Beetle (Coleoptera: Scirtidae) Facilitation of Larval Mosquito Growth in Tree Hole Habitats is Linked to Multitrophic Microbial Interactions

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Abstract Container-breeding mosquitoes, such as Aedes triseriatus, ingest biofilms and filter water column microorganisms directly to obtain the bulk of their nutrition. Scirtid beetles often co-occur with A. triseriatus and may facilitate the production of mosquito adults under low-resource conditions. Using molecular genetic techniques and quantitative assays, we observed changes in the dynamics and composition of bacterial and fungal communities present on leaf detritus and in the water column when scirtid beetles co-occur with A. triseriatus. Data from terminal restriction fragment polymorphism analysis indicated scirtid presence alters the structure of fungal communities in the water column but not leaf-associated fungal communities. Similar changes in leaf and water bacterial communities occurred in response to mosquito presence. In addition, we observed increased processing of leaf detritus, higher leaf-associated enzyme activity, higher bacterial productivity, and higher leaf-associated fungal biomass when scirtid beetles were present. Such shifts suggest beetle feeding facilitates mosquito production indirectly through the microbial community rather than directly through an increase in available fine particulate organic matter.

Introduction

Tree holes are small discrete ecosystems that contain heterotrophic communities driven by allochthonous inputs of soluble and particulate organic matter. In Eastern North American tree holes, larvae of the Eastern treehole mosquito, Aedes triseriatus (Say), are usually the dominant macroinvertebrate consumers. Leaves are a typical source of coarse particulate organic matter (CPOM) [48], but animal-derived detritus, such as invertebrate carcasses and fecal material, also supply energy to these systems [9, 50]. In addition, stemflow runoff brings dissolved organic carbon and nutrients into the habitats [5, 16, 19]. Success of A. triseriatus and similar mosquito species is characterized by high adult productivity and body weight at emergence, and is dependent on the nutrition obtained while in the larval stage [45]. The quantity and quality of leaf litter in tree holes are important determinants of successful mosquito production [34, 45], although in general larvae do not directly consume the material. Rather, microorganisms are the main nutritional resource for mosquito larvae, which feed by browsing on container surface microbial biofilms and filtering fine particulate organic matter (FPOM) and microorganisms from the water column [27]. Fungi associated with leaf detritus alone may account for around 10% of the detrital biomass in tree hole habitats [19, 20]. Hence, microbial degradation of detritus is a critical link between developing larvae and nutrients as processing by microorganisms incorporates nutrients and otherwise inaccessible carbon from organic detritus.
Despite previous studies addressing consumption of microorganisms by mosquito larvae in container habitats, microbial food resource dynamics remain ill-defined. Water column-associated bacteria exhibit negligible or inconsistent responses to mosquito presence and, thus far, there have not been any published studies of fungal communities associated with tree hole water columns. Furthermore, larval feeding effects on the composition of water column microbial communities are limited to bacteria and protists [8, 17, 19, 21, 44]. In contrast, there is convincing support for the importance of microbial communities associated with detritus in previous assessments of tree holes. Increased fungal enzyme activity, and decreased bacterial productivity and abundance are associated with mosquito feeding [16, 17, 19, 34]. Despite the response of microorganisms to larvae, the composition of microbial communities in tree holes has been addressed only recently [18, 36, 37, 49]. In those studies, larval feeding effects were evident on the taxa comprising fungal and bacterial communities associated with leaf detritus, with particular influence on Saccharomycetes, Dothideomycetes, and Chytridiomycota fungal taxa and on Alpha- and Betaproteobacteria.

In the Midwestern and Eastern USA, larval scirtid beetles (Helodes and Prionocyphon spp.) are shredders which often co-occur with A. triseriatus in tree holes [2, 32]. Scirtid feeding activity results in the skeletonization of leaf detritus and the associated conversion of CPOM to FPOM, including small leaf particles and feces. Several studies indicate that the presence of these beetles in tree holes conditionally facilitates the survival and development of A. triseriatus when leaf litter levels are low by improving the quality of resources available to A. triseriatus [3, 9–11, 31, 32]. Similar processing chain commensalsisms benefitting mosquitoes have been reported in pitcher plants. By increasing the conversion of coarse particulate plant material to fine particulates, the midge Metriocnemus knabi facilitates populations of the pitcher plant mosquito Wyeomyia smithii [15]. In addition to mosquito populations, processing chain benefits have been reported to facilitate populations of the ceratopogonid midge Culicoides gutti-pennis in the presence of Helodid beetles [33]. Increased conversion of leaf material into FPOM is the main mechanism cited for any benefit experienced by mosquitoes in the presence of scirtids [11, 12, 30, 31]. Indeed, Daugherty and Juliano [12] demonstrated that additions of scirtid fecal material and its associated microbiota to microcosms increased larval A. triseriatus development by providing additional nutrient resources; however, the source of this positive effect (feces, microbiota, or a combination) was not determined. Although the conversion of CPOM into FPOM may benefit foraging larvae by being ingested directly, it may also promote access to fungal material otherwise embedded in the leaf matrix. Furthermore, it is possible that any influence of scirtid feeding on the microbial community may affect mosquitoes given the direct utilization of microorganisms by larvae for nutrient acquisition. By altering the detritus such that the abundance of FPOM is greater, scirtid beetles modify the bottom-up influence of detritus on bacterial and fungal populations. Such blending of “bottom-up” influences and “top-down” trophic cascades are described for aquatic ecosystems, including tree hole container habitats, yet the ramifications of scirtid beetles on microorganisms is poorly understood [6, 7, 21, 22].

Although previous studies have postulated mechanisms for facilitation of mosquito larvae by scirtids, they are limited to measurements of macroscopic changes such as the abundance of FPOM, population densities of mosquitoes, and decomposition of scirtid carcasses [11, 12, 30, 32]. While all of these factors play a role in mosquito growth, these indirect assessments lack the intermediate microbial step linking the trophic levels. Elucidating the effect of scirtid-feeding activity on mosquito productivity requires an assessment of the changes such feeding renders on the microbial community.

In an effort to understand the ecological influences driving mosquito production, we sought to describe microbial mechanisms underlying facilitation of larval A. triseriatus growth and development by scirtid beetles via assessments of the microbial community associated with scirtid presence in microcosms simulating natural tree holes. To address the effects of macroinvertebrates on the structure of tree hole microbial communities, we used the culture-independent molecular technique, terminal restriction length polymorphism (T-RFLP) analysis. We postulated that detritus-associated fungal dynamics, rather than water column-associated, would be the most dramatically affected by scirtid presence as this location is the “center of activity” for foraging scirtid larvae [13]. Furthermore, we predicted that mosquito presence should have a dissimilar effect on microbial communities than that of scirtids due to the different feeding modes exhibited by these macroinvertebrates.

**Materials and Methods**

**Experimental Design**

The microbial dynamics underlying the facilitation of A. triseriatus development by scirtid beetle larvae were investigated by crossing beetle presence/absence with mosquito presence/absence in a multifactorial design with the following scirtid/A. triseriatus ratios: 0:0, 10:0, 10:40, and 0:40. The ratios were selected to reflect densities of the macroinvertebrates typically observed in natural tree holes.
Twelve replicates of each treatment combination were constructed, with one set of six replicates destructively sampled on day 20, approximately midway through the experiment. Because resource levels mediate the interaction between scirtids and mosquitoes such that facilitation is only apparent when resources are less abundant, low leaf litter rations were used for all replicate microcosms. Individual microcosms simulating natural tree holes received 1-g senescent oak leaf pack (Northern Red Oak, Quercus rubra) in 500 ml deionized water. Microcosms (20.3 cm height×7.6 cm inner diameter) were constructed similar to those described in previous tree hole studies [17, 20, 46]. A 3-ml microbial inoculum, consisting of homogenized tree hole water and particulates, was added to microcosms 3 days prior to the addition of macroinvertebrates. To insure that macroinvertebrates were not present, the microbial inoculum was visually inspected prior to addition.

A. triseriatus, obtained from our colony at Michigan State University, were added as newly hatched first instar larvae. Scritid beetle larvae, ranging from second to fourth instar, were obtained from local tree holes (Toumey Woodlot, E. Lansing, MI, USA) and replaced semi-weekly as necessary to maintain a constant population in scritid treatments. Replacement beetle larvae were taken from the original stock of scritids collected for this experiment and rinsed with sterile water before being added to microcosms. Dead beetles were removed to avoid confounding the effect of scritids by providing additional resources to mosquito or microbial populations. Due to the difficulty of identifying live beetle larvae, scritids were not identified to species [10, 12, 32].

Sampling

Leaf condition was assessed by taking dry weight measurements of leaf disks on days 20 and 40. Leaf disk samples (20 mm diameter) were procured from each microcosm on days 20 and 40 using a cork borer. Leaves were aseptically removed from microcosms and gently placed onto a sterile Petri dish prior to cutting to minimize disruption of the biofilm. Two leaf samples apiece were also collected for assessing bacterial abundance, bacterial productivity, fungal biomass, fungal degradation enzymes, and microbial community structure. Prior to analyses, leaf disk samples were sonicated for 12 min in an ice bath to obtain the loosely attached, surface-associated microorganisms, as these better represent those encountered by foraging mosquito larvae [18]. In addition to leaf samples, water samples were also taken from microcosms on each sample date. After stirring microcosms to homogenize water and particulates, 15-ml water column samples were removed from microcosms for the analyses indicated above.

Microcosms were checked daily for the presence of adult mosquitoes. Adults were collected and stored at 4°C until the microcosm was destructively samples. Larvae were gathered by decanting the remaining microcosm water through a fine mesh sieve and stored at 4°C. Where possible, we sexed and identified adults, larvae, and pupae to species before lyophilizing the mosquitoes to obtain dry mass measurements.

Bacterial Abundance and Productivity

Bacterial abundance on the leaf surface and in the water column was quantified from subsamples via direct microscopic counts of bacteria using the 4,6-diamidino-2-phenylindole fluorescent staining procedure [34, 38, 46]. Formalin (3% formaldehyde final concentration)-preserved samples were kept under dark conditions at 4°C until analysis, then filtered onto black Nucleopore filters (0.2-mm pore size; Costar, Cambridge, MA, USA). After filtering, samples were stained at a final concentration of 20 μg/ml for 5 min. Two filters and at least 20 fields per filter were counted (a minimum of 200 cells per filter) for each subsample at 1,000× using a Nikon E800 fluorescent microscope (Nikon, Inc. Melville, NY, USA).

Fungal Biomass

Fungal biomass was estimated using ergosterol, a sterol associated with fungal cell walls, as a surrogate measurement [28, 43]. Leaf disk subsamples were placed in high performance liquid chromatography (HPLC) grade methanol and stored in the dark at 4°C prior to extraction and quantification of ergosterol using HPLC and UV detection [16, 19].

Enzyme Activity

Fungal activity was estimated by measuring the degradation of xylose and cellobiose as surrogate measures of fungal enzymes. Leaf enzyme activity was assayed through incubations of leaf
disk subsamples with two methylumbelliferyl (MUF)-labeled substrates, 4-methylumbelliferyl-β-D-celllobioside and 4-methylumbelliferyl-β-D-xyloside, which estimate cellobiohydase and xylosidase activities, respectively. These substrates are analogous to plant polymers and thus provide an estimate of leaf-associated carbohydrate activity [19]. MUF was liberated upon enzymatic cleavage of the substrates during 1.5-h incubation at 22°C. Unbound MUF, which fluoresces at 360 nm, was measured using a 96-well Hoefer DyNA Quest 200 fluorometer (GE Healthcare, Little Chalfont, Buckinghamshire, UK).

Microbial Community Analysis

The compositions of leaf- and water column-associated microbial communities in microcosms were assessed using T-RFLP analysis, a culture-independent technique that estimates the diversity of microbial “species” present in samples [24, 26]. Six replicate microcosms per treatment were sampled for bacterial and fungal community analysis. Three of the microcosms were destructively sampled on day 20 and three on day 40. Leaf disk samples were sonicated as described above in sterile phosphate buffered saline to dislodge surface-associated microorganisms. Thereafter, DNA was immediately isolated from leaf sonicate and water samples using the MoBio UltraClean soil extraction kit (Carlsbad, CA, USA). Independent amplification of DNA for microbial communities in microcosms was assessed using universal eubacterial and fungal primers nu-SSU-0817-59F and nu-SSU-1536-39R (5′-CAGGCCTAACACATG CAAGTC-3′ and 5′-GGGCGGWGTGATCACAAGGC-3′) [25]. A 762-bp 18S rDNA fragment was amplified using universal fungal primers 18S rDNA forward primer 5′-SSU-0817-59F and 18S rDNA reverse primer 5′-ATGCAATGCYCTATCCCCA-3′ (6-FAM) at the 5′ end (IDT Technologies). Template DNA from bacteria (16S rDNA) and fungi (18S rDNA). A 1,300-bp 16S rDNA fragment was amplified using universal eubacterial primers 63F and 1387R (5′-CAGGCCTAACACATG CAAGTC-3′ and 5′-GGGCGGWGTGATCACAAGGC-3′) [25]. A 762-bp 18S rDNA fragment was amplified using universal fungal primers 18S rDNA forward primer 5′-SSU-0817-59F and 18S rDNA reverse primer 5′-ATGCAATGCYCTATCCCCA-3′ (6-FAM) at the 5′ end (IDT Technologies). Template DNA from bacteria (16S rDNA) and fungi (18S rDNA).

One-way analyses of variance (ANOVA) were used to assess mosquito response variables (survival, emergence time, and total biomass). Multivariate analysis of variance (MANOVA) was used to analyze related bacterial measurements including bacterial abundance and bacterial productivity in the water column and on leaf surfaces (Proc GLM, SAS v. 9.1, SAS Institute, Inc., Cary NC, USA). The activity of fungal degradation enzymes and fungal ergosterol were similarly analyzed in a separate MANOVA. Standardized correlation coefficients (SCCs) were calculated for each MANOVA to determine the magnitude of the contributions made to each variable to the respective measurements included in the analysis [41]. In addition, univariate analysis of variance (“protected” ANOVA) was conducted for dependent variables with significant MANOVA results followed by Bonferroni correction to control the experiment-wide error [39]. Treatment means were separated using Tukey’s “honestly significantly different” (HSD) test. Leaf mass lost was analyzed using a standard ANOVA. Means were square root transformed as needed to meet the assumptions of ANOVA.

T-RFLPs were aligned and peak areas determined using the GeneScan software package (Applied Biosystems). A sample was discarded if its combined peak area was less than 1,000 or if a normality plot indicated the presence of significant outliers. In addition, peaks that comprised <3% of the community or appeared in fewer than three samples were also discarded. We distinguished “real” peaks by separating signals from baseline noise in electropherograms. T-RFs differing by ± 1 bp were considered identical. Common peaks were aligned and binned using
the “T-RFLPs Stats” analysis tools (iBest, http://www.ibest.uidaho.edu/tools/trflpstats/index.php) [1]. The resulting matrices were converted to log ratios of the relative abundance of peak area in each sample, calculated as the peak area of a T-RF divided by total peak area of all fragments in a sample within a 50- to 800-bp range. Profiles from each of the four communities (water column bacteria and fungi and leaf-associated bacteria and fungi) were individually analyzed by principle component analysis (PCA) on the respective covariance matrices to reduce dimensionality. PCA ordinations were conducted using JMP® Statistical Discovery Software (SAS Institute, Inc., Cary NC, USA). Scores from PCs 1–3 were subjected to standard ANOVA for treatment comparisons.

Results

Leaf decomposition decreased significantly in the presence of scirtids ($F=11.96; df=1, 23; P=0.003$) such that microcosms containing scirtids had a lower proportion of leaf mass remaining than non-scirtid microcosm (Fig. 1; Tukey’s HSD, $\alpha=0.05$). Neither mosquito presence solely or with scirtids significantly affected leaf decomposition ($F=0.69; df=1, 23; P=0.417$; and $F=0.0; df=1, 23; P=0.962$, respectively). Mosquito survival and total biomass were highest in the presence of scirtid beetles, with correspondingly faster female development time (Table 1), although these differences were not statistically significant.

Scirtids, mosquitoes, and time had significant effects on measurements of bacterial community dynamics (MANOVA, Table 2). Macoinvertebrate treatment, time, and the interactions among the main effects were primarily explained by changes in the bacterial abundance on leaves (Table 2, SCC). In the water column, bacterial abundance was not significantly affected by the presence of either macoinvertebrate, although a significant decrease in abundance did occur from days 20 to 40 (Fig. 2a). In contrast, leaf surface bacterial abundance was depressed in the presence of macoinvertebrates (Table 3 and Fig. 2b). Under reduced-resource conditions, mosquito larvae significantly lowered bacterial abundance compared with no larvae conditions; however, scirtid presence significantly interacted with mosquito presence on day 20 such that microcosms containing both insects had a greater abundance of leaf-associated bacteria than those containing mosquito larvae alone (Table 3). Although scirtid beetles alone did not significantly decrease bacterial abundance at day 20; by day 40, microcosms containing only scirtids had a lower abundance of bacteria than microcosms with no macoinvertebrates present.

Significant reductions in water column bacterial productivity were evident in the presence of mosquito larvae and productivity remained unchanged whether scirtids were present or absent (Table 3, Fig. 2c). In addition, water column productivity also declined significantly from days 20 to 40 for microcosm treatments. Leaf-associated bacterial productivity was significantly lower in the presence of mosquito and scirtid beetle larvae compared with the no-macoinvertebrate treatment (Table 3, Fig. 2d). Productivity differences were not apparent among microcosms receiving macoinvertebrates, likely in response to the significant interaction between mosquito larvae and scirtid beetles.

The presence of scirtid beetles in microcosms had important effects on the activity and biomass of fungi in microcosms (MANOVA, Table 4). These effects were mainly explained by changes in the concentration of xylanase and cellulohydrolase (SCC, Table 4). Activity of fungal decomposition enzymes for these substrates was significantly greater ($F=9.28; df=1, 36; P=0.004$ and $F=8.57; df=1, 36; P=0.006$, respectively) in leaf samples from microcosms containing scirtids compared with microcosms without scirtid beetles (Fig. 3). In contrast, although fungal biomass (ergosterol concentration) in microcosm leaf samples was numerically greater in microcosms containing scirtids, this effect was not statistically significant. Enzyme activity and biomass, determined by ergosterol measurement, were not significantly influenced by the presence of mosquito larvae in microcosms; however, biomass was significantly influenced by the time main effect (MANOVA, Table 4). Fungal biomass was not significantly influenced by the presence of either A. triseriatus or scirtid larvae; however, the concentration of ergosterol was signif-
Significantly greater for all treatments on day 40 compared to day 20 \((F=12.84, df=1, 36; P=0.001; \text{Fig. 4})\).

T-RFLPs

MspI

In this study, 16S and 18S rRNA genes amplified from microcosms and digested with MspI were subjected to T-RFLP analysis to evaluate the role of macroinvertebrates in shaping tree hole microbial communities. T-RFLP analysis yielded 117 and 118 T-RFs associated with leaf and water column bacterial communities, respectively. PCA on transformed T-RF profiles revealed that the first three principle components (PCs) explained 41.3 and 42.7% of the total variability in the MspI dataset corresponding to bacterial communities in the water column and on leaf surfaces, respectively. PC1 scores for bacteria in the water column were significantly affected by time, exhibiting shifts in the community structure from day 20 to day 40 \((F=14.0, df=1, 23, P=0.002; \text{Fig. 5})\). Scirtid and mosquito presence also had significant effects on bacterial communities along PC1 \((F=16.9, df=1, 23, P=0.001; F=16.9, df=1, 23, P=0.019)\). Both time and mosquito presence significantly affected water column bacteria along PC2 \((F=9.3, df=1, 23, P<0.001; F=31.5, df=1, 23, P<0.001)\). There were no significant effects of treatment on PC3 scores. Both mosquito presence and time significantly affected PC1 scores for leaf surface bacterial communities \((F=45.5, df=1, 22, P<0.001; F=16.9, df=1, 22, P=0.001; F=16.9, df=1, 22, P=0.019)\). Neither insect presence nor time significantly affected PCs 2 and 3.

T-RFLP analysis yielded 99 T-RFs associated with leaf and 88 T-RFs associated with water column fungal communities. The first three PCs obtained from analysis of T-RF profiles explained 50 and 54.3% of the total variability in the MspI dataset corresponding to fungal communities in the water column and on leaf surfaces, respectively. Water column fungal communities exhibited significant shifts in response to mosquito presence and time along PC2 \((F=43.6, df=1, 22, P<0.001; F=12.4, df=1, 22, P=0.003; \text{Fig. 6})\). There were no significant effects of any factor on PC1. Mosquito presence, sample date, and the interactions of sample date with mosquito presence and with scirtid presence significantly affected water-associated fungi along PC3 \((F=3.3, df=1, 22, P=0.011; F=23.6, df=1, 22, P=0.048)\). Neither insect presence nor time significantly affected PCs 2 and 3.

### Table 2

| Model parameters | MANOVA | SCC |
|------------------|--------|-----|
|                  | Pillai's trace | \text{Bacterial abundance} | \text{Bacterial productivity} |
|                  | \text{Water} column | Leaf surface | \text{Water} column | Leaf surface |
| Scritid          | 0.34   | 3.22 | 4, 25 | 0.029*<sup>7</sup> | 0.21 | 1.778 | −0.252 | 1.248 |
| Mosquito         | 0.597  | 9.25 | 4, 25 | 0.000*<sup>7</sup> | −0.267 | 0.941 | −0.07 | 0.346 |
| Time             | 0.893  | 52.29 | 4, 25 | 0.000*<sup>7</sup> | −0.481 | 3.176 | 0.358 | −0.324 |
| Scritid \times mosquito | 0.446  | 5.04 | 4, 25 | 0.004*<sup>7</sup> | −0.157 | 1.649 | −0.134 | 1.293 |
| Scritid \times time | 0.596  | 9.25 | 4, 25 | 0.170 | −0.109 | 2.889 | 0.639 | −1.047 |
| Mosquito \times time | 0.476  | 5.69 | 4, 25 | 0.002*<sup>7</sup> | −0.339 | 3.145 | 0.31 | −0.488 |
| Scritid \times mosquito \times time | 0.327  | 3.04 | 4, 25 | 0.036*<sup>7</sup> | 0.832 | −2.561 | −0.662 | 0.898 |

Only the first canonical was significant

*\(P<0.05\), significance for MANOVA
Similarly, a shift in leaf-associated fungal communities in response to mosquito presence was significant along PC2 ($F=6.19$, $df=1, 23$, $P=0.024$). In general, however, no shifts were evident in leaf fungal communities in response to time or scirtid presence.

HhaI

In comparison to MspI, T-RFLP digestions using HhaI resulted in fewer T-RFs from water column- and leaf-associated bacterial communities (71 and 55, respectively). PC2 scores representing water column bacterial communities were significantly affected by mosquitoes ($F=6.2$, $df=1, 22$, $P=0.01$). Leaf surface bacterial communities represented by principal component axes were significantly affected by scirtids along PC1 ($F=5.1$, $df=1, 23$, $P=0.039$). The percentage of total variation explained by the first three PCs in the HhaI datasets corresponding to water column and leaf surface bacterial communities were 44.7% and 47%, respectively.

Similar to bacterial communities, fungal communities associated with water and leaf samples produced fewer T-RFs (46 and 41, respectively) when digested with HhaI compared with MspI. Water column fungal communities changed in response to the interaction of mosquito presence and time along PC2 ($F=11.5$, $df=1, 22$, $P=0.004$). PCs 1–3 explained 54% of the total variation associated with T-RFLP profiles in this community. Scirtids did not significantly affect changes in leaf-associated fungal communities, although these communities changed in response to the presence of mosquitoes (PC2: $F=45.5$, $df=1, 23$, $P<0.001$) and time (PC1: $F=24.3$, $df=1, 23$, $P<0.001$; PC2: $F=8.7$, $df=1, 23$, $P=0.010$). PCA analysis revealed that 56.2% of the variation in leaf fungal communities was explained by the first three principle components.

**Discussion**

This study evaluated the impact of scirtid beetle presence on microbial community dynamics in container habitats. We demonstrated that scirtid beetles increased the decay rate of leaf detritus, accelerated microbial activity on leaf
surfaces, and altered the community structure of microorganisms in water and on leaves. Shifts in T-RF profiles representing bacterial and fungal communities on these substrates were evident in the presence of mosquitoes and scirtids. Interactive effects between mosquitoes and scirtids were most pronounced in the leaf bacterial community, suggesting this component is important in facilitation, however, increased leaf carbohydrolytic activity and decay rates also implicate fungal activity in the interaction. Taken together, these results indicate accelerated leaf decay and microbial activity associated with beetle feeding activity may underlie scirtid-mediated facilitation of mosquito growth. The facilitative effect of scirtids is such that larger females are produced under low-resource conditions in the presence of scirtids [30]. In our study, we subjected these macroinvertebrates to similar low resource conditions and found that mosquito survival and total biomass were highest in the presence of scirtid beetles. Although not statistically significant, the trends evident in our results reflect the findings of previous studies [30] describing the facilitation of mosquitoes by scirtid beetles.

Enhancement of microbially associated decomposition enzymes, which convert particulate matter into soluble compounds, occurred in the presence of scirtids. Although scirtid-induced reductions in leaf mass have been documented previously [31], the current study is the first to indicate a microbial mechanism for leaf processing other than physical conversion of CPOM to FPOM. While the latter contributes to the pool of resources utilized by A. triseriatus, transitions between filtering (consuming FPOM and microorganisms) and browsing (consuming microbial biofilms) feeding modes are known to occur [27]. If browsing behavior is indeed more commonly exhibited by late-instar A. triseriatus as previously indicated, then we would expect that late-instar mosquito larvae would benefit from an increase in leaf-associated microbial biomass and fungal activity conferred by scirtid beetles.

### Table 3
Summary of ANOVA results for bacterial abundance and bacterial productivity values from microcosm water column and leaf material

| Model parameters | $F_{df}$ | $P$ value |
|------------------|---------|----------|
| Water column abundance (cells/ml) | | |
| Scirtid | 1.22, 28 | 0.278 |
| Mosquito | 1.37, 28 | 0.252 |
| Time | 15.26, 28 | <0.001* |
| Scirtid×time | 0.30, 28 | 0.857 |
| Mosquito×time | 0.71, 28 | 0.407 |
| Scirtid×mosquito | 0.78, 28 | 0.385 |
| Leaf surface abundance (cells/disk) | | |
| Scirtid | 8.90, 28 | 0.006* |
| Mosquito | 45.59, 28 | <0.001* |
| Time | 210.76, 28 | <0.001* |
| Scirtid×time | 4.74, 28 | 0.038 |
| Mosquito×time | 25.65, 28 | <0.001* |
| Scirtid×mosquito | 11.23, 28 | 0.002* |
| Water column productivity (nmol leucine/ml/h) | | |
| Scirtid | 0.73, 28 | 0.401 |
| Mosquito | 5.40, 28 | 0.028* |
| Time | 13.82, 28 | <0.001* |
| Scirtid×time | 0.47, 28 | 0.500 |
| Mosquito×time | 2.27, 28 | 0.143 |
| Scirtid×mosquito | 0.91, 28 | 0.348 |
| Leaf surface productivity (nmol leucine/disk/h) | | |
| Scirtid | 7.64, 28 | 0.010* |
| Mosquito | 11.73, 28 | <0.001* |
| Time | 2.95, 28 | 0.097 |
| Scirtid×time | 3.02, 28 | 0.093 |
| Mosquito×time | 0.99, 28 | 0.327 |
| Scirtid×mosquito | 13.02, 28 | <0.001* |

*p Values that are significant following Bonferroni adjustment

### Table 4
Multivariate analysis of variance (MANOVA) for the effect scirtid beetles, mosquito larvae, and sampling time on fungal biomass and leaf associated enzyme measurements, and the standardized canonical coefficients (SCC) of the first canonical variate

| Model parameters | Pillai’s trace | $F_{df}$ | $P$ | Ergosterol | Xylosidase | Cellulobioshydrase |
|------------------|---------------|---------|-----|-----------|-----------|-----------------|
| Scirtid | 0.227 | 3.320, 3, 34 | 0.031* | 0.241 | 0.559 | 0.555 |
| Mosquito | 0.057 | 0.680, 3, 34 | 0.571 | 0.911 | 0.592 | −0.858 |
| Time | 0.277 | 4.340, 3, 34 | 0.011* | 1.09 | 0.092 | 0.179 |
| Scirtid×mosquito | 0.104 | 1.320, 3, 34 | 0.285 | 0.567 | −0.718 | 1.322 |
| Scirtid×time | 0.104 | 1.320, 3, 34 | 0.284 | 0.547 | 0.552 | −0.021 |
| Mosquito×time | 0.054 | 0.650, 3, 34 | 0.59 | −0.654 | 0.655 | 0.475 |
| Scirtid×mosquito×time | 0.008 | 0.100, 3, 34 | 0.961 | 0.523 | 0.634 | 0.523 |

*P<0.05, significance for MANOVA
through direct ingestion of microbial cells exposed or created by beetle feeding, and perhaps through direct utilization of fungal enzymes- and leaf-degradation products. Additional research is needed to determine how leaf processing is partitioned between FPOM production through invertebrate feeding and microbial metabolic activity and the relative contribution each make to the production of *A. triseriatus*.

The presence of either macroinvertebrate depressed leaf-associated bacteria and their growth rates, however, the effect was most pronounced with mosquitoes alone. Productivity was higher when mosquitoes co-occurred with scirtids, suggesting that scirtid feeding may have interfered with mosquito browsing on leaf surfaces. Thus, it appears that reductions in bacterial biomass and productivity via scirtid grazing on leaf surfaces may play a role in the facilitation of mosquito growth by promoting fungal production. That mosquitoes drive down bacterial productivity on leaves is well-described by Kaufman et al. [16, 17, 19, 20]; however, this is the first study to indicate that another tree hole invertebrate may also affect bacterial productivity. Although a similar reduction in bacterial productivity in the presence of macroinvertebrates also occurred in the water column, there was no significant mosquito-scirtid interaction and thus a mosquito-grazing effect was not modified by the presence of scirtids as might be expected with increasing FPOM release. A corresponding change in bacterial abundance did not occur in response to the presence of either mosquitoes or scirtids and this is unsurprising in light of previous findings from tree holes. Kaufman et al. [17] suggested that such decoupling of bacterial abundance and growth rate may result from shifts in the structure of bacterial communities in response to macroinvertebrate presence. In this study, scirtids did affect water column bacterial community structure, as evidenced by T-RFLP results obtained from MspI digestions (Fig. 5), and is supportive of the idea that beetle presence alters bacterial groups. This effect is likely indirect, as scirtids do not graze in the water column and there is no indication that bacterial abundance or productivity in the water column increased when scirtids were present.
quito larvae generally had stronger influences on water column bacterial structure than scirtids, evidenced by T-RFLP results from both MspI and HhaI digestions, presumably because they directly feed upon that microbial resource. Bacterial community shifts in response to mosquito larvae have been described for pitcher plant and tree hole communities [8, 16, 23]; however, this is the first study to evaluate this response in the presence a facilitating organism.

On leaf surfaces, scirtids affected abundance and productivity of bacteria, while also altering the structure of bacterial communities. While mosquitoes altered MspI T-RF profiles of leaf bacterial communities, sciritid beetle presence was the only factor in this study associated with changes in T-RF profiles of leaf bacterial communities obtained from both MspI and HhaI digestions. Ostensibly, the more robust nature of sciritid feeding results in more dramatic changes in this microbial community than the “delicate” browsing by mosquito larvae. Similar to the changes in water column communities in response to mosquitoes, these changes likely reflect direct feeding by the beetles. In addition to the direct effect of sciritid feeding, shifts in T-RF profiles may be a response to differential enrichment of microbial communities due to increased leaf processing and macroinvertebrate waste, additions of macroinvertebrate-associated inocula, or the release of endophytic microorganisms or antimicrobial compounds during leaf processing. Although each of these indirect processes has the potential to alter microbial interactions, the effect of direct predation on microbial community structure has been demonstrated frequently in other systems, including mosquito habitats. In pitcher plants, another type of water-filled habitat, Peterson et al. [35] found that presence of a keystone predator, Wyomyia smithii, increased the diversity of bacterial assemblage present. It is apparent that mosquito larvae and sciritid beetles ingest microorganisms; thus, we expect that the feeding effects of these macroinvertebrates should be similar to the effects observed in other systems.

In contrast to bacterial communities, leaf surface fungal communities derived from MspI (Fig. 6) and HhaI T-RFs changed in response to mosquito larvae, but not the presence of sciritid beetle larvae. This is congruent with the findings of Kaufman et al. [16], who showed shifts in fungal community structure related to the presence or absence of A. triseriatus larvae in tree holes. Although leaf fungal communities were more active in their presence, sciritid beetles did not change the nature of these communities. Leaf fungal biomass in the presence of sciritids was statistically equivalent on a surface area basis (per leaf disk) compared with microcosms without sciritids, but this did not take into account the reduced leaf mass when beetles were present. Fungal biomass was presumably more concentrated on a per mass basis (same fungal mass, lower weight of each leaf disk) in the sciritid-present treatment and enzyme activity levels would have been similarly higher. We suggest that
scirtid influenced concentration of existing fungal community biomass/activity may facilitate A. triseriatus growth, but examination of fungal growth rates and more detailed analysis of fungal community structure (see below) is warranted in future studies.

Although leaf-associated fungal and water column bacterial communities in scirtid-only microcosms followed a similar trajectory to those with no macroinvertebrates, the influence of scirtid feeding on leaf bacterial dynamics is reflected in the differential shift of PC scores from microcosms with scirtids. That bacterial community changes evident on the leaf surface were influenced by scirtid as well as mosquito presence is unsurprising because scirtid feeding would necessarily result in the consumption of surface-associated bacterial biofilms along with leaf particulates. However, it is evident from the different principle component trajectories (Figs. 5 and 6) that these species have dissimilar effects on the microbiota. The effect of scirtids on bacteria in the water column suggests their movements (feeding and locomotion) along leaves may dislodge surface bacteria and FPOM into the water, thus altering the bacterial community in both locations. In addition, skeletonization of leaves should increase the surface area available for bacterial biofilm formation, further contribution to changes on the bacterial communities and the observed increase in bacterial productivity in microcosms containing scirtids and mosquitoes compared to those with only mosquitoes. Overall, mosquito presence had the most influence on changes in microbial community composition. Leaf surface- and water column-associated bacterial and fungal communities all changed significantly in response to mosquito presence in correspondence with our previous observations [16, 18, 47]. Although we did not assess water column fungal biomass in the present study, we suspect that mosquitoes change fungal compo-

Figure 5 PCA ordination of T-RFLP peak areas representing bacterial taxa from water (a and b) and leaf (c and d) samples taken from microcosms on day 20 (a, c) and day 40 (b, d). Treatments are represented by the following symbols: circles 0 macroinvertebrates; squares 10 scirtids, 0 mosquitoes; open triangles 0 scirtids, 40 mosquitoes; filled triangles: 10 scirtids, 40 mosquitoes. The percentage of variance explained by PC scores for each substrate (water or leaf) is indicated on the axes for each panel.
sition in the water column by reducing the taxa susceptible to larval digestion [14, 40]. Surprisingly, though scirtids may open additional niches for fungal colonization via shredding leaf material, scirtids did not appear to directly affect the structure of fungal communities through their feeding activity. The absence of scirtid-associated changes in the structure of leaf fungal communities may indicate these macroinvertebrates are not affecting the structure at all, or at least not at the level of resolution provided by T-RFLPs. Alternatively, changes may not be visible as a result of the technique used to harvest surface associated microorganisms from the leaf surfaces. Loosely attached fungi are represented in the sonicated leaf samples while fungi whose hyphae are emmeshed in the matrix of the decomposing leaf are not. Rather than accessing only loosely attached fungi, beetles consume whole leaf particles with the associated fungi; therefore, changes in the composition of fungal communities may be relatively uniform in contrast to the compositional changes in response to mosquitoes observed in this study and others [18]. Thus, scirtids may very well be altering leaf-associated fungal communities and increasing fungal-derived nutrients in the system via feeding on embedded fungal hyphae not measured here, however, that question should be addressed in future experiments.

The influence of time on microbial community dynamics is apparent in the high SCC values for this variable in Tables 2 and 3. Over the course of the experiment, bacterial

Figure 6 PCA ordination of T-RFLP peak areas representing fungal taxa from water (a and b) and leaf (c and d) samples taken from microcosms on day 20 (a, c) and day 40 (b, d). Treatments are represented by the following symbols: circles 0 macroinvertebrates; squares 10 scirtids, 0 mosquitoes; open triangles 0 scirtids, 40 mosquitoes; filled triangles 10 scirtids, 40 mosquitoes. The percentage of variance explained by PC scores for each substrate (water or leaf) is indicated on the axes for each panel.
abundance and productivity declined in the water column of microcosms lacking macroinvertebrates, as did the production of bacteria on leaf surfaces. In contrast, the abundance of bacteria on leaf surfaces increased with time in the no-macroinvertebrate microcosms (Fig. 2). Temporal succession is also evident in the structure of microbial communities in these microcosms (Figs. 5 and 6). Over time, PC scores from T-RF profiles shifted even in the absence of macroinvertebrates. However, the trajectories of PC scores representing microbial communities without macroinvertebrates generally differed in magnitude and direction from the trajectories of communities exposed to macroinvertebrate feeding, suggesting that these consumers play a more complex role in microbial dynamics than simply increasing the speed of microbial succession.

As shredders, sciritid feeding in tree holes accelerates detrital processing by reducing leaf material into finer fragments and variation in the condition of available resources is postulated to underlie the stochastic distribution of sciritid beetles occupying nearby tree hole such that tree holes with high quality litter resources are likely to harbor sciritid populations [29–31]. From a mosquito’s perspective, sciritid feeding alters detrital surfaces and water column content, thus directly impacting the primary larval feeding zones. Leaf surfaces in particular are a tough, non-nutritive vehicle upon which resides a film of rich nutrients incorporated as microbial biomass. Although bacteria in this biofilm are severely grazed down by mosquito larvae, fungal members of this biofilm are less susceptible because they are anchored into the matrix of decaying tissue itself. Scritid feeding may make this resource more available to larvae by exposing hyphae and/or releasing small, ingestible fragments containing fungal hyphae into the filter feeding zone. That scritid feeding altered water column fungi supports the latter mechanism but measurements of fungal biomass in the water column are necessary to further support this hypothesis. Because resource availability is the most important contributing factor to mosquito growth, sciritid-associated changes in the underlying structure of microbial communities and the spatial rearrangement of these communities may provide mechanisms by which sciritid beetle presence in tree holes alters mosquito success under nutrient-limited conditions.

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