of origin, with tissue-associated communities separating from luminal contents. The second principal coordinate, which accounted for 15% of the variation, separated communities by infection status (P < 0.00001), indicating that the proportional representation of the different types of bacteria also changed. The tissue samples were more widely separated along principal coordinate 2 than the samples from luminal contents, a finding consistent with
the idea that *C. rodentium* infection and associated inflammation most strongly affect tissue-associated bacteria. Collectively, Fig. 2A and B indicate that both the amounts and the types of gut bacteria detected differed due to the infection status and sample type.

Infected and uninfected animals from two additional litters were also available for comparison. Figure 2C shows females infected for 9 days and controls compared using unweighted UniFrac. Here, too, clustering by infection status and sample type is evident. Analysis after 14 days of infection (Fig. 2D and E) showed that distinctive communities persisted in infected versus uninfected animals up to this time point, even though the infection was beginning to resolve. The effects of infection were evident in both males (Fig. 2D) and females (Fig. 2E).

The temporal progression could also be assessed using unweighted UniFrac, comparing the uninfected (Fig. 3A) and infected (Fig. 3B) animals. For *C. rodentium*-infected animals from all three litters (Fig. 3B), communities from 9 days p.i. clustered separately from those from 14 days p.i., and the separation was highly significant (*P* < 10^-7 on PCA2). Separation over time was much less prominent for the uninfected controls (Fig. 3A), although it did achieve significance (*P* = 0.0002 on PCA1). The mice at the different time points were necessarily from different litters, and littermates are known to harbor relatively similar microbiomes (20), so the slight difference seen in the naive controls is not surprising. Together, these findings indicate that the temporal progression of the *C. rodentium* infection was accompanied by global changes in the composition of the gut microbiota.

Bacterial communities attached to the gut mucosa show changes in multiple lineages accompanying infection. We first present an analysis of bacterial taxa in communities adhered to the gut mucosa, since this is the where *C. rodentium* infection was most developed and where the greatest differences were detected in the UniFrac analysis (Fig. 4). Tissue-associated samples were pooled and analyzed at the phylum level for day 9 (Fig. 4A) or day 14 (Fig. 4B) samples. Samples were pooled over litters and genders of the mice. For all samples, the *Bacteroidetes* and *Firmicutes* comprised at least 85% of the sample. The next most abundant phyla were the *Proteobacteria* and the little-studied *Deferribacteres*. Infection resulted in an outgrowth of minor groups, including the *Deferribacteres*, some *Proteobacteria*, and *Tenericutes* (the *Tenericutes* are a newly formed phylum composed of what was previously the *Mollicutes* class of the *Firmicutes*). The extent of the expansion of the lower abundance phyla was notable at day 9 but more pronounced by day 14.

The deep-sequencing data allowed quantitative analysis of lineages at a finer taxonomic resolution (Fig. 4C and D). In each plot, the tissue-associated bacteria are separated into the cecum and proximal colon (sequences from the tissue-adhered samples from the distal colon were too few in number for analysis). Analysis of the cecal samples allows the minor classes to be identified primarily as *Bacteroidetes* and *Firmicutes* comprised at least 85% of the sample. The next most abundant phyla were the *Proteobacteria* and the little-studied *Deferribacteres*. Infection resulted in an outgrowth of minor groups, including the *Deferribacteres*, some *Proteobacteria*, and *Tenericutes* (the *Tenericutes* are a newly formed phylum composed of what was previously the *Mollicutes* class of the *Firmicutes*). The extent of the expansion of the lower abundance phyla was notable at day 9 but more pronounced by day 14.

The deep-sequencing data allowed quantitative analysis of lineages at a finer taxonomic resolution (Fig. 4C and D). In each plot, the tissue-associated bacteria are separated into the cecum and proximal colon (sequences from the tissue-adhered samples from the distal colon were too few in number for analysis). Analysis of the cecal samples allows the minor classes to be identified primarily as *Bacteroidetes* and *Firmicutes* comprised at least 85% of the sample. The next most abundant phyla were the *Proteobacteria* and the little-studied *Deferribacteres*. Infection resulted in an outgrowth of minor groups, including the *Deferribacteres*, some *Proteobacteria*, and *Tenericutes* (the *Tenericutes* are a newly formed phylum composed of what was previously the *Mollicutes* class of the *Firmicutes*). The extent of the expansion of the lower abundance phyla was notable at day 9 but more pronounced by day 14.

Exposure to *C. rodentium* results in dynamic changes in luminally dwelling microbiota. As discussed above, *C. rodentium* is an attaching-and-effacing bacterium that forms pedes-
tals on the luminal surface of intestinal epithelial cells. We describe above changes in the surface-attached microbial communities. We also sought to determine whether more global changes in microbial communities could also be identified in the luminal contents of the gut.

We found that the bacteria in luminal contents at all levels of infection differed significantly from tissue samples. The comparison between luminal contents and tissue samples achieved significance of $P < 0.0001$ on PCA1 and $P < 0.0028$ on PCA2 for day 9 females. The comparison between naive and infected achieved a significance of $P < 0.0001$ on PCA1 and $P < 0.0001$ on PCA2 for day 14 females.

**FIG. 2.** Global effects of *C. rodentium* infection, separated by time after infection and the sex of the mice compared. Points from naive (solid symbols) and infected (open symbols) animals are enclosed in ovals. Each litter is represented by a different symbol (circles, litter A; squares, litter B; triangles, litter C). Blue and yellow indicate tissue and luminal content samples, respectively. Axes show the principal coordinate number and the percent of variation explained. 

(A) Data for males at day 9 p.i., using unweighted UniFrac. The comparison between luminal contents and tissue samples achieved $P < 0.0007$ on PCA1; the comparison between naive versus infected achieved significance of $P < 0.0001$ on PCA2.

(B) Same as in panel A, but using weighted UniFrac. The comparison between luminal contents and tissue samples achieved $P < 0.0001$ on PCA1; the comparison between naive and infected mice achieved a significance of $P < 0.0001$ on PCA2.

(C) Data for day 9 females, unweighted UniFrac. The comparison between luminal contents and tissue samples achieved a significance of $P < 0.0001$ on PCA2; the comparison between naive and infected achieved a significance of $P < 0.0001$ on PCA1.

(D) Data for day 14 males, unweighted UniFrac. The comparison between luminal contents and tissue samples achieved a significance of $P < 0.0149$ on PCA1; the comparison between naive and infected achieved a significance of $P < 0.0001$ on PCA1.

(E) Data for day 14 females, unweighted UniFrac. The comparison between luminal contents and tissue samples achieved a significance of $P < 0.0001$ on PCA2; the comparison between naive and infected achieved a significance of $P < 0.0001$ on PCA1.
of the GI tract in uninfected and infected mice differed from the tissue-associated bacteria in the ratio of *Firmicutes* to *Bacteroidetes*: the ratio was <1 for all tissue-associated samples but >1 for luminal contents (compare Fig. 4 and 5). The representation of minor groups also varied, with the *Deferribacteres* group more abundant in the tissue-associated samples.

The microbial phyla of the lumen showed little difference associated with infection at day 9 (Fig. 5A), but by day 14 there was a notable expansion of minor phyla (Fig. 5B), including *Proteobacteria* and *Deferribacteres*. A finer analysis of taxa affected in the lumen of the cecum, distal colon, and proximal colon at the family level for day 9 (Fig. 5C) and day 14 (Fig. 5D) showed that many groups were altered in abundance by infection. This took place at both times and was not apparent in the phylum-level analysis because the composition actually changed within the *Firmicutes* phylum. Of these, the *Lactobacillaceae* were particularly strongly affected, showing considerably reduced abundance accompanying infection in all of the samples of luminal contents at both time points. Among other groups that were affected by the infection were the *Deferribacteraceae*, *Betaproteobacteria*, and *Alphaproteobacteria*, all of which increase both at day 9 and day 14 p.i., and *Mollicutes*, which increase substantially at day 14 p.i. Thus, these studies demonstrate previously unappreciated changes inboth luminal and tissue-adhered bacterial communities accompanying *C. rodentium* infection.

**Statistical analysis of affected lineages.** Because of the large number of samples analyzed by pyrosequencing, it was possible to carry out a rigorous statistical analysis of the lineages that were altered by *C. rodentium* infection, treating each mouse as a single measurement of proportions. This is in contrast to less rigorous studies in which many community samples are pooled and statistical analysis carried out on the pool (27), since in such a comparison animals could differ radically from each other, but this would be lost upon pooling.

Table 2 presents a nonparametric comparison in which the collection of proportions for each bacterial lineage, measured within each mouse, is compared between infected and uninfected mice. This shows that the *Deferribacteres* phylum and its single class and family consistently increased in abundance upon infection in both tissue-associated and luminal content samples. The *Proteobacteria* phylum increased in abundance in luminal contents in day 9 p.i. Several lineages of *Firmicutes* changed in abundance in both tissue-associated and luminal content samples. The *Lactobacillaceae* family decreased in abundance in the luminal contents of the distal colon at day 9 p.i. As a control, the uninfected control samples were compared over the two time points (days 9 and 14), and no significant changes in proportion were detected. Thus, thanks to the number of samples available for analysis, it was possible to document changes in microbiome composition that were consistently associated with infection over all animals tested.

**DISCUSSION**

We used deep sequencing of 16S rRNA gene segments to quantify the response of the gut microbiome to *C. rodentium* infection and the associated inflammatory response. This strategy allowed us for the first time to track an intestinal pathogen in space and time while monitoring the consequences for the full gut microbial community as well. We found distinctive communities in the three regions of the gut analyzed, and the communities differed between luminal and tissue-associated bacteria. In contrast to a previous report (27), we did not observe a general increase in *Enterobacteriaceae* lineages apart from *C. rodentium* itself, but we did detect alterations in abundance for *Proteobacteria*, *Deferribacteres*, *Clostridia*, and *Lactobacillaceae*. To our knowledge, this was the first use of 454/Roche deep sequencing to track the community-wide effects of an enteric pathogen in space and time. These findings specify bacterial groups that change their proportion in the community in response to *C. rodentium* infection and which may affect pathogenesis and clearance.

The 454 pyrosequencing approach used here differed from
many previously published studies. Sogin and coworkers emphasized the use of 454 sequencing to study the “rare biosphere” by sequencing deeply into a relatively small number of samples (16, 38). Others have used 454 pyrosequencing for resequencing of full genomes (28). Our approach relies on DNA bar coding to characterize large numbers of samples, but using comparatively modest numbers of 16S rRNA gene sequence reads, in the range of 1,200 per sample. This then allows statistical analysis over a large number of samples where each is treated as a separate measurement of proportions (Table 2). Similar approaches were used in other studies (8, 29, 40). Collecting data in this fashion allows the use of 454 pyrosequencing as a tool for hypothesis testing, in this case asking whether significant global differences could be detected consistently in *C. rodentium*-infected versus uninfected mice. A limitation is that rare bacteria will not be sampled efficiently, but effects on the most abundant groups are well resolved.

Early during infection, *C. rodentium* colonizes the cecum, but at the earliest time point of our analysis (day 9 p.i.) significant alterations in the composition of microbial communities accompanied infection in the proximal and distal colon as well. Given the capacity of bacteria to express antimicrobial products, changes in microbial diversity could be the result of direct competition between *C. rodentium* and resident bacterial communities (5). Alternatively, the robust innate and adaptive immune responses elicited by *C. rodentium* infection could have nonspecific effects on resident bacterial communities. Consistent with this possibility, we and others have observed elevated expression of IL-17, IL-22, and IFN-γ after infection (Fig. 1) (44). These proinflammatory cytokines can induce the expression of antimicrobial peptides in intestinal epithelial cells, including REGIII proteins (44), that could have potent nonspecific effects on *C. rodentium* and resident microbial communities. The observation that *C. rodentium* infection and...
the associated host inflammatory response correlate with alterations in attached and luminal microbial communities throughout the GI tract supports a model wherein *C. rodentium*-specific lymphocytes are primed in the draining lymph nodes and recirculate to the gut microenvironment, where they are widely distributed throughout the cecum and colon. As a result, the widespread *C. rodentium*-induced inflammatory response could explain the global changes in the microbiota we have observed after infection.

In a previous study of *C. rodentium*-infected animals from Lupp et al. (27), which relied on fluorescence in situ hybridization and small numbers of near-full-length 16S sequences (a total of 688 over multiple time points, treatments, and controls), significant evidence was obtained for a bloom of *Enterobacteriaceae*, and significant increases were seen here for this group in cecum-associated communities. However, in our data, this bloom could be attributed to the added *C. rodentium* itself, which is a member of the *Enterobacteriaceae*. Other members of the *Enterobacteriaceae* family could be quantified in our data and did not increase in abundance with infection. Thus, our data do not support the model of Lupp et al. that inflammation provokes a nonspecific outgrowth of *Enterobacteriaceae*. We note, however, that the gut microbiota present in mice prior to infection differed between the two studies, which may have affected the response to infection (discussed below).

We found that the *Lactobacillaceae* group was diminished in abundance during *C. rodentium* infection, a finding of interest because some *Lactobacillus* species have been studied as probiotics for use in the treatment of GI disease (reviewed in reference 41). *Lactobacillus* is a facultative anaerobic or mi-
croaerophilic gram-positive bacteria that convert sugars to lactic acid. The resulting acidification has been suggested to suppress the growth of some harmful bacterial lineages. We found that *C. rodentium* infection resulted in a reduction in the proportion of *Lactobacillus* in microbial communities. Whether or not the specific *Lactobacillus* species accompanying *C. rodentium* infection resulted in reduction of the commensal *Lactobacillaceae* is little studied.

An unexpected aspect of the data was the bloom of *Deferribacteres* accompanying *C. rodentium* infection. These bacteria are members of the eight bacteria comprising the Alter- erd Schaedler Flora, which is used to reconstitute germfree rodents with defined communities (35), but in general the group is little studied. *Deferribacteres* have been associated with periodontal diseases in humans (18, 19, 36). The linkage of this group is little studied.

### Statistical analysis of bacterial lineages changing in abundance during infection, treating each mouse as a data point

<table>
<thead>
<tr>
<th>Days p.i.</th>
<th>Location</th>
<th>Site</th>
<th>Level</th>
<th>Lineage</th>
<th>P</th>
<th>Change upon infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>Tissue</td>
<td>Cecum</td>
<td>Class</td>
<td><em>Clostridia</em></td>
<td>0.0253</td>
<td>Down</td>
</tr>
<tr>
<td></td>
<td>Proximal colon</td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0339</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0339</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Clostridiaceae</em></td>
<td>0.0339</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0275</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Clostridiaceae</em></td>
<td>0.0275</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0275</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Clostridiaceae</em></td>
<td>0.0275</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td>Luminal content</td>
<td>Phylum</td>
<td>Class</td>
<td><em>Lachnospiraceae</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Lactobacillaceae</em></td>
<td>0.0500</td>
<td>Down</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Other Firmicutes</em></td>
<td>0.0143</td>
<td>Up</td>
</tr>
<tr>
<td>14</td>
<td>Luminal content</td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0283</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0283</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0283</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0163</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0163</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Class</td>
<td><em>Deferribacteres</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Phylum</td>
<td>Family</td>
<td><em>Deferribacteraceae</em></td>
<td>0.0500</td>
<td>Up</td>
</tr>
</tbody>
</table>

**ACKNOWLEDGMENTS**

We thank Jeff Weiser and members of the Bushman and Artis laboratories for helpful discussions.

This study was supported in part by the Penn Genome Frontiers Institute and a grant with the Pennsylvania Department of Health. Research in the Artis lab is supported by the National Institutes of Health (NIH), AI61570, AI074878 (D.A.), T32 AI060516 Training in Bacteriology (D.A.H.), the Burroughs Wellcome Fund (Investigator in Pathogenesis of Infectious Disease Award [D.A.]), the Crohns and Colitis Foundation of America’s William and Shelby Model Family Foundation Research Award (D.A.), pilot grants from the University of Pennsylvania (University Research Foundation Award, Veterinary Center for Infectious Diseases Pilot Grant [D.A.]), and the Pilot Feasibility Program of the National Institute of Diabetes and Digestive and Kidney Diseases DK50306. We also acknowledge NIH instrument grant S10RR024525.

The Department of Health specifically disclaims responsibility for any analyses, interpretations, or conclusions.

**REFERENCES**


