Microbiota Diversification and Crash Induced by Dietary Oxalate in the Mammalian Herbivore Neotoma albigula

Aaron W. Miller, Colin Dale, M. Denise Dearing
Departments of Urology and Immunology, Cleveland Clinic, Cleveland, Ohio, USA; Department of Biology, University of Utah, Salt Lake City, Utah, USA

ABSTRACT Oxalate, broadly found in both dietary and endogenous sources, is a primary constituent in 80% of kidney stones, an affliction that has tripled in prevalence over the last 40 years. Oxalate-degrading bacteria within the gut microbiota can mitigate the effects of oxalate and are negatively correlated with kidney stone formation, but bacteriotherapies involving oxalate-degrading bacteria have met with mixed results. To inform the development of more effective and consistent bacteriotherapies, we sought to quantify the interactions and limits between oxalate and an oxalate-adapted microbiota from the wild mammalian herbivore Neotoma albigula (woodrat), which consumes a high-oxalate diet in the wild. We tracked the microbiota over a variable-oxalate diet ranging from 0.2% to 12%, with the upper limit approximating 10× the level of human consumption. The N. albigula microbiota was capable of degrading ~100% of dietary oxalate regardless of the amount consumed. However, the microbiota exhibited significant changes in diversity dynamically at the operational taxonomic unit (OTU), family, and community levels in accordance with oxalate input. Furthermore, a cohesive microbial network was stimulated by the consumption of oxalate and exhibited some resistance to the effects of prolonged exposure. This study demonstrates that the oxalate-adapted microbiota of N. albigula exhibits a very high level of degradation and tolerance for oxalate.

IMPORTANCE The bacteria associated with mammalian hosts exhibit extensive interactions with overall host physiology and contribute significantly to the health of the host. Bacteria are vital to the mitigation of the toxic effects of oxalate specifically as mammals do not possess the enzymes to degrade this compound, which is present in the majority of kidney stones. Contrary to the body of literature on a few oxalate-degrading specialists, our work illustrates that oxalate stimulates a broad but cohesive microbial network in a dose-dependent manner. The unique characteristics of the N. albigula microbiota make it an excellent source for the development of bacteriotherapies to inhibit kidney stone formation. Furthermore, this work successfully demonstrates methods to identify microbial networks responsive to specific toxins, their limits, and important elements such as microbial network cohesivity and architecture. These are necessary steps in the development of targeted bacteriotherapies.

KEYWORDS oxalate, urinary stone disease, woodrat

The mammalian gut is home to trillions of bacteria comprised of hundreds to thousands of interacting bacterial operational taxonomic units (OTUs) with myriad functions (1, 2). So extensive are the host-microbe interactions, that the microbiota is vital for the development of organs, the immune system, and for host metabolism (3, 4). As such, the gut microbiota plays a key role as a buffer to the negative effects of many dietary toxins (5–8). However, the extent of the toxin-buffering capacity and the effects of toxins on the whole microbiota are unknown. It is important to understand...
the interactions between dietary toxins and the gut microbiota for the development of targeted bacteriotherapies tailored to individual patients. Thus, to understand specific interactions between dietary toxins and the gut microbiota, it is necessary to focus on a sufficiently simplified system.

Oxalate is a simple organic acid that is broadly distributed among plants, is regularly consumed by mammalian herbivores and humans, and contributes to 80% of kidney stones (9–12). Despite the toxicity of oxalate to mammals, it cannot be degraded by mammalian enzymes (13). However, it can be metabolized by many gut bacteria, such as Oxalobacter formigenes and Lactobacillus acidophilus, among others (5, 14). For O. formigenes specifically, oxalate is a carbon and energy source for growth (5). The microbial degradation of oxalate in the mammalian gut primarily proceeds via a simple, two-step enzymatic reaction involving the enzymes oxalyl-coenzyme A (CoA) decarboxylase and formyl-CoA transferase to produce one molecule each of CO₂ and formate (15,16). Oxalate can also be toxic to some bacteria, inhibiting proliferation, and introduces a potential source of dynamic interactions within a complex gut microbiota (17, 18). Furthermore, powdered sodium oxalate can be added to a diet, allowing for the effects of a single dietary variable to be examined in animal studies (19, 20). Thus, the metabolism of dietary oxalate by the gut microbiota is a relatively simple process that excludes contributions from the host, introduces a potential source of dynamic interactions, and can be studied in isolation. However, mammals do produce oxalate as a terminal metabolite in the liver from certain dietary precursors, which can make its way into the gut, and must be accounted for in studies concerning the interaction between dietary oxalate and the gut microbiota (21, 22).

The mammalian herbivore Neotoma albigula (woodrat) naturally consumes a high 1.5% oxalate diet in the wild (8). This species harbors consortia of oxalate-degrading bacteria that are stimulated by the introduction of oxalate to the diet and can degrade large amounts of dietary oxalate even when provided at 9% of the diet by dry weight (19, 23, 24). This oxalate-degrading capacity is largely unique to N. albigula and thus makes for an excellent study system to examine the interactions between oxalate and the gut microbiota and inform the development of bacteriotherapy strategies to minimize renal oxalate exposure (23, 24).

The present study was designed to test the hypothesis that the gut microbiota of N. albigula harbors a distinct and cohesive microbial network involved in oxalate metabolism with a limited and quantifiable buffering capacity for the negative effects associated with oxalate exposure. Our objectives were to quantify the maximum tolerable dose of oxalate for N. albigula, quantify changes to gut microbiota diversity metrics across different dietary oxalate concentrations, and identify OTUs that exhibit a significant change in relative abundance associated with oxalate consumption.

RESULTS

Effect of oxalate on host physiology. A total of nine females and five males were used for the study (starting body masses of 171.15 ± 6.67 g and 195.22 ± 7.35 g, respectively). Seven out of the original 14 animals were removed from the trial starting after the first day of the 12% oxalate period, having lost >10% of their starting body mass, including three males and four females (Fig. 1). Animals remaining in the trial had a body mass change ranging from −4.3% to +8.8%. Those animals that persisted throughout the trial maintained −100% oxalate degradation with no significant increase in urinary oxalate across all oxalate treatments in the experiment (Fig. 2). Of the woodrats that were removed from the diet trial, five out of seven exhibited an anomalously low level of oxalate degradation or high level of urinary oxalate at least once during the trial prior to dropping out. For woodrats that completed the diet trial, only two of the seven exhibited at least one anomalous value for oxalate degradation or urinary oxalate (Table 1). For oxalate degradation, anomalous values ranged from 1.3- to 2.7-fold lower than the average. For urinary oxalate, anomalous values were 1.8- to 2.5-fold higher than the average. However, differences in anomalous values between animals that were removed and those that completed the trial were not significant.
Food intake, dry matter digestibility (DMD), fecal output, and fecal oxalate did not differ significantly over the course of the experiment. Dry matter digestibility is defined as (dry matter in the feed – dry matter in feces)/dry matter in the feed. Because there was no difference in these metrics over time, a global average is presented (see Table S1 in the supplemental material). Water intake and urine output did increase significantly with oxalate consumption (see Fig. S1 in the supplemental material).

Effect of oxalate on gut microbiota. Sequencing of fecal communities yielded 2,935,802 sequences after quality control. Across all samples, a total of 26,628 unique OTUs were defined. One animal was removed from the group that completed the trial as it did not produce a fecal sample for one of the time points. This resulted in a sample size of six animals with microbial inventories at each time point. Across OTUs, 93.4% were assignable to the phylum level, with 15.7% assignable to the genus level. The community was dominated by *Bacteroides*, particularly the S24-7 family (see Fig. S3 in the supplemental material).

Community membership and structure were most strongly influenced by interindividual variation (membership, \( P = 0.001, F = 3.402 \); structure, \( P = 0.001, F = 5 \)), and not by oxalate treatment (membership, \( P = 0.632, F = 0.9526 \); structure, \( P = 0.059, F = 1.43 \)) (Fig. 3). All \( \alpha \)-diversity metrics differed significantly across the diet trial, with a peak at the first 6% oxalate period and a trough at the end of the diet trial (Fig. 4).

![FIG 1](http://msphere.asm.org/) Kaplan-Meier survivor curve based upon the loss of 10% of body mass by *Neotoma albigula* on different oxalate diets.

**FIG 2** Oxalate handling for *N. albigula* in vivo for animals that completed the diet trial (\( n = 7 \)). (A) The amount of oxalate consumed correlated with the amount of oxalate degraded (estimated by the differential between oxalate consumed and the total oxalate excreted in the urine and feces). Significance was determined as a repeated-measure Pearson correlation: \( r = 0.9993, P < 0.001 \). (B) Oxalate degradation across the diet trial. Data were analyzed with a repeated-measure ANOVA (df = 6.28, \( P < 0.001 \)) and a post hoc Tukey’s analysis. Letters reflect statistical groups. (C) Daily urinary oxalate excretion for each dietary oxalate treatment. Data for the whole experiment were analyzed with a repeated-measure ANOVA (df = 6.28, \( P = 0.59 \)).
Coinciding with the trend in \(\beta\)-diversity, 8 of the 101 defined families exhibited a significant difference in the number of unique OTUs assigned to them, with a peak at the first 6% oxalate period and trough at the end of the diet trial (Table 2). We conducted paired \(t\) tests on the relative abundance of OTUs between the end of the first 6% oxalate period and the end of the diet trial to determine which OTUs dropped out of the community after the first 6% oxalate period. Analysis resulted in the identification of over 1,000 OTUs that saw a significant drop in relative abundance. However, significance was lost when correcting for false discoveries, indicative of a stochastic loss of OTUs (data not shown).

Repeated-measure Spearman correlations revealed that 1,004 OTUs out of the total 22,784 OTUs present in the six animals analyzed across the study exhibited a significant positive correlation with oxalate consumption after false-discovery rate (FDR) correction, while no OTUs exhibited a significant negative correlation, indicating that equivalent reductions in relative abundance were broadly distributed among the remaining OTUs (see Table S2 in the supplemental material). The SparCC analysis revealed a variable number of positive significant interactions between diet treatments ranging from 729 at the end of the diet trial to 1,682 after the first 6% oxalate period (Fig. 5).

**TABLE 1** Animals that exhibited an anomalously low value for oxalate degradation or high value for urinary oxalate excretion at some point in the diet trial and whether the animal was later removed

<table>
<thead>
<tr>
<th>Animal</th>
<th>Outlier diet (% oxalate)</th>
<th>Metric</th>
<th>Removal diet (% oxalate)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NALB 7</td>
<td>0</td>
<td>Urine oxalate</td>
<td>Not removed</td>
</tr>
<tr>
<td>NALB 10</td>
<td>0</td>
<td>Urine oxalate</td>
<td>12</td>
</tr>
<tr>
<td>NALB 8</td>
<td>3</td>
<td>Oxalate degradation</td>
<td>12</td>
</tr>
<tr>
<td>NALB 8</td>
<td>3</td>
<td>Urine oxalate</td>
<td>12</td>
</tr>
<tr>
<td>NALB 9</td>
<td>6</td>
<td>Urine oxalate</td>
<td>12</td>
</tr>
<tr>
<td>NALB 10</td>
<td>6</td>
<td>Oxalate degradation</td>
<td>12</td>
</tr>
<tr>
<td>NALB 12</td>
<td>6</td>
<td>Oxalate degradation</td>
<td>12</td>
</tr>
<tr>
<td>NALB 6</td>
<td>12</td>
<td>Oxalate degradation</td>
<td>Not removed</td>
</tr>
<tr>
<td>NALB 7</td>
<td>12</td>
<td>Urine oxalate</td>
<td>Not removed</td>
</tr>
<tr>
<td>NALB 13</td>
<td>12</td>
<td>Oxalate degradation</td>
<td>12</td>
</tr>
<tr>
<td>NALB 6</td>
<td>6 (2nd)</td>
<td>Urine oxalate</td>
<td>Not removed</td>
</tr>
</tbody>
</table>

*The diet based on percentage of oxalate on which the outlier occurred and the diet on which the animal was removed (if applicable) are indicated. The proportions of outliers between each group were compared with a Fisher’s exact test \((P = 0.184)\).*

![FIG 3](http://msphere.asm.org/) PCoA plots of the \(\beta\)-diversity (unweighted UniFrac) of individuals (left panel [Adonis \(P\) value of 0.001 and \(F\) value of 3.402]) and percentage of oxalate in the diet (right panel [Adonis \(P\) value of 0.632 and \(F\) value of 0.9526]). Feces were collected 4 to 5 days after the start of each diet treatment for microbial inventories. Different shapes and colors represent different animals or diets. Circles in the left panel show clustering of animals.
Furthermore, there was a significant increase in the relative abundance of the oxalate-degrading genes oxc and frc (Fig. S4).

**Effect of oxalate on renal histopathology.** Renal calcium deposition did not correlate with oxalate consumption in *N. albigula*. There were no significant differences in the surface area of kidney sections normalized to body mass between the wild-diet and 3% oxalate diets. There was a significant increase in the relative abundance of the oxalate-rich microbiota in *N. albigula* woodrats fed diets with different concentrations of oxalate. Treatments along the x axis are listed chronologically. Different letters indicate significant differences as determined by a Tukey’s post hoc analysis. (A) Species richness (repeated-measure ANOVA, *df* = 5.24, *P* < 0.001); (B) Species evenness (repeated-measure ANOVA, *P* = 0.006); (C) Shannon’s index (repeated-measures ANOVA, *P* < 0.001); (D) phylogenetic diversity (repeated-measures ANOVA, *P* < 0.001).

**TABLE 2** Taxonomic families (out of 101) that exhibit significant differences in number of unique OTUs among oxalate diets

<table>
<thead>
<tr>
<th>Family</th>
<th>No. of OTUs on diet shown:</th>
<th>FDR</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0% oxalate</td>
<td>3% oxalate</td>
</tr>
<tr>
<td>Lactobacillaceae</td>
<td>19 ± 1.7 A</td>
<td>18.8 ± 2.4 C</td>
</tr>
<tr>
<td></td>
<td>(1.89E-3)</td>
<td>(1.88E-3)</td>
</tr>
<tr>
<td>S24-7</td>
<td>625 ± 155.1 A</td>
<td>1,199.5 ± 147.1 BC</td>
</tr>
<tr>
<td></td>
<td>(1.92E-1)</td>
<td>(4.44E-1)</td>
</tr>
<tr>
<td>Ruminococcaceae</td>
<td>510.5 ± 54.1 A</td>
<td>450 ± 19.1 A</td>
</tr>
<tr>
<td></td>
<td>(1.09E-1)</td>
<td>(9.20E-2)</td>
</tr>
<tr>
<td>Coriobacteriaceae</td>
<td>6.2 ± 1 A</td>
<td>9.2 ± 1.4 AB</td>
</tr>
<tr>
<td></td>
<td>(3.47E-4)</td>
<td>(2.62E-4)</td>
</tr>
<tr>
<td>Lachnospiraceae</td>
<td>382.8 ± 36.6 AB</td>
<td>259.8 ± 25.6 BC</td>
</tr>
<tr>
<td></td>
<td>(1.37E-1)</td>
<td>(5.57E-2)</td>
</tr>
<tr>
<td>Streptococcaceae</td>
<td>7.3 ± 1.5 AB</td>
<td>10.7 ± 2.2 B</td>
</tr>
<tr>
<td></td>
<td>(0.001101295)</td>
<td>(0.000659257)</td>
</tr>
<tr>
<td>Rikenellaceae</td>
<td>68.8 ± 9.3 AB</td>
<td>71.3 ± 6.2 AB</td>
</tr>
<tr>
<td></td>
<td>(4.90E-2)</td>
<td>(3.63E-2)</td>
</tr>
<tr>
<td>Mogibacteriaceae</td>
<td>6 ± 0.7 A</td>
<td>5.7 ± 1.1 A</td>
</tr>
<tr>
<td></td>
<td>(4.53E-4)</td>
<td>(2.35E-4)</td>
</tr>
</tbody>
</table>

*Columns reflect the mean number of unique OTUs ± standard error. Significance was calculated with a repeated-measure ANOVA (FDR corrected). Letters reflect statistical grouping as determined by an FDR-corrected, post hoc Tukey’s analysis. The relative abundance of each family and time point is listed in parentheses after the unique number of OTUs.*
fed and high-oxalate-diet-fed animals (Table 3). All kidneys, whether from animals feeding on the wild diet or the high-oxalate diet, exhibited some calcium deposition (Fig. 6). However, there was no significant increase in the percentage of the total kidney surface area covered by calcium deposits for *N. albigula* feeding on a 1.5% oxalate diet versus 6 or 12% oxalate (Table 3).

**DISCUSSION**

Diet is one of the primary drivers influencing the form and function of the mammalian gut microbiota (25–28). Dietary toxins in particular help to shape and are shaped by the gut microbiota, with considerable implications for the health and physiology of the host (5–8, 19). The results of our study strongly support but do not explicitly confirm the hypothesis that microbial oxalate metabolism in the *N. albigula* gut involves a distinct and cohesive, functionally resilient microbial network with a limited oxalate-buffering capacity. To test the hypothesis, we created artificial conditions for the gut microbiota of *N. albigula* that are well outside what is normally seen in the wild.

**TABLE 3** Surface area of kidney sections normalized to body mass and surface area covered by calcium deposits in *Neotoma albigula* woodrats freshly caught in the wild or given a high-oxalate diet

<table>
<thead>
<tr>
<th>Metric</th>
<th>Result for woodrat group</th>
<th>t score</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface area of kidney section (cm²/kg body mass)</td>
<td>Wild caught: 8.7 ± 1.1</td>
<td>8 ± 0.5</td>
<td>3.955</td>
</tr>
<tr>
<td>% of surface area covered by calcium deposits</td>
<td>0.014 ± 0.007</td>
<td>0.007 ± 0.003</td>
<td>3.722</td>
</tr>
</tbody>
</table>

Shown is the surface area of kidney sections normalized to body mass and surface area covered by calcium deposits in *Neotoma albigula* woodrats freshly caught in the wild or given a high-oxalate diet (6 or 12%). Significance was determined by a t test (df = 8).
to specifically stimulate the microbial network that benefits from oxalate exposure and quantify the oxalate-buffering capacity of the gut microbiota as a whole. Prior to the experimental period, animals were under captive conditions, feeding on a very-low-oxalate (0.2%) diet for 7 months. These conditions reduced the gut microbiota α-diversity and helped the microbiota acclimate away from oxalate exposure, thus setting the conditions necessary to quantify the effect of oxalate exposure (7). Following the acclimation period, oxalate, which is one of the primary toxins found in the wild *N. albigula* diet, was reintroduced and brought up to levels 8-fold higher than what is naturally seen (8). Evidence that supports our oxalate metabolic network hypothesis include the following: (i) the microbiota exhibited a two-phase, dynamic change in diversity where oxalate shifted from resource to poison for the gut microbiota, (ii) >1,000 microbial OTUs exhibited a statistically significant positive correlation to oxalate over the course of the diet trial, with the number of interactions among them increasing and decreasing in accordance with oxalate input (Fig. 5), and (iii) despite an overall drop in diversity beginning after the first 6% oxalate period, oxalate metabolism operated at maximum capacity throughout the trial. Each of these points will be discussed in detail below.

Oxalate, from the perspective of the gut microbiota, is both a resource and a poison. On the one hand, oxalate can be used as a carbon and energy source for many OTUs inhabiting the gut, sometimes exclusively, as is the case with *O. formigenes* (8, 14, 29–32). However, oxalate is also known to inhibit the growth of some bacteria that are known oxalate degraders and to induce the differential regulation of over 300 genes (17, 18). In the present study, both aspects of oxalate appear to impact the gut microbiota, inducing a two-phase dynamic shift in the gut microbiota. In the first phase, which lasted up to the first 6% diet, oxalate acted as a resource, enabling the community to support greater diversity (Fig. 4). The increase in diversity may at least partially reflect a return to historical conditions given a reintroduction of a resource with which the gut microbiota had been historically adapted and not just the introduction of a novel resource in general. In previous studies, the gut microbiota has been partially restored in captive animals when their natural diet was reintroduced (7). Similar increases in α-diversity have also occurred with the reintroduction of creosote, a toxin present in the natural diet of other woodrat species, *N. lepida* and *N. bryanti* (33). In contrast, when oxalate or creosote is introduced into the diets of animals without prior experience to the toxins, the gut microbiota exhibits significant decreases in

![FIG 6](http://msphere.asm.org/)

FIG 6 Von Kossa staining of kidney sections from freshly trapped woodrats on a 1.5% oxalate diet (A) and captive animals on a 12% oxalate diet (B). Black areas represent calcium deposits within the tissue.
diversity (20, 33). In the second phase of gut microbiota change, which occurs when oxalate reaches 12% of the diet, the level of oxalate exposure exceeds the buffering capacity of the gut microbiota and thus acts as a poison. This level of oxalate consumption is approximately $10\times$ the typical daily consumption for humans (34) (Fig. 2). The effects of oxalate as a poison are reflected in the body mass loss of the *N. albignula* hosts (Fig. 1) and in the α-diversity crash that began at 12% oxalate and persisted to the end of the diet trial (Fig. 4). Additionally, even the Oxalobacteraceae, some of which are oxalate-degrading specialists, exhibited a significant decline in relative abundance during the 12% period (see Fig. S2 in the supplemental material). Similar drops in α-diversity have been demonstrated after exposure to antibiotics, which are explicitly used to reduce microbial populations (35, 36).

Over the course of the diet trial, some OTUs exhibited statistically significant changes in relative abundance, while others did not. It is expected that if each animal harbored the exact same microbiota at the start of the trial, then consistent OTU-level shifts in relative abundance would occur and a clear diet effect would be apparent. Indeed, 1,004 out of 22,784 OTUs (3.8%) and 8 out of 101 families (7.9%) exhibited significant positive correlations to oxalate exposure over the course of the experiment (Table 2; and Table S2). However, there were no OTUs that were negatively correlated to oxalate exposure, meaning that the reciprocal declines in relative abundance for other taxa were broadly distributed in a stochastic manner. Our results indicate that the correlation between the relative abundance of OTUs and oxalate consumption followed by cooccurrence analysis is an effective means to identify and track bacteria that respond to oxalate, even with the sample size of six animals. As shown in the principal coordinate analysis (PCoA) plots of β-diversity (Fig. 3), microbial communities clustered significantly by animal, but not by diet. The β-diversity results indicate that the stochastic shifts in OTUs that are dependent on the unique microbiota of each animal at the start of the experiment outweighed the statistically significant OTU responses to oxalate. The taxa that positively responded to oxalate exposure were primarily composed of taxa expected to be a part of an oxalate metabolic network. They included potential oxalate degraders from the Oxalobacteraceae, Lactobacillus, and Bifidobacterium, as well as taxa that harbor a complete oxalate metabolic pathway, such as the S24-7 family (5, 14, 37). In fact, given the number and relative abundance of S24-7 members responding to oxalate, they may be even more important to oxalate degradation than the Oxalobacteraceae (Table 2; Table S2). Other taxa that responded to oxalate exposure do not have an obvious association with oxalate metabolism. These bacteria may benefit indirectly from oxalate metabolism, perhaps by taking advantage of the by-products produced, formate and CO₂. Formate and CO₂ may be used in downstream pathways such as acetogenesis, methanogenesis, or sulfate reduction (38). Indeed, many of the OTUs with significant correlation to oxalate exposure can potentially engage in these metabolic activities (Table S3). However, it is impossible to determine what functions these bacteria engaged in for the present study. The multi-layer network analysis restricted to OTUs that positively correlated to oxalate exposure (Fig. 5) shows a growth of the overall network with increasingly fewer isolated interactions that plateaued between the first 6% and 12% oxalate diets with a collapse by the end of the trial. This is further indication of a cohesive microbial network that specifically responds to oxalate exposure and has a limited buffering capacity. Network analysis was validated by correlating the number of cooccurring pairs at each time point to oxalate degradation. This validation step independently ensures that the endpoint of the network analysis produces biologically meaningful results rather than producing an arbitrary subset of OTUs present in the microbial inventories. As expected, there was a significant correlation with the biggest deviations from the trendline occurring at the two 6% oxalate periods. The physiological relevance of our results was validated through quantitative PCR (qPCR) analysis of the oxalate-degrading genes, *oxc* and *frc*, which increased in relative abundance between the 0% and 6% oxalate periods. For qPCR analysis, primers were designed based on the *oxc* and *frc*
genes from the Oxalobacteraceae, which exhibited a significant response to oxalate according to microbial inventories generated by 16S rRNA sequencing (Fig. S4).

Despite the fact that the oxalate-buffering capacity of the microbiota was exceeded, oxalate metabolism continued to operate at maximum capacity during the 12% and the second 6% oxalate periods (Fig. 2). While the proportion of animals that both were removed and exhibited anomalous values for oxalate excretion/degradation was higher than for those animals that did not drop out, the difference was not significant (Table 1). In examining the network analysis, we see also that the number of interactions after the first 6% diet (1,682) is similar to that after the 12% diet (1,575), in contrast to the overall α-diversity crash (Fig. 4). These results are indicative of a functionally resilient microbial network with a high residual capacity for oxalate metabolism.

Over the course of the diet trial, 50% of the animals lost >10% of their body mass (Fig. 1). However, there was no significant change in food intake (Table S1). Furthermore, in contrast to the significant drop in overall α-diversity between the first and second 6% oxalate periods, there were no significant declines in OTU relative abundance, indicative of stochastic population reductions. These results, combined with the persistently high oxalate degradation and microbial network resilience, suggest that the loss of body mass in the animal hosts was not a direct result of oxalate exposure, but rather could result from the stochastic changes in the gut microbiota and their effect on host physiology. However, further examination of the causes behind the mass loss was out of the scope of the experiment, and it is impossible to know for sure given the data.

During the diet trial, oxalate was added to the diet as sodium oxalate, thus in effect creating two stressors to the host and microbiota. While there is considerable evidence for the effect of sodium on host physiology, little is known about its effect on the microbiota (39–41). The animals here exhibited a significant increase in water intake and urine output in accordance with the oxalate diet, indicative of an effect of sodium (Fig. S1). Furthermore, the genus Allobaculum, which exhibited a significant correlation to the diet here, also significantly increases in relative abundance with sodium chloride intake (40). However, in mammalian herbivores, sodium is the most limiting mineral for growth and reproduction (42). Sodium is efficiently absorbed, with very little making it to the microbiota of the distal colon (43). Many of the OTUs stimulated by oxalate here have previously been associated with oxalate or negatively associated with kidney stone disease. Oxalobacter is an oxalate specialist (5). Bacteria belonging to the taxa Oxalobacteraceae, Bifidobacterium, Clostridium, S24-7, Lactobacillus, and Clostridiales are known to harbor oxalate-degrading genes (8, 37). Additionally, bacteria from the Oxalobacteraceae, Parabacteroides, Desulfovibrio, and Bacteroides taxa have been negatively associated with kidney stone disease (12, 44, 45). Finally, the taxa stimulated by oxalate here exhibited an 82% overlap with taxa that cooccur with Oxalobacter at the family level in woodrats and Sprague-Dawley rats, as determined in another study (46). Therefore, while some of the changes to bacterial taxa may result from exposure to sodium, our results suggest that the changes to the microbiota in this study were primarily driven by oxalate.

Our results indicate that the physiology of the host-microbe holobiont of N. albigula is sufficient to inhibit the negative effects of oxalate consumption, even at extreme levels. A diet of 12% oxalate is lethal for many mammals, including cattle and sheep (47). However, even on a 6% or 12% oxalate diet, there was no significant accumulation of renal calcium deposition in N. albigula. This is in contrast to Sprague-Dawley rats fed a high-glyoxalate diet, which produces excess endogenous oxalate, which exhibited significant increases in renal calcium deposition and the development of renal calculi (48).

Conclusions. The microbiota of N. albigula is unique in its capacity for oxalate metabolism, which can be exploited for the development of personalized bacteriotherapies. The results of the present study have broad implications for the development of bacteriotherapies that target oxalate for elimination. Here we have shown that (i)
TABLE 4 Timeline for the diet trial to quantify the maximum tolerable dose of oxalate and its effect on the gut microbiota

<table>
<thead>
<tr>
<th>Diet</th>
<th>No. of days on diet</th>
<th>Day of fecal collection^a</th>
</tr>
</thead>
<tbody>
<tr>
<td>0% oxalate</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>3% oxalate</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>6% oxalate</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>12% oxalate</td>
<td>14</td>
<td>4</td>
</tr>
<tr>
<td>6% oxalate (2nd)</td>
<td>5</td>
<td>5</td>
</tr>
</tbody>
</table>

^aFeces were collected for microbial inventories at the specified time points.

cohesive microbial networks respond to a specific oxalate challenge, (ii) stochastic effects on the microbial community unique to each individual’s microbiota can mask the effect of oxalate on the microbial networks, (iii) the buffering capacity of the N. albigula microbiota can be exceeded given a high enough level of oxalate exposure, and (iv) the N. albigula oxalate metabolic network is resilient even despite an overall community-level crash. This work can inform strategies for the development of personalized bacteriotherapies for patients suffering from recurrent episodes of calcium oxalate stone formation. Furthermore, the methods employed here can be used to identify microbial networks responsive to other toxins, along with their limits. Thus, this work adds to the toolbox for the development of targeted bacteriotherapies.

MATERIALS AND METHODS

Oxalate diet trials. Fourteen adult N. albigula woodrats were collected from Castle Valley, UT (38.63’N, 109.41’W) in September 2014, using Sherman live traps. Animals were transported to the animal facility at University of Utah, where they were housed in individual cages (48 by 27 by 20 cm) with a 12-h/12-h light/dark cycle, at 28°C and 20% humidity. Animals were maintained on a 0.2% oxalate, high-fiber rabbit chow (Harlan Teklad formula 2031; Envigo, Denver, CO) for 7 months prior to experimentation, which reduces the overall diversity of the microbiota of these animals but maintains their native microbiota (49). Thus, captive maintenance on a low-oxalate diet provides a baseline microbiota to which the addition of oxalate can be compared. All methods were approved by the IACUC under protocol no. 12-12010.

Woodrats were placed in a diet trial in which the oxalate concentration of the food was gradually increased over time along a gradient from 0.2% (for days 1 to 5; herein referred to as 0%), 3% (days 6 to 10), 6% (days 11 to 15), 12% (days 16 to 30 days), and then decreased to 6% (days 31 to 36) (Table 4). The oxalate concentration of the diet was adjusted by adding the appropriate amount of sodium oxalate (Fisher Scientific, Pittsburgh, PA) into the powdered rabbit chow on a dry weight basis. Animals were removed from the diet trial if they lost >10% of their starting body mass. The maximum tolerable dose of oxalate was defined by the point at which 50% of the animals were removed from the diet trial due to body mass loss. The multiday schedule for each concentration was chosen to ensure that the gut microbiota had time to respond to the specific diet (19, 50). The 14-day period for the 12% oxalate diet was employed to evaluate the potential for chronic exposure to a high concentration of oxalate. We did not go above 12%, as this concentration is known to be lethal to other mammalian herbivores and is found in some plants native to the same habitat as N. albigula (47). During the trial, we measured body mass and food intake on a daily basis, and from these data we estimated DMD and oxalate consumption.

For the duration of the diet trial, animals were placed in metabolic cages to separate urine and feces into sterile 50-ml conical tubes to allow for quantification along with food and water intake (given ad libitum). Acidified 24-h urine collections were frozen (−20°C) and feces dried overnight (45°C) prior to oxalate assays. Prior to drying, a subsample of feces was collected from each animal every 4 to 5 days on each diet treatment. Samples were frozen at −80°C for microbial analyses. Oxalate excretion was also quantified from urine and feces for the same time periods (Table 4).

Oxalate extractions and assays were performed as previously described (19, 20). Briefly, acidified urine was thawed and centrifuged to remove precipitates. The pH of the urine was brought up to 7 with NaOH, and 0.1 g of CaCl₂ was added to precipitate calcium oxalate. Samples were centrifuged and decanted. A volume of deionized water equal to the starting volume of urine was added to calcium oxalate precipitate, and the solution was titrated as described below. Dried fecal samples were ground to a powder and acidified with HSO₆₄ for 15 min to extract oxalate. Precipitates were filtered out with a grade 4 Whatman filter, and the pH was raised to 7 with NaOH. Calcium oxalate was precipitated with the addition of CaCl₂ and a volume of deionized water equal to the volume of the filtrate was added prior to titration. Calcium oxalate solutions were acidified and heated to 80°C prior to titration with 0.01 M KMnO₄. Titration volumes were then compared to standards that have undergone the same procedures with known amounts of oxalate added. These methods allow for the recovery of 90 to 110% of urinary and fecal oxalate (19, 20). Oxalate degradation was defined as oxalate consumed minus oxalate excreted, which is a conservative estimate (19, 20). Data were evaluated with repeated-measure analysis of variance (ANOVA) with a post hoc Tukey’s analysis when applicable of animals that made it to the end of the trial. To determine if animals that were removed from the trial for excessive loss of mass exhibited
a difference in oxalate degradation or urinary oxalate excretion from those that persisted, box and whisker plots were generated from all data for each time point and anomalous values were defined as being outside 1.5× the interquartile range above the upper quartile or below the lower quartile. The proportions of anomalous values in each group were compared with a Fisher’s exact test.

**Microbial inventories.** For microbial inventories, DNA was extracted from 180 to 220 mg of fecal samples with the QIAamp DNA stool minikit (Qiagen, Germantown, MD). Microbial inventories were generated from feces of the seven woodrats that persisted through the entire trial. Extracted DNA was sent to Argonne National Laboratory (Chicago, IL) for sequencing of the V4 region of the 16S rRNA gene with primers 315F and 806R (51). The DNA was barcoded with 12-bp sequences, and samples were multiplexed for a single-lane run on an Illumina MiSeq with paired-end sequencing of 150 bp each, as described previously (52).

The resulting data were demultiplexed and quality controlled with default parameters in QIIME (52). Microbial OTUs were assigned de novo with UCLUST at a sequence identity cutoff of 97%. Chloroplast and mitochondrion sequences, as well as sequences with fewer than 10 representations across the whole data set, were removed. Following processing, the data set was normalized with the DESEQ2 algorithm, which executes a negative binomial Wald test and maintains rare taxa (53, 54).

From the normalized OTU table, we calculated the following α-diversity metrics: species richness (Margalef’s), evenness (equitability), Shannon index, and phylogenetic diversity. The α-diversity metrics were analyzed with a repeated-measure ANOVA and post hoc Tukey’s analysis or a paired t test with false-discovery rate (FDR) correction where applicable. Furthermore, unweighted and weighted UniFrac analyses were performed to compare community membership and structure, respectively (55). Comparisons between animals and diet were made with Adonis after 999 permutations. The total number of unique OTUs in each bacterial family for each diet was quantified and analyzed with a repeated-measure ANOVA across time points (FDR corrected) followed by a post hoc Tukey’s analysis (R statistical package). All analyses were performed in QIIME unless otherwise noted.

Those OTUs exhibiting a significant increase in abundance relative we defined as an oxalate metabolic network, which was specifically tracked across the diet trial. To quantify the network, we performed a repeated-measure Spearman correlation analysis (FDR corrected) for OTU relative abundance and oxalate consumption using R statistical software. Spearman correlations were chosen because they are more appropriate for nonlinear correlations, and we expected a plateauing or decline of microbial network bacteria at the 12% oxalate level. Significant, positive Spearman correlations were used to produce a restricted list of OTUs. From this list, the SparCC algorithm was used for cooccurrence analyses of the OTUs among animals and time points. This algorithm minimizes spurious associations and only strong correlations ($P < 0.05$) were used in downstream analyses (56). Cooccurrence networks were visualized in Cytoscape (57).

**Validation of network analysis.** Network analysis results were validated in two ways. First, the number of cooccurring pairs within the network was quantified for presence/absence in the microbiota of each animal and time point. This was then correlated to oxalate degradation at the same time point. This ensures that the biological significance inferred from network analysis correlated to the expected biological outcomes and that the analytical methods to identify OTUs responding to oxalate were valid. Second, we conducted qPCR analysis of the oxalate-degrading genes oxc and frc for the microbiota after the 0.2% oxalate diet and after the first 6% oxalate diet, where the biggest difference in microbial diversity was quantified. This analysis ensures that the microbial function expected to increase the most for the 0.2% oxalate diet and after the first 6% oxalate diet, where the biggest difference in microbial function expected to increase the most.

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**Histopathology.** A total of eight N. albigena woodrats were euthanized under isoflurane for the histopathological analysis of kidneys. As a control, four animals in captivity for one night after trapping while feeding on their natural diet of *Opuntia* cactus (1.5% oxalate) were sacrificed to quantify normal renal calcium deposition for *N. albigena*. To maximize the probability of detecting renal calcium oxalate deposition for animals in the present study, two animals that dropped out while on 6% oxalate and two animals that dropped out on 12% oxalate were chosen for renal histopathology. Kidneys were fixed in formalin, which was replaced with 70% ethanol after 24 h and kept at 4°C until they were processed for histology. Prior to embedding in paraffin, kidneys were longitudinally bisected along the center line of the kidney. Longitudinal sections of the center of the kidney were stained with the Von Kossa stain, in which silver nitrate is used to replace calcium and becomes opaque. Kidney sections were visualized on an Olympus BX41 microscope at ×100 magnification. Images were taken with a Diagnostic Instruments color mosaic camera (18.2) and Spot version 4.6 software. Total kidney surface area and the surface area of calcium deposits were quantified with ImageJ software.

**Accession number(s).** Sequence reads are available at the Sequence Read Archive under accession no. SRR5261472.
SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00428-17.

FIG S1, PDF file, 0.1 MB.
FIG S2, PDF file, 0.1 MB.
FIG S3, PDF file, 0.1 MB.
FIG S4, PDF file, 0.05 MB.
TABLE S1, PDF file, 0.1 MB.
TABLE S2, PDF file, 0.2 MB.
TABLE S3, PDF file, 0.1 MB.

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A.W.M. carried out experiments and analyses. All authors participated in the design of the study and statistical analyses, helped to draft the manuscript, and read and approved the final manuscript.

REFERENCES


