Manuscript Draft: Aim3 Leaf Traits

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5 1 Abstract

2 Introduction

7 3 Methods

8 3.1 Field

- Growth and host plant inoculation seven tropical tree species was conducted at the greenhouses in the Gamboa
- Research Station, Smithsonian Tropical Research institute, Republic of Panama. The species, *Theobroma ca-*
- an cao, Dypterix sp., Lacmellea panamensis, Apeiba membranacea, Heisteria concinna, Chrysophyllum caim
- 12 ito, and Cordia alliodora were chosen due to their variance in leaf traits (J.Wright unpublished data) and the
- availability of seeds in January- April 2019. Seeds of tree species were collected from the forest floor and
- grown in the greenhouse. Seedlings were kept in a chamber made out of PVC and clear plastic to prevent in-
- oculation from spore fall inside the greenhouse. NEEDS INFORMATION ON THE SOIL MIXTURE AND
- ¹⁶ AUTOCLAVING PROTOCOL. Seedlings reached a minimum of 4 true leaves before endophyte inoculation.
- 17 Then 10 individual plants of each species were exposed to 10 nights of spore fall to achieve a high endophyte

load (E+) and 10 homologous plants were kept inside the greenhouse plastic chamber to maintain a low endophyte load (E-) (Fig. ? MAKE A DIRAGRAM?). Plants exposed to spore fall were placed near (~10 m) the forest edge at dusk (~18:OO hours) and returned to the greenhouse at dawn (~07:00 hours) (Bittleston et al. 2011).

2 3.1.1 Leaf trait measurements

Three mature leaves were haphazardly collected from each of the individual plants in each treatment (E+, E-) within 7-10 days after inoculation treatment. Anthocyanin (ACI) content and leaf thickness (LT) were measured while the leaf was still attached to the plant. We measured anthocyanin content with ACM-200 plus 25 (Opti-Sciences Inc. Hudson, New Hampshire, U.S.A.) on three haphazardly selected locations (working from the petiole out to the leaf tip) on the leaf surface of three haphazardly selected leaves for a total of nine measurements per plant (Tellez et al., 2022). The ACM-200 calculates an anthocyanin content index (ACI) value from the ratio of % transmittance at 931 nm/% transmittance at 525 nm (opti-sciencesinc?). On compound leaves (i.e., Dypterix sp.) we measured at three different leaflets. Leaf thickness (µm) was measured with a Mitutoyo 7327 Micrometer Gauge (Mitutoyo, Takatsu-ku, Kawasaki, Japan) in sthe same manner as the anthocyanin measurements, taking care to avoid major and secondary veins. After anthocyanin and leaf thickness measurements were completed, we removed the leaves from their stems, placed them inside a plastic bag (i.e. Ziploc), place in an ice chest and moved them to the lab for further measurements. Leaf punch strength (LPS) was measured with an Imada DST-11a digital force gauge (Imada Inc., Northbrook, IL, United States) by conducting punch-and-die tests with a sharp-edged cylindrical steel punch (2.0 mm diameter) and a steel die with a sharp-edged aperture of small clearance (0.05 mm). The leaf punch measurements were taken by puncturing the leaf lamina at the base, mid-leaf and tip on both sides of the mid-vein, avoiding minor leaf veins when possible (Tellez et al., 2022). Once leaf toughness was measured, we used a 7 mm diameter punch hole to puncture disks for leaf mass per area (LMA) measurements. We collected one three disks per

leaf (see Supplementary material for details). The disk punches dried at 60 °C for 48-72 hours. before being weighed.

43 3.1.2 Leaf tissue preparation for molecular work

The selected leaves were also used to profile endophyte community composition, abundance, and richness via amplicon sequencing (Illumina MiSeq). The leaf tissue remaining after the leaf trait measurements had the main vein and margins excised so that only the lamina remained. The lamina was haphazardly cut into 2 x 2 mm segments, enough to obtain a total of 16, and surface sterilized by sequential rinsing in 95% ethanol (10 s), 0.5 NaOCl (2 mins) and 70% ethanol (2 mins), as per (Arnold et al., 2003; Higgins et al., 2014; Tellez et al., 2022). After, leaves were air-dried briefly under sterile conditions. Sixteen leaf segments per leaf, a total of forty-eight leaf segments per plant, were plated in 2% malt extract agar (MEA), sealed with Parafilm M (Bemis Company Inc., U.S.A.) and incubated at room temperature. The cultured leaf segments were used to estimate endophyte colonization of E+ and E- leaves. The presence or absence of endophytic fungi in the leaf cuttings was assessed 7 days after plating. The remaining sterilized leaf lamina was preserved in sterile 15 mL tubes with ~ 10 mL CTAB solution (1 M Tris—HCl pH 8, 5 M NaCl, 0.5 M EDTA, and 20 g CTAB). Leaf tissue in CTAB solution was used for amplicon sequencing (described in detail below). All leaf tissue handling was performed in a biosafety cabinet with all surfaces sterilized by exposure to UV light for 30 minutes and cleaned sequentially in between samples with 95% ethanol, 0.5% NaOCl and 70% ethanol to prevent cross contamination.

59 3.2 Amplicon sequencing

Leaf tissue in CTAB solution was stored for 2 months at room temperature prior to being placed at -80 C for 3 months before extracting DNA. In preparation for DNA extraction, we decontaminated all instru-

ments, materials, and surfaces with DNAway (Molecular BioProducts Inc., San Diego, CA, United States), 95% Ethanol, 0.5 % NaOCl, and 70 % Ethanol, and subsequently treated with UV light for 30 minutes in biosafety cabinet. We then transferred 0.2 - 0.3 g of leaf tissue into duplicate sterile 2mL tubes, resulting in 2 subsamples. Total genomic DNA from subsamples was extracted as described in U'Ren & Arnold (2017). In brief, we added two sterile 3.2 mm stainless steel beads to each tube and proceeded to lyophilize samples for 72 hours to fully remove CTAB content from tissue. After this period, we submerged the sample tubes in liquid nitrogen for 30s and proceeded to homogenize samples to a fine powder for 45 s in FastPrep-24 Tissue and Cell Homogenizer (MP Biomedicals, Solon, OH, USA). Afterwards, we repeated the decontamination procedure described before and used QIAGEN DNeasy 96 PowerPlant Pro-HTP Kit (U'Ren & Arnold, 2017) (QIAGEN, Valencia, CA, USA). After all genomic DNA was extracted, we pooled the subsamples for each individual sample before amplification. We used sterile equipment and pipettes with aerosol-resistant tips with filters in all steps before amplification. We followed a two-step amplification approach previously described by Sarmiento et al. (2017) and U'Ren & Arnold (2017). We used primers for the fungal ITSrDNA region, ITS1f (5'-CTTGGTCATTTAGAGGAAGTAA-3') and ITS4 (5'-TCCTCCGCTTATTGATATGC-3') with modified universal consensus sequences CS1 and CS2 and 0-5 bp for phase-shifting. Every sample was amplified in two parallel reactions containing 1-2 μL of DNA template (U'Ren & Arnold, 2017; see 77 also Tellez et al., 2022). We visualized PCR (PCR1) reactions with SYBR Green 1 (Invitrogen, Carlsbad, CA, USA) on 2% agarose gel (Oita et al., 2021). Based on the electrophoresis band intensity, we combined parallel PCR1 reactions and diluted 5 µL of amplicon product with molecular grade water to standardize to a concentration of 1:15 (Sarmiento et al., 2017 for details; Tellez et al., 2022). We included DNA extraction blanks and PCR1 negatives in this step. We used a separate set of sterile pipettes, tips, and equipment to reduce contamination. We used a designated PCR area to restrict contact with pre-PCR materials (Oita et al., 2021).

We used 1 µL of PCR1 product from samples and negative control for a second PCR (PCR2) with barcode

adapters (IBEST Genomics Resource Core, Moscow, ID, USA). Each PCR2 reaction (total 15 μ L) contained 1X Phusion Flash High Fidelity PCR Master Mix, 0.075 μ M of barcoded primers (forward and reverse pooled at a concentration of 2 μ M) and 0.24mg/mL of BSA following Sarmiento (2017) and U'Ren & Arnold (2017). Before final pooling for sequencing, we purified the amplicons using Agencourt AMPure XP Beads (Beckman Coulter Inc, Brea, CA USA) to a ratio of 1:1 following the manufacturer's instructions. The products were evaluated with Bio Analyzer 2100 (Agilent Technologies, Santa Clara, CA, USA) (Tellez et al., 2022). We quantified the samples through University of Arizona Genetics Core, and subsequently diluted them to the same concentration to prevent over representation of samples with higher concentration, see (CITATION). Amplicons were normalized to 1 ng/ μ L, then pooled 2 μ L of each for sequencing. No contamination was detected visually or by fluorometric analysis. To provide robust controls we combined 5 μ L of each PCR1 negative and the DNA extraction blanks and sequenced them as samples. Ultimately, we combined samples into a single tube with 20 ng/ μ L of amplified DNA with barcoded adapters for sequencing on the Illumina MiSeq platform with Reagent Kit v3 (2 × 300 bp) following protocols from the IBEST Genomics Resource Core at the University of Idaho, USA. Again, we included the DNA extraction blanks and two PCR1 negatives and sequenced with samples. Sequencing yielded 3,778,081 total ITS1 reads.

on 3.2.1 Mock Communities

We processed and sequenced 12 mock communities following the methods described above. This allowed us to assess the quality of our NGS data set. We used two mock communities that consisted of PCR product from DNA extractions of 32 phylogenetically distinct fungi, representing lineages that are typically observed as endophytes: Ascomycota, Basidiomycota, Zygomycota and Chytridiomycota (Oita et al., 2021; see Daru et al., 2019 for details). In brief, we used six mock communities with equimolar concentrations of DNA from all 32 fungal taxa and another six mock communities with tiered concentrations of DNA from the same fungal taxa (Daru et al., 2019). Each mock community was sequenced five times (i.e., five replicates) (Oita

et al., 2021). The read abundance from the equimolar and tiered communities was positively associated with the expected read number (with replicates as a random factor: R2Adj = 0.87, P = XXXX, see Supplementary Material). Using mock communities allowed us to evaluate the sequencing effectiveness in communities with known composition and structure (Bowman & Arnold, 2021). Henceforth, we used read abundance as a relevant proxy for biological OTU abundance (U'Ren et al., 2019).

14 3.2.2 Bioinformatic analyses

We used VSEARCH (v2.14.1) for de novo chimera detection, dereplication and sequence alignment. 115 VSEARCH is an open-source alternative to USEARCH that uses an optimal global aligner (full dynamic 116 programming Needleman-Wunsch), resulting in more accurate alignments and sensitivity (Rognes et al., 117 2016). For mock communities and experimental samples, we used forward reads (ITS1) for downstream bioinformatics analyses due to their high quality, rather than reverse reads (ITS4). Following Sarmiento 119 et al. (2017), we concatenated all reads in a single file and used FastQC reports to assess Phred scores 120 above 30 and determine the adequate length of truncation. We processed 892,713 of sequence reads 121 from mock communities and 3,778,081 from experimental samples. We truncated mock community and experimental sample reads to a length of 250 bp with command fast_trunclen and filtered them at a 123 maximum expected error of 1.0 with command fast_maxee. We then clustered unique sequence zero 124 radius OTUs (that is, zOTUs; analogous to amplicon sequence variants (Callahan et al., 2016)), by using 125 commands derep_fulllength and minseqlength set at 2. Sequentially we denoised and removed 126 chimeras from read sequences with commands cluster_unoise, and uchime3_denovo, respectively (see 127 Supplementary YYY for details). Finally, we clustered zOTUs at a 95% sequence similarity with command 128 usearch_global and option id set at 0.95. After which, 3,035,960 sequence reads from experimental samples remained.

Taxonomy was assigned with the Tree-Based Alignment Selector Toolkit [v2.2; Carbone et al. (2019)] by
placing unknowns within the Pezizomycotina v2 reference tree (Carbone et al., 2017). ITS sequences were
blasted against the UNITE database by the ribosomal database project (RDP) classifier. A total of 2147 OTUs
hits were obtained and are composed of 68.6% Ascomycota, 26.8% Basidiomycota, <0.05% Chytridiomycota,
<0.05% Glomeromycota, <0.05% Mortierellomycota, <0.05% Rozellomycota, 0.05% Kickxellomycota, and
4.2 % BLAST hit misses. Only OTUs representing Ascomycota were used for downstream statistical analyses since foliar endophyte communities in tropical trees are dominated by Ascomycota (Arnold & Lutzoni,
2007).

For each OTU identified, we removed laboratory contaminants from experimental samples by substracting the 139 average read count found in control samples from the DNA extraction and PCR steps. Our analysis of mock 140 communities allowed use to identify and remove false OTUs from experimental samples, those with fewer than 10 reads, and remove 0.1% of the read relative abundance across all samples (Oita et al., 2021). Removed reads represent the frequency of reads classified as contamination in the mock communities relative to the 143 expected read count. Three experimental samples from *Theobrona cacao* (n=2) and *Apeiba membranacea* (n=1) were removed from all analyses due to incomplete entries. After pruning taxa with zero reads from experimental samples, we identified 260 OTUs found exclusively in control (E-) plants (n=78) and deemed 146 them as artifacts resulting from the greenhouse conditions. Consequently, these were consistently eliminated 147 from treatment (E+) plants across all species. We converted reads for each fungal OTU to proportions of 148 total sequence abundance per sample to reduce differences in sampling effort, following previous studies (Weiss et al. (2017); McMurdie & Holmes (2014)). We then removed singletons and obtained an average 150 of 2,464,558 sequence reads in 529 Ascomycota OTUs across 156 experimental samples of 7 tree species. 151 All analyses post taxonomic assignment were performed in R [v. 4.3.2; R Core Team (2023)] using the 152 phyloseq package (McMurdie & Holmes, 2013) and custom scripts (see Supplementary Material).

4 3.2.3 Ant-endophyte interaction assays

A fresh fourth leaf was used in ant assays. To assess leaf-cutter ant damage, we introduced one detached leaf per plant per treatment to an actively foraging leaf-cutter ant colony for a two-hour assay. We presented leaf-cutter ant colonies with a choice of an E+ or an E- leaf on one disposable plastic plate next to an active nest trail. Carefully, we collected and placed debris from the trail leading up to the plate to lure ants into the plate. We initiated the ant assay as soon as one ant entered the plate and explored the leaf contents (for ~ 10-20 seconds). Every five minutes we took a digital photo of the choice arena until about 75% of the leaf content of one of the leaves was consumed. We used the digital photo at time zero and at the end of trial to quantify the leaf area removed using ImageJ [v1.52r; Schneider et al. (2012)]. Ant recruitment was estimated by counting individuals in the choice arena throughout trial event.

164 3.2.4 Pathogen assays

For the pathogen assays, we introduced an agar plug inoculated with hyphae of *Calonectria* sp. (P+ treatment), and an agar plug without the pathogen (P- control) to similarly aged/sized leaves within 10-14 days
after endophyte inoculations (CITATION). Leaves with the P+ or P- treatment were misted with sterile
water two times a day (morning and afternoon) to maintain moisture. After four days, we removed the plugs
and took digital photos to analyze leaf area damage using ImageJ [v1.52r; Schneider et al. (2012)].

3.2.5 Statistical Analyses

We explored how leaf functional traits and foliar fungal symbionts correlated to herbivory and pathogen damage on leaves. We present the analyses at the leaf and at the plant level. Leaf functional traits were measured and are presented in their raw form, at the leaf level, while FEF data was explored and is presented

at the plant level. In analyses where leaf functional traits and FEF are combined we used averages of the leaf functional traits.

To test for H2, we used a general linear mixed model (GLM) with XXXX as the response variable. To determine which fixed effects to include in the models we used the vif function in *R* to calculate the variance inflation factor for all explanatory variables (ACI, LT, LPS and LMA) (R Core Team, 2023). We then created a correlation matrix with cor function to assess correlations among covariates. We opted to maintain explanatory variables pertaining to physical barriers (LT, LPS and LMA) and exclude ACI from subsequent linear models due to high collinearity with LPS (0.54) and LMA (0.73). Every variable kept exhibits some degree of collinearity and this is well recorded in the literature (CITE HERE).

Additionally, Principal Component Analysis (PCA) was used to reduce dimensions among covariates and reveal underlying interactions that could influence fungal endophyte abundance, diversity and community composition in seedlings. The PCA was computed using the prcomp function in R statistical software (R Core Team, 2023). A complete PCA was computed with variables ACI, LT, LPS, and LMA (FIGURE 2a?). We then proceeded to compute a PCA with the data from leaves of plants used in the ant (n = 210) and pathogen assays (n = 192).

The PCA revealed how covariates (LMA, ACI, Thickness and Toughness) interact. I overlapped tree species groups on the PCA axes to show how the variance in the data is explained by PC1 (60%) and PC2 (27%) (Fig. 4). This is indicative of correlation among covariates. Thickness and toughness were orthogonal to each other in PCA, indicative of low correlation.

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- 5 Figures
- 253 **5.1 Figure 1**
- 254 **5.2 Figure 2**
- 255 **5.3 Figure 3**

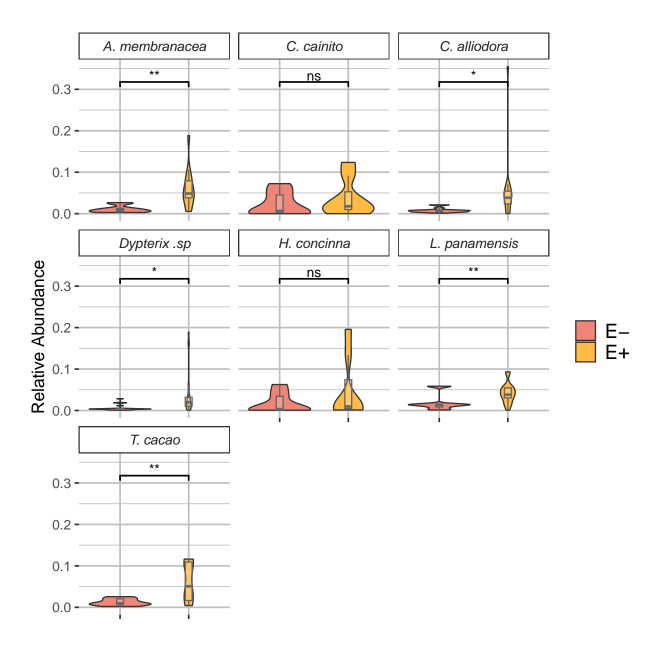


Figure 1: Relative abundance (RA) of Ascomycota OTUs of seven tree species used in the study. Violin plots show the distribution of the RA. The horizontal line within the embedded boxplots represents the median, the box represents the interquartile range (IQR), and the whiskers represent the 1.5xIQR. Outliers are represented by dots. Significance levels are represented by asterisks [p = 0.05 (*), p = 0.01 (*), and $p^* = 0.001$ (***)].

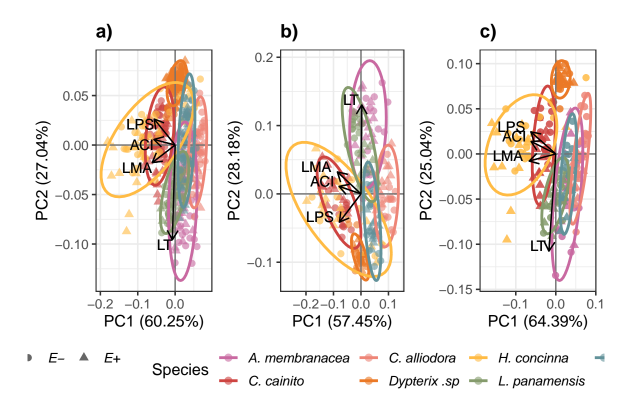


Figure 2: a) Principal Component Analysis (PCA) of leaf functional traits from all tree species separated by *E*- and *E*+ treatment. b) PCA of leaf functional traits of plants solely used in ant herbivory assays. c) PCA leaf functional traits of plants used solely in pathogen damage assays. Colors represent individual species. Circle and triangles represent *E*- and *E*+ treatments, respectively. Colored ellipses correspond to tree species and represent 95% confidence intervals.

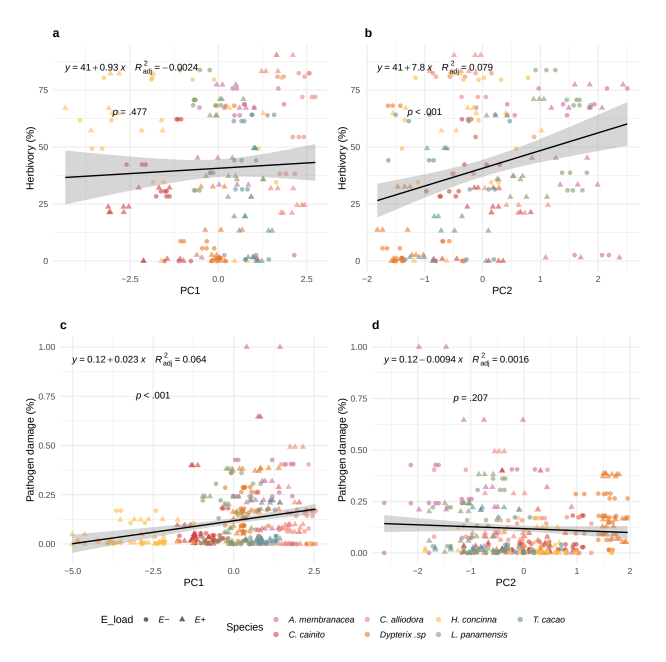


Figure 3: Simple linear regressions of herbivory and pathogen damage on PC1 and PC2 axes from PCA's of leaf traits for ant herbivory and pathogen damage assays. Linear regression of a) percent herbivory damage and PC1 axis (R^2 -adjusted= -0.0024, p=0.447); b) percent herbivory damage and PC2 axis (R^2 -adjusted= 0.079, p=<0.001); c) percent pathogen damage and PC1 axis (R^2 -adjusted= 0.064, p=<0.001); and d) percent pathogen damage and PC2 axis (R^2 -adjusted= 0.0016, p=0.207). Colors represent individual species. Circle and triangles represent E- and E+ treatments, respectively.