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Abstract Microbial communities typically vary in composition and structure over space and time. Little is known about the inherent characteristics of communities that govern various drivers of these changes, such as random variation, changes in response to perturbation, or susceptibility to invasion. In this study, we use 16S ribosomal RNA gene sequences to describe variation among bacterial communities in the midguts of cabbage white butterfly (*Pieris rapae*) larvae and examine the influence of community structure on susceptibility to invasion. We compared communities in larvae experiencing the same conditions at different times (temporal variation) or fed different diets (perturbation). The most highly represented

phylum was *Proteobacteria*, which was present in all midgut communities. The observed species richness ranged from six to 15, and the most abundant members affiliated with the genera *Methylobacteria*, *Asaia*, *Acinetobacter*, *Enterobacter*, and *Pantoea*. Individual larvae subjected to the same conditions at the same time harbored communities that were highly similar in structure and membership, whereas the communities observed within larval populations changed with diet and over time. In addition, structural changes due to perturbation coincided with enhanced susceptibility to invasion by *Enterobacter* sp. NAB3R and *Pantoea stewartii* CWB600, suggesting that resistance to invasion is in part governed by community structure. These findings along with the observed conservation of membership at the phylum level, variation in structure and membership at lower taxonomic levels, and its

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relative simplicity make the cabbage white butterfly larval community an attractive model for studying community dynamics and robustness.

Introduction

Microbial communities are dynamic, often experiencing changes in composition and structure. Changes can result from alterations in nutrient availability, physical aspects of the environment, and proximity to other organisms [1–4]. Many communities, such as those inhabiting lakes, soil, insects, humans, and other animals, experience temporal changes associated with factors such as season, nutrient availability, and host development [5–10].

In addition to cyclical or programmed influences, sudden disturbances can also alter community composition. In animal-associated communities, a common disturbance is change in host diet. Plant- vs nonplant-based diets, differences in plant species, fiber content and type, and fat source all have been implicated in changes in gut community composition [11–15]. The addition of antibiotics to diet and intravenous administration also alter community composition [16–18]. While the impact of antibiotics on the human gastrointestinal microbial community has attracted interest as a medical issue, antibiotics also provide a tool for exploring the ecology of animal gut-associated communities.

One of the most important ecological processes is invasion, arising from both natural and anthropogenic introductions. The process of biological invasion can be modeled in gut communities. Biological invasion theory attempts to predict invasion patterns, characteristics of successful invaders, characteristics of communities susceptible to invasion, consequences of invasion, and processes driving establishment [19, 20]. General theories of invasibility are needed, both to develop a more proactive and predictive approach to the increased frequency of biological invasions and to help guide the strategies and success of deliberate introductions.

In some situations, microbial invasions are associated with damage to an ecosystem (i.e., reference [21] and [22]), and in others, invasion is desired. For example, bacteria comprising probiotic preparations and disease-suppressive bacterial biocontrol agents for crop health must invade a community to provide a benefit [23, 24]. The ability to resist invasion by exogenous bacteria, also known as the barrier effect or colonization resistance, is a central attribute of the microbial communities in the human gastrointestinal tract and vagina [25–27]. Despite the importance of invasions, community susceptibility to them is not well understood [28, 29].

Here, we present the lepidopteran midgut as a potential model system for studying ecological processes in animal-associated gastrointestinal communities. Recently, the midgut of *Lymantria dispar*, the gypsy moth, was shown to

contain a simple bacterial community of approximately seven members [30]. Because of its relative simplicity, the community in the lepidopteran midgut is attractive as a potential model system. The lepidopteran species used in this study, *Pieris rapae*, the cabbage white butterfly, has several practical attributes, including ease of care, handling, and manipulation, and a short life cycle that facilitates multigenerational studies. Although there has been much work examining the digestive physiology of the cabbage white butterfly (i.e., the maintenance of a slightly alkaline gut pH and the presence of endopeptidases), there has been little work exploring the gut microbiota [31, 32]. A previous microbiological study of this insect was limited to culture-based analysis of the adult alimentary tract [33].

The goals of this study were to (1) characterize the species richness and composition of the midgut bacterial community of cabbage white butterfly larvae using culture-independent methods, (2) investigate the effects of time, diet, and antibiotics on this community, and (3) explore the cabbage white butterfly midgut as a model for studying aspects of inherent community features, such as robustness. Robustness is a comprehensive term used to describe the extent to which a community exhibits temporal stability (constancy in structure over time; [34]), resistance (ability to resist change following perturbation; [35, 36]), and resilience (ability to return to an initial structure following perturbation; [37]). Two aspects of robustness, temporal stability and resistance (to dietary perturbation and invasion), were examined in this study.

Methods

Treatment and Rearing of Larvae

Cabbage white butterfly eggs were obtained from Carolina Biological Supply Company (Burlington, NC, USA) and soaked in a solution of 1% Tween and 2% bleach for 3 min and then rinsed in sterile distilled water. This treatment was used to reduce the presence of fungi and bacteria on the surface of the egg capsule. The eggs were then dried in a sterile hood and placed in a sterilized Petri dish with either unamended sterilized standard gypsy moth artificial diet (MP Biomedical, Irvine, CA, USA), sterile artificial diet amended with a penicillin and streptomycin cocktail (Sigma-Aldrich Biotechnology, St. Louis, MO, USA) at 10 units per milliliter and 10 µg/ml, respectively, and sterilized artificial diet amended with sinigrin (Sigma-Aldrich Biotechnology), a major phytochemical component of Brussels sprouts, at a concentration of 3.0 mg/ml, which is comparable to the concentration in fresh Brussels sprouts [38] or Brussels sprout leaves. Larvae were reared to fourth instar in growth chambers with 16 h/8 h (light/dark) photoperiods at 27 °C. Petri dishes containing larvae were

opened only inside a sterile hood for feeding and cleaning purposes.

Preparation of Diet

A denatured wheat germ diet used to rear many Lepidoptera, such as gypsy moth, was prepared as directed by the manufacturer's instructions and autoclaved (MP Biomedicals, Irvine, CA, USA). Diet was cooled, and treatments were incorporated (compounds and concentrations described above). Diet samples were plated periodically to confirm the absence of culturable bacteria and fungi. Conventionally raised Brussels sprouts were obtained from a local grocery store. Brussels sprout leaves were separated and sonicated for 60 s in 5% bleach, washed in 5% bleach for 5 min, and rinsed twice in sterile distilled water for 2 min. Efforts to culture bacteria from treated Brussels sprout leaves were unsuccessful, thereby confirming effectiveness of bleach treatment. Leaves were then dried in a sterile hood and stored for up to 5 days in parafilm-sealed sterile Petri dishes at 4 °C until use.

Sampling and Dissection

Fourth instar larvae were used in all experiments. Larvae were placed in sterilized Petri dishes and starved for 4 to 6 h before dissection to reduce food content in the midgut and the presence of transient bacteria. Dissections were performed as described previously [30], and guts were stored at −20 °C prior to DNA isolation or used immediately when bacterial cultivation was necessary.

Bacterial Cultivation

Bacteria were isolated from midguts as described previously [30]. Briefly, midguts were sonicated for 60 s and plated on 1/10th strength tryptic soy agar or 1/10th tryptic soy agar amended with 25 µg/ml nalidixic acid when appropriate.

Analysis of 16S Ribosomal RNA Genes

DNA was extracted from pools of ten or from individual midguts as described previously [30]. Individual guts were sampled to determine intrinsic variability between insects reared under the same conditions at the same time. Amplification of 16S ribosomal RNA (rRNA) genes was performed as described previously using primers 27F and 1492R [39]. Polymerase chain reaction (PCR) products were then ligated into pGEM-T (Promega Corporation, Madison, WI, USA) according to the manufacturer's directions. *Escherichia coli* JM109 (Promega Corporation) was transformed with the ligation mix according to the manufacturer's instructions. Clones were grown in Luria–Bertani broth containing 100 µg/mL ampicillin. The 16S

rRNA genes were amplified in PCR reactions containing standard vector primers M13F and M13R. PCR products were purified using the AMPure magnetic bead system (Agencourt Bioscience, Beverly, MA, USA). Sequencing reactions were conducted as described previously using the BigDye V3.1 reaction mix (Applied Biosystems, Foster City, CA, USA) and primers 27F for onefold coverage or 27F and 907R for twofold coverage of ~500–900 bp of the 5' end of the 16S rRNA gene [30, 39]. The reverse sequences obtained using the 907R primer differed from the forward sequences, on average, by less than 1%; therefore, reverse sequences were not included in the analysis of most of the experiments. Products were then purified using the CleanSEQ magnetic bead system (Agencourt Bioscience) and analyzed at the University of Wisconsin–Madison Biotechnology Center.

Phylogenetic Analysis and Temporal Stability of Community

Sequences were initially analyzed in SeqMan (DNASTAR, Madison, WI), rapidly aligned using the align tool of the Greengenes web application (www.greengenes.lbl.gov) [40] and then manually aligned further using ARB (www.arb-home.de) [41, 42]. Bellerophon (<http://foo.maths.uq.edu.au/~huber/bellerophon.pl>) [43] was used to detect chimeras using the Huber–Hugenholtz correction; poor quality and chimeric sequences were removed from the group to be analyzed further. Distance matrices were generated using the Jukes–Cantor correction in ARB and used in the subsequent analyses. DOTUR (<http://schloss.micro.umass.edu/software/dotur.html>) [44] was used to assign sequences to operational taxonomic units (OTUs) and to calculate diversity indices for each of the 14 libraries constructed. Sequences that were in the same OTU_{0.03} (identity ≥97%) were considered to be from the same species, and sequences with an identity of less than 97% were considered to be different species [44]. Sequences that were in the same OTU_{0.20} (identity ≥80%) were considered to be from the same phylum [44]. BLASTN (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>) [41] was used to compare sequences in OTUs to sequences in GenBank and to assign identity. The sequences were assigned to phylogenetic divisions and/or species based on those results. An estimate of library coverage was determined using the formula

$$C = [1 - (n_1/N)] \times 100,$$

where C is the percent coverage, n_1 is the number of sequences appearing once, and N is the total number of sequences in the library [45] for each library. Descriptive characteristics of each library are summarized in Table 1. SONS (<http://schloss.micro.umass.edu/software/sons.html>) [46] was used to calculate indices that measure similarity

Table 1 Characteristics of 16S rRNA gene libraries constructed from midgut communities in cabbage white butterfly larvae reared

Experiment	Library name	Diet	Date of sampling	Sample composition	No. of sequences	No. of OTUs observed ^a	Chao estimate OTUs (95% confidence interval)	Good's coverage	Shannon–Weaver diversity index (95% confidence interval)
Effect of diet	BL-1	Artificial	1/05	Pool—10 guts	79	7	7.4 (7.0–20)	97%	1.2 (0.9–1.4)
	BL-2	Sinigrin	1/05	Pool—10 guts	59	9	10 (9.1–20)	93%	1.7 (1.4–1.9)
	BL-3	Brussels sprouts	1/05	Pool—10 guts	69	7	7.3 (7.0–16)	97%	0.7 (0.4–1.0)
Effects of antibiotics	BL-4	Artificial	11/05	Pool—10 guts	99	11	14 (11–34)	96%	1.7 (1.6–1.9)
	BL-5	PenStrep	11/05	Pool—10 guts	100	15	43 (22–120)	93%	2.0 (1.8–2.2)
	BL-6	PenStrep » artificial	11/05	Pool—10 guts	98	10	13 (10–33)	97%	1.4 (1.2–1.6)
Larva-to-larva variation	BL-7	Artificial	12/05	Individual gut	69	9	38 (18–115)	94%	0.8 (0.4–1.1)
	BL-8	Artificial	12/05	Individual gut	48	6	9.0 (6.4–31)	98%	0.9 (0.6–1.2)
	BL-9	Artificial	12/05	Individual gut	56	8	16 (10–52)	98%	0.9 (0.6–1.3)
Batch-to-batch variation	BL-10	Artificial	3/04	Pool—10 guts	136	6	7.0 (6.1–20)	99%	0.9 (0.6–1.1)
	BL-11	Artificial	11/04	Pool—10 guts	146	9	9.5 (9.0–17)	99%	0.6 (0.5–0.8)
Larva-to-larva variation	BL-12	Artificial	8/06	Individual gut	77	8	8.0 (8.0–8.0)	99%	1.6 (1.4–1.8)
	BL-13	Artificial	8/06	Individual gut	82	8	14 (12–33)	96%	1.4 (1.2–1.6)
	BL-14	Artificial	8/06	Individual gut	76	10	14 (11–34)	95%	1.7 (1.5–1.9)

Artificial sterile artificial diet, *Sinigrin* sterile artificial diet amended with sinigrin, *PenStrep* sterile artificial diet amended with penicillin and streptomycin, *PenStrep* » artificial PenStrep transferred to sterile artificial diet

^a All sequences within each OTU differ by a sequence divergence of 3% or less

in structure (θ , Yue and Clayton index) and the ratio of shared to unshared species (Jaccard's index). The Yue and Clayton index, θ , measures structural similarity by calculating proportions of the community represented by shared and unshared species and placing more weight on shared species that are similar in abundance than those of dissimilar abundance [47]. We used the Lehman and Tilman measure of total community temporal stability, S_T , as

$$S_T = \frac{\sum \text{abundance}}{\sqrt{\sum \text{var} + \sum \text{cov}}}$$

[48]. Instead of using absolute abundances as was previously described [48], we calculated S_T using relative abundances (proportions of libraries represented by each

OTU_{0.03}). Sums were calculated for the mean proportion, and variance for each OTU_{0.03} represented in at least one of the libraries as well as the covariance for each pair of OTU_{0.03}s.

Susceptibility to Invasion

To assess the effects of diet and antibiotics on invasion, we placed larvae in sterile Petri dishes without food for 4 to 6 h, after which they were fed a diet disk inoculated with approximately 1.0×10^7 colony-forming units (CFUs) of one bacterial strain (Table 2). Each strain was chosen because of its relationship with the cabbage white butterfly midgut community. *Pantoea stewartii* CWB600 is native to the cabbage white butterfly midgut environment but is exogenous

Table 2 Bacterial strains used in this study

Strains	Description	Source/reference	Medium
<i>Pantoea stewartii</i> CWB600	Isolate from midguts of cabbage white butterfly larvae fed Brussels sprouts	This study	1/10th TSB
<i>Bacillus cereus</i> UW85	Zwittermicin-producing isolate from soil with biocontrol activity	[21]	1/10th TSB
<i>Enterobacter</i> sp. NAB3R	Rif ^R spontaneous mutant of isolate from midguts of gypsy moth larvae fed sterile artificial diet	[70]	1/10th TSB
<i>Pantoea</i> sp. CWB304	Nal ^R spontaneous mutant of isolate from midguts of cabbage white butterfly larvae fed artificial diet	[61]	1/10th TSB + Nal

Rif rifampicin, Nal nalidixic acid, 1/10th TSB one tenth strength tryptic soy broth

to the sterile artificial diet community (Table 2). This strain was also naturally resistant to penicillin and streptomycin (data not shown). *Pantoea* sp. CWB304 is native to the communities in larvae reared on sterile artificial diet community and Brussels sprouts and served as a colonization control [49] (Table 2). Two strains were chosen because they were completely exogenous to the cabbage white butterfly midgut community. *Enterobacter* sp. NAB3R was isolated from the gypsy moth and *Bacillus cereus* UW85 was originally isolated from soil [30, 50]. Interestingly, *B. cereus* UW85 also produces zwittermicin, a broad-spectrum antibiotic [51]. Dissections were performed as previously described 24 h after the bacterial feeding [30]. Experiments were conducted at least three times, and bacterial counts were log transformed to reduce heteroscedasticity and analyzed by analysis of variance. When transformations were inappropriate, the Kruskal–Wallis and Mann–Whitney tests were used to determine significance. Statistical computations were conducted using Minitab Statistical Software (Minitab, Inc., State College, PA) and R version 2.4.1 (<http://www.r-project.org/>) [52].

Nucleotide Accession Numbers

16S rRNA gene sequences from this study were deposited in GenBank and are available through the accession numbers DQ342363–DQ343128, DQ349068–DQ349097, DQ537959–DQ538132, EU352364–EU352599, and EU984512.

Results

Species- and Phylum-Level Composition

The bacterial community in the cabbage white butterfly midgut contained 10^3 – 10^6 CFUs per gut (data not shown) and is dominated by organisms whose 16S rRNA genes affiliated with those of the Proteobacteria with additional sequences from the Firmicutes and Bacteroidetes (Fig. 1). The most abundant sequences affiliated with the genera *Methylobacterium*, *Asaia*, *Enterobacter*, *Acinetobacter*, and *Pantoea* (Table 3). Less prevalent sequences affiliated with the genera *Escherichia*, *Roseomonas*, *Lactobacillus*, *Enterococcus*, *Staphylococcus*, *Acidovorax*, *Pseudomonas*, *Rhizobium*, *Bacillus*, *Imtechium*, *Moraxella*, *Ralstonia*, *Hymenobacter*, *Flavobacterium*, *Propionibacterium*, *Nevskia*, *Corynebacterium*, and *Comamonas* (Supplementary Table 1). Eight species affiliates (OTUs that affiliated with specific species) were considered predominant because they were present in at least 50% of the libraries and/or represented the most sequences in at least one library (Table 3). Seven of the eight dominant species affiliates grouped with Proteobacteria and one grouped with Bacteroidetes. Other phyla represented among

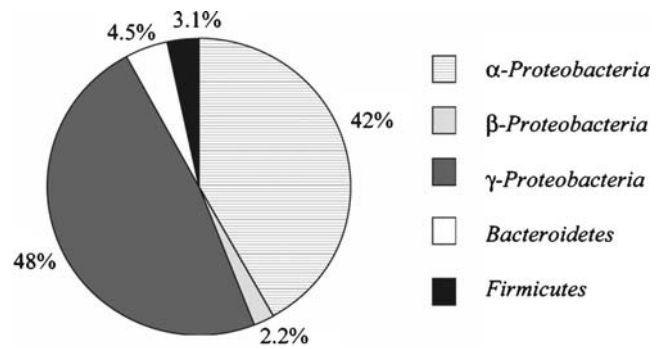


Figure 1 Average distribution of clones affiliated with the α -, β -, and γ -Proteobacteria subphyla and other phyla identified in cabbage white butterfly midgut bacterial communities. Graph excludes Actinobacteria and Chloroflexi, which were each detected once in one sample (0.13%)

the less prevalent species affiliates included Firmicutes, Actinobacteria, and Chloroflexi (Fig. 1). Observations at the phylum level and subphylum level revealed differences in the distribution between the α - and γ -Proteobacteria subphyla (Fig. 1 and Table 3). Three of the seven Proteobacteria affiliates that were dominant community members were α -Proteobacteria, and four were γ -Proteobacteria.

Batch-to-Batch and Larva-to-Larva Variation, and Temporal Stability

In order to assess stability of the gut flora in an insect colony over time and compare batch-to-batch variation, midgut communities from the lab colony larvae were sampled at two dates, March 2004 (BL-10) and November 2004 (BL-11). The communities were similar in both structure and membership at the species level (Fig. 2a and Table 4). In contrast, the structures of communities from larvae reared from commercially obtained eggs sampled in January (BL-1) and November 2005 (BL-4) were not similar in structure or membership at the species level (Fig. 2a, Table 4, and Supplementary Table 1).

There was high similarity among communities in the midguts of the individual larvae that were fed the same diet, sampled at the same time, and sampled on an individual basis (BL-7, BL-8, and BL-9 sampled in December 2005 and BL-12, BL-13, and BL-14 sampled in August 2006). Species affiliates detected in at least two of the three communities sampled in December 2005 included *Methylobacterium*, *Asaia*, *Acinetobacter*, *Ralstonia*, and *Staphylococcus* affiliates as well as an OTU that affiliated closely with the Cytophaga–Flavobacterium–Bacteroides division (Supplementary Table 1). At least two of the three communities sampled in August 2006 contained the same *Methylobacterium*, *Ralstonia*, and *Staphylococcus* affiliates listed above as well as *Propionibacterium* and *Corynebacterium* affiliates (Supplementary Table 1). These communi-

Table 3 Predominant species in the larval midgut community of the cabbage white butterfly

Phylogenetic division	Representative clone GenBank accession no.	Closest culturable match GenBank; accession no.; % identity	Proportion of libraries containing OTU (%)	Representation of OTU in library (%)	Proportion of libraries dominated by OTU (%)
$\tilde{\alpha}$ Proteobacteria	DQ342928	Phenanthrene-degrading bacterium; AY177358; 99%	91	34	36
	DQ342958	<i>Roseomonas gilardii</i> AY150045; 100%	64	3	0
	DQ342728	<i>Asaia</i> sp. SF2.1; AB025929; 99%	55	5	9
γ -Proteobacteria	DQ342888	<i>Enterobacter cloacae</i> ; AB244457; 99%	73	24	27
	DQ342721	<i>Acinetobacter</i> sp. pheno2; APH278311; 99%	73	10	18
	DQ342869	<i>Escherichia coli</i> O157:H7; 99%	64	3	0
	DQ342866	<i>Pantoea</i> sp. PPE7; AY501386; 97%	36	7	9
Bacteroidetes	DQ342843	Flavobacteriaceae bacterium YMS-2; EF017801; 99%	64	4	0

OTUs were present in $\geq 50\%$ of libraries or were the most abundant OTU in a library

ties also shared similar levels of diversity and were almost identical structurally (Fig. 2a, Table 4).

As controls, the temporal stabilities of communities sampled at the same time were calculated. As expected, when the temporal stability of communities in larvae sampled from the same batch was calculated, the stability was high ($S_T=19.6$ and 21.6 for BL 7–9 and BL 12–14, respectively). The temporal stability of all communities,

across all batches (BL 1–9, 12–14) was relatively low ($S_T=2.39$).

Resistance to Diet- and Antibiotic-Induced Changes

Larvae were reared from hatching on sterile artificial diet (BL-1; the control community to which others would be compared), sterile artificial diet amended with

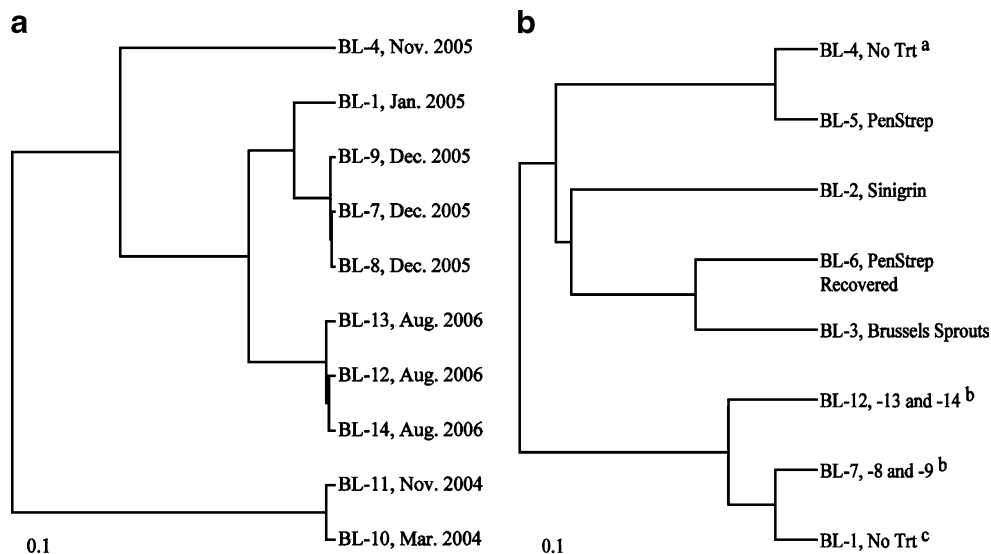


Figure 2 Comparisons of community structure as determined by Yue and Clayton (θ) indices at OTU_{0.03}. **a** Batch-to-batch and larva-to-larva structural variation. Unweighted pair-group method using arithmetic mean (UPGMA) clustering of communities from larvae reared on sterile artificial diet only. Library names are followed by date of sampling. **b** Effects of diet on community structure. UPGMA clustering of communities in larvae reared on sterile artificial diet (No

Trt), Brussels sprouts, sinigrin, penicillin and streptomycin (*PenStrep*), or penicillin and streptomycin and then transferred to sterile artificial diet (*PenStrep Recovered*). Reference bar length corresponds to a distance of 0.10 (distance = $1 - \theta$). No treatment control for antibiotics experiment (a); sequences from libraries constructed from individual larvae reared on sterile artificial diet were pooled and treated as one library (b); no treatment control for diet experiment (c)

Table 4 Pairwise comparisons of structural similarity between batches of larvae and individual larvae fed sterile artificial diet

Libraries compared	Yue and Clayton index ^a	Jaccard's index
Batches of larvae		
BL-1 (Jan. 2005)×BL-4 (Nov. 2005)	0.24 (0.06) ^b	0.24
BL-10 (Mar. 2004)×BL-11 (Nov. 2004)	0.97 (0.01)	0.12
Combined BL-7, 8 and 9 (Dec. 2005)×combined BL-12,13, and 14 (Aug. 2006) ^c	0.65 (0.06) ^b	0.32
Individual larvae		
BL-7 (larva 1-Dec. 2005)×BL-8 (larva 2-Dec. 2005)	0.99 (0.01)	0.23
BL-7 (larva 1-Dec. 2005)×BL-9 (larva 3-Dec. 2005)	0.99 (0.01)	0.5
BL-8 (larva 2-Dec. 2005)×BL-9 (larva 3-Dec. 2005)	0.98 (0.01)	0.4
BL-12 (larva 1-Aug. 2006)×BL-13 (larva 2-Aug. 2006)	0.96 (0.05)	0.45
BL-12 (larva 1-Aug. 2006)×BL-14 (larva 3-Aug. 2006)	0.98 (0.02)	0.58
BL-13 (larva 2-Aug. 2006)×BL-14 (larva 3-Aug. 2006)	0.98 (0.02)	0.43

Larvae within a batch hatched from eggs from the same source were reared at the same time and on the same diet and sampled at the same time

^a Yue and Clayton index provided with 95% confidence interval in parentheses

^b The value differs significantly from 1.0. Similarity increases as value approaches 1.0 for both indices

^c Libraries BL-7, BL-8, and BL-9 and BL-12, BL-13, and BL-14 were constructed from individual midguts. The resulting sequences were combined into two groups based on batch (Dec. 2005 or Aug. 2006) and then compared to each other

sinigrin (BL-2), or Brussels sprouts (BL-3). All communities contained sequences whose closest cultured matches in GenBank were *Methylobacterium* sp. PB133 (99% similarity) and *Roseomonas gilardii* strain ATCC49956 (100% similarity). The addition of sinigrin to artificial diet, or substituting Brussels sprouts for artificial diet, resulted in a different midgut community composition and structure (Fig. 2b, Table 5, and Supplementary Table 1). The compositions of the communities in the larvae reared on Brussels sprouts or sinigrin were more similar to each other than either was to those in larvae reared on unamended diet (Supplementary Table 1). For example, addition of sinigrin to artificial diet or substitution with Brussels sprouts resulted in the presence of an *Enterobacter* species that was not detected

in the communities of larvae fed unamended diet and in the tenfold reduction of the representation of a *Methylobacterium* species that had been the dominant member of the control communities (Supplementary Table 1). Despite differences in total composition, there was overlap among some members of the three communities (Supplementary Table 1).

Resistance to antibiotic-induced changes was assessed by rearing larvae on untreated sterile artificial diet, or sterile artificial diet amended with antibiotics, and sampling the communities. Two experiments were conducted. Trial 1 revealed the communities in these larvae were similar in composition, but the addition of antibiotics resulted in the detection of previously unobserved species (e.g., an affiliate

Table 5 Pairwise comparisons of structural similarity of communities from larvae fed different diets

Libraries compared	Yue and Clayton index ^a	Jaccard's index
BL-1 (unamended) × BL-2 (sinigrin)	0.08 (0.04) ^b	0.22
BL-1 (unamended) × BL-3 (Brussels sprouts)	0.05 (0.03) ^b	0.36
BL-2 (sinigrin) × BL-3 (Brussels sprouts)	0.42 (0.11) ^b	0.28
BL-4 (unamended) × BL-5 (penicillin and streptomycin)	0.88 (0.06)	0.5
BL-4 (unamended) × BL-6 (penicillin and streptomycin recovered)	0.26 (0.06) ^b	0.57
BL-5 (penicillin and streptomycin) × BL-6 (penicillin and streptomycin recovered)	0.37 (0.08) ^b	0.53

Unamended larvae were fed unamended sterile artificial diet, sinigrin larvae were fed sterile artificial diet amended with sinigrin at 3.0 mg/mL, Brussels sprouts larvae were fed Brussels sprouts, penicillin and streptomycin larvae were fed sterile artificial diet amended with penicillin and streptomycin at 10 units/ml and 10 mg/ml, respectively, penicillin and streptomycin recovered larvae were fed penicillin and streptomycin and then transferred to unamended sterile artificial diet 24 h before they molted to fourth instar

^a Yue and Clayton index provided with 95% confidence interval in parentheses

^b The value is significantly different from 1.0. Similarity increases as value approaches 1.0 for both indices

of *Pantoea* sp. PPE7, Supplementary Table 1), and an increase of the Chao1 estimated richness from 14 (11–34) to 43 (22–120) (BL-4 and BL-5; Table 1). Testing whether communities could recover by transferring larvae from diet that contained antibiotics to diet that did not contain antibiotics (BL-6) resulted in a community that was significantly different from the communities in larvae fed only antibiotic-containing diet ($\theta=0.26$, s.e.=0.06) and in larvae fed only unamended diet ($\theta=0.37$, s.e.=0.08; Table 5). The recovery period also resulted in the increased detection of a *Pantoea* species that was previously detected at low levels only in the communities of larvae fed antibiotics (Fig. 3) and in the twofold reduction of an *Acinetobacter* species that represented about 30% of the antibiotic and control communities (Supplementary Table 1).

In trial 2, the addition of antibiotics increased the Chao1 estimated species richness from 8.25 (8.0–12.8) to 15.3 (12.5–34.1), similar to what was observed in trial 1, although less dramatic. Unlike the treatment of communities sampled in trial 1, in trial 2, antibiotics significantly altered the structure of community ($\theta=0.07$, s.e.=0.02). The dominant member of the control community was an *Asaia* species, but the dominant member in the microbial community within larvae fed antibiotics was an *Acidovorax* species. Additionally, antibiotics in artificial diet reduced the abundance of culturable bacteria by 1,000-fold (Fig. 4).

Resistance to Invasion

Establishment by *Pantoea stewartii* CWB600 was enhanced in cabbage white butterfly larval midguts by feeding antibiotics (Mann–Whitney test, $P<0.01$; Fig. 5a). When

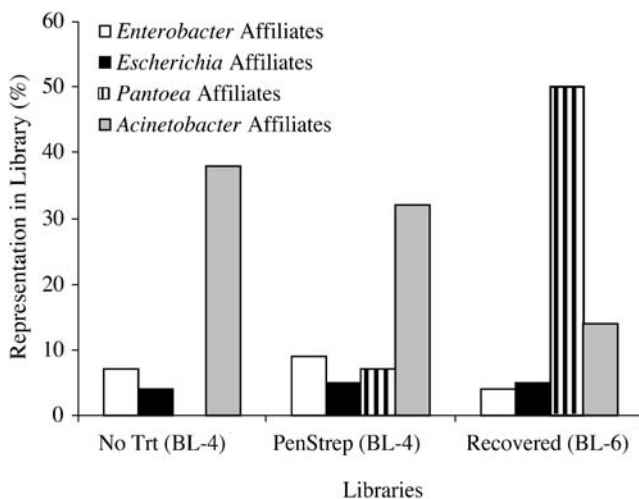
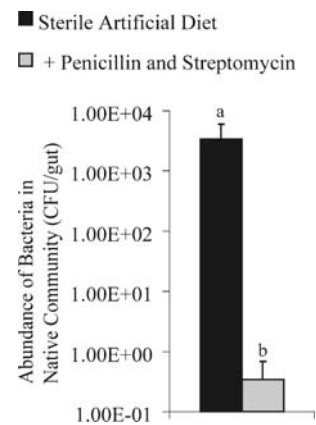


Figure 3 Distribution of predominant γ -Proteobacteria in libraries constructed from larvae reared on unamended sterile artificial diet (BL-4), sterile artificial diet amended with penicillin and streptomycin (BL-5), or sterile artificial diet amended with penicillin and streptomycin and then transferred to unamended artificial diet (BL-6)

Figure 4 Effect of antibiotics on abundance of culturable bacteria. Midguts from cabbage white butterfly larvae fed untreated sterile artificial diet ($n=24$) or penicillin and streptomycin ($n=29$) were plated on one tenth strength tryptic soy agar. Error bars are standard error mean. Different letters indicate significant difference (Kruskal–Wallis and Mann–Whitney; $P<0.05$)



compared to feeding on untreated sterile artificial diet and feeding on Brussels sprouts, feeding on sinigrin increased community susceptibility to establishment by *Enterobacter* sp. NAB3R (Mann–Whitney test, $P<0.01$; Fig. 5b). A similar pattern was observed when *Pantoea* sp. CWB304 was introduced to communities of larvae fed the diets above, although establishment was not increased significantly (Mann–Whitney test, $P=0.13$ and 0.27 , respectively, Fig. 5b).

Discussion

Here, we explore robustness of the cabbage white butterfly midgut bacterial community as a potential model system for studying both specific aspects of microbial ecology and general theories of biological invasion. We calculated variation in community structure over time, examined the effects of diet and antibiotics on this community, and explored the community's susceptibility to invasion.

Steinhaus observed that the alimentary tract of adult cabbage white butterflies contained *Enterobacter* spp. (called *Aerobacter* spp. in 1941), as well as a *Flavobacterium* sp. [33]. We likewise detected these members and further determined that the larval stage of this insect contains several additional genera. The community is similar in composition to those found in other insects that have recently been characterized using culture-independent techniques. For example, *Enterobacter* and *Pantoea* have been identified in numerous insects including gypsy moths (Lepidoptera: Lymantriidae), ant lions (Neuroptera: Myrmeleontidae), biting midges (Diptera: Ceratopogonidae), stable flies (Diptera: Muscidae), grasshoppers (Orthoptera: Acrididae), mosquitoes (Diptera: Culicidae), and thrips (Thysanoptera: Thripidae) as well as the cabbage white butterfly (Lepidoptera: Pieridae) [30, 53–56]. Other members of the cabbage white butterfly larval midgut community that are also found in other insects are *Bacillus*, *Acinetobacter*, *Lactobacillus*, and *Pseudomonas* spp. [57–60].

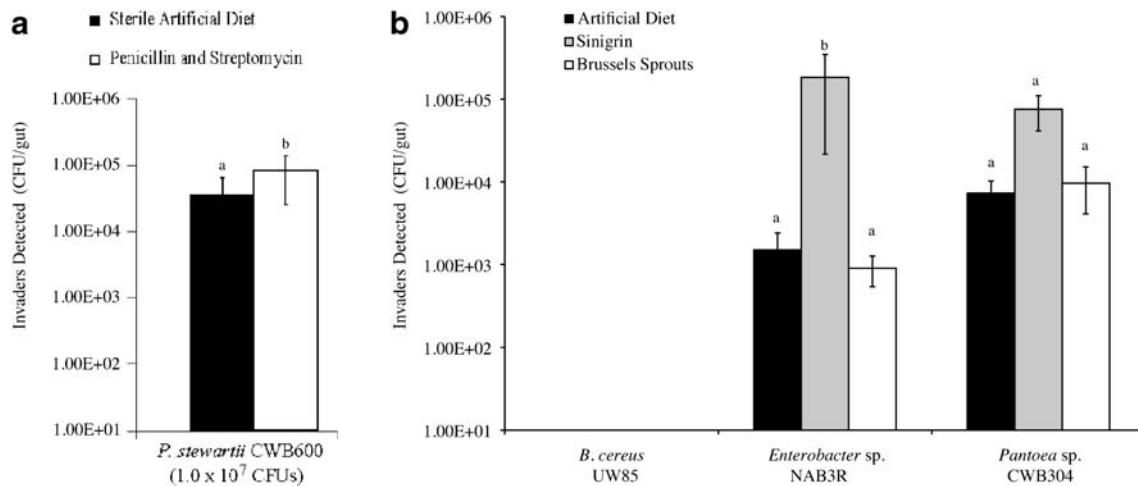


Figure 5 Effects of antibiotics and diet on establishment of invaders in cabbage white butterfly midgut communities. **a** Larvae fed unamended sterile artificial diet ($n=36$) or penicillin and streptomycin (100 units/ml and 100 $\mu\text{g}/\text{ml}$, respectively; $n=33$) from hatching were fed *Pantoea* sp. CWB600. **b** Larvae fed unamended sterile artificial diet ($n=27$) or

sterile artificial diet amended with sinigrin (3.0 mg/ml; $n=18$) or Brussels sprouts ($n=23$) from hatching were fed *Pantoea* sp. CWB304. Error bars are standard error mean. Different letters indicate that values differ significantly (Kruskal–Wallis and Mann–Whitney; P values <0.05

In addition to identifying the compositional similarities that exist between the bacterial communities in cabbage white butterfly and other insects, this study also revealed that the cabbage white butterfly midgut bacterial community exhibits temporal instability at the species level and conservation of membership at the phylum level. Similar patterns of change and conservation of membership that are linked to a specific phylogenetic level occur in other communities as well. The human colon, for example, experiences changes in membership at the species level throughout the lifetime of an individual, between individuals, and in response to diet, but phylum-level and often genus-level memberships are consistent [61–67]. This sort of phylum-level stability is also exhibited in the gypsy moth, in which species composition changes with diet, while the γ -Proteobacteria and Firmicutes phyla are present in under all conditions [30]. This trend toward consistency at higher taxonomic levels and flexibility at lower levels may indicate that the overall function of a community is often more important than the presence of particular members. This explanation has been suggested for reported variations among human gastrointestinal microbial communities [68]. In this case, multiple members within certain groups of bacteria, i.e., the α -Proteobacteria, might be equally able to meet particular functional requirements.

Functional requirements of communities may also play a role in the observation that host plants and phytochemicals, specifically Brussels sprouts and sinigrin, alter community structure in cabbage white butterfly larvae midguts. Our hypothesis was that amending sterile artificial diet with sinigrin, a major component of Brussels sprouts, would alter the community to resemble that in larvae fed Brussels

sprouts. In the wild, the cabbage white butterfly midgut community is consistently exposed to sinigrin, which is a glucosinolate—a class of phytochemicals whose breakdown products have antimicrobial properties [69–73]. Sinigrin resulted in a community that was more similar to the community Brussels sprouts-fed larvae than the sterile artificial diet-fed larvae, but was significantly different from both. Relatives of several cabbage white butterfly community members are able to degrade sinigrin and utilize the end products of insect-mediated sinigrin degradation suggesting the possibility that certain community members assist in degradation of sinigrin and benefit from it [74–80]. Future work including the functional analyses of the community in larvae-fed sinigrin would likely reveal that some proportion of this community is able to degrade and/or utilize sinigrin.

The presence of antibiotics in the artificial diet perturbed the community structure and resulted in changes in susceptibility to invasion, as did the presence of sinigrin. Our results provide experimental support for the view that perturbation can increase the susceptibility of communities to invasion [81, 82]. Both theoretical treatments and correlative analyses have argued for the importance of this relationship, but more widespread acceptance and implementation of corresponding management tactics have lagged pending more direct evidence. The manipulative experiments on insect gut communities described here will hopefully help bridge these various approaches and scales. It is not yet possible to quantify the generality of our results, but it is informative that similar conclusions emerge from studies of macroscale (e.g., lagoon, grassland) [81–84] and microscale (e.g., gut microbial) communities.

These results reinforce the importance of community structure and microbe–microbe interactions and support the view that the presence or absence of only a few members may influence community resistance and susceptibility to invasion. This raises another point of interest that has been explored in macroscale communities—the impact of invaders on existing community interactions [85, 86]. Because of this community's relatively low richness, it would be amenable to studies that seek to introduce an exogenous species and monitor the resulting changes in the community.

Our experiments with antibiotics indicate that perturbation can alter the relative abundance of various community members. Specifically, we detected community members after antibiotic treatment that were not detectable before treatment. The antibiotic might have reduced the population sizes of some members that normally dominate the community, enabling rare members to fill the vacated niche. For example, a *Pantoea* affiliate that was not detected in the artificial diet community (trial 1, BL-4) and represented a small fraction of the community in antibiotic-fed larvae became the numerically dominant member (51%) of the recovered community (Fig. 3). Similarly, the population of this *Pantoea* affiliate increased to 20% of the community following larval consumption of Brussels sprouts. Antibiotic treatment also increased the number of species present at low abundance, producing higher species richness. The results of both studies indicate that this community may have low resistance to structural change when confronted with antibiotic exposure, and trial 1 suggests that the community may lack short-term structural resilience after antibiotic exposure. A longer recovery period might reveal that the community has the capability to return to a structure similar to its native state as is the case for other microbial communities (i.e., [18]). The differences we observed between our antibiotic trials, as well as the differences among batches of larvae suggest that the assembly and reassembly processes are complex and include both elements of randomness and underlying species interactions, host–microbial relationships, and external drivers not yet understood. These features make the cabbage white butterfly community a rich opportunity for studying secondary succession, the process by which a community reestablishes following a disturbance [87].

Model systems have proven to be essential to understanding microbial interactions [88]. For example, study of the squid–*Vibrio* symbiosis led to the discovery of bacterial quorum sensing [89–92]. The termite hindgut has been a fruitful source of information about metabolic processes in communities, including the demonstration of microbe-regulated oxygen and hydrogen gradients and the linking of function and spatial organization to specific organisms [93, 94]. Each of these systems has characteristics that make it amenable to the kinds of studies that led to

development of new principles in microbial ecology. We present the cabbage white butterfly larval community as a relatively simple, easily manipulatable, multispecies community in which to test ecological hypotheses about interspecies interactions and community robustness.

In this study, we initiated the evaluation of robustness of the bacterial community in the cabbage white butterfly larval midgut by measuring temporal stability, resistance, and resilience. Further exploration of robustness will generate principles that govern the dynamics of this community and perhaps others. Understanding the determinants of robustness will also require development of improved statistical tools to quantify it. Because the lepidopteran larval gut presents a community that is relatively simple, tractable, and easy to manipulate, it is ideal for building and testing statistical models and investigating ecological events such as succession and invasion and the basis for robustness.

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