

# Leaf Mycobiome and Mycotoxin Profile of Warm-Season Grasses Structured by Plant Species, Geography, and Apparent Black-Stroma Fungal Structure

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ABSTRACT Grasses harbor diverse fungi, including some that produce mycotoxins or other secondary metabolites. Recently, Florida cattle farmers reported cattle illness, while the cattle were grazing on warm-season grass pastures, that was not attributable to common causes, such as nutritional imbalances or nitrate toxicity. To understand correlations between grass mycobiome and mycotoxin production, we investigated the mycobiomes associated with five prominent, perennial forage and weed grasses [Paspalum notatum Flügge, Cynodon dactylon (L.) Pers., Paspalum nicorae Parodi, Sporobolus indicus (L.) R. Br., and Andropogon virginicus (L.)] collected from six Florida pastures actively grazed by livestock. Black fungal stromata of Myriogenospora and Balansia were observed on P. notatum and S. indicus leaves and were investigated. High-throughput amplicon sequencing was applied to delineate leaf mycobiomes. Mycotoxins from P. notatum leaves were inspected using liquid chromatography-mass spectrometry (LC-MS/MS). Grass species, cultivars, and geographic localities interactively affected fungal community assemblies of asymptomatic leaves. Among the grass species, the greatest fungal richness was detected in the weed S. indicus. The black fungal structures of P. notatum leaves were dominated by the genus Myriogenospora, while those of S. indicus were codominated by the genus Balansia and a hypermycoparasitic fungus of the genus Clonostachys. When comparing mycotoxins detected in P. notatum leaves with and without M. atramentosa, emodin, an anthraquinone, was the only compound which was significantly different (P < 0.05). Understanding the leaf mycobiome and the mycotoxins it may produce in warm-season grasses has important implications for how these associations lead to secondary metabolite production and their subsequent impact on animal health.

**IMPORTANCE** The leaf mycobiome of forage grasses can have a major impact on their mycotoxin contents of forage and subsequently affect livestock health. Despite the importance of the cattle industry in warm-climate regions, such as Florida, studies have been primarily limited to temperate forage systems. Our study provides a holistic view of leaf fungi considering epibiotic, endophytic, and hypermycoparasitic associations with five perennial, warm-season forage and weed grasses. We highlight that plant identity and geographic location interactively affect leaf fungal community composition. Yeasts appeared to be an overlooked fungal group in healthy forage mycobiomes. Furthermore, we detected high emodin quantities in the leaves of a widely planted forage species (*P. notatum*) whenever epibiotic fungi occurred. Our study demonstrated the importance of identifying fungal communities, ecological roles, and secondary metabolites in perennial, warm-season grasses and their potential for interfering with livestock health.

**KEYWORDS** mycotoxin, mycobiome, warm-season grasses, forage, fungi, endophytes, hypermycoparasite, *Myriogenospora*, *Balansia*, *Clonostachys* 

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Plants harbor a diverse array of microorganisms, including species from the kingdom Fungi (1). The leaf surface and interior provide habitats for epibiotic and endophytic fungi, respectively. While often not causing observable symptoms to the plant hosts, the impacts of these leaf-inhabiting fungi on their host plants are dynamic, ranging from beneficial to neutral to detrimental (2).

Of all leaf-inhabiting fungi, the role that C3 grass-associated endophytes (for example, *Epichloë* spp.) play has been well characterized. These vertically transmitted fungi obtain photosynthates from the plant and produce toxins which deter herbivores (2). While fungally derived metabolites can protect plants against herbivores, they also sometimes cause severe health problems when consumed by livestock (3). In temperate regions where tall fescue (*Festuca arundinacea* Schreb.) grass is a major pasture forage species, fescue toxicosis is a significant problem. This disease is the result of ingestion of ergot alkaloids released by *Epichloë coenophiala* in grazing livestock. The U.S. beef cattle industry alone loses an estimated \$2 billion annually from fescue toxicosis-related sequelae, including abortion and weight loss (4).

Perennial, warm-season, C4 grasses contribute significantly to feeding livestock in the southern regions of the United States (5). Considering nutritive value and soil adaptability, a variety of perennial grass forage species are grown across Florida; however, *Paspalum notatum* Flügge (bahiagrass), which is native to South America (approximately -20 to -35 degrees latitude), is one of the most important perennial cultivated forage species. It is easy to establish, can be used for soil stabilization, has good persistence, requires minimal maintenance, and has low susceptibility to diseases and pests (6). Cultivars 'Pensacola' and 'Argentine' are among the most widely planted varieties, but several improved cultivars are increasingly being planted (7).

Depending upon pasture management, less desirable grasses can contaminate pastures and are regarded as weeds. For example, *Paspalum nicorae* Parodi (Brunswickgrass) is considered a weed, since it is much less palatable to cattle and can contaminate *P. notatum* seed production fields (8). Originally from tropical Asia, *Sporobolicus indicus* (smutgrass) is regarded as one of the most troublesome pasture grass weeds (9) since it is adapted to a variety of soil types and often outcompetes other pasture forages (10). *S. indicus* is considered a weed grass due to its low nutritive value and unpalatable nature (11). Although native to the Americas, *Andropogon virginicus* (broomsedge) has infiltrated millions of acres of pastureland across the southeastern United States (12), and it competes for resources with preferred forage grasses (9, 11). *S. indicus* and *A. virginicus* often coinhabit *P. notatum* pastures and, while less palatable, can occasionally be consumed by livestock.

Unlike C3 grasses, perennial, warm-season, C4 grasses have had little delineation of their leaf-associated endophytic and epibiotic fungal communities. Recently, Depetris et al. (13) reported various culturable endophytes from Paspalum spp. and demonstrated their positive effects on plant host performance. Many fungi of the Clavicipitaceae family can live as endophytes inside asymptomatic plant tissues (defined here as apparently healthy plant tissues without visible fungal structures) for part or all of their life and may occasionally produce visible reproductive structures on the plant surface (14). While many plant-inhabiting fungi are regarded as endophytes in the Clavicipitaceae, some members of this family adopt an epibiotic existence whereby they live superficially on the exterior of host plants (15). Ergot alkaloids, a group of mycotoxins often produced by clavicipitaceous fungi, have been reported from several C4 and C3 grass species (16). For example, ergot alkaloids were detected in situ from three Balansia epichloë-infected grass leaves, including Sporobolus from the southeastern United States (17). Compared to clavicipitaceous fungi living within C3 grasses, genera of Clavicipitaceae that occur on C4 grass leaves (including Myriogenospora and Balansia) remain relatively understudied (18, 19). Besides the mostly endophytic genus Balansia, the epibiotic genus Myriogenospora has been reported multiple times within warm-season grasses, including P. notatum; it forms black stromata superficially on the plant leaves (defined as symptomatic here), causing a "tangle-top" symptom (18, 20). Despite its frequent occurrence and phylogenetic similarity to other mycotoxinproducing fungi, the ability of Myriogenospora to produce mycotoxins is unknown (21).

Understanding the potential factors that influence mycotoxin production in pastures is critically important for plant and animal health.

Phyllosphere microbiomes, including the epibiotic and endophytic microbial communities, can be impacted by various abiotic and biotic factors, including plant host genetics (e.g., species and genotype), plant traits (e.g., leaf chemical composition), management (e.g., chemical applications), seasonality, and geographic location (22-26). Core microbial members commonly present across multiple plant species and geologic areas may reflect their indispensable interactions with plants (27, 28). In contrast to the core microbes, indicator microbial species found abundantly in particular plants may reflect specific associations and adaptations of certain plant-microbe pairs (25). Overall, by affecting plant traits and plant-microbial interactions, phyllosphere microbiomes can have a direct impact on and potential application for use in agricultural and environmental health (27, 29, 30). Culture-based detection methods have revealed a high abundance of fungi belonging to Ascomycota that are associated with warmseason grasses (13). Recently, culture-independent methods, which can detect more complete assemblies of fungi, have been applied to the biofuel crop Panicum virgatum L. (switchgrass), a warm-season, C4 grass. By investigating the mycobiome of *P. virgatum* grown in temperate fields, the grass genotype and seasonality were revealed to significantly affect leaf mycobiome assemblies (31, 32). Additionally, leaf mycobiome had intricate interactions with the bacterial communities living therein (32). However, warm-season grasses growing in warmer regions have not been subjected to culture-independent mycobiome detection. Climate change and anthropogenic activities, including agrotechnology, have raised the concerns of changing plant-fungal interaction and the emergence of new diseases (33, 34). These knowledge gaps call for investigations in the mycobiome and potential mycotoxin production in the perennial, warm-season grasses.

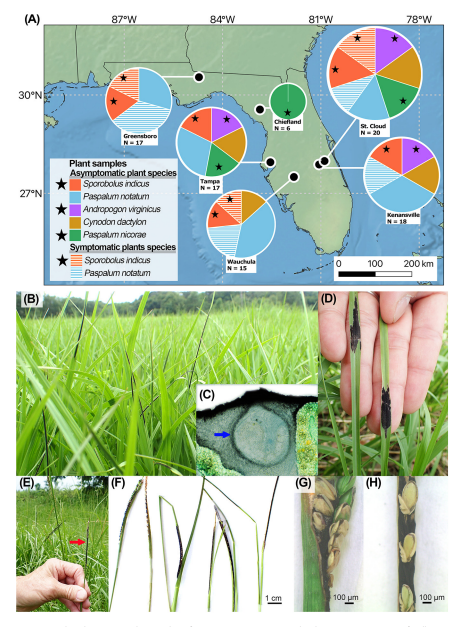
To further our understanding of the effect that plant taxonomy (i.e., species and cultivar) has on microbial community and mycotoxin profiles of warm-season grass-associated fungi across different Florida locations, we investigated the leaf-inhabiting fungal community of two improved perennial forage species, *P. notatum* and *Cynodon dactylon*, and three weed species, *P. nicorae*, *S. indicus*, and *A. virginicus*, which often contaminate improved forage pastures and hay fields across six locations in Florida. Our objectives were to (i) investigate the fungal community assemblies across Florida pastures for cooccurring forage and weed species using high-throughput amplicon sequencing, (ii) cross-compare the foliar fungal communities between grass samples both with and without obvious fungal stromata in *P. notatum* and *S. indicus*, and (iii) detect mycotoxins present in the important forage *P. notatum* via liquid chromatography-tandem mass spectrometry (LC-MS/MS). Our results revealed the core and unique fungal communities across warm-season pastures and the mycotoxin composition of one critical forage species (*P. notatum*) in Florida, providing key insights relevant to plant and cattle health, pasture management, and the emerging concern and potential impact of mycotoxins on livestock production.

#### **RESULTS**

In addition to being on the leaf, black fungal structures were also associated with the inflorescence of *P. notatum* (Fig. 1E to H) and were confirmed to be *Myriogenospora atramentosa* by its nuclear ribosomal internal transcribed spacer (nrITS) region (Sanger sequencing, GenBank accession numbers ON678271 to ON678272; specimen code at the University of Florida Herbarium, FLAS-F-63886). DNA amplicon sequencing that targets fungal ITS1 was further applied to identify and compare the mycobiomes living in and on symptomatic and asymptomatic leaf samples. Fungal ITS1 metabarcoding resulted in merged read numbers ranging from 9,805 to 54,194 per grass sample, yielding 989 amplicon sequence variants (ASVs).

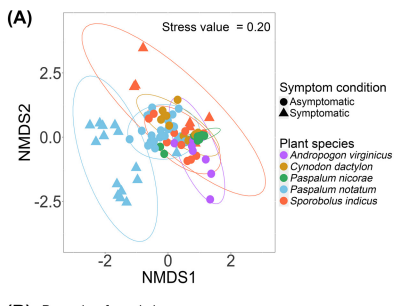
Comparison of the fungal composition in asymptomatic with that in symptomatic tissues. The mycobiome of grass samples bearing black fungal stromata, especially symptomatic P. notatum, was distinct from that of the asymptomatic ones (Fig. 2A). In the asymptomatic leaves of P. notatum and S. indicus, taxa in the class Dothideomycetes (S. indicus, S1.1%; S1.1%; S2.1.1%; S3.1% had the greatest abundance, followed by Tremellomycetes (S3. indicus, S3.1% to S4.1%; S5. indicus, S4.2%; S5. indicus, S7. notatum, S8.3% (Fig. 2B), S8.3% (Fig. 2B), S9. indicus, S9. indicus,

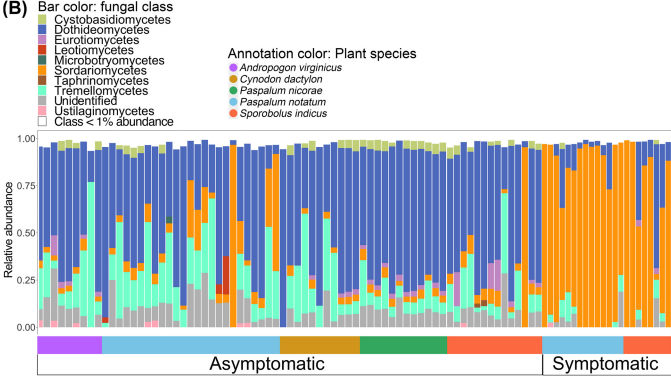
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**FIG 1** Sampling locations and examples of symptomatic grasses in Florida pastures. (A) Map of collection sites across Florida, USA. Each pie chart corresponds to the sampling of a given site. The color indicates plant species, and the pattern indicates symptom pathology. The size of the pie chart corresponds to the number of samples at each site. Stars indicate plant species considered weeds. (B) *Paspalum notatum* leaves with black fungal structures produced by *Myriogenospora atramentosa*. (C) Cross section of *P. notatum* leaf bearing the superficial fungal structure of *M. atramentosa*. Blue arrow identifies the black *M. atramentosa* perithecium. The surrounding green part is *P. notatum*'s leaf tissue. (D) *Sporobolus indicus* leaf blades with fungal structures produced by *Balansia epichloë*. (E to H) *M. atramentosa* on the seed head of *P. notatum* (red arrow).

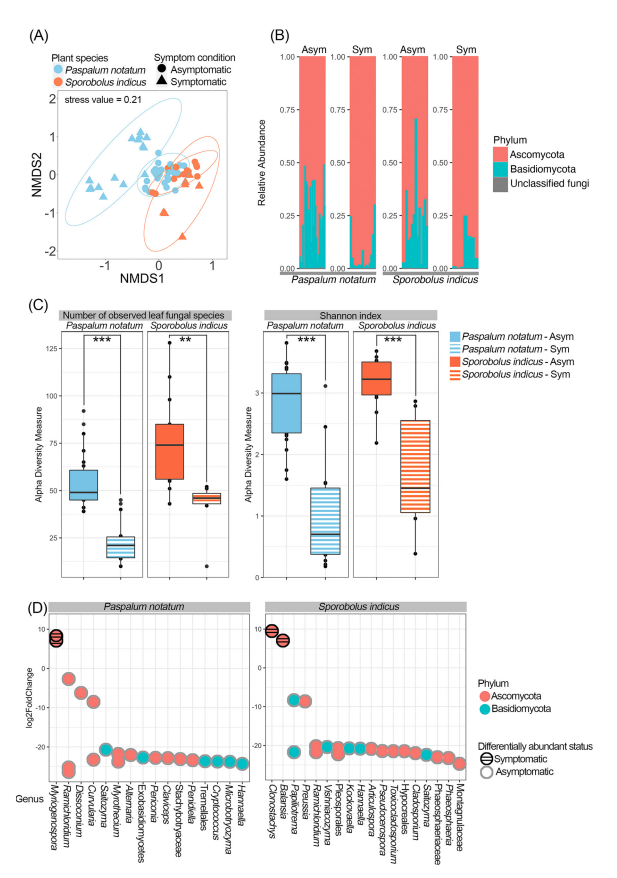
regardless of plant species. For both plant species with symptoms, Sordariomycetes were the most prevalent fungal class (*S. indicus*,  $62.8\% \pm 34.9\%$ ; *P. notatum*,  $82\% \pm 18.8\%$ ), compared to other classes living in and on the same tissue (Fig. 2B). Both *P. notatum* (P=0.001) and *S. indicus* (P=0.006) hosted fungal communities distinct between symptomatic and asymptomatic samples (Fig. 3A), with a greater shift of fungal communities in *P. notatum* than in *S. indicus*. The relative abundance of Basidiomycota was significantly greater in asymptomatic (P=0.001) and P=0.0010. Indicus = P=0.0011 has in symptomatic (P=0.0011) and P=0.0012. Indicus = P=0.0013 leaves for both P=0.0013 and P=0.0014 and P=0.0014





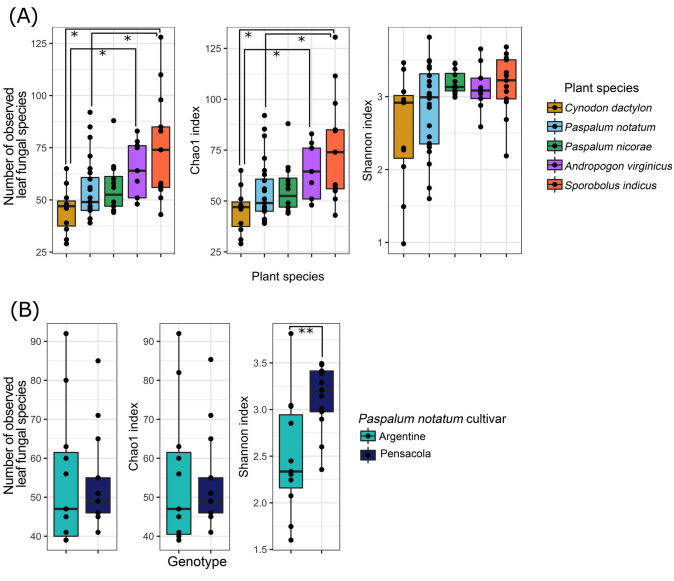
**FIG 2** (A) Nonmetric multidimensional scaling (NMDS) plot of fungal beta-diversity (Bray-Curtis dissimilarity) of all grass samples. (B) Stack barplots of fungal community at class level for asymptomatic and symptomatic samples of the five forage grass species evaluated.

0.046) (Fig. 3B). Despite being relatively more abundant in symptomatic samples, the phylum Ascomycota dominated the grass-inhabiting fungal communities (Fig. 3B) regardless of the symptom condition (asymptomatic, P.  $notatum = 93.2\% \pm 9.3\%$ , S.  $indicus = 91.3\% \pm 9.5\%$ ; symptomatic, P.  $notatum = 76.2\% \pm 15.1\%$ , S.  $indicus = 77.9\% \pm 17.7\%$ ). Fungal richness (observed number of species and Chao1, S. indicus, P = 0.002; P. notatum, P < 0.001) and Shannon index (S. indicus and P. notatum both with P values of <0.001) were greater in asymptomatic leaves than in symptomatic leaves (Fig. 3C). Two ASVs, both assigned to Myriogenospora, were significantly more abundant (false-discovery rate [FDR] < 0.05) in symptomatic P. notatum (Fig. 3D). A Clonostachys ASV classified as Clonostachys miodochialis and a Balansia ASV were also



**FIG 3** (A) Nonmetric multidimensional scaling (NMDS) plot of fungal beta-diversity (Bray-Curtis dissimilarity) of the symptomatic and asymptomatic *Paspalum notatum* and *Sporobolus indicus*. (B) Stack barplots of fungal phyla in asymptomatic and symptomatic *P. notatum* and (Continued on next page)

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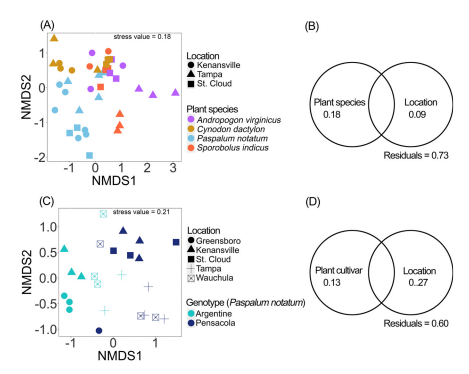
**FIG 4** (A) Fungal alpha-diversity of asymptomatic grasses across plant species. (B) Fungal alpha-diversity of asymptomatic grasses of the two *Paspalum notatum* cultivars 'Argentine' and 'Pensacola'. \*, P < 0.05; \*\*, P < 0.01.

significantly more abundant in symptomatic *S. indicus* samples (Fig. 3D). A total of 19 ASVs spanning Ascomycota and Basidiomycota were more abundant in the asymptomatic samples of *P. notatum* and *S. indicus* (Fig. 3D).

Fungal diversity and composition in and on asymptomatic leaves across grass species, cultivar, and locality. We then focused on asymptomatic grass samples to assess factors impacting mycobiomes. We estimated the number of observed species, Chao1 index, and Shannon index of asymptomatic tissues of each plant species. Fungal diversity was not the same across a given plant species (number of observed species, P = 0.002; Chao1, P = 0.002, Shannon index, P = 0.04). Of all pairwise plant species comparisons, significant differences in fungal richness were observed in three pair-species comparisons (Fig. 4A). No significant differences in Shannon diversities of leaf fungi were found across all of the pair-plant species comparisons (Fig. 4A). Among the five examined grass species, *S. indicus* had the greatest

## FIG 3 Legend (Continued)

S. indicus. Each bar corresponds to one sample. (C) Diversity measurements of asymptomatic versus symptomatic plants of P. notatum and S. indicus. (D) Significantly differentially abundant ASVs (Wald test, false-discovery rate [FDR] of <0.05) between symptomatic and asymptomatic P. notatum and S. indicus. Every circle corresponds to one ASV. ASV, amplicon sequence variant \*\*, P < 0.01; \*\*\*, P < 0.001.



**FIG 5** (A) Nonmetric multidimensional scaling (NMDS) plot of fungal beta-diversity (Bray-Curtis dissimilarity) of four asymptomatic grasses at three locations. (B) Variation partitioning of plant species and location effects on mycobiome composition. (C) Nonmetric multidimensional scaling (NMDS) plot of fungal beta-diversity (Bray-Curtis dissimilarity) of two *Paspalum notatum* cultivars. (D) Variation partitioning of *P. notatum* cultivar and location effects on mycobiome composition.

fungal diversity richness (Fig. 4A). This trend was observed across all sampling sites except for Kenansville, where the observed fungal species in *Andropogon virginicus* exceeded those in *S. indicus*.

The leaf mycobiomes of both P. notatum and S. indicus revealed similar richnesses and Shannon diversities across sampling sites (P > 0.05). The mycobiome of P. notatum was assessed for plant cultivar effect. While similar observed numbers of species and Chao1 measurements were detected for the two cultivars, the Shannon index suggested a significantly higher diversity of leaf fungi present in 'Pensacola' than in 'Argentine' (Fig. 4B).

To assess the plant identity and geography effects on mycobiome composition, we focused on three locations with four grass species sampled to avoid sampling bias. Plant species and location had an interactive effect on the beta-diversity of fungal communities living in and on grass leaves (permutational multivariate analysis of variance [PERMANOVA], plant species, location, and plant species-location had a P value of <0.01) (Fig. 5A). Therefore, these two factors were assessed separately. PERMANOVA suggested a significant difference for fungal beta-diversity among plant species (P, St. Cloud = 0.03, Kenansville = 0.01, and Tampa = 0.01) (Fig. 5A). Additionally, PERMANOVA suggested significant differences of fungal beta-diversity across locations for P. notatum and S. indicus (P < 0.001 for both plant species). Variance partitioning analysis revealed a slightly greater plant species effect (0.18) than location effect (0.09) (Fig. 5A and B). In comparison, the two P. notatum cultivars had significant cultivar, location, and cultivar-location effects (PERMANOVA, cultivar, location, and cultivar-location all with P values of <0.01). Variance partition analysis detected a greater impact of location effect (0.27) than cultivar effect (0.13) on mycobiome assembly (Fig. 5C and D).

Fungal indicators and core mycobiome of different plant species using asymptomatic grass samples. Asymptomatic leaf mycobiomes were investigated to find fungal members unique to certain grass species as "indicators" or present across different grasses as "cores" in the Florida pastures. Indicator ASVs are defined as ASVs found significantly more often with one plant species than with others. Among the four plant species, *S. indicus* had the highest number of indicator ASVs (60) followed by *A. virginicus* (28). *C. dactylon* and *P. notatum* only had 14 and 11 indicator ASVs, respectively. These ASVs were distributed across 48

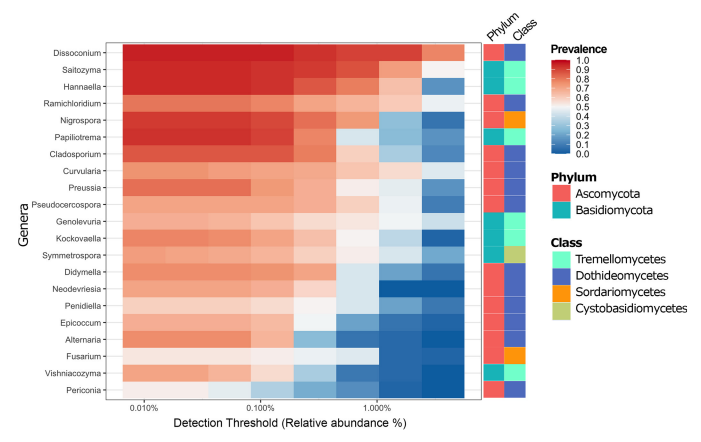


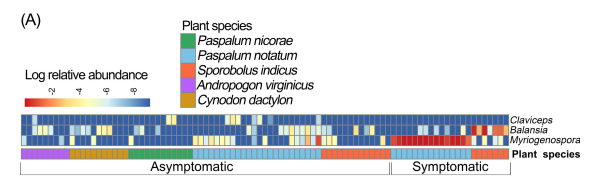
FIG 6 Core fungal genera (i.e., relative abundance of >0.01% in more than half of the samples) across all asymptomatic plant species.

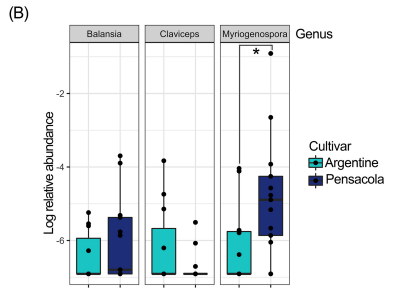
genera (see Fig. S1 in the supplemental material). Only 33 were identified to include a single ASV as an indicator. Seven ASVs belonging to the genus *Ramichloridium* were identified as indicator ASVs in *S. indicus*, *P. notatum*, and *C. dactylon*.

To identify the core mycobiome shared across plant species, a genus with at least 0.01% relative abundance in over half of the asymptomatic samples was defined as a core genus of the mycobiome. Twenty-one genera were detected as being part of a core mycobiome within the asymptomatic leaf samples, with 12 being in the class Dothideomycetes and six in the Tremellomycetes, followed by two in the Sordariomycetes and one in the Cystobasidiomycetes (Fig. 6). All seven Basidiomycota fungal genera detected as core members across Florida pastures have yeast forms, highlighting the potential importance and ubiquity of yeast fungi in pasture grasses (Fig. 6).

Distribution of Clavicipitaceae fungi across plant species and locations. Because Clavicipitaceae are of interest in the grass mycobiome and could be major mycotoxin producers, we examined their relative abundance across symptomatic and asymptomatic leaves. A total of three genera (Myriogenospora, Balansia, and Claviceps) of the family Clavicipitaceae were detected from all grass samples (Fig. 7A), while symptomatic P. Notatum and S. Notatum and Notatian a greater abundance of Notatian Notatian and Notatian Notatian and Notatian Notatian

**Mycotoxin diversity and quantity in** *Paspalum notatum.* Forty-five metabolites (see Table S1 in the supplemental material) were examined for the widely planted forage *P. notatum.* The goal was to evaluate if the epibiotic *M. atramentosa* is responsible for mycotoxin production. Six mycotoxins (alternariol methyl ether, beauvericin,





**FIG 7** (A) Abundance of fungal genera of Clavicipitaceae across Florida pastures. (B) Clavicipitaceae of cultivars 'Argentine' and 'Pensacola' of the asymptomatic *Paspalum notatum*. \*, P < 0.05.

dihydrolysergol, beta-zearalenone, emodin, fumonisin B1) were detected in at least one leaf sample of P. notatum (Fig. 8). Comparing these metabolites among mixed, symptomatic, and asymptomatic samples of P. notatum, emodin was significantly greater in symptomatic samples than in mixed or asymptomatic samples (P = 0.01). This suggests that the black fungal stromata associated with the symptomatic samples were associated with emodin production.

#### **DISCUSSION**

By using a culture-independent, high-throughput amplicon sequencing approach, we showed that the community of leaf fungi was structured by a combination of factors including plant species, cultivar, and sampling location, as well as the presence of fungal structures on leaves (symptomatic versus asymptomatic leaves). An unexpectedly high relative abundance of basidiomycetous yeasts was detected from the asymptomatic grasses, highlighting their overlooked ecological importance in warm-season grasses. The metabolite emodin was significantly more abundant in *P. notatum* with the epiphytic fungus *Myriogenospora atramentosa*. Our study underlines the complexity of the fungal community associated with Florida forages and begins to elucidate its ecological function, potential concern, and opportunity.

Fungal community assembly and diversity of warm-season grasses. Fungal communities of plants are often affected by complex factors. Similar to a previous study showing a more significant effect of grass host species than of geological factors like elevation (35) on the composition of the fungal community, our results suggest that leaf mycobiomes are likely manipulated by grass species and are less affected by

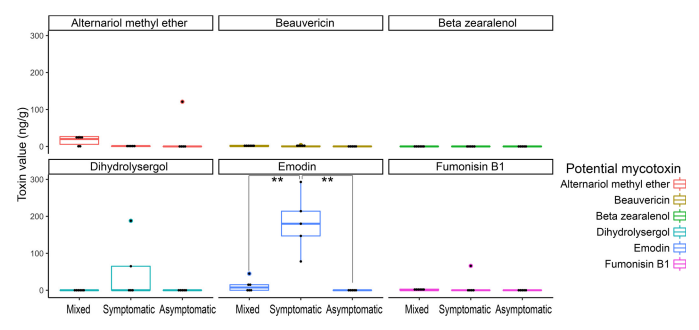


FIG 8 Six mycotoxins with nonzero values in at least one sample of Paspalum notatum. \*\*, pairwise Wilcox test, false-discovery rate (FDR) of <0.01.

specific grass cultivars and regional-scale location. Studies of temperate grasses also revealed complex factors involved in shaping the foliar mycobiome. In prairie systems, seasonality, geographic separation, and host species played major roles in shaping the foliar grass mycobiome (36). In grassland farms, the mycobiome of different grass species responded distinctively to land management and within-field microclimate (37, 38). Of the five grass species investigated in this study, the two weedy species S. indicus and A. virginicus harbored significantly higher richness of leaf fungi than did the two forage grasses. Fungal richness may be determined by the outcome of plant-fungus interactions, especially for plant defense responses. Foliar endophytes have been shown to promote plant resistance to biological and environmental stress (39, 40) and to promote increased growth rates (41), leading to higher competitiveness with local plants. Future research can manipulate the fungal richness of these grasses and test for their competitiveness in warm-season pastures. In comparing the leaf mycobiomes between two P. notatum cultivars, despite no richness differences detected, 'Pensacola' P. notatum had a significantly higher Shannon diversity of leaf fungi, which measured both richness and evenness (Fig. 4B). Ploidy differences of 'Argentine' (tetraploid) and 'Pensacola' (diploid) (7) could lead to plant trait differentiation and may result in recruiting and retaining different groups and a diversity of mycobiomes through plant traitfungal trait interactions (42).

Compositional differences of the fungal community in symptomatic versus asymptomatic samples were more significant in *P. notatum* than in *S. indicus* (Fig. 3A), suggesting that *M. atramentosa* and *B. epichloë* led divergent mycobiome responses. Interestingly, along with the black-stroma producer genus *Balansia*, an ASV of the genus *Clonostachys* also had higher abundance in symptomatic leaves. *Clonostachys* was previously reported as a hypermycoparasite on *Epichloë typhina* (43), a species that belongs to the same family as *Balansia*. *Clonostachys* has biocontrol features to deter phytopathogenic fungi and a high resistance to mycotoxins produced by *Fusarium* (44). The host-parasite association between *Clonostachys* and Clavicipitaceae has yet to be elucidated. Hypermycoparasites are often neglected in studies focusing on plant-associated fungi. Our finding highlights that hypermycoparasites should be considered for fungal metabolite accumulation and hold potential for bioprospecting fungal discovery.

**The dominant fungal taxa living in/on warm-season grasses.** Our results revealed high fungal diversity for those samples from apparently healthy (asymptomatic) grasses. Studies of grass-associated endophytic fungi, especially those detected by culture-based methods, often detected Ascomycota only (13, 45). Culture-independent methods recovered more

tremellomycetous (Basidiomycota) fungi (26, 46), likely due to their low rates of growth or inability to grow on artificial media. While fungi of the phylum Ascomycota dominated the community assembly here, a surprisingly high Basidiomycota biodiversity was detected (Fig. 2). Fungal members of Ascomycota and Basidiomycota probably interact in the phyllosphere. For example, Tremellomycetes (Basidiomycota) and Dothideomycetes (Ascomycota) have been reported to increase together in flower tissues during the ripening stage of wheat (46) and were detected as "hubs" of grass-mycobiome networks (31), suggesting possible cooperation of these two classes in the natural environment. Diverse reproduction strategies have been observed across different taxa of Tremellomycetes: some taxa produce complex fruiting bodies, some exist in the yeast form, and some are dimorphic (47). The core genera detected here (e.g., Saitozyma and Hannaella, Table S2, Fig. 6) prefer the yeast form. Although basidiomycetous yeasts are occasionally isolated from plants and soil (48), the roles this fungal group plays in fungal-plant interactions, soil biology, transmission, and ecology are largely unexplored. Some yeasts, including Cryptococcus of Tremellomycetes, have been proposed as agents for biocontrol against phytopathogens (49-51). Among dothideomycetous lineages, Dissoconium and Ramichloridium were the core genera detected in the asymptomatic leaves for all of the grass species sampled in this study (Fig. 6). Genomic analysis of one Ramichloridium species revealed its potential mechanism to be well adapted to extreme environments, including those with long periods of desiccation, low nutrients, and high solar intensity on the leaf surface (52). In addition to being the core genus, Ramichloridium had seven ASVs detected as indicators of three plant species (see Fig. S1 in the supplemental material), suggesting members of Ramichloridium had specific interactions with individual plant species. Dissoconium and Ramichloridium were both isolated from C4 grasses previously and demonstrated antifungal abilities (53, 54). Members of the tremellomycetous yeasts and Dothideomycetes were also identified as core in Panicum virgatum grown in a temperate climate (31). Together with our finding, tremellomycetous yeasts and Dothideomycetes are the core mycobiome of warm-season grasses regardless of the climate factor, highlighting their potential benefit to plants as fungal generalists. Future studies should be carried out to understand their ecology (e.g., trophic mode), physical association (e.g., epibiotic or endophytic), potential impact on plant host development, and fitness, as well as antagonistic effects on other organisms (54).

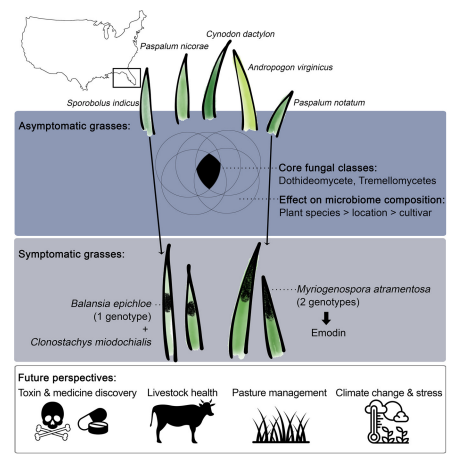
Culture-dependent and culture-independent methods can detect complementary microbial members of grass endophytes and epibionts (55, 56). For example, in *Paspalum*, the eurotiomycetous fungi such as *Penicillium* and *Aspergillus* were detected via culture-dependent methods (13), yet neither of these two genera was relatively abundant in the *P. notatum* samples investigated in this study using culture-independent methods. However, the culture-independent methods are generally thought to capture more comprehensive mycobiomes, especially those fungi that engage in interactions with plants or other microbial consortia (57).

Mycotoxins in Paspalum notatum and Myriogenospora. Plant-associated fungi can produce a chemically diverse variety of mycotoxins. These compounds can benefit the plant by making it less palatable, or even toxic, as a defense mechanism against grazing herbivores (58), thus allowing infected plants to thrive. Some of the best known mycotoxicoses in livestock are fescue foot and ryegrass staggers caused by clavicipitaceous fungi in cool-season grasses (59). Alkaloids, especially indole-diterpenes and ergot alkaloids, are the main types of toxins that cause the clinical signs seen in these syndromes (14). One ergot alkaloid precursor, dihydrolysergol, was detected in two out of the five symptomatic P. notatum samples (Fig. 8). Because the quantity of dihydrolysergol was not significantly different among symptom statuses, the black stroma (M. atramentosa) was unlikely to be the main source of dihydrolysergol production. If accidentally consumed by livestock, these toxins could raise health concerns, including average daily gain effects, reduced reproductive capacity, and hyperthermia (Table S3) (59-61). Such symptoms have recently been observed and were of concern in cattle operations in the southern part of the United States; however, the causal agents require future investigation (62). Albeit at a much lower level than on the symptomatic leaves, fungi of Clavicipitaceae were present across Florida pastures, even in asymptomatic plants (Fig. 7A).

This is unexpected for *Claviceps* and *Myriogenospora* as the former is an ovarian parasite and the latter is a leaf epibiont. According to an inoculation experiment, *Claviceps* can infect leaves, suggesting that a *Claviceps*-leaf association is possible (63). To our knowledge, *Myriogenospora* has not been confirmed microscopically inside the plant tissue as an endophyte. *Myriogenospora* reads of asymptomatic samples might be derived from early fungal infection without visible signs of fungi.

Although fungi of Clavicipitaceae such as Balansia epichloë that were detected in the samples of this study are well known for producing alkaloids, other fungi in the Dothideomycetes as well as Sordariomycetes, such as the core genera Alternaria and Fusarium, are also capable of producing other types of mycotoxins (Fig. 6) (17, 64). Toxicity symptoms could be triggered by changing environmental conditions such as fluctuation of temperature (65, 66). Previous research reported high zearalenone in Paspalum notatum (67), but similar concentrations were not recovered here (Fig. 8). Instead, we detected alternariol methyl ether in P. notatum (Fig. 8). Alternariol methyl ether was mostly produced by fungi of Alternaria (68), one of the core fungal genera examined here across Florida pastures. Finally, given the high consistency of emodin detected in grass samples with Myriogenospora atramentosa fruiting bodies (Fig. 8) and the fact that emodin can be produced by both fungi and plants (69), it is likely that M. atramentosa is responsible for producing or triggering the plant to produce emodin in vivo within P. notatum. Emodin is an anthraquinone also found in rhubarb which has been shown to have largely beneficial properties, including antimicrobial abilities, herbivore resistance, and properties of medicinal value (e.g., candidate as an antitumor drug) (69-71), highlighting the bioprospecting potential of this forage-associated mycotoxin. There is evidence that emodin can be hepatotoxic and nephrotoxic and lead to reproductive toxicity with high doses and chronic use (70), however, so caution must be exercised as a more complete picture surrounding the potential therapy/toxicity tradeoffs of this compound emerges. While previous research indicated seed heads from the cultivar 'Argentine' to be more susceptible than 'Pensacola' to ergot caused by Claviceps paspali (72, 73), we detected similar relative abundances of *Claviceps* in the leaves of both cultivars (Fig. 7B).

The fungal family Clavicipitaceae contains many plant-associated fungi capable of producing ergot alkaloids, especially in the Clavicipitaceae clade A (74) which includes Verticillium, Balansia, Epichloë, Claviceps, and Myriogenospora. Because clavicipitaceous fungi's main hosts are Poaceae, morning glories and sedges which are of essential use for humans in developing agricultural systems, these fungi and their mycotoxins have been widely studied (75). However, Myriogenospora represents an exception. Despite the wide host range of Myriogenospora on forage grasses and on crops such as sugar cane and lemongrass (19-21, 76), its capability of producing secondary metabolites including mycotoxins is unknown. In this study, ergot alkaloids were not detected in Myriogenospora-infected grass, suggesting a loss of function in the evolutionary history of the Clavicipitaceae clade A. Atkinsonella hypoxylon (75), an epibiotic fungus closely related to Myriogenospora, was predicted to produce ergot alkaloid according to its genome content. However, ergot alkaloid was undetectable in plant tissues infected by A. hypoxylon. The mycotoxin profile potentially produced in affected host plants needs to be more thoroughly studied by sampling M. atramentosa in vivo with different plant species or in vitro by culturing it in fungal medium as well as sampling across seasons. Other types of alkaloids, including indole-diterpenes, which cause livestock illness, should be investigated to eliminate food safety and animal health concerns of Myriogenospora-infected plants. Two different genotypes of M. atramentosa, confirmed with MiSeq and Sanger sequencing, were detected in our sampling (Fig. S2), agreeing with the work of Glenn et al. (19) that cryptic biodiversity is present within this fungal species. While many plant-associated clavicipitaceous fungi can transmit vertically with seeds, such as Claviceps and Epichloë, M. atramentosa is not known to be specifically vertically or horizontally transmitted (19). The seed heads were, however, occasionally found covered with M. atramentosa tissue (Fig. 1E to H). Whether the M. atramentosa-infected seeds remain viable is unknown and will require further investigation. Despite the many concerning aspects of clavicipitaceous fungi, they often provide a natural defense for plants against herbivore predation, holding promising applications in the realm of sustaining agriculture where insect resistance is a sought-after quality in crop systems (77).



**FIG 9** Schematic summary of warm-season grass leaf mycobiomes, mycotoxins, and future perspectives. Five perennial warm-season grasses were sampled. In the asymptomatic leaves, tremellomycetous yeasts and Dothideomycetes were revealed to be the core fungal classes across Florida pastures. The stroma-bearing (symptomatic) leaves of the forage grass *Paspalum notatum* were enriched by two *Myriogenospora atramentosa* genotypes. In comparison, the stroma-bearing leaves of the weedy grass *Sporobolus indicus* were enriched by *Balansia epichloë* and a mycoparasitic fungus, *Clonostachys* sp. *M. atramentosa* likely produced or triggered the production of emodin, an anthraquinone compound, in *P. notatum*. This study highlights the diverse leaf mycobiomes and potential production of secondary metabolites of warm-season grasses, holding potential concerns and opportunities. We call for interdisciplinary research that monitors animal health considering future climate change and pasture management strategies to better understand warm-season grass mycobiomes.

**Conclusions.** A holistic view of the foliar mycobiome of Florida warm-season grasses is illustrated in this study (Table S2, Fig. 9). We revealed that the weed Sporobolus indicus harbors the highest richness of fungi, posing potential concern about livestock grazing in pastures with cooccurring species of grass forage. Fungal assemblies are affected by both the host plant identity (i.e., species and cultivar) and the geographic location. Despite mycotoxins being a concern for livestock health, they also present a potential biocontrol tool for plants to combat insect predation and impacts from climate change as well as discovering drugs used to treat human diseases. We demonstrated that high-throughput amplicon sequencing is a powerful approach to delineate the mycobiome of healthy grasses. Additionally, unknown taxa (e.g., two Myriogenospora atramentosa genotypes) were corroborated, and overlooked fungal diversity (e.g., Clonostachys miodochialis accompanying Balansia epichloë stromata) was uncovered (Fig. 9). Significant knowledge gaps remain in understanding what mechanisms are involved in warm-season grasses and their interaction with leaf fungi, what environmental drivers can manipulate such interactions, and how these associations lead to secondary metabolite production and their subsequent impact on animal health. Future examination can incorporate host genetic differences, pasture management style, and other environmental factors such as weather data and soil nutrient level to better understand the complex factors shaping the mycobiome of warm-season grasses. We

advise that a multidisciplinary approach be taken in development of new research to be conducted on grass mycobiomes to further decipher plant-fungus interactions, determine their impacts on palatability and animal health, and elucidate the potential application of these fungal resources.

#### **MATERIALS AND METHODS**

Sample collection for fungal community assessment. We sampled five perennial warm-season grasses: P. notatum, C. dactylon, P. nicorae, S. indicus, and A. virginicus. P. notatum, P. nicorae, and A. virginicus belong to the subfamily Panicoideae while S. indicus and C. dactylon belong to Chloridoideae of the Poaceae family. The growth type of P. notatum, P. nicorae, and C. dactylon is rhizomatous in comparison to the bunch-type S. indicus and A. virginicus. Samples were collected from six pastures in Florida (Greensboro, St. Cloud, Kenansville, Tampa, Wauchula, and Chiefland; latitude and longitude range between 27°29′56" to 30°32′33"N and 84°43′08" to 80°53'13"W, respectively [Fig. 1]). Samples were collected in the summer of 2017 (between May and August) to avoid seasonal variation. All the collection sites have humid subtropical climates (78) but span four plant hardiness zones with average minimum winter temperatures ranging from 15°F to 35°F (79). Forage and weed species coinhabited all but the Chiefland site (Fig. 1A). All grass leaves sampled remained green without yellowing or signs of herbivory. Samples considered "asymptomatic" were leaves from plant individuals without any sign of fungal symptoms, while "symptomatic" samples were leaves with visible black stromata. The fungal stromata of symptomatic samples were included for all analysis. These symptomatic samples were collected from four and three out of six sampling sites for P. notatum and S. indicus, respectively (Fig. 1A to D). Three replicates were collected from each plant species at individual locations. Each replicate contained five leaf pieces. Samples were placed in sealed plastic bags, set in an ice-filled cooler, and transported back to the lab, where they were removed, placed into microcentrifuge tubes, and stored immediately in a low-temperature freezer (-80°C) until the DNA extraction step.

DNA extraction, library preparation, and sequencing. To extract DNA from leaf material, a cetyltrimethylammonium bromide (CTAB) protocol (80) was used with minor modifications. Specifically, 10 zirconia beads (2.0-mm diameter; BioSpec Products Inc., Bartlesville, OK, USA) and two stainless steel beads (2.3-mm diameter; BioSpec Products) were placed in 2-mL tubes, and samples were ground using a MiniG tissue homogenizer (SPEX SamplePrep, Metuchen, NJ, USA). Amplicon sequencing libraries were prepared with ITS1F and ITS2 primers (81, 82), targeting the ITS1 region of fungal nuclear ribosomal DNA (nrITS). To amplify the target region and to attach a barcode sequence for each sample, a three-step PCR approach was performed, as described previously (83). Briefly, the ITS region was amplified using standard primer sets (ITS1F, 5'CTTGGTCATTTAGAGGAAGTAA3'; ITS2, 5'GCTGCGTTCTTCATCGATGC3') with 10 cycles (81, 82). The PCR product was then used as the DNA input for a second 10-cycle PCR with the same primer sets (ITS1F/ITS2) combined with Illumina adaptors. The third PCR then added a unique barcode to each sample, which enabled multiplexing. The PCR products generated from the third round of PCR were cleaned up with 1:1 (vol/vol) Agencourt AMPure XP beads prior to pooling and sequencing. The concentration and purity of the final PCR product were assessed by Nanodrop (ThermoFisher Scientific) and gel electrophoresis. The PCR products were pooled with equimolar ratios of DNA and sequenced by the Illumina MiSeq 250 pair-end platform at the Duke Genome Center of Biology (Durham, NC, USA). The raw reads were submitted to the National Center for Biotechnology Information (NCBI) Sequence Read Archive (BioProject identifier [ID], PRJNA733319). Whenever a Sanger sequencing verification step was involved, we used ITS1F and LR3 (84) primer sets and followed the work of Chen et al. (83) for PCR preparation and sequencing.

Bioinformatics and statistics. Reads generated with the Illumina MiSeq were demultiplexed, and primers were trimmed with the program Cutadapt (85). The paired-end reads were quality filtered and merged, followed by amplicon sequence variant (ASV) identification in DADA2 (86). The taxonomy was determined with the UNITE database (87) using the RDP naive Bayesian classifier algorithm (88) implemented in DADA2. Key taxa were checked twice by manually performing a BLAST search (BLASTN) with the sequences against NCBI GenBank. Data analyses were conducted in R unless otherwise noted (89). Alpha-diversity was calculated based on a data set rarefied to 9,805 reads per sample, and singletons were retained in this data set. For the remaining analyses, read count of each ASV was normalized to the proportion of all reads per sample (90). Fungal community assessment was conducted in Phyloseq (91). The R package ranacapa (92) was used to examine the species accumulation curve for sampling completeness. To visualize the community assemblies shaped by locality and plant identity, nonmetric multidimensional scaling (NMDS) analysis based on beta-diversity (Bray-Curtis dissimilarity) (93) was performed. The function ADONIS, which conducts permutational multivariate analysis of variance (PERMANOVA), was performed to assess community differentiation (93) with 999 permutations. Alpha-diversity was measured with observed species and Chao1 index for richness and the Shannon index, which considers both richness and evenness (93). Wilcoxon rank sum tests (for two treatments) or Kruskal-Wallis tests (for multiple treatments) were performed for data skewed from normal distribution; otherwise, Welch's t tests (for two treatments) and ANOVA (for multiple treatments) were conducted. A post hoc Tukey honestly significant difference (HSD) test was performed for pairwise comparisons. For multiple comparisons, P values were adjusted with false-discovery rate (FDR) correction.

To test if the relative abundance of certain ASVs was significantly more abundant in symptomatic or asymptomatic *P. notatum* and *S. indicus*, we performed pairwise comparisons with the Wald test using package DESeq2 (94). Members of core mycobiomes were detected at the genus level using the R package "microbiome" (95). A core member was defined by its relative abundance of >0.01% in more than half of the samples. Because not all plants were sampled at every site (e.g., only one plant species, *P. nicorae*, was sampled

at Chiefland), to understand how plant identity and geographic location contributed in shaping mycobiome composition, we focused on a sub-data set containing four plant species (P. notatum, C. dactylon, S. indicus, and A. virginicus) which were sampled at three locations (Kenansville, Tampa, and St. Cloud) to avoid sampling bias. Indicator species were identified by the indicspecies package (96). To test for plant cultivar effect on mycobiome assembly, two P. notatum cultivars ('Argentine' and 'Pensacola') were investigated. To determine the relative degree of host (i.e., at species or cultivar level)-versus-location effect, we performed variance partitioning with redundancy analysis (RDA), using the Varpart function in the Vegan package (93).

To reconstruct phylogeny, sequences were first aligned with MAFFT (97). The phylogeny was reconstructed in RAxML with a maximum likelihood algorithm (98).

Mycotoxin analysis. The Greensboro site (Fig. 1) was revisited in the summer of 2018 to collect Paspalum notatum samples for mycotoxin detection. Each sample contained at least 50 individual leaf pieces. We collected three sample types. In addition to the "asymptomatic" and "symptomatic" samples as defined above, "mixed" samples were collected without careful examination in the field. Each of the three sample types had 5 to 6 replicates. Samples were placed in an ice-filled cooler until return to the lab and then freeze-dried (FreeZone2.5; Labconco, MO, USA) on the same collection day.

Forty-five mycotoxins (see Table S1 in the supplemental material) were examined via multiple reaction monitoring of two transitions (quantitative and qualitative) per compound on an LC-MS/MS system. Each sample was ground to pass an 0.5-mm screen, then extracted in triplicate by adding 4 mL of a 79:20:1 (vol/vol) acetonitrile-water-acetic acid extraction solution per gram of material, and then rotated in the dark for 90 min (99). Samples were centrifuged at 3,000 rpm for 5 min and then diluted 1:1 with 20:79:1 (vol/vol/vol) acetonitrile-water-acetic acid. Mycotoxins were detected on an AB/Sciex 3200 QTrap LC-MS/MS system (Applied Biosystems, Foster City, CA, USA) via electrospray ionization, with separation performed using a Perkin-Elmer (Waltham, MA, USA) series 200 high-pressure liquid chromatograph (HPLC) connected to a Gemini C<sub>18</sub> column (150 by 4.6 mm, 5 µm; Phenomenex [Torrance, CA, USA]) with a 4- by 3-mm security guard cartridge with similar packing (99). Mobile phases consisted of 5 mM ammonium acetate and methanol-water-acetic acid in a ratio of 10:89:1 (vol/vol/vol) (A) or 97:2:1 (B) and were run in a gradient program at 1 mL/min. Ammonium acetate (HPLC grade) and methanol and acetic acid (LC-MS grade) were purchased from commercial sources. The 18-m $\Omega$  water was obtained from an Elga Ultra PureLab water system (Cary, NC, USA). The presence of a mycotoxin was confirmed when the signal was equal to or greater than a signal-to-noise (S/N) ratio of 3:1 (limit of detection [LOD]), and both quantitative and qualitative transitions were present. Mycotoxins were quantified for each sample against standard curves established using certified analytical standards purchased from commercial sources. To analyze mycotoxin content, the concentrations of different types of mycotoxins among the three sample types (symptomatic, asymptomatic, and mixed) of P. notatum were compared with the Kruskal-Wallis test followed by a pairwise Wilcox test.

Data availability. All sequences are available at NCBI GenBank. The raw reads were submitted to the Sequence Read Archive (BioProject identifier [ID], PRJNA733319). Sanger sequencing accession numbers are ON678271 to ON678272. The fungal specimen was deposited at the University of Florida Herbarium under no. FLAS-F-63886.

# **SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

SUPPLEMENTAL FILE 1, PDF file, 1.1 MB.

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F.M., K.-H.C., C.M., and A.B. collected grass samples for molecular work and mycotoxin detection. K.-H.C. analyzed the data and led the writing of the manuscript. F.M. conducted library preparation toward sequencing. J.D. measured the mycotoxins. H.-L.L. designed and supervised the project. All authors contributed to the writing and editing of the manuscript.

#### REFERENCES

- 1. Vandenkoornhuyse P, Quaiser A, Duhamel M, Van AL, Dufresne A. 2015. The importance of the microbiome of the plant holobiont. New Phytol 206:1196-1206. https://doi.org/10.1111/nph.13312.
- 2. Rodriguez RJ, White JF, Jr, Arnold AE, Redman RS. 2009. Fungal endophytes: diversity and functional roles. New Phytol 182:314-330. https://doi .org/10.1111/j.1469-8137.2009.02773.x.
- 3. Bezerra da Rocha ME, Freire FDCO, Maia FEF, Florindo Guedes MI, Rondina D. 2014. Mycotoxins and their effects on human and animal health. Food Control 36:159-165. https://doi.org/10.1016/j.foodcont.2013.08.021.
- 4. Kallenbach RL. 2015. Bill E. Kunkle interdisciplinary beef symposium: coping with tall fescue toxicosis: solutions and realities. J Anim Sci 93:5487-5495. https://doi.org/10.2527/jas.2015-9229.

- Alison MW, Pitman WD, Aiken GE, Han K-J. 2018. Potential for use of a perennial C3 grass in a warm-temperate region with pastures dominated by C4 grasses. *In* Tadele Z (ed), Grasses as food and feed. IntechOpen, London, United Kingdom.
- Blount AR, Acuña CA. 2009. Bahiagrass. In Singh RJ (ed), Genetic resources, chromosome engineering, and crop improvement series: forage crops. CRC Press. Boca Raton. FL.
- Bunnell BT, Baker RD, McCarty LB, Hall DW, Colvin DL. 2003. Differential response
  of five Bahiagrass (*Paspalum notatum*) cultivars to metsulfuron. Weed Technol
  17:550–553. https://doi.org/10.1614/0890-037X(2003)017[0550:DROFBP]2.0.CO;2.
- Blount A, Wallau M, Sellers B, Hancock D, Dillard L, Dubeux J, Mackowiak C, Vendramini J, Cooper C. 2022. Brunswickgrass (Paspalum nicorae): a weed contaminant in southern pastures and Bahiagrass seed production fields. Publication SS-AGR-405. Agronomy Department, UF/IFAS Extension, University of Florida, Gainesville, FL. https://doi.org/10.32473/edis-ag408-2022.
- Rana N, Wilder BJ, Sellers BA, Ferrell JA, MacDonald GE. 2012. Effects of environmental factors on seed germination and emergence of Smutgrass (Sporobolus indicus) varieties. Weed Sci 60:558–563. https://doi.org/10.1614/ WS-D-11-00208.1.
- Rana N, Sellers BA, Ferrell JA, MacDonald GE, Silveira ML, Vendramini JM. 2013. Impact of soil pH on Bahiagrass competition with giant Smutgrass (Sporobolus indicus var. pyramidalis) and small smutgrass (Sporobolus indicus var. indicus). Weed Sci 61:109–116. https://doi.org/10.1614/WS-D-12-00070.1.
- Butler TJ, Redmon LA, Stritzke JF, Goad CL. 2006. Using prescribed fire, tillage, and fertilizer to manage Broomsedge-infested pastures. Forage Grazinglands 4:1–7. https://doi.org/10.1094/FG-2006-0424-01-RS.
- Peters EJ, Lowance SA. 1974. Fertility and management treatments to control Broomsedge in pastures. Weed Sci 22:201–205. https://doi.org/10 .1017/S0043174500036900.
- Depetris MB, Acuña CA, Gutierrez S, Marcón F, Felitti SA. 2020. Fungal endophyte diversity in *Paspalum* and effects on plant growth. Grass Forage Sci 75:316–325. https://doi.org/10.1111/gfs.12494.
- Schardl CL, Young CA, Faulkner JR, Florea S, Pan J. 2012. Chemotypic diversity of epichloae, fungal symbionts of grasses. Fungal Ecol 5:331–344. https://doi.org/10.1016/j.funeco.2011.04.005.
- White JF, Glenn AE. 1994. A study of two fungal epibionts of grasses: structural features, host relationships, and classification in the genus *Myriogenospora* (Clavicipitales). Am J Bot 81:216–223. https://doi.org/10.1002/j.1537-2197.1994.tb15432.x.
- Pažoutová S, Odvody GN, Frederickson DE, Chudíčková M, Olšovská J, Kolařík M. 2011. New *Claviceps* species from warm-season grasses. Fungal Divers 49:145–165. https://doi.org/10.1007/s13225-011-0102-4.
- Bacon CW, Hinton DM, Mitchell TR, Palencia ER. 2018. In situ ergot alkaloid detection in three Balansia epichloe-infected grass species. J Appl Microbiol 125:976–985. https://doi.org/10.1111/jam.13941.
- Rykard DM, Luttrell ES, Bacon CW. 1982. Development of the conidial state of *Myriogenospora atramentosa*. Mycologia 74:648–654. https://doi.org/10.1080/00275514.1982.12021562.
- Glenn AE, Rykard DM, Bacon CW, Hanlin RT. 1998. Molecular characterization of *Myriogenospora atramentosa* and its occurrence on some new hosts. Mycol Res 102:483–490. https://doi.org/10.1017/S0953756297005236.
- Chen K-H, Blount A, Justesen B, Walter JH, Wallau M, Liao H-L. 2019. First report of the association of *Myriogenospora atramentosa* with the plant genus *Hemarthria*. Plant Health Prog 20:244–246. https://doi.org/10.1094/PHP-07-19 -0043-BR.
- Cruz-Laufer AJ, Mardones M, Piepenbring M. 2019. Systematics, taxonomy, and distribution of species of *Myriogenospora* G.F. Atk. (Clavicipitaceae, Hypocreales, Ascomycota). Check List 15:735–746. https://doi.org/10.15560/15.5.735.
- Sapkota R, Knorr K, Jørgensen LN, O'Hanlon KA, Nicolaisen M. 2015. Host genotype is an important determinant of the cereal phyllosphere mycobiome. New Phytol 207:1134–1144. https://doi.org/10.1111/nph.13418.
- Knorr K, Jørgensen LN, Nicolaisen M. 2019. Fungicides have complex effects on the wheat phyllosphere mycobiome. PLoS One 14:e0213176. https://doi.org/10.1371/journal.pone.0213176.
- Noble AS, Noe S, Clearwater MJ, Lee CK. 2020. A core phyllosphere microbiome exists across distant populations of a tree species indigenous to New Zealand. PLoS One 15:e0237079. https://doi.org/10.1371/journal.pone.0237079.
- Al Ashhab A, Meshner S, Alexander-Shani R, Dimerets H, Brandwein M, Bar-Lavan Y, Winters G. 2021. Temporal and spatial changes in phyllosphere microbiome of *Acacia* trees growing in arid environments. Front Microbiol 12:656269. https://doi.org/10.3389/fmicb.2021.656269.
- Yao H, Sun X, He C, Maitra P, Li X-C, Guo L-D. 2019. Phyllosphere epiphytic and endophytic fungal community and network structures differ in a tropical mangrove ecosystem. Microbiome 7:57. https://doi.org/10.1186/s40168-019-0671-0.

- Toju H, Peay KG, Yamamichi M, Narisawa K, Hiruma K, Naito K, Fukuda S, Ushio M, Nakaoka S, Onoda Y, Yoshida K, Schlaeppi K, Bai Y, Sugiura R, Ichihashi Y, Minamisawa K, Kiers ET. 2018. Core microbiomes for sustainable agroecosystems. Nat Plants 4:247–257. https://doi.org/10.1038/s41477-018-0139-4.
- Berg G, Rybakova D, Fischer D, Cernava T, Vergès M-CC, Charles T, Chen X, Cocolin L, Eversole K, Corral GH, Kazou M, Kinkel L, Lange L, Lima N, Loy A, Macklin JA, Maguin E, Mauchline T, McClure R, Mitter B, Ryan M, Sarand I, Smidt H, Schelkle B, Roume H, Kiran GS, Selvin J, de Souza RSC, van Overbeek L, Singh BK, Wagner M, Walsh A, Sessitsch A, Schloter M. 2020. Microbiome definition re-visited: old concepts and new challenges. Microbiome 8:103. https://doi.org/10.1186/s40168-020-00875-0.
- Baldrian P. 2017. Forest microbiome: diversity, complexity and dynamics. FEMS Microbiol Rev 41:109–130. https://doi.org/10.1093/femsre/fuw040.
- Castañeda LE, Miura T, Sánchez R, Barbosa O. 2018. Effects of agricultural management on phyllosphere fungal diversity in vineyards and the association with adjacent native forests. PeerJ 6:e5715. https://doi.org/10.7717/ peerj.5715.
- 31. VanWallendael A, Benucci GMN, da Costa PB, Fraser L, Sreedasyam A, Fritschi F, Juenger TE, Lovell JT, Bonito G, Lowry DB. 2022. Host genotype controls ecological change in the leaf fungal microbiome. PLoS Biol 20: e3001681. https://doi.org/10.1371/journal.pbio.3001681.
- Bowsher AW, Benucci GMN, Bonito G, Shade A. 2021. Seasonal dynamics of core fungi in the switchgrass phyllosphere, and co-occurrence with leaf bacteria. Phytobiomes J 5:60–68. https://doi.org/10.1094/PBIOMES -07-20-0051-R.
- 33. Nnadi NE, Carter DA. 2021. Climate change and the emergence of fungal pathogens. PLoS Pathog 17:e1009503. https://doi.org/10.1371/journal.ppat.1009503.
- Anderson PK, Cunningham AA, Patel NG, Morales FJ, Epstein PR, Daszak P. 2004. Emerging infectious diseases of plants: pathogen pollution, climate change and agrotechnology drivers. Trends Ecol Evol 19:535–544. https://doi.org/10.1016/j.tree.2004.07.021.
- 35. Kivlin SN, Mann MA, Lynn JS, Kazenel MR, Taylor DL, Rudgers JA. 2022. Grass species identity shapes communities of root and leaf fungi more than elevation. ISME Commun 2:25. https://doi.org/10.1038/s43705-022-00107-6.
- Bailes G, Thomas D, Bridgham SD, Roy BA. 2020. Drivers of grass endophyte communities in prairies of the Pacific Northwest, USA. bioRxiv. https://doi.org/10.1101/2020.02.23.95348.
- Schiro G, Colangeli P, Müller MEH. 2019. A metabarcoding analysis of the mycobiome of wheat ears across a topographically heterogeneous field. Front Microbiol 10:2095. https://doi.org/10.3389/fmicb.2019.02095.
- 38. Wemheuer B, Thomas T, Wemheuer F. 2019. Fungal endophyte communities of three agricultural important grass species differ in their response towards management regimes. Microorganisms 7:37. https://doi.org/10.3390/microorganisms7020037.
- Mejía LC, Rojas El, Maynard Z, Bael SV, Arnold AE, Hebbar P, Samuels GJ, Robbins N, Herre EA. 2008. Endophytic fungi as biocontrol agents of *Theo-broma cacao* pathogens. Biol Control 46:4–14. https://doi.org/10.1016/j.biocontrol.2008.01.012.
- 40. Busby PE, Peay KG, Newcombe G. 2016. Common foliar fungi of *Populus trichocarpa* modify *Melampsora* rust disease severity. New Phytol 209: 1681–1692. https://doi.org/10.1111/nph.13742.
- 41. Kleczewski NM, Bauer JT, Bever JD, Clay K, Reynolds HL. 2012. A survey of endophytic fungi of switchgrass (*Panicum virgatum*) in the midwest, and their putative roles in plant growth. Fungal Ecol 5:521–529. https://doi.org/10.1016/j.funeco.2011.12.006.
- 42. Wipf HML, Coleman-Derr D. 2021. Evaluating domestication and ploidy effects on the assembly of the wheat bacterial microbiome. PLoS One 16: e0248030. https://doi.org/10.1371/journal.pone.0248030.
- 43. Górzyńska K, Ślachetka M, Ryszka P, Turnau K, Płachno BJ, Lembicz M. 2018. Incidence, identification, and mycoparasitic activity of *Clonostachys epichloë*, a hyperparasite of the fungal endophyte *Epichloë typhina*. Plant Dis 102:1973–1980. https://doi.org/10.1094/PDIS-02-18-0320-RE.
- Kosawang C, Karlsson M, Jensen DF, Dilokpimol A, Collinge DB. 2014.
   Transcriptomic profiling to identify genes involved in *Fusarium* mycotoxin deoxynivalenol and zearalenone tolerance in the mycoparasitic fungus *Clonostachys rosea*. BMC Genomics 15:55. https://doi.org/10.1186/1471-2164-15-55.
- Higgins KL, Arnold AE, Coley PD, Kursar TA. 2014. Communities of fungal endophytes in tropical forest grasses: highly diverse host- and habitat generalists characterized by strong spatial structure. Fungal Ecol 8:1–11. https://doi.org/10.1016/j.funeco.2013.12.005.
- Rojas EC, Sapkota R, Jensen B, Jørgensen HJL, Henriksson T, Jørgensen LN, Nicolaisen M, Collinge DB. 2020. Fusarium head blight modifies fungal endophytic communities during infection of wheat spikes. Microb Ecol 79:397–408. https://doi.org/10.1007/s00248-019-01426-3.

- 47. Liu X-Z, Wang Q-M, Theelen B, Groenewald M, Bai F-Y, Boekhout T. 2015. Phylogeny of tremellomycetous yeasts and related dimorphic and filamentous basidiomycetes reconstructed from multiple gene sequence analyses. Stud Mycol 81:1–26. https://doi.org/10.1016/j.simyco.2015.08.001.
- 48. Landell MF, Brandão LR, Barbosa AC, Ramos JP, Safar SVB, Gomes FCO, Sousa FMP, Morais PB, Broetto L, Leoncini O, Ribeiro JR, Fungsin B, Takashima M, Nakase T, Lee C-F, Vainstein MH, Fell JW, Scorzetti G, Vishniac HS, Rosa CA, Valente P. 2014. *Hannaella pagnoccae* sp. nov., a tremellaceous yeast species isolated from plants and soil. Int J Syst Evol Microbiol 64:1970–1977. https://doi.org/10.1099/ijs.0.059345-0.
- Kasfi K, Taheri P, Jafarpour B, Tarighi S. 2018. Identification of epiphytic yeasts and bacteria with potential for biocontrol of grey mold disease on table grapes caused by *Botrytis cinerea*. Span J Agric Res 16:e1002. https://doi.org/10.5424/sjar/2018161-11378.
- Ferraz P, Cássio F, Lucas C. 2019. Potential of yeasts as biocontrol agents of the phytopathogen causing cacao witches' broom disease: is microbial warfare a solution? Front Microbiol 10:1766. https://doi.org/10.3389/ fmicb.2019.01766.
- Freimoser FM, Rueda-Mejia MP, Tilocca B, Migheli Q. 2019. Biocontrol yeasts: mechanisms and applications. World J Microbiol Biotechnol 35: 154. https://doi.org/10.1007/s11274-019-2728-4.
- Wang B, Liang X, Gleason ML, Zhang R, Sun G. 2017. Genome sequence of the ectophytic fungus *Ramichloridium luteum* reveals unique evolutionary adaptations to plant surface niche. BMC Genomics 18:729. https://doi .org/10.1186/s12864-017-4118-3.
- Maia NDC, Souza PNDC, Godinho BTV, Moreira SI, de Abreu LM, Jank L, Cardoso PG. 2018. Fungal endophytes of *Panicum maximum* and *Pennisetum* purpureum: isolation, identification, and determination of antifungal potential. Rev Bras Zootec 47:e20170183. https://doi.org/10.1590/rbz4720170183.
- 54. de Gama DS, Santos ÍAFM, de Abreu LM, de Medeiros FHV, Duarte WF, Cardoso PG. 2020. Endophytic fungi from *Brachiaria* grasses in Brazil and preliminary screening of *Sclerotinia sclerotiorum* antagonists. Sci Agric 77: e20180210. https://doi.org/10.1590/1678-992x-2018-0210.
- Oita S, Carey J, Kline I, Ibáñez A, Yang N, Hom EFY, Carbone I, U'Ren JM, Arnold AE. 2021. Methodological approaches frame insights into endophyte richness and community composition. Microb Ecol 82:21–34. https://doi.org/ 10.1007/s00248-020-01654-y.
- Anguita-Maeso M, Olivares-García C, Haro C, Imperial J, Navas-Cortés JA, Landa BB. 2019. Culture-dependent and culture-independent characterization of the olive xylem microbiota: effect of sap extraction methods. Front Plant Sci 10:1708. https://doi.org/10.3389/fpls.2019.01708.
- Chaudhry V, Runge P, Sengupta P, Doehlemann G, Parker JE, Kemen E.
   2021. Shaping the leaf microbiota: plant–microbe–microbe interactions. J
   Exp Bot 72:36–56. https://doi.org/10.1093/jxb/eraa417.
- 58. Fuchs B, Krischke M, Mueller MJ, Krauss J. 2017. Herbivore-specific induction of defense metabolites in a grass–endophyte association. Funct Ecol 31:318–324. https://doi.org/10.1111/1365-2435.12755.
- Tor-Agbidye J, Blythe LL, Craig AM. 2001. Correlation of endophyte toxins (ergovaline and lolitrem B) with clinical disease: fescue foot and perennial ryegrass staggers. Vet Hum Toxicol 43:140–146.
- Gallagher RT, Campbell AG, Hawkes AD, Holland PT, McGaveston DA, Pansier EA, Harvey IC. 1982. Ryegrass staggers: the presence of Lolitrem neurotoxins in perennial ryegrass seed. N Z Vet J 30:183–184. https://doi .org/10.1080/00480169.1982.34936.
- Bennett JW, Klich M. 2003. Mycotoxins. Clin Microbiol Rev 16:497–516. https://doi.org/10.1128/CMR.16.3.497-516.2003.
- 62. Gott P, Johns A, Stam A, Miller B, Weaver T, Bell B, Hendel E, Hofstetter-Schahs U, Murugesan G. 2018. PSXVI-31 intervention strategy for zearale-none's negative effects on performance of cow-calf pairs supplemented with liquid feed in South Central Florida a field study. J Anim Sci 96:383. https://doi.org/10.1093/jas/sky404.840.
- Mantle P. 2020. Comparative ergot alkaloid elaboration by selected plectenchymatic mycelia of Claviceps purpurea through sequential cycles of axenic culture and plant parasitism. Biology 9:41. https://doi.org/10.3390/biology9030041.
- Young CA, Schardl CL, Panaccione DG, Florea S, Takach JE, Charlton ND, Moore N, Webb JS, Jaromczyk J. 2015. Genetics, genomics and evolution of ergot alkaloid diversity. Toxins (Basel) 7:1273–1302. https://doi.org/10 .3390/toxins7041273.
- Spiers DE, Zhang Q, Eichen PA, Rottinghaus GE, Garner GB, Ellersieck MR. 1995. Temperature-dependent responses of rats to ergovaline derived from endophyte-infected tall fescue. J Anim Sci 73:1954–1961. https://doi.org/10.2527/1995.7371954x.

- Medina Á, Rodríguez A, Magan N. 2015. Climate change and mycotoxigenic fungi: impacts on mycotoxin production. Curr Opin Food Sci 5:99–104. https://doi.org/10.1016/j.cofs.2015.11.002.
- Gott P, Stam A, Johns A, Hendel E, Mendoza S, Hofstetter-Schahs U, Bell B, Murugesan G. 2018. 424 mycotoxin survey of southern US pasture grasses. J Anim Sci 96:209. https://doi.org/10.1093/jas/sky404.453.
- Dall'Asta C, Cirlini M, Falavigna C. 2014. Mycotoxins from Alternaria. Adv Mol Toxicol 8:107–121. https://doi.org/10.1016/B978-0-444-63406-1.00003-9.
- Izhaki I. 2002. Emodin a secondary metabolite with multiple ecological functions in higher plants. New Phytol 155:205–217. https://doi.org/10 .1046/i.1469-8137.2002.00459.x.
- Dong X, Fu J, Yin X, Cao S, Li X, Lin L, Huyiligeqi Ni J. 2016. Emodin: a review of its pharmacology, toxicity and pharmacokinetics. Phytother Res 30:1207–1218. https://doi.org/10.1002/ptr.5631.
- 71. Sevilla P, Rivas JM, García-Blanco F, García-Ramos JV, Sánchez-Cortés S. 2007. Identification of the antitumoral drug emodin binding sites in bovine serum albumin by spectroscopic methods. Biochim Biophys Acta 1774:1359–1369. https://doi.org/10.1016/j.bbapap.2007.07.022.
- Burton GW. 1955. Breeding Pensacola Bahiagrass, *Paspalum notatum*:
   I. method of reproduction. Agron J 47:311–314. https://doi.org/10.2134/agronj1955.00021962004700070008x.
- Rios E, Blount A, Harmon P, Mackowiak C, Kenworthy K, Quesenberry K. 2015. Ergot resistant tetraploid Bahiagrass and fungicide effects on seed yield and quality. Plant Health Prog 16:56–62. https://doi.org/10.1094/PHP -RS-14-0051.
- 74. Sung G-H, Sung J-M, Hywel-Jones NL, Spatafora JW. 2007. A multi-gene phylogeny of Clavicipitaceae (Ascomycota, Fungi): identification of localized incongruence using a combinational bootstrap approach. Mol Phylogenet Evol 44:1204–1223. https://doi.org/10.1016/j.ympev.2007.03.011.
- 75. Florea S, Panaccione DG, Schardl CL. 2017. Ergot alkaloids of the family Clavicipitaceae. Phytopathology 107:504–518. https://doi.org/10.1094/PHYTO -12-16-0435-RVW.
- Shivas R, White J, Irwin J, Blatch A, Whittle P. 1999. First record in Australia of *Myriogenospora atramentosa* on lemongrass and sugarcane. Australas Plant Pathol 28:336. https://doi.org/10.1071/AP99053.
- 77. Kauppinen M, Saikkonen K, Helander M, Pirttilä AM, Wäli PR. 2016. *Epichloë* grass endophytes in sustainable agriculture. Nat Plants 2:15224. https://doi.org/10.1038/nplants.2015.224.
- 78. Olson DM, Dinerstein E, Wikramanayake ED, Burgess ND, Powell GVN, Underwood EC, D'Amico JA, Itoua I, Strand HE, Morrison JC, Loucks CJ, Allnutt TF, Ricketts TH, Kura Y, Lamoreux JF, Wettengel WW, Hedao P, Kassem KR. 2001. Terrestrial ecoregions of the world: a new map of life on Earth. Bioscience 51:933–938. https://www.sciencebase.gov/catalog/item/508fece8e4b0a1b43c29ca22.
- Agricultural Research Service. 2005. USDA plant hardiness zone map: United States. Agricultural Research Service, US Department of Agriculture, Washington, DC, and Oregon State University, Corvallis, OR. https://planthardiness.ars.usda.gov/.
- Liao H-L, Chen Y, Bruns TD, Peay KG, Taylor JW, Branco S, Talbot JM, Vilgalys R. 2014. Metatranscriptomic analysis of ectomycorrhizal roots reveals genes associated with *Piloderma–Pinus* symbiosis: improved methodologies for assessing gene expression *in situ*. Environ Microbiol 16:3730–3742. https://doi.org/10.1111/1462-2920.12619.
- 81. White TJ, Bruns T, Lee S, Taylor JW. 1990. Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics, p 315–322. *In* Innis MA, Gelfand DH, Sninsky JJ, White TJ (ed), PCR protocols: a guide to methods and applications. Academic Press, New York, NY.
- Gardes M, Bruns TD. 1993. ITS primers with enhanced specificity for basidiomycetes application to the identification of mycorrhizae and rusts. Mol Ecol 2:113–118. https://doi.org/10.1111/j.1365-294X.1993.tb00005.x.
- 83. Chen K-H, Liao H-L, Arnold AE, Bonito G, Lutzoni F. 2018. RNA-based analyses reveal fungal communities structured by a senescence gradient in the moss *Dicranum scoparium* and the presence of putative multi-trophic fungi. New Phytol 218:1597–1611. https://doi.org/10.1111/nph.15092.
- Vilgalys R, Hester M. 1990. Rapid genetic identification and mapping of enzymatically amplified ribosomal DNA from several *Cryptococcus* species. J Bacteriol 172:4238–4246. https://doi.org/10.1128/jb.172.8.4238-4246.1990.
- Martin M. 2011. Cutadapt removes adapter sequences from high-throughput sequencing reads. EMBnet J 17:10–12. https://doi.org/10.14806/ej.17.1.200.
- Callahan BJ, McMurdie PJ, Rosen MJ, Han AW, Johnson AJA, Holmes SP. 2016. DADA2: high-resolution sample inference from Illumina amplicon data. 7. Nat Methods 13:581–583. https://doi.org/10.1038/nmeth.3869.
- 87. Nilsson RH, Larsson K-H, Taylor AFS, Bengtsson-Palme J, Jeppesen TS, Schigel D, Kennedy P, Picard K, Glöckner FO, Tedersoo L, Saar I, Kõljalg U,

- Abarenkov K. 2019. The UNITE database for molecular identification of fungi: handling dark taxa and parallel taxonomic classifications. Nucleic Acids Res 47:D259–D264. https://doi.org/10.1093/nar/gky1022.
- Wang Q, Garrity GM, Tiedje JM, Cole JR. 2007. Naive Bayesian classifier for rapid assignment of rRNA sequences into the new bacterial taxonomy. Appl Environ Microbiol 73:5261–5267. https://doi.org/10.1128/AEM.00062-07.
- 89. R Core Team. 2019. R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. https://www.R-project.org/.
- Weiss S, Xu ZZ, Peddada S, Amir A, Bittinger K, Gonzalez A, Lozupone C, Zaneveld JR, Vázquez-Baeza Y, Birmingham A, Hyde ER, Knight R. 2017. Normalization and microbial differential abundance strategies depend upon data characteristics. Microbiome 5:27. https://doi.org/10.1186/s40168-017-0237-y.
- 91. McMurdie PJ, Holmes S. 2013. phyloseq: an R package for reproducible interactive analysis and graphics of microbiome census data. PLoS One 8: e61217. https://doi.org/10.1371/journal.pone.0061217.
- Kandlikar GS, Gold ZJ, Cowen MC, Meyer RS, Freise AC, Kraft NJB, Moberg-Parker J, Sprague J, Kushner DJ, Curd EE. 2018. ranacapa: an R package and Shiny web app to explore environmental DNA data with exploratory statistics and interactive visualizations. F1000Res 7:1734. https://doi.org/10.12688/f1000research.16680.1.

- 93. Dixon P. 2003. VEGAN, a package of R functions for community ecology. J Veg Sci 14:927–930. https://doi.org/10.1111/j.1654-1103.2003.tb02228.x.
- 94. Love MI, Huber W, Anders S. 2014. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol 15:550. https://doi.org/10.1186/s13059-014-0550-8.
- 95. Lahti L. 2019. Microbiome R package. http://microbiome.github.io.
- Cáceres MD, Legendre P. 2009. Associations between species and groups of sites: indices and statistical inference. Ecology 90:3566–3574. https://doi.org/10 .1890/08-1823.1.
- 97. Katoh K, Standley DM. 2013. MAFFT multiple sequence alignment software version 7: improvements in performance and usability. Mol Biol Evol 30:772–780. https://doi.org/10.1093/molbev/mst010.
- 98. Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30:1312–1313. https://doi.org/10.1093/bioinformatics/btu033.
- 99. Sulyok M, Berthiller F, Krska R, Schuhmacher R. 2006. Development and validation of a liquid chromatography/tandem mass spectrometric method for the determination of 39 mycotoxins in wheat and maize. Rapid Commun Mass Spectrom 20:2649–2659. https://doi.org/10.1002/rcm.2640.