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Between two trees: Environmental effects of *I. micheliana* and *A. latifolia* on leaf litter ants in a coffee agroecosystem

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Abstract

Shade trees provide important ecological services that support productivity in coffee agroforestry systems. Processes such as biological nitrogen fixation play a key role in this. Less is known, however, about potential indirect mechanisms by which nitrogen fixation supports coffee productivity. One potential route for this to occur is by providing ecological benefits to other above- and belowground organisms that enrich the overall function of agroecosystems. A useful lens with which to evaluate the ecological benefits to these communities under shade trees is to assess how ground-dwelling ant communities respond to the quality of leaf litter from established nitrogen (N)-fixing tree species. Here, we use two trees commonly planted in coffee agroecosystems: Inga micheliana, an N-fixing species, and Alchornea latifolia, a non-N-fixing species. In this study, we set out to answer the following questions: (1) How does the leaf litter environment differ between I. micheliana and A. latifolia? (2) Do differences in environmental factors between I. micheliana and A. latifolia correlate with differences in ant abundance and species richness? and (3) Do differences in environmental factors between I. micheliana and A. latifolia correlate with differences in ant community composition? Twenty-eight randomly selected sites (14 I. micheliana and 14 A. latifolia) were established within a 45-ha plot in a shaded organic coffee farm in Chiapas, Mexico. Three 1-m² quadrats within a 5-m radius from the base of the selected trees were established, and the leaf litter within the quadrats was removed and sieved. Ant specimens were extracted from leaf litter collected from quadrats using the mini-Winkler method and identified to genus and species, or morphospecies, level. Results indicate that *I. micheliana*, the N-fixing species, has a lower C:N ratio than A. latifolia. Differences in C:N ratios are correlated with ant abundance but not with ant species richness. Distance to edge (in meters) has significant effects on leaf litter ant abundance, richness, and species composition. Results suggest that there may be unaccounted feedbacks from N- and non-N-fixing vegetation to brown food webs enabling them to sustain similar ground-dwelling ant communities.

KEYWORDS

agroecosystem, ants, brown food web, coffee, Formicidae, Inga, leaf-litter, nitrogen

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INTRODUCTION

Coffee agroecosystems have been established in biodiversity hotspots around the world and constitute the livelihood of millions of traditional farmers in the tropics (Perfecto et al., 2014; Perfecto & Vandermeer, 2015). Ecological literature in the last three decades has evidenced the array of coffee management practices and their impacts on biodiversity, from low-management (shade-grown) styles that promote high levels of biodiversity (Philpott, Arendt, et al., 2008) to intensified agricultural plantations (sun-coffee) that result in low biodiversity levels (Armbrecht et al., 2005; Mas & Dietsch, 2003; Moguel & Toledo, 1999; Perfecto et al., 1996). Planned biodiversity (e.g., coffee plants and shade trees), in conjunction with associated biodiversity, facilitates a set of valuable ecosystems services (e.g., biological pest control) and ecosystem functions (e.g., formation of soil organic matter [SOM]) (Barnes et al., 2017; Jha et al., 2011; Moguel & Toledo, 1999; Tscharntke et al., 2005).

Shade trees play an important role in coffee agroecosystems. They maintain soil moisture, control weeds, alleviate pest breakouts (Morris & Perfecto, 2016; Soto-Pinto et al., 2002), and provide a good source of fuelwood, fruits, and construction material (García-Barrios et al., 2009; Peeters et al., 2003; Valencia et al., 2014). Nonetheless, shade trees are not always present in coffee systems (i.e., sun-coffee agroecosystems). In shade-coffee farms, tree species composition can vary widely, and it is highly dependent on the farmer's management practice (Valencia et al., 2014). Farmers in Central America manage highly weathered and nutrient-poor soils in coffee agroecosystems by favoring N-fixing trees, particularly those from the *Inga* genus (Grossman et al., 2006; Romero-Alvarado et al., 2002; Valencia et al., 2014). The practice varies according to region and country, especially when use of non-N-fixing trees is also common given their multiple uses (e.g., timber, fruits, light shade, etc.) (Peeters et al., 2003). Extensive efforts have been directed at understanding the role that N-fixing trees play in coffee production and maintenance of biodiversity; nonetheless, many of these studies have focused on interactions taking place in the arboreal component of the agroecosystem (Barrios et al., 2018; Hajian-Forooshani et al., 2016; Philpott et al., 2004; Philpott & Bichier, 2012). Less is known about how N-fixing trees influence ground level and belowground food webs in coffee agroecosystems.

Due to the capacity of species in the genus *Inga* for biological nitrogen fixation (BNF) (Pennington, 1997), the trees generally have a lower carbon-to-nitrogen ratio (herein C:N ratio) in their foliage, which produces high-quality leaf litter and has the potential to accelerate loss of lignin and soluble C in the leaf litter (Talbot & Treseder, 2012). High-quality leaf litter, either due to the identity of leaves

(i.e., specific C:N ratio) or addition of resources (e.g., necromass), can increase microbial activity and accelerate decomposition (Clay et al., 2013; Shik & Kaspari, 2010; Talbot & Treseder, 2012; Zhang & Zak, 1995). Differing C:N ratios have been shown to correlate with altered microbial communities and to affect potential synergisms in decompositions rates (Chapman et al., 2013), ultimately influencing nutrient cycling. Changes in microbial communities due to nutrient availability could potentially lead to increases in decomposition rates, which in turn could cause the loss of habitat space for soil arthropods, including leaf litter ants (Shik & Kaspari, 2010). Nevertheless, increased decomposition rates could also lead to increases in soil macrofauna abundance and richness, especially for fungi grazers (i.e., Collembola), increasing availability of resources (e.g., prey) for ground-dwelling ants. Although past studies have shown the effects that C:N ratio has on microbial communities and feedback loops between them (Beare et al., 1992; Coleman, 2011), it remains elusive how the C:N ratio affects ground-dwelling ant communities.

In this study, we investigate how leaf litter from Inga micheliana (a N-fixing species) and Alchornea latifolia (a non-N-fixing species) influences leaf litter ant communities. Ants (Hymenoptera: Formicidae) are highly abundant and diverse in tropical ecosystems (Hölldobler & Wilson, 1990) and are regularly used to assess ecosystem responses to land management (Offenberg, 2015). They also serve as ecosystem engineers (e.g., through soil bioturbation) (Lobry de Bruyn, 1999; Nkem et al., 2000; Vandermeer & Perfecto, 2007), biocontrol agents (Morris & Perfecto, 2016; Perfecto & Castiñeiras, 1998; Philpott & Armbrecht, 2006; Vandermeer et al., 2010), and are good indicators of ecosystem health (Benckiser, 2010). Ants respond to changes in local factors such as tree species richness, tree abundance, and leaf litter biomass (Armbrecht et al., 2005; De la Mora et al., 2013; Philpott & Armbrecht, 2006). Furthermore, shaded-coffee farms have been shown to support a high diversity of ants and other leaf litter arthropods (Perfecto et al., 1996, 2014; Philpott et al., 2004). However, little is known about how ground-dwelling ant species respond to changes in the C:N ratio of leaf litter in tropical agroecosystems (Philpott & Armbrecht, 2006).

We examined the abundance, richness, and community composition of ground-dwelling ant communities in leaf litter from *I. micheliana* and *A. latifolia* in a shaded-coffee agroecosystem. We specifically asked: (1) How does the leaf litter environment differ between *I. micheliana* and *A. latifolia*? (2) Do differences in environmental factors between *I. micheliana* and *A. latifolia* correlate with differences in ant abundance and species richness? and (3) Do differences in environmental factors between *I. micheliana* and *A. latifolia* correlate with differences in ant community

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composition? Since *I. micheliana* carries out BNF (Pennington, 1997; Romero-Alvarado et al., 2002), we predicted a lower C:N ratio in the leaf litter that accumulates beneath its crown as compared to *A. latifolia*, which does not fix N. Furthermore, we predicted that increased nutrient availability influences ground-dwelling ant abundance and richness, with greater ant abundance and richness in leaf litter with a lower C:N ratio.

METHODS

Study site

We collected samples from a 45-ha plot in Finca Irlanda, an organic shaded-coffee farm in the Soconusco region of Chiapas, Mexico, located approximately at 92°20′29″ W and 15°10′6″ N. Altitude in the plot ranges from 900 to 1200 m above sea level (asl) (Li et al., 2016). The region is characterized by two distinct seasons: a wet season from mid-late May to October and a dry season from November to April. Mean annual rainfall is 4500 mm (De la Mora et al., 2013; Philpott, Perfecto, & Vandermeer, 2008).

Local site selection and environmental factors

To avoid potential competitive exclusion of grounddwelling ants (Ennis, 2010) by keystone ant species, Azteca sericeasur, trees were selected that met the following criteria: (1) free of A. sericeasur nests for the last 3 years; (2) neighboring trees within a 10-m radius had to be free of A. sericeasur; (3) the paired trees were a minimum of 10 m apart to avoid shared leaf litter content and a maximum of 100 m from each other to ensure sampling was done in a similar area of the 45-ha plot and sampled during the same day (e.g., sampling in overgrown or managed area is not representative of the coffee farm as a whole) (De la Mora et al., 2013); (4) diameter at breast height (dbh) had to be equal to or greater than 30 cm. Using a georeferenced map of the 45-ha plot in Finca Irlanda, we selected 28 trees for sampling: 14 of I. micheliana, the most abundant N-fixing species, and 14 of A. latifolia, the most abundant non-N-fixing species (Li et al., 2016). At each tree, we set four 5-m-long transects from the base of the tree in a cross pattern following all cardinal directions. Along the NS transect, we established three 1-m² quadrats for arthropod extraction, for a total of 84 quadrats: one at the base of the tree and two at 5 m from the base of the tree. All transects were used to measure local environmental site factors (e.g., leaf litter depth).

Local environmental site information (from now on "local factors") was collected for each tree site to capture potential effects on ant abundance, richness, and species composition. A total of 13 local factors were measured: dbh (in centimeters), number of coffee plants within the four 5-m-long transects, mean leaf litter depth (in millimeters) within all 5-m-long transects, leaf litter depth (in millimeters) within each 1-m² quadrat, slope cardinality (in degrees), altitude (in meters above sea level), distance to edge (in meters) (edge of trails and roads in coffee farm), mean percent crown cover, pH, percent soil humidity, total percent nitrogen (% N), total percent carbon (% C), and C:N ratio. Measurements of leaf litter depth were taken at base of the tree, 2.5 m. and 5 m from the base of the tree in all four cardinal directions for total of 10 data points. Leaf litter depth within 1-m² quadrats was measured at the center and all four corners. Measuring along transects and within quadrats provided a measure of the mean leaf litter volume under the tree crown. Slope and altitude of sites were determined with a Garmin 72H model (www.garmin.com). To determine mean percent crown cover, four measurements in cardinal directions were taken with a spherical crown densiometer, Model A (Forestry Suppliers), at the base of the tree. Mean soil pH was determined by colorimetric method (Lovibond Soil pH Test Kit, MPN number 694, www.forestry-suppliers.com). Briefly, we collected a core from the soil surface using a 2-mL vial tube at each of the three 1-m² quadrats established within the NS transects of selected trees for arthropod extraction. We then homogenized each core and proceeded following manufacturers' guidelines. Soil samples were processed in the field laboratory within 6 h of being collected. Soil humidity was assessed through the gravimetric method by taking an additional 4-6 g of soil from each of the 1-m² quadrats in NS transects of selected trees. Each sample was weighed and then dried for a minimum of 72 h at 50°C until no further mass loss was recorded. All 13 local factors were measured on the same date as ant sampling occurred for each tree pair (see Ant sampling protocol). For C and N analyses, we collected leaf litter from an additional fourth 1-m² quadrat in proximity to NS transect (n = 28). Collecting from an additional quadrat ensured that leaf litter was not disturbed by previous measurements and sampling efforts (see below for details).

Ant sampling protocol

Leaf litter was collected from the three 1-m² quadrats established at each tree and sifted from these quadrats using the Winkler method (Agosti et al., 2000). All leaf litter

inside the quadrat was sifted and placed in mini-Winkler extractors and left for 72 h to collect ants and other leaf litter arthropods in containers with 70% ethyl alcohol. Mini-Winkler extractors were equipped with a 50–75 W incandescent light bulb to increase extraction efficiency. Ants were separated from other arthropods and organisms and placed in vials with 70% ethyl alcohol for further identification. Specimens were identified to species and morphospecies levels using the *Identification Guide to the Ant Genera of the World* (Bolton, 1994).

Leaf litter chemical analyses: C:N ratio

All dried leaf litter inside the fourth 1-m² quadrat was collected and dried for a minimum of 72 h at 50°C until a constant mass was obtained. Following that, we homogenized and subsampled 20 g of leaf litter and placed it in plastic bags (Ziploc, 16.5 × 14.9 cm) for chemical analyses. Subsamples were pulverized using a Krups brand coffee grinder (model GVX212) in the finest setting. Approximately 0.2–0.3 g of ground sample was analyzed for C and N content using a LECO Trumac CN combustion analyzer (LECO Corporation, Saint Joseph, MI). Chemical analyses were conducted at the laboratory of Dr. Jennifer Blesh at the University of Michigan (Ann Arbor, MI, USA). Data from this fourth quadrat are considered representative of the chemical composition of leaf litter of *I. micheliana* and *A. latifolia* in this study and statistical analyses.

Statistical analyses

Ant species abundance, richness, and community composition

To assess the effectiveness of our sampling efforts at capturing ant richness, we computed species accumulation curves (SACs) for observed species richness. We used the "BiodiversityR" package (Kindt, 2022; Kindt & Coe, 2005) with parameters set to 100 permutations with the "exact" method and second-order jackknife with 95% confidence intervals (CIs). Additionally, mean ant abundance and richness were tested for significant differences between treatments using paired Student's *t* tests, where ant abundance was the count of individuals and richness was the number of unique species.

Local environmental factors

To address our first question, we computed paired t tests for all the local factors of I. micheliana and A. latifolia

sites: dbh, altitude, slope cardinality, distance to edge, number of coffee plants within all 5-m-long transects, leaf litter depth, leaf litter depth within 1-m² quadrats, pH, percent soil humidity, total % N, total % C, and C:N ratio. We computed simple linear regression to explore the correlation between significantly different local factors and ant abundance and richness in the leaf litter of both tree species.

Predicting ant abundance, richness, and community composition

To address our second question, we computed generalized linear mixed models (GLMMs). We computed the variance inflation factors (VIFs) with the vif function in the "car" (Fox et al., 2022) to examine multicollinearity among all local factors: dbh, number of coffee plants within the four 5-m-long transects, mean leaf litter depth within all 5-m-long transects, leaf litter depth within each 1-m² quadrat, slope cardinality, altitude, distance to edge (edge of trails and roads in coffee farm), mean percent crown cover, pH, percent soil humidity, % N, % C, and C:N ratio. Computation was performed by building two initial GLMMs for abundance and richness using the glmer function in "lme4" package and optimized parameters with the bobyga method (Bates et al., 2015). Local factors with a VIF greater than 5 were considered highly correlated. Although total % N was highly correlated (VIF > 5) with C:N ratio in both initial GLMMs, we still considered it relevant as it has been reported to be a significantly positive predictor of leaf litter predators (Kaspari & Yanoviak, 2009). To ensure the best fit possible, we scaled and centered the continuous local factors using the scale function in R software (R Core Team, 2022) and computed the GLMMs with a Poisson distribution for ant abundance and species richness. Due to the high number of local factors, we did not model interaction effects, this allowed us to avoid convergence issues in our GLMMs. We consider our models a reasonable subset of the truly maximal model. All local factors were set as fixed effects in the models. We set sampling dates (eight dates during June-July 2016) and site identification (ID) as random effects in our models for predicting ant abundance to capture changes in other environmental changes not measured throughout the season. We eliminated site ID as a random effect to avoid model singularity in our GLMM for predicting ant species richness. To select the most parsimonious model, we used the "buildmer" package (Voeten, 2022) to perform a stepwise backward elimination of insignificant variables until we reached a model that maximized model fit. The same optimizer parameters were applied. Best-fit model

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residuals were visually evaluated to corroborate fit using function gqPlot from the car package (Fox et al., 2022). Marginal and conditional R^2 values for best-fit models were computed with "MuMin" package (Bartoń, 2022). We removed an outlier A. latifolia site with more than 2000 ant individuals from analyses. Because all sites were uniquely paired, the corresponding paired individual of I. micheliana was also removed from all analyses. For all response variables and local factors, apart from leaf litter depth, total % N, total % C, and C:N ratio, we report data from a total of 78 quadrats (13 I. micheliana sites \times 3 quadrats and 13 A. latifolia sites \times 3 quadrats). Data for leaf litter depth, total % N, total % C, and C:N ratio are reported on a per site basis (n = 26). Mean values and standard errors for local factors were compared and tested with paired Student's t tests.

Finally, we computed a permutational multivariate analysis of variance (PERMANOVA) using adonis2 function from the "vegan" package (Oksanen et al., 2022) to understand how differences in local factors between I. micheliana and A. latifolia correlate with differences in ant community composition (9999 permutations, method = "bray"). We used a permutational analysis of multivariate dispersion (PERMDISP) as companion to PERMANOVA to corroborate homogeneous dispersion of variances, a necessary assumption for this test, and exclude the possibility that any significant differences between leaf litter ant communities were caused by heterogeneous dispersion of variances. The PERMDISP was computed with betadisper and tested for significance with permutest functions in the vegan package (Oksanen et al., 2022). To visualize how local factors correlate with leaf litter ant communities, we computed a distance-based redundancy analysis (dbRDA) by applying the capscale function to a Bray-Curtis similarity matrix (Oksanen et al., 2022). The dbRDA and corresponding visualization appropriately illustrate the underlying patterns of compositional differences, as it is considered analogous to PERMANOVA with non-Euclidean distance matrices (Legendre & Anderson, 1999). We used the simper function in vegan to discriminate which species contribute the most to compositional differences between groups (Oksanen et al., 2022). All statistical analyses were performed using R statistical software (v4.2.1, R Core Team, 2022).

RESULTS

Sampling effort to capture ant abundance, richness, and community composition

Our sampling of 78 quadrats (39 per tree species) resulted in a total of 6574 ant individuals from 8 subfamilies, 34 genera, and 67 morphospecies and species (Appendix S1: Table S1).

SACs estimated asymptotes indicate that our sampling effort captured the mean richness of leaf litter ant species (Figure 1). The overlap of the CIs of the SACs revealed no significant difference in observed species richness between leaf litter types. We corroborated this with paired t tests (t=0.049, $\mathrm{df}=38$, p=0.96). For overall ant abundance, no significant differences between tree species were observed as well (paired t tests, t=-0.73, $\mathrm{df}=38$, p=0.47). We found no significant differences in abundance for species present at more than five tree sites, except for *Nylanderia* sp1. (p=0.02). Among species that were present in less than five tree sites, we found significant differences for *Solenopsis zeteki* (p=0.01) and *Solenopsis* sp1. (p=0.01).

Effects of local factors on ant abundance and richness

Several local factors were found to be significantly different between I. micheliana and A. latifolia sites (Table 1). dbh (in centimeters) and C:N ratio were significantly greater under A. latifolia than I. micheliana, while total % N and number of coffee plants were greater under the I. micheliana trees (Figure 2, Table 1). Quadrats at host tree sites did not differ for any of the other measured local factors. Exploratory simple linear regressions revealed, overall, no significant trends between significantly different local factors and ant abundance and richness (Figure 3). We only observed a significant correlation for dbh and ant richness at I. micheliana quadrats (p = 0.01) (Figure 3e).

GLMMs revealed how local factors correlate with leaf litter ant abundance and richness (Table 2). The initial GLMM for predicting leaf litter ant abundance reports that distance to edge (p = 0.001); mean leaf litter depth within 1-m² quadrats (p = 0.002); pH (p < 0.001); percent soil humidity (p < 0.001); total % N (p < 0.001); total % C (p < 0.001); and C:N ratio (p < 0.001) are significant predictors (marginal R^2 : 0.44 and conditional R^2 : 0.99) (Figure 4, Table 2). The most parsimonious model for ant abundance revealed that from the initial 13 local factors, only distance to edge (fixed effect) (p = 0.0371) and sampling date (random effect) were significant in predicting leaf litter ant abundance (marginal R²: 0.32 and conditional R^2 : 0.97). The GLMM model with all the factors for predicting leaf litter ant richness revealed that only distance to edge (p = 0.003) was significant (marginal R^2 : 0.24 and conditional R^2 : 0.57). The final parsimonious model for predicting richness of leaf litter ants revealed that distance to edge (fixed effect) (p = 0.003) and sampling date (random effect) are significant factors (marginal R^2 : 0.08 and conditional R^2 : 0.44). Distance to edge was the only local factor that had a significant

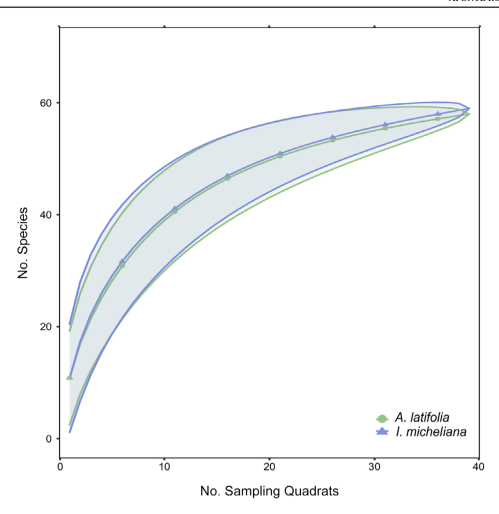


FIGURE 1 Species accumulation curves of leaf litter ant communities under *A. latifolia* and *I. micheliana*. Shaded areas represent 95% CIs.

TABLE 1 Values (mean \pm SE) of local factor characteristics at *I. micheliana* and *A. latifolia* sites in coffee agroecosystem in Chiapas, Mexico.

| Site characteristics | Inga micheliana | Alchornea latifolia | t | p |
|---|--------------------|---------------------|-------|-----------|
| dbh (cm) | 32.6 ± 1.12 | 59.97 ± 1.92 | 12.04 | <0.001*** |
| Altitude (m) | 1039.77 ± 4.58 | 1032.39 ± 2.60 | -1.33 | 0.19 |
| Slope cardinality (°) | 194.39 ± 14.68 | 224.00 ± 15.05 | 1.72 | 0.09 |
| Distance to edge (m) | 12.71 ± 1.20 | 17.54 ± 2.61 | 1.40 | 0.17 |
| Leaf litter depth (mm) | 53.07 ± 3.19 | 58.95 ± 3.70 | 1.66 | 0.11 |
| Leaf litter 1-m ² quadrat (mm) | 57.62 ± 2.69 | 60.48 ± 4.25 | 0.54 | 0.59 |
| Crown cover (%) | 66 ± 5.11 | 72.00 ± 4.12 | 0.79 | 0.44 |
| No. coffee plants | 30.07 ± 1.63 | 23.15 ± 1.84 | -2.71 | 0.01** |
| Percent soil pH | 6.00 ± 0.15 | 6.36 ± 0.11 | 1.72 | 0.09 |
| Percent soil humidity | 83.82 ± 7.38 | 81.51 ± 3.20 | -0.29 | 0.77 |
| C:N ratio | 20.27 ± 0.31 | 25.28 ± 0.68 | 8.02 | <0.001*** |
| Total % N | 2.32 ± 0.03 | 1.92 ± 0.06 | -7.20 | <0.001*** |
| Total % C | 46.61 ± 0.36 | 47.01 ± 0.29 | 0.83 | 0.41 |

Note: Values show mean, standard error, and results from paired t tests (n = 78, df = 38). Values in boldface are significant. **p < 0.01; ***p < 0.001.

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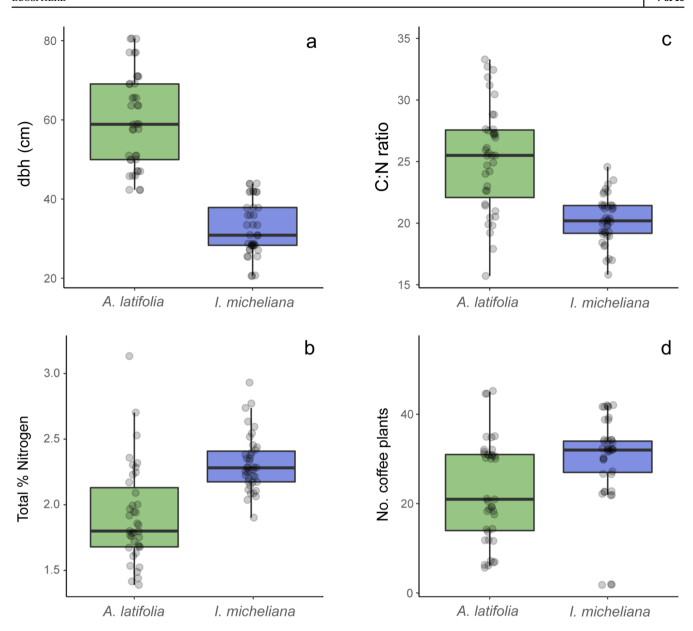


FIGURE 2 Boxplots of significantly different local factors at *I. micheliana* and *A. latifolia* sites: (a) dbh (p < 0.001); (b) total % N (p < 0.001); (c) C:N ratio (p < 0.001); and (d) number of coffee plants (p = 0.01). All significant values were computed with paired t tests. Lines within boxes represent median values, boxes enclose 25th and 75th percentiles, and whiskers enclose 5th and 95th percentiles.

negative correlation with both ant abundance and species richness in initial and final (parsimonious) GLMMs (Figures 4 and 5, Table 2). We summarized the log-mean estimates from the initial GLMMs with 95% CIs, to better illustrate the significant negative correlations for mean 1-m² quadrat leaf litter depth and mean soil and significantly positive correlations with mean soil pH, percent soil humidity, C:N ratio, total % N, and total % C with ant abundance (Figure 4). The log-mean estimate visualization of the GLMM for ant richness highlights only that distance to edge has a significant negative correlation.

The PERMDISP tests revealed no significant differences in the centroids of the dispersions of leaf litter ant

communities from *I. micheliana* and *A. latifolia* (F=0.4, p=0.54). Visualization of species composition with dbRDA showed overlap in the community structure as well as the associations between local factors with ant communities in *I. micheliana* and *A. latifolia* leaf litter (Figure 6). The axes CAP1 and CAP2 of the dbRDA explain 21.2% and 19%, respectively, of 5.9% of the constrained variance (Figure 6). Results from PERMANOVA revealed significant correlations for tree species ($R^2=0.03$, p<0.0001); dbh ($R^2=0.02$, p=0.02); altitude ($R^2=0.02$, p=0.02); slope cardinality ($R^2=0.02$, p=0.01); distance to edge ($R^2=0.03$, p=0.002); and percent soil humidity ($R^2=0.02$, p=0.03) (Table 3). Marginally significant correlations were observed for mean 1-m² leaf litter depth

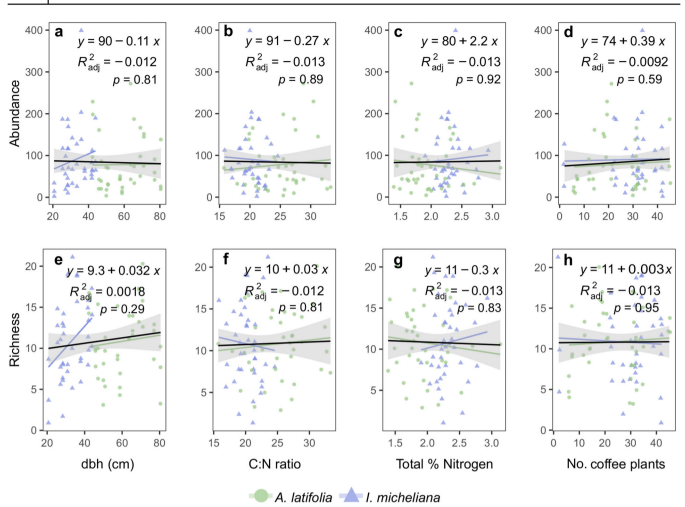


FIGURE 3 Simple linear regressions between significantly different local factors at *I. micheliana* and *A. latifolia* quadrats and ant abundance and richness in the leaf litter. Regression lines in black represent all data points (n = 78). Linear equation, r^2 value, and p value correspond to all data points. Shaded areas represent 95% CIs.

 $(R^2=0.02,\ p=0.06)$ and total % N $(R^2=0.02,\ p=0.06)$ (Table 3). The species with the greatest cumulative contributions to compositional differences between groups are: Solenopsis terricola (0.17), Pheidole protensa (0.30), Solenopsis picea (0.37), Eurhophalotrix sp1. (0.44), S. zeteki (0.51), Solenopsis sp1. (0.58), Gnamptogenys striatula (0.62), Strumigenys gundlachi (0.66), Hypoponera nitidula (0.69), and Eurhophalotrix sp2. (0.72).

DISCUSSION

Our study investigated two tree species of distinct chemical compositions and leaf trait morphologies and found that leaf litter from *I. micheliana* had significantly lower C:N ratio than that of *A. latifolia* (Figure 2c, Table 1). Tree species also differed significantly in dbh, number of coffee plants, total % N, and C:N ratio (Figure 2, Table 1). Results align well with reports from other studies showing that N-fixing trees produce leaf litter with low C:N

ratio that can contribute to greater accumulation of SOM and higher decomposition rates. High inputs of N to soil may result in increased coffee production and agroecosystem sustainability (Leblanc et al., 2006; López-Rodríguez et al., 2015). The results from our initial GLMMs partially support our prediction that C:N ratio in leaf litter is an important predictor of leaf litter ant abundance, yet there was no significant difference in leaf litter ant abundance between tree species (data not shown). In other words, it is possible that the statistical significance in the model is not biologically relevant. The differences detected in this study do not support the hypothesis that C:N ratio is correlated to leaf litter ant richness (Figure 4, Table 2). Other local factors that were significant predictors variables of leaf litter ant abundance were distance to edge, mean 1-m2 quadrat leaf litter depth, percent soil humidity, mean soil pH, total % N, and total % C (Table 2). On the other hand, in the GLMM for predicting ant richness, only distance to edge was a significant negative predictor variable (Table 2).

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TABLE 2 Generalized linear mixed models (GLMMs) for predicting leaf litter ant abundance and richness.

| | C | GLMM: Ant abundance | | | GLMM: Ant richness | | | |
|--|----------|---------------------|------------------|---------|--------------------|------|----------------|---------|
| Coefficient | Log-mean | SE | 95% CI | p | Log-mean | SE | 95% CI | p |
| (Intercept) | 4.47*** | 0.29 | 3.90 to 5.03 | < 0.001 | 2.27*** | 0.13 | 2.02 to 2.53 | < 0.001 |
| Tree species | -0.30 | 0.43 | -1.14 to 0.53 | 0.475 | 0.23 | 0.18 | -0.11 to 0.58 | 0.188 |
| dbh (cm) | -0.03 | 0.22 | -0.46 to 0.41 | 0.909 | 0.10 | 0.08 | -0.06 to 0.27 | 0.210 |
| Altitude (m asl) | -0.12 | 0.15 | -0.41 to 0.16 | 0.400 | -0.03 | 0.05 | -0.14 to 0.08 | 0.580 |
| Slope cardinality (°) | -0.20 | 0.17 | -0.54 to 0.13 | 0.239 | -0.00 | 0.07 | -0.14 to 0.14 | 0.993 |
| Distance to edge (m) | -0.49** | 0.15 | −0.79 to −0.20 | 0.001 | -0.19** | 0.06 | −0.32 to −0.07 | 0.003 |
| Mean 1-m² quadrat litter depth (mm) | -0.07** | 0.02 | −0.11 to −0.02 | 0.002 | -0.04 | 0.05 | -0.14 to 0.06 | 0.428 |
| Mean leaf litter depth (mm) | 0.29 | 0.18 | -0.06 to 0.65 | 0.107 | 0.11 | 0.08 | -0.04 to 0.26 | 0.142 |
| No. coffee plants | 0.16 | 0.15 | -0.15 to 0.46 | 0.314 | 0.07 | 0.06 | -0.05 to 0.19 | 0.242 |
| Percent crown cover | 0.12 | 0.17 | -0.22 to 0.45 | 0.486 | -0.01 | 0.07 | -0.14 to 0.13 | 0.939 |
| Mean percent soil pH | -0.05*** | 0.01 | -0.08 to -0.02 | 0.001 | 0.02 | 0.04 | -0.05 to 0.10 | 0.553 |
| Mean percent soil humidity | 0.35*** | 0.03 | 0.30 to 0.40 | < 0.001 | 0.08 | 0.05 | -0.01 to 0.18 | 0.090 |
| C:N ratio | 0.59*** | 0.10 | 0.39 to 0.80 | < 0.001 | 0.06 | 0.21 | -0.35 to 0.46 | 0.783 |
| % N | 0.80*** | 0.12 | 0.57 to 1.02 | < 0.001 | -0.05 | 0.21 | -0.45 to 0.36 | 0.820 |
| % C | 0.12*** | 0.04 | 0.05 to 0.19 | 0.001 | 0.05 | 0.08 | -0.09 to 0.20 | 0.476 |

Note: Random effects for GLMM: ant abundance: $\sigma^2 = 0.01$; $\tau_{00} = 0.24_{\text{ID}}$; $\tau_{00} = 0.24_{\text{Sampling_date}}$; ICC = 0.97; $N = 9_{\text{Sampling_date}}$; $N = 26_{\text{ID}}$; observations = 78; marginal $R^2 = 0.441$; conditional $R^2 = 0.985$. Random effects for GLMM: ant richness: $\sigma^2 = 0.09$; $\tau_{00} = 0.07_{\text{Sampling_date}}$; intraclass correlation coefficient (ICC) = 0.43; $N = 9_{\text{Sampling_date}}$; observations = 78; marginal $R^2 = 0.245$; conditional $R^2 = 0.572$. Values in boldface are significant.

p < 0.01: *p < 0.001.

Even though t tests results also confirm that dbh, total % N, and number of coffee plants significantly different between I. micheliana and A. latifolia, it is not enough to be significantly correlated with leaf litter ant species richness between the groups (Figure 2, Table 3). Again, this could be due to the possibility that the statistical difference of these local factors is not enough to be biologically relevant for the ant community (Figure 1, Table 2). In a similar study, Murnen et al. (2013) reported a small increase in ant colony growth and species richness after adding necromass to leaf litter and increasing its nutrient quality. Unfortunately, no data for % N or other nutrients are reported in this study. In general, we see no significant differences for ant species richness under I. micheliana compared to A. latifolia leaf litter, as shown by the SACs (Figure 1). Although we observe high overlap in species composition (Figure 4), PERMANOVA results highlight that between group differences are correlated with distance to edge, dbh, altitude, slope cardinality, and percent soil humidity. We focus on results from our initial GLMMs to highlight the effects of the different local factors on ant abundance and richness (Figure 4, Table 2). Our final parsimonious models also support the same general conclusion that distance to edge is an important predictor of ant abundance and richness (Figure 5).

Although we did not collect data on soil macrofauna, it is likely that higher N content and lower C:N ratio in leaf litter could potentially increase their abundance and richness, thus increasing availability of resources (e.g., prey) for ground-dwelling ants. This pathway could result in increased abundance of generalist species (e.g., *Solenopsis* spp.) at the cost of lower species richness. Few other studies have investigated the specific role of C:N ratio from leaf litter in ant communities. Hence, comparison across studies is difficult and at times speculative. Studies focused on comparing decomposition of leaf litter from pioneer and old-growth forests do report significant differences, where old-growth forest litter hosts a greater abundance and diversity of leaf litter and soil invertebrate communities (Laird-Hopkins et al., 2017).

Distance to edge was a significant predictor variable for ant abundance and richness (Figure 4, Table 2). It is also strongly correlated to species composition (Table 3). Here, distance to edge referred to the distance to the nearest walking path (1–2 m wide) in the coffee farm, not the edge of a forest or agricultural system like it is commonly utilized (Majer et al., 1997; Silva et al., 2011). The light gap created by trails in the coffee farms has similar effects to gaps in natural forests (Majer et al., 1997; Perfecto & Vandermeer, 1996), potentially allowing colonization of niche space and increased food resources

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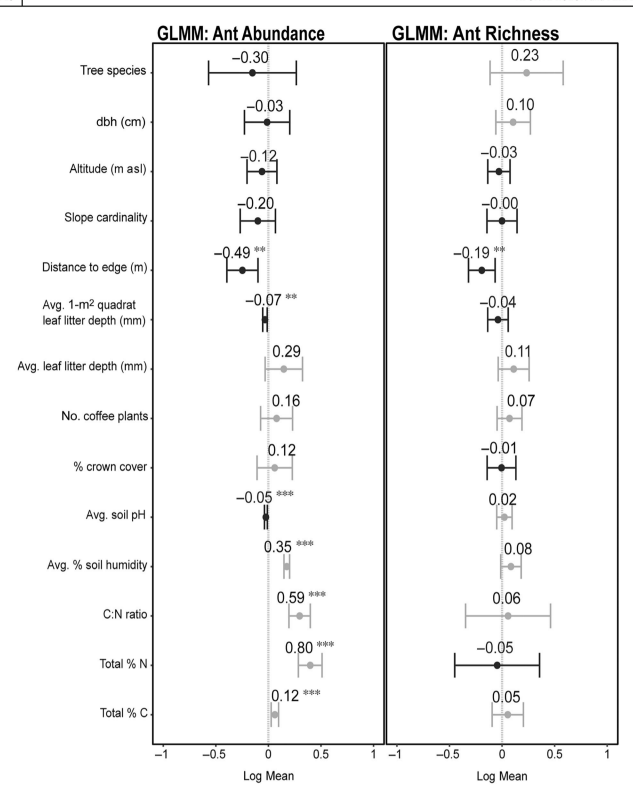


FIGURE 4 The effect of local factors on ant abundance and species richness in *I. micheliana* and *A. latifolia* leaf litter. A positive value indicates that local factors are positively correlated with ant abundance or species richness. Positive correlations are in gray circles and negative correlations are in black circles. Values represent log-mean estimates of generalized linear mixed models (GLMMs). Bars represent 95% CIs. Asterisks denote significance of effect size (**p < 0.01; ***p < 0.001). Avg. average.

(e.g., food disposal by farm workers; Aponte Rolón, personal observations) for genera like *Solenopsis*, *Pheidole*, and *Wasmannia auropunctata*, which have a high

capacity for recruitment of workers and can build large colonies (Hölldobler & Wilson, 1990). Nevertheless, the presence of these dominant species can have a negative ECOSPHERE 11 of 16

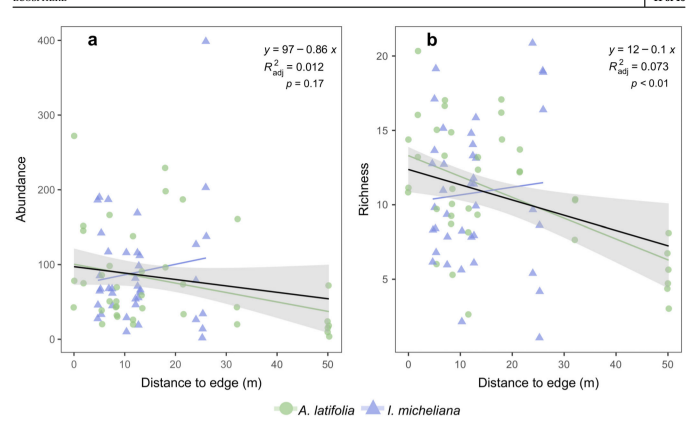


FIGURE 5 Simple linear regressions between distance to edge (in meters) and *I. micheliana* and *A. latifolia* quadrats' ant abundance and richness in the leaf litter. Regression lines in black represent all data points (n = 78). Linear equation, r^2 value, and p value correspond to all data points. Shaded areas represent 95% CIs.

effect on other ants, therefore reducing species richness (Ennis & Philpott, 2017).

Our results contrasted with other studies that report a strong correlation of ant abundance and richness with organic matter mass and leaf litter depth (De la Mora et al., 2013; Sabu et al., 2008). Rather, our results supported reports from Shik and Kaspari (2010), which showed no differences in leaf litter ant abundance or richness in experimental plots, and another study by Kaspari et al. (2010), which reported homogeneous ant species richness along a topographic and nutrient gradient (e.g., nitrogen [N], phosphorus [P], and potassium [K] additions). Results reported by Shik and Kaspari (2010) partially support the "more food, less habitat" hypothesis, where leaf litter decomposes faster due to increased microbial activity hence preventing observable differences in ant abundance and richness. Similar results were reported by Murnen et al. (2013) when they compared ant communities among forest, sun-coffee, and shaded-coffee habitats and found that habitat type did influence ant abundance and richness, but not food addition. Schmitt et al. (2020) examined the decomposition of I. micheliana leaves but found no difference; it was the presence of A. sericeasur that changed the leaf litter ant community composition. This points towards higher order ecological interactions influencing ants in the leaf litter.

Overall, we found that the leaf litter produced by I. micheliana (the N-fixing tree) had a statistically significant lower C:N ratio and dbh, and a higher total % N and number of surrounding coffee plants than A. latifolia (the non-N-fixing tree) (Figure 2, Table 1). However, these differences do not seem to be biologically important to distinguish the ant community living in leaf litter under these tree species. More specifically, we could not detect any significant differences in the leaf litter ant abundance and richness. In contrast, we see differences in species composition under these tree species that are correlated with multiple local factors. With GLMMs, we found that distance to edge, mean 1-m² quadrat leaf litter depth, pH, percent soil humidity, C:N ratio, total % N, and total % C were significant predictors of ant abundance. Abundance is positively correlated with percent soil humidity, C:N ratio, total % N, and total % C, while distance to edge, mean 1-m² quadrat leaf litter depth is negatively correlated (Figures 4 and 5). Finally, distance to edge (i.e., distance to a trail or road) was a significant negative predictor of both ant abundance and species richness. Leaf litter ants decline in abundance and richness as distance to edge increases. A slightly different set of local factors correlates with ant species composition, as shown by PERMANOVA results (Table 3) and dbRDA

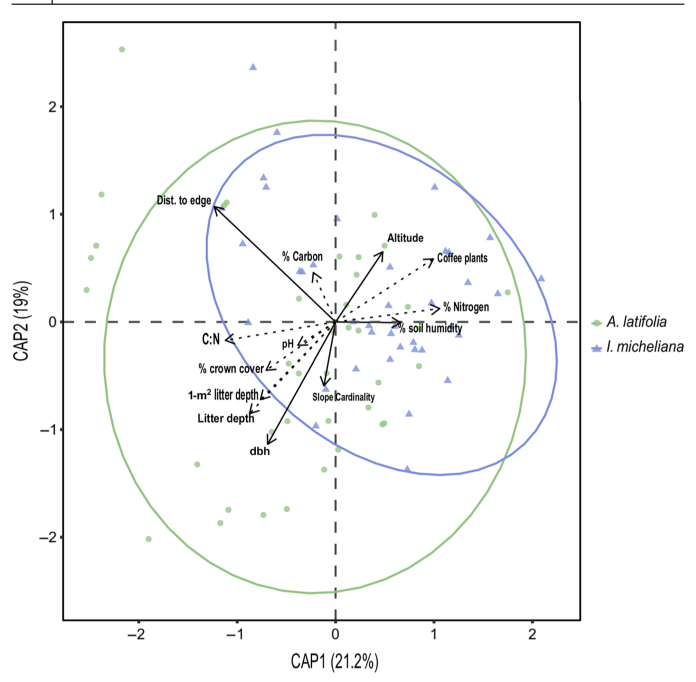


FIGURE 6 Leaf litter ant community composition associated with local factors. Ant community variation within and between sample quadrats (n = 78) under *I. micheliana* and *A. latifolia* trees from distance-based redundancy analysis models constrained by local factors. Solid lines represent significant associations (p < 0.05). Each point represents leaf litter ant community sampled at quadrats.

(Figure 6). Compositional differences between leaf litter ant communities are significantly correlated with dbh, altitude, slope cardinality, soil percent humidity, and distance to edge (Figure 6, Table 3). Regardless of local factors' statistically significant differences, only distance to edge contributes to biologically important differences in leaf litter ant abundance, richness, and species composition.

Further studies should focus on manipulative experiments that isolate the effects of leaf litter type and get at the potential priority effects on ant community

composition. As well as focus on comparing changes in microbial activity due to leaf litter type and its effects on ant community composition. Results suggest that a shaded-coffee agroecosystem can support high levels of ant biodiversity regardless of which tree species, *I. micheliana* or *A. latifolia*, is planted by farmers. Although from this study, it is not clear at what scale (e.g., quadrat) local factors most contribute to observable biological differences in leaf litter ant abundance, richness, and species composition.

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TABLE 3 Permutational multivariate analysis of variance of local factors and leaf litter ant community composition.

| Characteristic | df | SS | R^2 | F | p |
|---|----|--------|-------|-------|-----------|
| Tree species | 1 | 0.786 | 0.034 | 2.934 | <0.001*** |
| dbh (cm) | 1 | 0.510 | 0.022 | 1.905 | 0.02* |
| Altitude (m asl) | 1 | 0.510 | 0.022 | 1.902 | 0.02* |
| Slope cardinality (°) | 1 | 0.543 | 0.024 | 2.027 | 0.01** |
| Distance to edge (m) | 1 | 0.661 | 0.029 | 2.467 | 0.002** |
| Mean 1-m ² quadrat litter depth (mm) | 1 | 0.430 | 0.019 | 1.604 | 0.061 |
| Mean leaf litter depth (mm) | 1 | 0.364 | 0.016 | 1.360 | 0.144 |
| No. coffee plants | 1 | 0.352 | 0.015 | 1.314 | 0.173 |
| Percent crown cover | 1 | 0.333 | 0.015 | 1.245 | 0.218 |
| Mean percent soil pH | 1 | 0.089 | 0.004 | 0.331 | 0.996 |
| Mean percent soil humidity | 1 | 0.465 | 0.020 | 1.736 | 0.034* |
| C:N ratio | 1 | 0.230 | 0.010 | 0.857 | 0.631 |
| % N | 1 | 0.425 | 0.019 | 1.585 | 0.061 |
| % C | 1 | 0.300 | 0.013 | 1.119 | 0.315 |
| Residuals | 63 | 16.875 | 0.738 | | |
| Total | 77 | 22.872 | 1.000 | | |

Note: Values in boldface are significant.

AUTHOR CONTRIBUTIONS

Bolívar Aponte Rolón and Ivette Perfecto are responsible for the conception of research questions and experimental design. Bolívar Aponte Rolón is responsible for fieldwork, data collection, and analysis. Bolívar Aponte Rolón produced the first manuscript draft. Both authors agree on the final manuscript.

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CONFLICT OF INTEREST STATEMENT

The authors declare that they have no conflict of interest.

DATA AVAILABILITY STATEMENT

All data and source code (Aponte Rolón, 2023) are available from Zenodo: https://doi.org/10.5281/zenodo.7573987.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article. How to cite this article: Aponte Rolón, Bolívar, and Ivette Perfecto. 2023. "Between Two Trees: Environmental Effects of *I. Micheliana* and *A. Latifolia* on Leaf Litter Ants in a Coffee Agroecosystem." *Ecosphere* 14(2): e4442. https://doi.org/10.1002/ecs2.4442