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Abstract

Background: The native microflora associated with mosquitoes have important roles in mosquito development and vector competence. Sequencing of bacterial V3 region from 16S rRNA genes across the developmental stages of *Culex* mosquitoes (early and late larval instars, pupae and adults) was used to test the hypothesis that bacteria found in the larval stage of *Culex* are transstadially transmitted to the adult stage, and to compare the microbiomes of field-collected versus laboratory-reared mosquitoes.

Results: Beta diversity analysis revealed that bacterial community structure differed among three life stages (larvae, pupae and adults) of *Culex tarsalis*. Although only ~2 % of the total number of bacterial OTUs were found in all stages, sequences from these OTUs accounted for nearly 82 % of the total bacterial sequences recovered from all stages. *Thorsellia* (*Gammaproteobacteria*) was the most abundant bacterial taxon found across all developmental stages of field-collected *Culex* mosquitoes, but was rare in mosquitoes from laboratory-reared colonies. The proportion of *Thorsellia* sequences in the microbiomes of mosquito life stages varied ontogenetically with the greatest proportions recovered from the pupae of *C. tarsalis* and the lowest from newly emerged adults. The microbiome of field-collected late instar larvae was not influenced significantly by differences in the microbiota of the habitat due to habitat age or biopesticide treatments. The microbiome diversity was the greatest in the early instar larvae and the lowest in laboratory-reared mosquitoes.

Conclusions: Bacterial communities in early instar *C. tarsalis* larvae were significantly more diverse when compared to late instar larvae, pupae and newly emerged adults. Some of the bacterial OTUs found in the early instar larvae were also found across developmental stages. *Thorsellia* dominated the bacterial communities in field-collected immature stages but occurred at much lower relative abundance in adults. Differences in microbiota observed in larval habitats did not influence bacterial community profiles of late instar larvae or adults. However, bacterial communities in laboratory-reared *C. tarsalis* larvae differed significantly from the field. Determining the role of *Thorsellia* in mosquitoes and its distribution across different species of mosquitoes warrants further investigation.

Keywords: Thorsellia, Outdoor mesocosms, Bacteria, Biopesticide, Transstadial transmission

Background

Bacteria ingested by the immature stages of mosquitoes (Culicidae) are generally known to provide nutrition [1, 2], facilitate successful development [3, 4], and influence vector competence [5–8]. The presence of certain bacteria has been also shown experimentally to provide

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immunity against pathogens, and reduce vector competence [9–11]. Therefore, interest has grown in understanding the obligate and facultative roles of the gut microflora of mosquitoes, their interaction with parasites and their manipulation to devise alternative and more sustainable vector control strategies (e.g., paratransgenic control) [7, 9, 12–16].

To date, the majority of the studies on microbiomes of mosquitoes have focused largely on investigating the gut microflora (and/or symbionts) associated only with adult mosquitoes (e.g., 9). The focus on adult mosquitoes is justified, on one hand, because adult mosquitoes

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transmit pathogens directly or cause nuisance to humans and animals. On the other hand, the interactions of bacteria and immature stages of mosquitoes will potentially influence the microbiome of the adult mosquitoes. Moreover, previous studies focused on the dominant tropical disease vector mosquito species such as Anopheles gambiae, and not much was known about North American native mosquito species such as Culex tarsalis. Few studies have considered transstadial transmission (larvae to pupae to adult) of microbial communities because the entire midgut of culicid mosquitoes is generally thought be replaced during development such that entire bacterial communities associated with the larval midgut are eliminated prior to eclosion [2, 17]. However, circumstantial evidence in Anopheles mosquitoes suggested that some bacteria species found in larval stages persist through metamorphosis and are transferred to adults [18, 19]. Recently, these contradictory observations have been reconciled in studies from Anopheles stephensi where the Malpighian tubules function as a "refugium" for bacteria during metamorphosis [20, 21]. In addition, most other studies were based on laboratory-reared mosquitoes, fed standardized diets, and raised under controlled environmental conditions for several generations. However, these artificial conditions are likely to restrict and/or alter the host microbiomes of lab-raised insect populations relative to natural populations (e.g. Drosophila; [22]).

The Gram-negative genus Thorsellia (Gammaproteobacteria) is the most abundant bacterial group found among the gut microflora of field-collected late instar larvae of Culex spp. (3rd and 4th instars; [23]). Thorsellia anophelis was first described from adult Anopheles arabiensis in Kenya [24, 25] and is the type specimen for a new family of the Bacteria, Thorselliaceae [26]. Thorsellia anophelis has been reported to be the predominant bacterial species found in adult Anopheles gambiae sensu lato [27] and has also been found in Anopheles culicifacies [28]. It has been hypothesized that the bacterium is acquired during larval feeding, and is then transferred transstadially to adult Anopheles, although the evidence is somewhat conflicting [18, 19, 27]. Rani and colleagues recovered this bacterium in both larvae and adults of Anopheles stephensi, but not in the pupal stage [18], whereas Briones and colleagues did not recover the bacterium from either larval or pupal stages of Anopheles [27]. However, others found Thorsellia to be the 5th most abundant genus in Anopheles gambiae larvae, and the 10th most abundant genus in pupae [19]. Direct evidence supporting the transstadial transmission of Thorsellia and other bacterial species in the microbiome of *Culex* mosquitoes is lacking.

It was unknown whether succession or other factors such as use of pesticides causing changes in bacterial

communities of the larval developmental sites also affect the microbiome in mosquitoes. Pesticides (i.e., malathion, permethrin, atrazine and glyphosate) alter bacterial communities in the larval environment [29]. Removal of mosquito larvae using biopesticides such as *Bacillus thuringiensis* subsp. *israelensis* (*Bti*) was reported to lessen grazing pressure on bacterial communities [30–32] and increase the diversity of bacteria in the habitats [33].

In this study, we: 1) identified bacteria found in the guts of different developmental stages of *C. tarsalis*; 2) compared the microbiomes of field-collected *C. tarsalis* larvae with laboratory-reared individuals; and 3) assessed the effects of habitat age and manipulating larval mosquito density (using *Bti* applications) on the microbial communities found in late-instar mosquito larvae.

Results

Bacterial taxa in developmental stages of C. tarsalis

A total of 14,634 OTUs (4,609,186 bacterial sequences) were generated from 41 Culex mosquito samples (39 field-collected and two laboratory-reared). Proteobacteria (56 %), Bacteroidetes (15 %), Cyanobacteria (14 %), Firmicutes (7 %), Actinobacteria (2 %), and Spirochaetes (2 %) were the most abundant bacterial phyla found in developmental stages of Culex mosquitoes (Additional file 1: Table S1). Unclassified sequences accounted for ~0.6 % of sequences, whereas, bacterial sequences unclassified to phyla accounted for 0.8 % of sequences. Another 30 phyla accounted for the remaining 1 % of the bacterial communities. Although the relative abundance of bacterial taxa changed across the mosquito developmental stages, Proteobacteria were the dominant bacteria found in the guts of C. tarsalis (Fig. 1).

A total of 235 bacterial OTUs, comprising 82 % of the total bacterial sequences, were found among the three life stages of field-collected larvae (early and late instars), pupae, and adults. However, only 27 OTUs classified into seven phyla, representing 66 % of all the sequences recovered from the field-collected mosquitoes, were shared among the samples from all developmental stages (Additional file 2: Table S2). Proteobacteria, Cyanobacteria, Bacteroidetes and Firmicutes were the most abundant phyla shared among the different stages (Fig. 1). Thorsellia (40 %), Cyanobacteria (18.0 %) and Dysgonomonas (13 %) were the most abundant genera shared across developmental stages. A Thorsellia OTU (#2664) was found in all developmental stages of field-collected larvae, pupae, and adults of C. tarsalis, and was the most abundant bacterial species across all mosquito life stages (Additional file 2: Table S2). Phylogenetic analysis using longer 16S rRNA gene sequences (1070 bp) of *Thorsellia* obtained from C. tarsalis larvae revealed that this species is closely related to T. anophelis described from

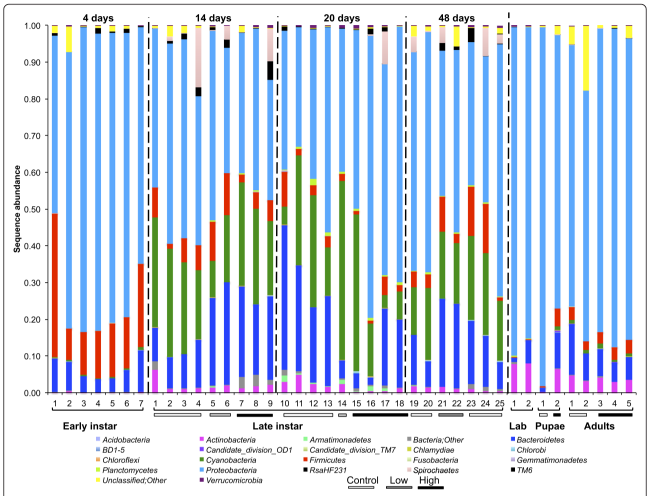


Fig. 1 Proportional sequence abundance of bacterial phyla across *Culex* developmental stages. The samples were taken 4, 14, 20 and 48 days after the onset of the experiment. Only phyla with an average abundance ≥ 0.01 % were included. Numbers on the x-axis represent sample identification numbers. Stages of mosquitoes are denoted by Early instar, Late instar, Lab = lab reared late instar, Pupae, and Adults. Light bar refers to control; gray refers to low *Bti* and dark bar for high *Bti* treatments

Anopheles gambiae sensu stricto but may also contain a separate strain (Fig. 2). The two *Thorsellia* sequences from *Culex* mosquitoes were both 99 % similar to *T. anophelis*, but with individual differences. Out of the six clones sequenced, five clones were identical to clone #6, whereas clone #7 was unique but had nucleotide differences that

are supported by the other *Thorsellia* species (data not shown).

Early instar larvae (not identified to species)

A total of 5,888 OTUs (805,169 sequences from 7 samples) in 34 bacterial phyla were recovered from early (1st

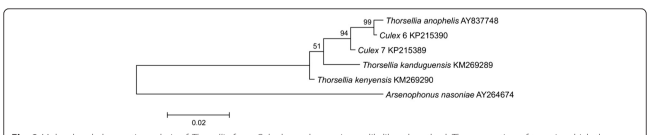


Fig. 2 Molecular phylogenetic analysis of *Thorsellia* from *Culex* larvae by maximum likelihood method. The proportion of trees in which the associated taxa clustered together is shown next to the branches (1000 bootstraps). The tree is drawn to scale, with branch lengths measured in the number of substitutions per site

and 2nd) instars of *Culex* larvae. *Proteobacteria* (73 %), *Firmicutes* (18 %), and *Bacteroidetes* (7 %) dominated the early instar *Culex* larvae (Additional file 1: Table S1). Overall, bacterial taxa in *Gammaproteobacteria* (43 %), *Betaprotobacteria* (26 %) and *Bacilli* (13 %) were the three most abundant classes found in the early stages of the mosquito life cycle (Additional file 1: Table S1). *Thorselliaceae* (27.2 %) and *Comamonadaceae* (21.2 %) were the two most abundant families found in early instar larvae (Fig. 3). *Thorsellia* was the most abundant (19 %) genus found in the early stages followed by an unclassified taxon of *Gammaproteobacteria* (8.8 %) and *Aeromonas* (7 %) (Additional file 3: Table S3).

Late instar C. tarsalis larvae Field-collected C. tarsalis larvae

Overall, 34 bacterial phyla (10,115 OTUs from 25 samples) were found in field-collected late larval (3rd and 4th) instars. *Proteobacteria* accounted for nearly half (49 %) of the sequences followed by *Cyanobacteria* (20 %), *Bacteroidetes* (18 %), and *Firmicutes* (6 %) (Fig. 1). *Spirochaetes* and *Actinobacteria* accounted for nearly

3 % and 2 %, respectively, of the sequences recovered from late-instar larvae. *Thorselliaceae* (43.9 %) and *Porphyromonadaceae* (24.9 %) were the two most abundant families found in late-instar larvae (Fig. 2). *Thorsellia* was the most abundant genus in field-collected late instar larvae (27 %) followed by *Cyanobacteria* (19 %) and *Dysgonomonas* (*Bacteroidetes*; 11 %) (Additional file 4: Table S4).

Laboratory-reared larvae

The late-instar larvae from a laboratory colony were dominated by *Proteobacteria* (87 %) followed by *Actinobacteria* (8 %) and *Bacteroidetes* (4 %). Overall, 786 OTUs (319,786 sequences from 2 samples) were obtained from the laboratory-reared colonies. *Enterobacteriaceae* (77.8 %) was the most abundant family found in late instar laboratory-reared mosquitoes (Fig. 4). *Rahnella* (*Gammaproteobacteria: Enterobacteriaceae*) (64 %), unclassified *Enterobacteriaceae* (12 %), and unclassified *Microbacteriaceae* (*Actinobacteteria*) (8 %) dominated the bacterial communities in the larvae from the laboratory-reared colony (Additional file 5: Table S5).

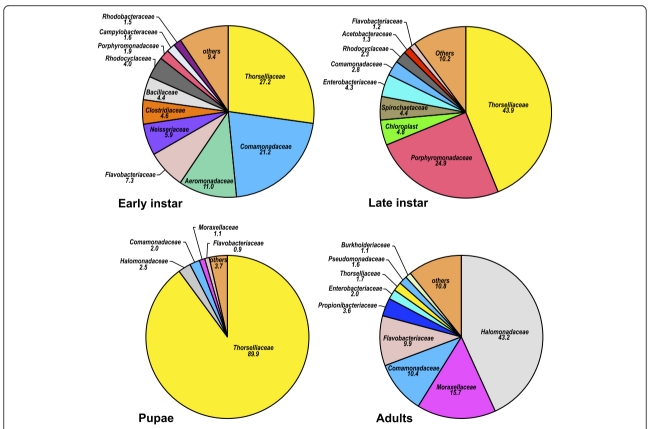


Fig. 3 Family-level abundance of bacterial communities. Family-level abundance (%) of bacterial communities in field-collected early (1st and 2nd) instar larvae, late (3rd and 4th) instar larvae, pupae and adults of *C. tarsalis*. Only sequences classified to family level were included. Because the treatments effects on the gut bacterial community structure within each stage were not significantly different, mosquitoes from all treatments were included in this figure

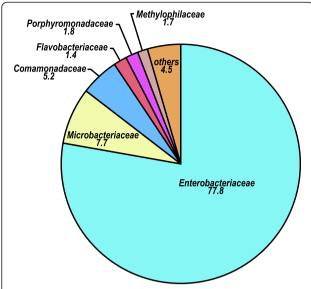


Fig. 4 Family-level abundance (%) of bacterial communities in laboratory-reared late instar *C. tarsalis*. Only sequences classified to family level were included. These samples were from a laboratory *C. tarsalis* colony

In addition to the two OTUs of Rahnella, we found 24 other bacterial OTUs, including one OTU of each of Thorsellia (OTU# 2464), Spirochaeta, Aeromonas, Enterobacteriaceae, Rhodocyclaceae, Bacillaes, Microbacteriaceae, Hydrogenophaga, Halomonadaceae, Alteromonadales, Flavobactrium, Diaphorobacter, Cloacibacterium, Aquabacterium, Propionibacterium, Sphingomonadales, Dysgomonas, and two OTUs of Cyanobacteria, Gammaproteobacteria and Clostridium that were also shared among other samples.

In comparison to late instar larvae collected from the field, nearly 62 % (484/786) of all the OTUs found in lab were also found in the field-collected larvae, whereas only 5 % (484/10,115) of the OTUs found in the field-collected larvae were found in the lab colony.

C. tarsalis pupae

Twenty phyla (represented by 638 OTUs from two samples) were recovered from *C. tarsalis* pupae among which *Proteobacteria* (96 %), *Bacteroidetes* (1.8 %), *Actinobacteria* (0.8 %), *Firmicutes* (0.7 %), and *Cyanobacteria* (0.1 %) were the dominant phyla (Additional file 1: Table S1). Unclassified bacteria accounted for 0.7 % of sequences. *Thorselliaceae* dominated the bacterial community at the family level (Fig. 3). Members of *Gammaproteobacteria* genera [*Thorsellia* (89.9 %), an unclassified taxon in *Halomonadaceae* (2.4 %), an unclassified taxon in *Alteromondales* (2.1 %), and *Actinobacter* (0.9 %)] dominated the bacterial communities in *C. tarsalis* pupae (Additional file 6: Table S6). However, the composition of the bacterial communities in the two samples varied

considerably. Pupae collected from untreated control contained a greater percentage (93.2 % of 101,093 sequences) of *Thorsellia* and the bacterial community was similar to late-instar larvae. The bacterial community in pupae collected from the high *Bti* treatment contained a much smaller (1 % of 7,835) relative abundance of *Thorsellia* and was more similar to that in adults.

Newly emerged non-blood-fed adult C. tarsalis

Overall, 277 genera representing 831 OTUs (47,700 sequences from five samples) were found in newly emerged non-blood-fed adult C. tarsalis. The most abundant bacterial phyla found in eclosing C. tarsalis included Proteobacteria (83.0 %), Bacteroidetes (8.2), Actinobacteria (4.0 %), Firmicutes (3.3 %), and Cyanobacteria (1.0 %) (Additional file 1: Table S1). Another 22 bacterial phyla contributed for < 1 % of the total sequences recovered from adults. At the family level, the bacterial communities of adult C. tarsalis were dominantly enriched with an unclassified taxon of Halomonadaceae (43 %), Moraxellaceae (10.9 %), Commamonadaceae (7.2 %), and Flavobacteriaceae (6.9 %) (Fig. 3). Three Gammaproteobacteria members [Halomonas (29.9 %), an unclassified taxon in Altermonadales (19 %), and Acinetobacter (10.4 %)] were the dominant genera found in adult mosquitoes (Additional file 5: Table S5). Thorsellia accounted for 1.3 % of all the sequences recovered from adult mosquitoes (Additional file 7: Table S7).

Diversity of bacterial communities across developmental stages of *Culex*

Beta diversity of bacterial communities differed significantly among mosquito developmental stages as assessed by multi-response permutation procedure (MRPP) on the Bray-Curtis and UniFrac distance matrices (Fig.s 5 and 6: MRPP analysis: A: 0.32, p < 0.001 for both). Bacterial diversity in early-instar *Culex* larvae from the first sampling date separated significantly from that of the late-instar larvae, pupae, and adults. Bacterial communities from lab-reared mosquitoes differed significantly from the bacterial communities in field-collected samples (Fig. 5). Only two samples of pupae and two samples of lab-reared larvae were analyzed in this study. Bacterial communities from one of the two pupal samples grouped tightly with adult samples, whereas the other sample was closely clustered to samples of fieldcollected late-instar larvae, mainly due to the differences in abundance of Thorsellia (Fig. 6). Bacterial communities from late-instar Culex larvae collected from the field were significantly separated from the other developmental stages (PC 1) and were dominated by Thorsellia and Cyanobacteria.

Alpha diversity of bacterial communities in early instar *C. tarsalis* larvae was significantly higher than the

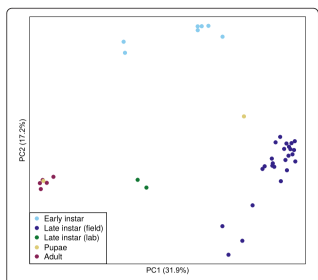


Fig. 5 Principal coordinate analysis (based on Bray-Curtis distances) of bacterial communities in *C. tarsalis*. Bacterial communities in early and late instars, pupae, and adults sampled from outdoor mesocosms, and in laboratory-reared late larval instars of *C.tarsalis* were significantly separated. Pupae and adults were collected only from high *Bti* treatment and untreated control (see material and methods for details). Late instar larvae were sampled from all treatments

bacterial diversity found in field-collected late larval instars and adults, and in laboratory-reared late-instar larvae (p < 0.001, Fig. 7). The diversity of bacterial communities in pupae varied considerably between the two samples and did not differ from the other developmental stages. The laboratory-reared larvae contained significantly less diverse bacterial communities than did larvae collected from the field (Fig. 7).

Influences of *Bti* and habitat age on gut bacterial communities of late instar mosquitoes

The bacterial communities in the late instar *Culex* larvae were not influenced significantly by age of the habitat or the larvicide treatments (Fig.s 8 and 9; MRPP $A \le 0.1$), although there was noticeable separation of bacterial communities from high *Bti* treatments from the other treatments. The bacterial communities in these larvae were dominated by OTUs of *Thorsellia*, *Cyanobacteria*, *Porphyromonadaceae*, *Aeromonadaceae*, *Rhodocyclaceae*, *Clostridiales* and *Spirochaeta* in descending order of abundance (Fig. 8).

Discussion

Marker gene sequence analysis revealed the dynamics of bacterial community structure across developmental stages of *Culex tarsalis* (Fig. 1) and suggested transstadial transmission of some bacterial groups in *Culex* mosquitoes from North America. Overall, 235 bacterial OTUs in the phyla *Proteobacteria*, *Cyanobacteria*,

Bacteroidetes, Actinobacteria, Firmicutes, Spirochaetes, and RsaHF231 persisted from the early larval instars to newly emerged adults of *C. tarsalis*. The relative abundance of taxa in bacterial microbiome differed among the developmental stages (Fig. 5) but, within 3rd and 4th instars, did not differ significantly across sampling dates or in response to changes in the microbial community of the feeding zone that occurred following the application of a high dose of biopesticides. Our study provides additional evidence that, contrary to a commonly accepted hypothesis [17], metamorphosis does not completely eliminate gut microbiota. In *Culex pipiens* mosquitoes, some bacterial communities from the larval stages were shown to be sequestered in meconial peritrophic membrane and can be passed to the adult stages [17].

Thorsellia was found across all stages of C. tarsalis collected from the field but was rare in mosquitoes from laboratory colonies. However, the proportion of Thorsellia sequences in field-collected mosquitoes decreased during development. In adults derived from fieldcollected larvae, Thorsellia only accounted for 3.3 % of all sequences identified to the genus level, as compared to the early (31 %), and late (52.3 %) instar larvae, and pupae (93 %). This genus accounted for only 0.09 % of sequences identified to genus level from a lab colony of late C. tarsalis instar larvae suggesting that this bacterium is associated more strongly with natural aquatic habitats. The possible symbiotic role played by the *Thor*sellia to the different species of mosquitoes is currently unknown. Briones et al. reported a dominance of T. anophelis in the guts of adult Anopheles gambiae [27]. Likewise, Wang and colleagues found Thorsellia to be the dominating species in newly emerged adults making up 2/3 of their bacteria [19]. In this study, Thorsellia sequences obtained from C. tarsalis adults only accounted for a very small proportion of the bacterial sequences. That said, Thorsellia was found to be the primary resident in the guts of early and late larval instars of fieldcollected C. tarsalis (this study and 23). Overall, nearly 44 % of the sequences identified to genus level in the present study were Thorsellia sequences. Our finding indicates the persistence of taxa from this bacterial genus across all developmental stages of *C. tarsalis*.

Early instar larval mosquitoes harbored significantly more diverse bacterial communities (in three of four diversity indices) than the late-instar larvae collected from the field, and pupae and adult stages. The early instar larvae were sampled four days after the onset of the experiment and the bacterial communities in the guts of these mosquitoes mirrored the higher bacterial diversity observed in the water column on this sampling date [33]. It is unknown whether the reduction of bacterial communities especially during the late stages of development was due to the dominance of phytoplankton

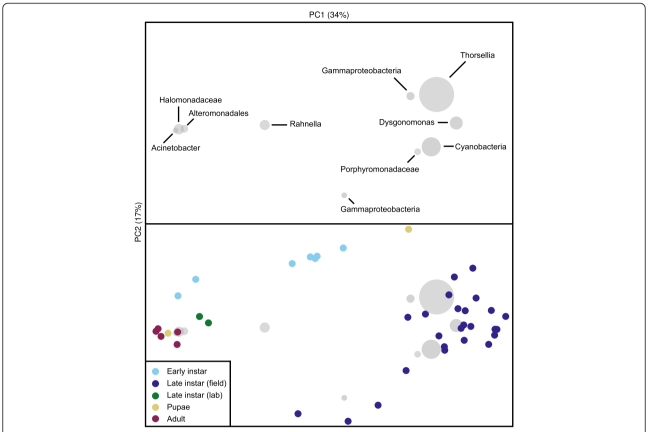


Fig. 6 Principal coordinate analysis ordination (based on weighted UniFrac distances) of bacterial communities sampled from different stages of *C. tarsalis*. Top panel: The ten most abundant bacterial taxa found in the different mosquito life stages as they relate to the ordination space in the bottom panel. Bottom panel: Mosquito life stages projected on bacterial community profiles of samples collected in this study. "Lab" is late instar larvae from a laboratory colony of *C. tarsalis*

(particularly *Cyanobacteria*) in the water column (the feeding zone of *Culex* larvae) or the reduction of some bacterial taxa from the gut of the mosquitoes during metamorphosis. In addition, the early instar larvae likely included *Culex stigmatosoma* and *Culex quinquefasciatus*, and the greatest diversity observed in this stage might be because different mosquito species were present in the samples.

Late instar *Culex* larvae maintained fairly stable bacterial communities in their gut regardless of changes of microbial communities in the feeding zones of developmental sites. Duguma *et al.* also found that the bacterial communities within mosquito larvae did not change significantly during succession in two bioremediation treatments [23]. As revealed by PCoA ordinations, marked differences in the diversity of bacterial taxa were found between late instars, pupae and adults. Although the majority of the microbes found in the feeding stages (i.e., larvae) were lost likely during metamorphosis, our study suggests that certain species of bacteria were found among life stages investigated and warrant further experimental evaluation of these microbiomes whether

they are obligate or facultative symbionts and persist through the different mosquito life stages.

The microbiome communities in laboratory-reared mosquitoes differed significantly from those found in mosquitoes in nature. We found that bacterial communities from field-collected Culex larval guts were significantly more diverse than their lab-reared counterparts. On average, 21 % of the sequences per sample from field-collected larvae were Cyanobacteria, whereas this taxon was virtually absent from the guts of lab-reared mosquitoes (Fig. 1). Similarly Bacteroidetes accounted 17 % of sequences found in the larvae collected from field whereas this bacterium only accounted 3 % of the sequences recovered from the lab-reared colony. The bacterial communities from laboratory-reared C. tarsalis larvae were primarily dominated by Rahnella (Gammaproteobacteria: Enterobacteriaceae), which occurred rarely in the field-collected larvae. Rani et al. also reported a significant reduction of bacterial diversity in lab-reared Anopheles stephensi [18]. It is well documented that gut microbial consortia play a significant role in shaping the physiology of insects [34, 35].

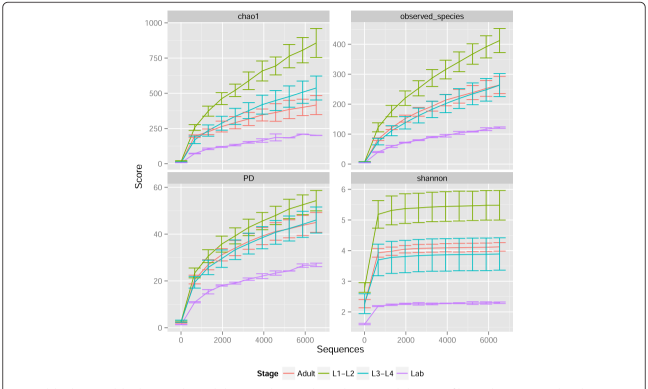


Fig. 7 Alpha diversity. Alpha diversity (Chao1, Phylogenetic diversity, observed species and Shannon) of bacterial communities (based on OTUs) in different *Culex* mosquito stages: early-instar larvae, late-instar larvae (lab-reared or field-collected) and adults. Pupal samples were not included in the alpha diversity plots shown below because there was a large variation in the number of sequences between the two samples analyzed

Bacteria found in lab-reared fruit flies were either rare or absent from wild *Drosophila* populations [22] posing a fundamental question on the extrapolations of many lab-reared host insect-microbiome models to the natural populations.

The high *Bti* treatment (equivalent to 48 kg Vectobac *G* ha⁻¹) that reduced late (3rd and 4th instars) larval mosquito abundance by >50 % for a month, changed the microbiota (bacterial communities and phytoplankton) and nutrient concentrations in the water column [33]. However, neither the differences in planktonic bacterial communities caused by this larvicide treatment nor the differences in succession of bacterial communities across time influenced the bacterial communities in the gut of *C. tarsalis*. Interestingly, the water column bacterial diversity was significantly higher in the high *Bti* mesocosms and significantly separated from the other treatments on this date [33].

Although the majority of late instar larvae were removed from the water column by the high *Bti* treatment, the late instar larvae that were analyzed in this study were likely unaffected by the *Bti* application at the time of sampling. Nine weeks (44 days) after the *Bti* application, larval mosquito abundance in the high *Bti* treated mesocosms was significantly higher than in the low *Bti* and untreated control mesocosms [33]. Assessing the

gut microbiota of mosquitoes surviving the biopesticide treatment has important implications because it might help understand the resistance mechanism and other physiological attributes provided by microbiota to the larval mosquitoes [35].

C. tarsalis is one of the most important vectors of arboviruses (e.g., western equine encephalomyelitis, St. Louis encephalitis and West Nile viruses) in North America. Interestingly, this mosquito species lacks endosymbionts such as Wolbachia, which is ubiquitous among several other arthropods including congeners [23,36]. Most other notable symbionts of mosquitoes such as Asaia and Spiroplasma are also lacking from C. tarsalis. The present study and a previous study [23] showed the dominance of the promising potential symbiont candidate, Thorsellia, in C. tarsalis collected from natural habitats. Both studies also reported the first evidence of this bacterium in a North American mosquito species. The function of this potential symbiont in the nutrition, physiology and vector competence of this mosquito species warrants further investigation.

Conclusions

The metagenomic analysis of the bacteria microbiome (both cultivable and uncultivable) revealed significant differences among three life stages (larva, pupa and

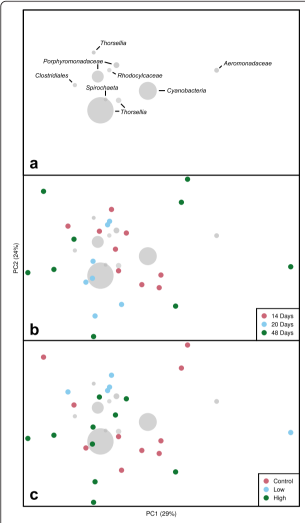


Fig. 8 Principal coordinate analysis ordination (based on weighted UniFrac distances) of bacterial communities sampled from *C. tarsalis* late-instar larvae. Top panel (**a**): 10 most abundant taxa found in field-collected late-instar *C. tarsalis* larvae as they relate to the ordination space in the middle and bottom panels. Middle panel (**b**): bacterial community profiles from mosquitoes sampled in different habitat ages. Bottom panel (**c**): bacterial community profiles from mosquitoes sampled in three *Bti* treatments. Neither *Bti* treatment nor sampling date significantly influenced the bacterial communities in late-instar larvae of *C. tarsalis*

newly emerged adults) of *C. tarsalis* collected from the field, and larvae from a laboratory colony. The greatest diversity was observed in the early instar larvae and the lowest bacterial diversity was found in the lab colony. *Thorsellia* (*Gammaproteobacteria*) dominated the active feeding stages (larvae) collected from the field and persisted through non-feeding developmental stages suggesting, respectively, that these bacteria are ingested from the aquatic habitats and transstadial transmission is possible between stages. However, it is unknown whether the persistence of *Thorsellia* in the gut of larval

mosquitoes is due to its resistance to digestion or is the result of continuous ingestion from the environment. The physiological role of this dominant bacterium within mosquitoes and whether it invades tissues also warrants further investigation.

The higher bacterial diversity in early instar larvae compared to the later developmental stages might have been due to either species-specific microbiomes that differed among C. tarsalis and two other congeners in the early -instar samples or other factors associated with the age of the habitat. The bacterial diversity was highest in the water column early in succession in the mesocosms. The gut bacterial community within late instar larval mosquitoes in outdoor mesocosms was conserved and did not change concomitantly with changes in the bacterial community present in the larval feeding zone across time as well as following the manipulation of larval density using a biopesticide treatment. The gut bacterial communities in pupae and adults were less diverse than in larvae. The mechanism(s) causing the loss of gut bacteria diversity during ontogeny require further study.

Finally, the bacterial communities found in laboratory-reared larvae were significantly less diverse compared to those found in field-collected larvae and only constituted a very small percentage (5 %) of the bacterial microbiomes found in the field-collected counterparts. *Thorsellia* was rare in mosquitoes from the laboratory colony. The low microbiome diversity found in the laboratory mosquitoes could have important implications for the interpretation of results and applicability to natural populations from mosquito-microbiomes studies using laboratory-reared mosquito larvae.

Methods

Study site and experimental design

The study was conducted in 1 m² experimental outdoor mesocosms at the Aquatic and Vector Control Research Facility of the University of California Riverside Agricultural Experiment Station. Detailed description of the study site and experimental protocol are given in a previous study [33] but briefly, each of twelve mesocosms were filled with 300 liters of water from an irrigation reservoir and enriched with both organic (50 g of alfalfa rabbit pellets) and inorganic nutrients (40 g of ammonium sulfate) on September 28, 2012. The nutrients were added to stimulate mosquito and microbial colonization. On October 2, three treatments (two application rates of Bti [Vectobac G: high = 4.81 g/mesocosm and low = 0.06 g/mesocosm] and an untreated control) were assigned to the mesocosms in a completely randomized experimental design. The mean water level was adjusted to 30 cm on October 1, and adjusted again to 30 cm on October 5, and no water was added afterwards. The mean water level in the 12 mesocosms was

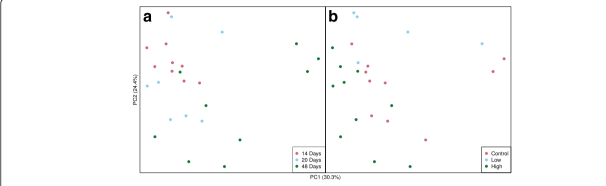


Fig. 9 Principal coordinate analysis of bacterial communities in *C. tarsalis* late-instar larvae. Late-instar larvae collected from three *Bti* treatments (a), and three sampling dates (b)

28 (± 0.32 ; mean \pm SE) cm on October 12, and 25 (± 0.28) cm on October 19 and 17 (± 0.65) cm on November 16.

Mosquito sampling for bacterial DNA extraction

Five early instar *Culex* larvae per mesocosm were collected on October 1 (4 days after starting the experiment), placed in 95 % ethanol in 15 mL sterile centrifuge tubes and stored at -20 °C until DNA extraction. The identification of these early instars was not possible using morphology but late (3rd and 4th) instar *Culex* larvae collected the following day was comprised of *C. stigmatosoma* (74 %), *C. quinquefasciatus* (23 %), and *C. tarsalis* (3 %). The proportion of *C. tarsalis* in the mesocosms increased over time while *C. stigmatosoma* tended to be more abundant early in experiment.

Five late (3rd and 4nd) instar larvae of *C. tarsalis* larvae per mesocosm were sampled on three dates: October 12, October 18, and November 15 (i.e., 14, 20 and 48 days, respectively from the onset of the experiments) to determine the influences of *Bti* treatments and habitat age on bacterial community structure associated with *Culex* larvae. These dates correspond to 10, 16 and 44 days after *Bti* treatments, respectively. Two groups of five late instar *C. tarsalis* larvae from a laboratory colony were also sampled to compare bacterial communities of laboratory-reared mosquitoes with field-collected mosquitoes. *C. tarsalis* colonies have been maintained in the laboratory for > 5 years and the larvae were fed a mixture of yeast and ground rat chow [37].

For pupae and adult mosquitoes, late instar larvae were collected from the mesocosms, morphologically identified to *C. tarsalis* under a dissecting microscope, and then reared to pupae or to adulthood. Water from the corresponding mesocosms was used to rear the larvae. Immediately after pupation, two groups of five pupae from the high *Bti* treatment and the control were preserved in 95 % ethanol and placed in a -20 °C freezer until DNA extraction. The remaining pupae were then transferred to rearing cages and, immediately after emergence, five groups of

five adult *C. tarsalis* (three from high *Bti* and two from control treatments) were placed in 95 % ethanol in 15 mL sterile tubes and kept in -20 °C freezer until DNA extraction.

DNA extraction, PCR and Illumina library preparation

The procedures of DNA extraction and amplification of the V3 hypervariable region of 16S rRNA genes using polymerase chain reaction (PCR) were similar to those used for late larval instars in previous study [23]. In addition to late instar larvae, early (1st and 2nd) instar larvae, pupae and adults of C. tarsalis were also sampled in the present study. All mosquito samples within the 15-mL tubes were sonicated for 3 min in iced sterile water according to a previous study [23]. DNA was extracted from only three (out of five) pooled, intact mosquitoes in 1.5 mL microcentrifuge tubes per replicate mesocosm using a Qiagen kit as described in [23]. PCR and Illumina library preparation were also carried out according to a previous study [23]. After libraries were prepared and quantified using an Agilent Bioanalyzer, all samples were normalized to 10 nM using Tris-HCl (10 mM, pH 8.5) and combined to create two multiplexed samples. The multiplexed samples were then subjected to a 2 × 150 base paired-end sequencing on a MiSeq Illumina platform at GENOSEQ (Sequencing and Genotyping Core) of the University of California Los Angeles, Los Angeles. Overall, a total of 41 mosquito samples [two late-instar samples from laboratory colonies, seven samples of field-collected early instars, 25 samples of late instars (taken on three sampling dates), two samples of pupae and five samples of adult mosquitoes] were submitted for sequencing.

Thorsellia cloning and phylogeny

PCR in 25 µL reactions was performed with Illustra PuRe Taq Ready-To-Go PCR Beads (GE Healthcare, Uppsala, Sweden), 0.4 µM each of forward and reverse *Thorsellia* primers 207f (5'-GCACTAGGATGAACCCAGG-3') and

reverse primer 1277r (5'-CTTTATGAGTTCCGCTT ACCC-3'), and 2 μ L of DNA from larvae of *C. tarsalis*. These primers were designed for investigating *Thorsellia* in *Anopheles gambiae* sensu latu in a previous study [38]. The PCR program was 98 °C for 5 min followed by 30 cycles of [95 °C for 30 s, 55 °C for 30 s and 72 °C for 1 min] and a final step of 72 °C for 10 min. Amplification products (1.1 kb) were cloned into TOPO 2.1 (Invitrogen) and sequenced at Macrogen (South Korea).

Sequences were aligned with available GenBank accessions for three described Thorsellia species (T. anophelis [AY837748], T. kanduguensis [KM269289], and T. kenyensis [KM269290]) and a single outgroup taxon (Arsenophonous nasoniae [AY264674]) using MEGA version 6 [39]. Positions containing gaps or missing data were eliminated, resulting in a final dataset of 1070 positions. MEGA was again used to infer phylogentic relationships among the sequences using the Maximum Likelihood method based on the Tamura-Nei model [40]. The initial tree for the heuristic search was obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites [5 categories (+G, parameter = 0.05)].

Sequence analysis, alignment, taxonomy assignment and statistical analysis

Analysis of the sequence reads was carried out using QIIME [41] version 1.7.0 and AXIOME version 1.6.0 [42] pipelines. Clustering of sequences to operational taxonomic units (OTUs) was carried out using cd-hit-est (multi-threaded version) with 97 % sequence identity [43]. Taxonomy assignment was conducted using the RDP classifier v2.2 with a confidence level of 0.6, and trained against the SILVA v111 16S/18S database [44]. All sequences that classified to Eukaryota were discarded. The statistical analyses of the sequences were carried out using procedures described in [33]. Briefly, beta diversity analysis using principal coordinate analysis (PCoA) based on Bray-Curtis dissimilarity distance matrix was carried out to assess the significance of differences among samples from different mosquito developmental stages, between Bti treatments and untreated controls, and among sampling dates. Beta diversity analyses were performed on OTU tables that were randomly subsampled (without replacement) down to the sample with the lowest number of sequences. Analyses including all stages were subsampled down to 6,539 sequences per sample, and analyses on late instar samples were subsampled down to 72,987 sequences per sample. The significance of the separation of sample groups in Bray-Curtis and UniFrac ordinations was assessed by MRPP [45] via QIIME. MRPP returns a within-group homogeneity value of A and a p value, which represents the probability of the observed differences between the groups occurring by chance. A values closer to 1 indicate increased sample similarity within-group, and an A value of 0 indicates the within-group similarity expected by chance. Alpha diversity measures based on phylogenetic distances were compared among samples within the mosquito developmental stages.

Availability of supporting data

The data set supporting the results of this article is available in the ENA repository under project accession number PRJEB6788. Two representative clone sequences were deposited in GenBank under accession numbers KP215389 and KP215390. The data set supporting the results of this article are included within the article and its additional files.

Additional files

Additional file 1: Table S1. *Bacteria* phyla and percent (by number of sequences) abundance in *Culex* life cycle stages.

Additional file 2: Table S2. Shared bacterial OTUs across developmental *stages of C. tarsalis* collected from the field. Bacterial OTUs represented by at least 1 sequence per sample were included.

Additional file 3: Table S3. The most abundant bacterial OTUs (≥1 %) found in field-collected early-instar *Culex* larvae.

Additional file 4: Table S4. The most abundant bacterial OTUs (≥1 %) found in field-collected late-instar *C. tarsalis* larvae.

Additional file 5: Table S5.The most abundant bacterial OTUs (≥1 %) found in laboratory-reared late-instar *C. tarsalis* larvae.

Additional file 6: Table S6. The most abundant bacterial OTUs (≥0.5 %) found in *C. tarsalis* pupae (reared from late-instar larvae collected from the field).

Additional file 7: Table S7.The most abundant bacterial OTUs (≥1 %) found in *C. tarsalis* adults (reared from late-instar larvae collected from the field).

Competing interest

The authors have declared no conflict of interests.

Authors' contributions

DD and WEW conceived the experiment. DD and WEW designed and conducted the field experiment. DD and PRJ conducted the molecular laboratory procedures. DD, MWH and JDN analyzed the data. RS contributed reagents. OT contributed the phylogenetic data and analysis. All authors contributed to writing the manuscript and have approved its submission.

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