

SHORT COMMUNICATION

Links between plant and fungal communities across a deforestation chronosequence in the Amazon rainforest

This article has been corrected since Advance Online Publication and a corrigendum is also printed in this issue

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Understanding the interactions among microbial communities, plant communities and soil properties following deforestation could provide insights into the long-term effects of land-use change on ecosystem functions, and may help identify approaches that promote the recovery of degraded sites. We combined high-throughput sequencing of fungal rDNA and molecular barcoding of plant roots to estimate fungal and plant community composition in soil sampled across a chronosequence of deforestation. We found significant effects of land-use change on fungal community composition, which was more closely correlated to plant community composition than to changes in soil properties or geographic distance, providing evidence for strong links between above- and below-ground communities in tropical forests.

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Land-use change, such as deforestation, is one of the greatest threats to biodiversity worldwide (Thomas *et al.*, 2004), particularly within tropical ecosystems (Sala *et al.*, 2000; Gibson *et al.*, 2011). In tropical ecosystems, the responses of soil fungal biodiversity to land-use change are only beginning to be explored, and responsiveness has been shown in some cases (for example, Fracetto *et al.*, 2013) but not in others (for example, Leal *et al.*, 2009). Fungal communities can be structured by nutrient availability (Rousk *et al.*, 2010) and plant community composition (Carney and Matson, 2006; Peay *et al.*, 2013), suggesting that shifts in soil fungal communities in response to land-use change (Castro *et al.*, 2008; Fracetto *et al.*, 2013) could result from alterations in soil properties and/or plant communities. However, the relative strength of these factors in determining fungal community composition in tropical systems has not been examined.

We used an established chronosequence of land use in the Brazilian Amazon rainforest (Rodrigues *et al.*, 2013) that included a primary forest, secondary forest and pasture. We sampled soil using a spatially explicit design within single hectare plots for a total of nine samples per site. To examine changes in fungal community composition we targeted the ITS1 region of the rRNA gene using the fungal-specific primer pair ITS1F-ITS2 with a two-stage PCR designed for paired-end Illumina sequencing. Amplicons were sequenced on an Illumina HiSeq 2000 (Illumina, San Diego, CA, USA) at the University of Oregon, quality-filtered to remove low-quality bases and putative chimeras, and clustered into operational taxonomic units at 97% sequence similarity using the program UCHIME (Edgar, 2010) in the QIIME platform (Caporaso *et al.*, 2010). Plant community composition was determined by direct PCR (Phire Direct PCR, ThermoScientific, Waltham, MA, USA) and Sanger sequencing of the chloroplast *trnL* intron region (Taberlet *et al.*, 1991, 2007) of roots from the same soil cores collected for microbial community analysis. Root sequences were grouped by plant genera on the basis of top BLAST matches to the GenBank database (Supplementary Table 1). Unique root sequences were deposited in GenBank under accession numbers KF661456–KF661514, and ITS1

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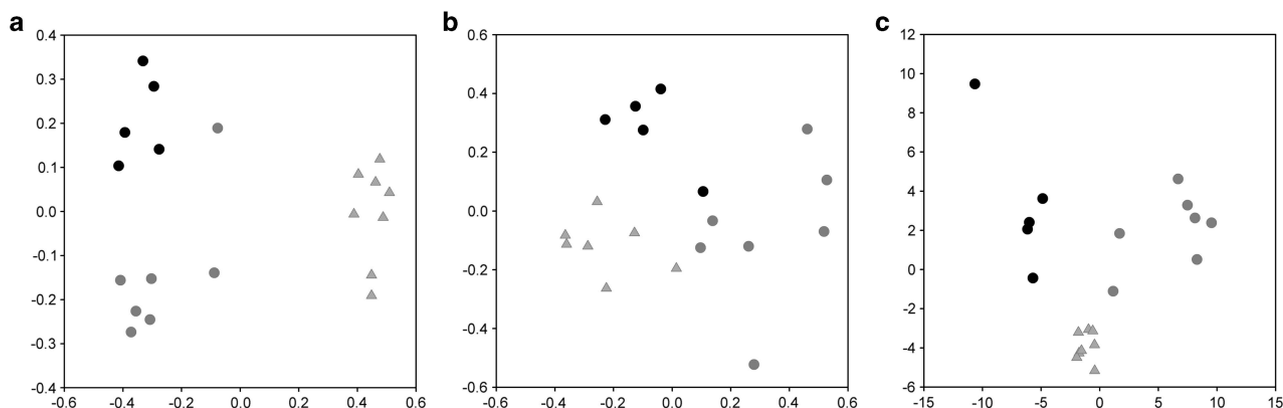


Figure 1 Nonmetric multidimensional scaling ordinations of (a) fungal communities, (b) plant communities and (c) soil properties in matched soil cores. Land-use types are as follows: primary forest (black circles), pasture (triangles) and secondary forest (gray circles).

sequences were deposited in MG-RAST under ID 4536676.3. Soil properties were determined by quantifying micro- and macronutrients with protocols modified for tropical soil (van Raij *et al.*, 2001; Supplementary Table 2). Analyses were undertaken on matched samples that had sufficient data for fungi, roots and soil properties for a total of 20 samples across the three land uses. A more detailed description of the methods can be found in the Supplementary Material.

Using an operational taxonomic unit community matrix rarefied to 10 000 sequences per sample, fungal community composition (Bray–Curtis) was compared among the sites using PERMANOVA (Anderson, 2001) and visualized using nonmetric multidimensional scaling. The fungal community composition was significantly different among the three land uses (that is, primary forest, secondary forest and pasture; $R^2 = 0.32$, $F_{2,17} = 4.14$, $P = 0.001$; Figure 1a), driven mostly by decreased relative abundance of Basidiomycota in the pasture and secondary forest (Supplementary Figure 1). PERMANOVA analysis also showed significant differences in plant community composition ($R^2 = 0.41$, $F_{2,17} = 5.92$, $P = 0.001$) and measures of soil properties ($F_{2,17} = 25.6$, $R^2 = 0.75$, $P = 0.001$; Figures 1b and c).

The relative contribution of variation in pairwise geographic distance, environmental distance (Hellinger-transformed Euclidean distance) and plant community dissimilarity (Bray–Curtis) to variation in fungal community composition was determined using multiple regression on distance matrices (Lichstein, 2006). We found that plant community composition was the strongest driver of differences in fungal community composition (Table 1). Although we found significant differences among the sites (Supplementary Table 2), fungal richness was not significantly correlated with plant richness ($F = 1.68$, $P = 0.21$).

Plant identification through targeting of roots within soil cores provides the means to isolate the plant species most likely to interact with the fungal

Table 1 Multiple regression on distance matrices comparing the relationship between fungal community similarity and geographic distance, plant community similarity and environmental similarity on the basis of soil properties (Model: $F = 57.02$, $R^2 = 0.48$, $P = 0.001$)

	Coefficient	P-value
Plant community	0.169	0.002
Geographic	0.006	0.027
Environmental	0.005	0.078

Coefficient values are partial regression coefficients, which indicate the rate of change in community similarity determined by a given independent variable while other independent variables are held constant. Bolded values indicate significance at $\alpha = 0.05$.

community in a given sampling location. We used this novel approach to characterize plant composition among plots and found that the plant community was a stronger predictor of the fungal community than were soil properties or geographic distance. These results are consistent with those of Mitchell *et al.* (2010), who showed that plant community composition was a better predictor of microbial community composition than was soil chemistry across a plant successional gradient in a boreal ecosystem. Our findings support those of Peay *et al.* (2013), who found a significant correlation between plant community composition and fungal community composition in three tropical forests, and further show that fungal community composition is more closely linked to plant community composition than to either environmental variability or geographic distance.

Our results also provide evidence that effects of deforestation on fungal communities are likely mediated in large part by changes in plant communities. Fungal communities in secondary forests were more similar to primary forests than were pasture communities (Figure 1), suggesting that the recovery of fungal communities following pasture abandonment is closely tied to that of plants. A clearer understanding of interactions between plant and fungal communities could prove useful to

conservation and restoration biology, as it could identify management strategies that better promote both reforestation and the recovery of microbially mediated ecosystem functions in degraded areas (van der Heijden *et al.*, 2008).

Conflict of Interest

The authors declare no conflict of interest.

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Supplementary information accompanies this paper on The ISME Journal website (<http://www.nature.com/ismej>)