



Phylogenetic diversity of Amazonian tree communities

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ABSTRACT

Aim To examine variation in the phylogenetic diversity (PD) of tree communities across geographical and environmental gradients in Amazonia.

Location Two hundred and eighty-three *c.* 1 ha forest inventory plots from across Amazonia.

Methods We evaluated PD as the total phylogenetic branch length across species in each plot (PD_{ss}), the mean pairwise phylogenetic distance between species (MPD), the mean nearest taxon distance (MNTD) and their equivalents standardized for species richness (ses.PD_{ss}, ses.MPD, ses.MNTD). We compared PD of tree communities growing (1) on substrates of varying geological age; and (2) in environments with varying ecophysiological barriers to growth and survival.

Results PD_{ss} is strongly positively correlated with species richness (SR), whereas MNTD has a negative correlation. Communities on geologically young- and intermediate-aged substrates (western and central Amazonia respectively) have the highest SR, and therefore the highest PD_{ss} and the lowest MNTD. We find that the youngest and oldest substrates (the latter on the Brazilian and Guiana Shields) have the highest ses.PD_{ss} and ses.MNTD. MPD and ses.MPD are strongly correlated with how evenly taxa are distributed among the three principal angiosperm clades and are both highest in western Amazonia. Meanwhile, seasonally dry tropical forest (SDTF) and forests on white sands have low PD, as evaluated by any metric.

Main conclusions High ses.PD_{ss} and ses.MNTD reflect greater lineage diversity in communities. We suggest that high ses.PD_{ss} and ses.MNTD in western

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Amazonia results from its favourable, easy-to-colonize environment, whereas high values in the Brazilian and Guianan Shields may be due to accumulation of lineages over a longer period of time. White-sand forests and SDTF are dominated by close relatives from fewer lineages, perhaps reflecting ecophysiological barriers that are difficult to surmount evolutionarily. Because MPD and ses.MPD do not reflect lineage diversity *per se*, we suggest that PDss, ses.PDss and ses.MNTD may be the most useful diversity metrics for setting large-scale conservation priorities.

Keywords

Amazon basin, Eudicots, Magnoliids, Monocots, phylogenetic diversity, species richness.

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INTRODUCTION

A central task of biology is to quantify biodiversity and how it varies geographically (Myers *et al.*, 2000). Elucidating and understanding the patterns of diversity is particularly important within the tropics, because of their high species richness and the pressing need to develop and apply effective conservation strategies in the face of massive habitat alteration. While the species diversity of specific areas can be measured using different indices (e.g. species richness, Fisher's alpha), these ecological metrics may fail to account for the evolutionary, or lineage, diversity of communities. As a result, some authors have advocated developing and implementing metrics, such as phylogenetic diversity, which quantify the lineage diversity of communities (Vane-Wright *et al.*, 1991; Faith, 1992).

Phylogenetic diversity (PD) is generally estimated as the total branch length of a phylogeny representing the species in a community (PD_{ss}; Faith, 1992). Alternative metrics to represent the evolutionary diversity in communities are available, such as the mean phylogenetic distance between all species and the mean phylogenetic distance between each species and its closest relative (MPD and MNTD respectively; Webb *et al.*, 2002; Helmus *et al.*, 2007; Cadotte *et al.*, 2010). All these metrics are often correlated with species richness (SR; the total number of species in a community), and thus SR can sometimes be used as a proxy for PD (Polasky *et al.*, 2001; Rodrigues & Gaston, 2002). However, some areas contain significantly greater or less PD than expected given their SR (Sechrest *et al.*, 2002; Forest *et al.*, 2007), and null model approaches have been developed to estimate PD while controlling for variation in SR (Kembel *et al.*, 2010). These standardized metrics may add complementary information about the evolutionary history and conservation significance of sites (Winter *et al.*, 2013). The availability of these recently developed PD metrics, in conjunction with the advent of standardized floristic sampling across Amazonia (Malhi *et al.*, 2002; Phillips & Miller, 2002) and a robust angiosperm phylogeny (Bremer *et al.*, 2009), now make it possible to examine how PD varies at large spatial scales across the world's most species-rich forest (Gentry, 1988; ter Steege *et al.*, 2013; see also Chave *et al.*, 2007).

Previous research has shown tree species diversity in 1 ha plots across the Amazon to be highest in its western and central regions and lowest in the east, on the Guianan and Brazilian shields (ter Steege *et al.*, 2003). Because PD is correlated with SR, we would expect that PD is greatest in the western and central Amazon, but this has yet to be thoroughly tested (although see Chave *et al.*, 2007). In addition, numerous factors may drive spatial variation in PD and whether communities show greater or less PD than expected given their SR. For example based on variation in substrate age, one might hypothesize that tree communities on the Guiana and Brazilian Shields, which overlay substrates of ancient Pre-Cambrian origin (Quesada *et al.*, 2011), might

have higher PD than expected given their relatively low SR. This high PD would reflect accumulated lineage diversity over tens of millions of years, with many deep phylogenetic branches separating species from these older diversification events (Swenson, 2009). In contrast, tree communities of western Amazonia overlying Pliocene and Pleistocene sediments from the Andes (Hoorn *et al.*, 2010; Quesada *et al.*, 2011) might be expected to show lower PD than expected given their high SR because of the dominance of recent evolutionary radiations of certain clades within which phylogenetic branches are short (Richardson *et al.*, 2001; Erkens *et al.*, 2007).

Soil fertility and precipitation seasonality also vary across Amazonia. Overall, the relatively young soils of western Amazonia are fertile in comparison with the highly weathered soils of central and eastern Amazonia and the Guianan and Brazilian Shields, whereas the poorest soils are found beneath white-sand forests that occur sporadically in small to large patches throughout the northern part of the basin (Quesada *et al.*, 2011). In addition, the dry season varies from being essentially absent in the north-west to lasting 5–6 months in the south-east and some northern areas (Sombroek, 2001), where moist forests give way to savannas and seasonally dry tropical forest (SDTF). Some of these environmental conditions may represent ecophysiological barriers that few lineages have been able to overcome (Anacker & Harrison, 2012; Miller *et al.*, 2013). Thus, an additional hypothesis to the one above, based on substrate age, is that tree communities in areas of the Amazon with greater ecophysiological barriers to growth (i.e. potentially more stressful environments) will show the lowest phylogenetic diversity (Qian *et al.*, 2013).

We used a network of 283 forest inventory plots (RAINFOR; Malhi *et al.*, 2002) to quantify the PD of tree communities and examine its spatial and environmental variation across Amazonia. We rarefied all plots to the same number of individuals, and then calculated (1) the total phylogenetic branch length of all species occurring in each plot, PD *sensu stricto* (PD_{ss}; Faith, 1992), (2) the mean pairwise phylogenetic distance between species (MPD; Webb, 2000; Webb *et al.*, 2002), and (3) the mean nearest taxon distance (MNTD; Webb, 2000). We also calculated standardized versions of these metrics that account for variation in SR. We then tested the hypothesis, based on substrate age, that tree communities in the Guiana and Brazilian Shields will show the greatest PD, whereas those in the western Amazon will show lower PD. And while our sample size outside of typical terra firme and floodplain moist forest is limited, we conducted a preliminary test of the hypothesis that tree communities in potentially more stressful environments, namely white sands, savannas, and SDTFs, will show the lowest PD. By examining the phylogenetic diversity of tree communities throughout Amazonia, we aim to provide insights into its biogeographical history and to inform the setting of conservation priorities.

METHODS

Tree community plot data

In this study, we used a total of 283 inventory plots of the RAINFOR forest plot network curated at Forest-Plots.net (see Table S1 in Supporting Information). Plots are generally one hectare in size (mean \pm SD = 1.1 ± 0.6 ha) and with all trees ≥ 10 cm diameter at breast height (DBH) sampled. We restricted analyses to old-growth forest plots and excluded plots with limited species identifications. Each plot was treated as a community and classified into three main biomes (Fig. 1): tropical moist forest, TMF ($n = 265$ plots), seasonally dry tropical forest, SDTF ($n = 13$), and savanna, S ($n = 5$). Fourteen plots were from the northern Andes (Colombia and Venezuela), outside the Amazon basin, but were included because of their close phylogeographical connection to Amazonia. SDTF plots are located from Bolivia to Venezuela, whereas savanna plots are only from Brazil and are separated by a maximum of 250 km.

The 265 tropical moist forest plots were further classified by the maximum age of the underlying geological formation. The Guiana and Brazilian Shields represent the oldest geological formations in Amazonia (TMF.o: > 500 Ma), followed by formations of central and eastern Amazonia (TMF.i: 20–100 Ma) located between the Shields, whereas areas near to the Andes (western Amazonia and northern Andes) are dominated by younger sediments (TMF.y: < 20 Ma; Quesada *et al.*, 2011) deposited mainly during the Pliocene and the

Pleistocene (Hoorn *et al.*, 2010) (Fig. 1). All TMF plots were also classified by forest type: montane forest, flooded forest, terra firme forest, and white-sand forest. Terra firme and flooded forests were sampled for each substrate age category, whereas montane forests were only sampled in western Amazonia on young substrates and white-sand forests were not sampled on substrates of intermediate age (see Table S1).

In total, the initial dataset included 183,908 individual trees sampled in Bolivia, Brazil, Colombia, Ecuador, French Guiana, Guyana, Peru, Surinam and Venezuela. To ensure a standardized nomenclature across plots based on the APG-III classification (Bremer *et al.*, 2009), the Taxonomic Name Resolution Service version 3.0 was used (<http://tnrs.iplantcollaborative.org>; accessed on 01/03/2013). Tree ferns and gymnosperms only occur in significant numbers in montane plots, and they are exceedingly rare in lowland forest, which is the focus of this study. These very rare species represent 0.018% of all individual trees in our lowland plots and are essentially stochastically sampled in any given 1 ha plot (tree ferns and gymnosperms were found in a total of nine and two lowland plots respectively). Given this stochasticity and the strong effect of tree ferns and gymnosperms on phylogenetic diversity metrics (they are subtended by very long phylogenetic branches; Faith *et al.*, 2004; Kembel & Hubbell, 2006; Chave *et al.*, 2007), we excluded them from phylogenetic diversity calculations. We also excluded all individuals not identified to a named species (13.6% of individuals). To determine if unidentified individuals could be biasing results, we assessed the correlation between the PD metrics and the proportion of unidentified individuals in each plot.

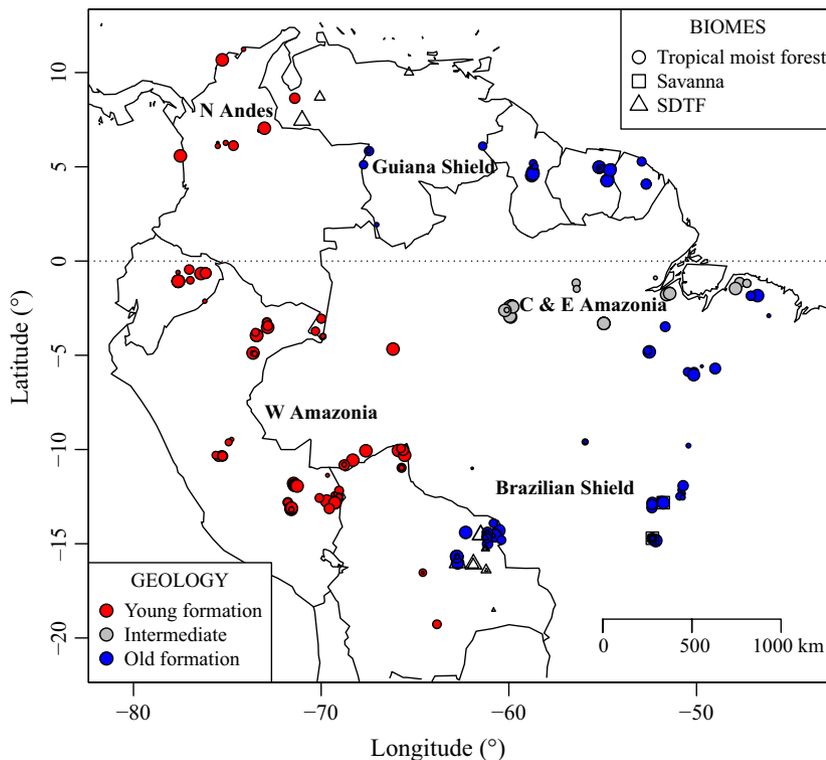


Figure 1 Location of 283 permanent RAINFOR plots classified by geological formation and biome in South America. Circle size represents species richness per plot (9 to 99 species for 249 rarefied individuals). Geographical regions used in the text are indicated in bold.

The final dataset contained a total of 157,340 individuals, belonging to 3868 species, 732 genera and 126 families of angiosperms.

Phylogenetic trees

A phylogenetic tree of the whole species pool (see Fig. S1) was generated using Phylomatic in PHYLOCOM version 4.2 (Webb *et al.*, 2008). This tool provides a phylogenetic hypothesis for the relationships among taxa by matching the list of species with up-to-date family and genus names, and tip labels of a provided megatree (Webb & Donoghue, 2005). In this case, the topology of R20120829.new provided at <http://phylodiversity.net/phyloomatic/> was used. An ultrametric phylogeny including branch length in millions of years (Ma) was obtained using *bladj* in PHYLOCOM. This command fixes the root node (angiosperms, 179 Ma) and other nodes to specified ages based on Wikström *et al.* (2001). Inconsistencies in syntax between internal node labels of the phylogeny and the ages file were modified manually to ensure a better performance of the node calibration using *bladj* (Gastauer & Meira-Neto, 2013). To determine if PD metrics are affected by phylogenetic resolution, we compared our results generated using the PHYLOCOM phylogeny with those using a phylogeny of Amazonian tree genera generated from DNA sequences of *rbcL* and *matK* plastid genes (K. G. Dexter & J. Chave, unpublished data). Full details of the temporally-calibrated, ultrametric phylogeny construction can be found in the Supporting information.

Phylogenetic diversity metrics

We used the PHYLOCOM phylogeny, which includes all genera in our dataset, to calculate six metrics that evaluate the evolutionary history present in communities: (1) the total phylogenetic branch length of all species occurring in a given community, i.e. phylogenetic diversity *sensu stricto* (PD_{ss}; Faith, 1992); (2) mean pairwise phylogenetic distance between species in terms of branch length (MPD; Webb, 2000; Webb *et al.*, 2002); (3) mean nearest taxon distance (MNTD; Webb, 2000; Webb *et al.*, 2002) and (4, 5 & 6) their equivalents, standardized for species richness (ses.PD_{ss}, ses.MPD, and ses.MNTD). For each community, these standardizations were accomplished by randomly drawing the same number of species from the phylogeny as present in the community, repeating this 1000 times, calculating PD_{ss}, MPD and MNTD for each randomization, taking the difference between the observed value of PD_{ss}, MPD, and MNTD and the mean of the random values, and dividing these differences by the standard deviation across the randomizations. These derived metrics therefore represent standardized effect sizes (ses) and are designated as such. ses.MPD and ses.MNTD are equivalent to the inverse of the NRI and NTI indices of Webb (2000). We consider the total phylogenetic branch length

(PD_{ss}) in communities (Faith, 1992; Forest *et al.*, 2007) and its deviation from expectation given species richness (ses.PD_{ss}) to be the most straightforward measures of evolutionary diversity in communities with respect to conservation prioritization. Lastly, we included the MPD, MNTD, ses.MPD, and ses.MNTD metrics of PD because of their history of use in the literature (e.g. Forest *et al.*, 2007; Gonzalez *et al.*, 2010; Fine & Kembel, 2011); MPD measures phylogenetic structure at deep nodes and MNTD at shallow nodes (Webb, 2000).

Data assessment and analysis

To minimize the effects of variation in sampling effort (i.e. plot size) and tree density, we used a rarefaction procedure that standardized all plots to 249 individuals, which was the lowest observed number of individual trees (≥ 10 cm DBH) among all plots. Values for PD_{ss}, MPD, MNTD, ses.PD_{ss}, ses.MPD, ses.MNTD and SR (the total number of species) for each rarefied community were calculated using the package *picante* (Kembel *et al.*, 2010) in the R STATISTICAL SOFTWARE version 2.15.1. PD metrics can also be sensitive to the most basal clades in a phylogeny (Swenson, 2009), so we classified taxa into one of the three major angiosperm clades (Magnoliids including Chloranthales, Monocots, and Eudicots), and the percentage of species in each clade was calculated. The mean across 100 rarefactions of the PD metrics, SR, and the proportion of major clades were used in subsequent analyses.

The values of PD_{ss}, MPD, MNTD, ses.PD_{ss}, ses.MPD and ses.MNTD were compared among communities growing on substrates of different geologic ages and forest types using *F*-tests and Tukey tests. We additionally compared all communities in potentially more stressful environments (white-sand forests, savannas and SDTF) vs. all in potentially less stressful environments (terra firme and montane forests) using a *t* test. Flooded forests were excluded from the analysis of stressful habitats because intensity and length of flooding is known to vary among plots, but we lack precise information on this. We also assessed the correlation of PD metrics with SR, the proportions of species in major clades, and the latitude and longitude of plots.

We assessed if there was any bias to the phylogenetic diversity metrics with respect to unidentified individuals by examining the correlation between percentage of unidentified individuals in plots and the various PD metrics. We also re-analysed a subset of the data ($n = 117$ plots each with >500 trees), rarefying the plots to 500 individuals per sampling unit, in order to test the effect of sample size in the rarefaction procedure on estimating phylogenetic diversity. Finally, we re-analysed a subset of the data ($n = 257$ plots), including plots that have more than 80% of species and individuals sampled in the sequenced-based genus-level phylogeny, in order to test the effect of phylogenetic resolution on estimating phylogenetic diversity. The random resolution of species-level relationships within genera in the

genus-level phylogeny was repeated for each set of rarefied communities.

RESULTS

Species richness and major angiosperm clades

Terra firme moist forests of intermediate and young geological formations have the highest species richness (SR), with an average of 88 and 72 species respectively (for 249 rarefied individuals; Table 1). Flooded moist forest communities in western and central Amazonia had greater SR than flooded and terra firme forests on the Guiana and Brazilian Shields, whereas the lowest SR was found in white-sand forests of the Guiana Shield and Andean montane forests (Table 1). SDTF and savannas show intermediate values of SR, resembling values of forest types on old geological formations.

On average, 85.8% of species per plot belong to Eudicots, 11.1% to Magnoliids and 3.1% to Monocots. Early diverging clades such as Magnoliids and Monocots tend to have a higher percentage of species on young geological formations than on intermediate and old formations, whereas Eudicots show the opposite pattern (Table 1). SDTF shows the lowest percentage of Magnoliid and Monocot species, and the greatest of Eudicots, but the abundance of these clades in savannas is more similar to the values typical of the moist forest plots.

Phylogenetic diversity metrics

Species richness strongly correlates with PDss ($r = 0.98$, $P < 0.001$; Fig. 2a) and MNTD ($r = -0.89$, $P < 0.001$; Fig. 2c), following a power relationship (PDss = $230.6 \cdot \text{SR}^{0.7}$, $r^2 = 0.96$; MNTD = $361.3 \cdot \text{SR}^{-0.4}$, $r^2 = 0.79$, both $P < 0.001$), which was a better fit than a linear relationship for both metrics (PDss = $1160.0 + 37.1 \cdot \text{SR}$, $r^2 = 0.92$; MNTD = $121.0 - 0.6 \cdot \text{SR}$, $r^2 = 0.71$, both $P < 0.001$). A much weaker correlation was observed between species richness and MPD ($r = 0.38$, $P < 0.001$; see Fig. S2). In contrast, the percentage of species in Magnoliids + Monocots (i.e. = 1–Eudicots) correlates strongly with MPD ($r = 0.88$, $P < 0.001$; Fig. 2b), which is driven mostly by variation in the relative abundance of Magnoliids ($r = 0.88$, $P < 0.001$) rather than Monocots ($r = 0.27$, $P < 0.001$). These correlations reflect the fact that communities with more Magnoliids and Monocots have a more even distribution of species across the three major angiosperm clades (see Fig. S3); a perfectly even split (1/3 in each clade) would give the highest value for MPD. The correlations of the percentages of species in major clades with PDss ($r_{1\text{-Eudicots}} = 0.52$, $r_{\text{Magnoliids}} = 0.48$, $r_{\text{Monocots}} = 0.26$, all $P < 0.001$) and MNTD ($r_{1\text{-Eudicots}} = 0.39$, $r_{\text{Magnoliids}} = 0.40$, $r_{\text{Monocots}} = 0.10$, all $P < 0.001$) were weaker (see Fig. S2). Both MPD and MNTD were strongly correlated with their standardized equivalents (MPD and ses.MPD: $r = 0.94$,

Table 1 Community composition and diversity across forest types, showing proportional representation of major clades and mean values of species richness (SR) and phylogenetic diversity. Phylogenetic diversity *sensu stricto* (PDss), mean pairwise phylogenetic distance between species (MPD) and mean nearest taxon distance (MNTD) are given in millions of years (Ma), whereas ses.PDss, ses.MPD and ses.MNTD are standardized metrics without units

Biome (max. geological age)	Forest type	No. of plots	Sample area (ha)	ID to spp (%) [*]	Species (mean, %)			Mean diversity values						
					Magnoliids	Monocots	Eudicots	PDss (Ma)	MPD (Ma)	MNTD (Ma)	ses. PDss	ses. MPD	ses. MNTD	
Tropical moist forest (< 20 Ma)	Flooded	12	17	86	16	6	78	72	3963	260	74	-1.24	0.48	-0.99
	Montane	16	16	80	12	1	87	29	2180	255	105	-0.41	0.03	-0.47
	Terra firme	86	95	85	14	5	81	78	4148	256	74	-1.42	0.15	-1.10
	White sand	4	4	83	10	5	85	42	2839	254	106	0.06	-0.12	-0.07
Tropical moist forest (20–100 Ma)	Flooded	2	2	73	9	0	90	72	3478	242	62	-3.44	-1.59	-2.69
	Terra firme	39	54	85	12	1	87	88	4200	248	64	-2.86	-0.90	-2.11
Tropical moist forest (> 500 Ma)	Flooded	17	16	89	5	2	93	34	2368	238	111	-0.59	-1.22	-0.23
	Terra firme	85	94	87	10	3	87	56	3310	247	85	-1.07	-0.91	-0.79
	White sand	4	4	87	7	0	93	22	1608	233	97	-1.90	-1.41	-1.53
Savanna	Savanna	5	4	100	5	2	93	47	3105	239	88	-0.28	-1.56	-0.63
SDTF	Dry forest	13	14	96	2	3	95	34	2195	224	89	-2.12	-2.42	-1.30
Total		283	320	86	11	3	86	63	3510	249	81	-1.41	-0.57	-1.06

^{*}The mean proportion of individuals identified to species.

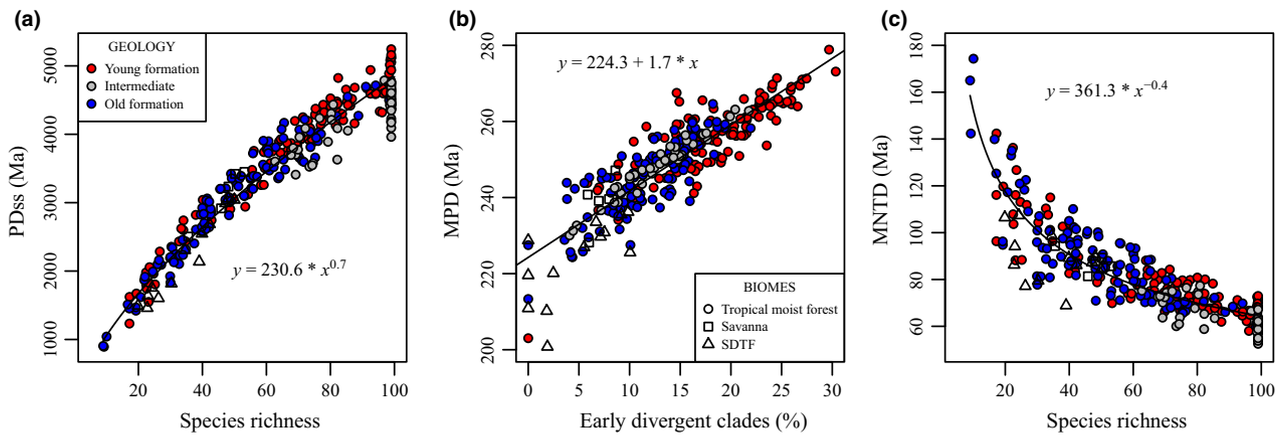


Figure 2 Relationship between (a) phylogenetic diversity *sensu stricto* and species richness, between (b) mean pairwise phylogenetic distance between species and the proportion of species of Magnoliids and Monocots (= 1–Eudicots), and between (c) mean nearest taxon distance and species richness. Tropical moist forest biome is classified based on maximum age of geological formations (young: < 20 Ma; intermediate: 20–100 Ma, old: > 500 Ma; Quesada *et al.*, 2011).

$P < 0.001$; MNTD and ses.MNTD: $r = 0.71$, $P < 0.01$; see Fig. S2), and show similar results with respect to the plot groupings we considered (e.g. substrate age, forest type). We therefore present results for only ses.MPD and ses.MNTD below. Interestingly, ses.PDss and ses.MNTD are strongly positively correlated with each other ($r = 0.95$, $P < 0.001$; see Fig. S2), whereas neither shows a strong relationship with ses.MPD (ses.PDss and ses.MPD: $r = 0.23$, $P < 0.001$; ses.MNTD and ses.MPD: $r = 0.09$, $P = 0.14$; see Fig. S2).

Geographical and environmental patterns

All of the PD metrics show non-random spatial distributions across Amazonia (Fig. 3a–d). While PDss shows weak relationships with both latitude and longitude ($r_{\text{Latitude}} = 0.14$, $P < 0.05$; $r_{\text{Longitude}} = -0.16$, $P < 0.05$) and ses.PDss and ses.MNTD show weak latitudinal gradients (ses.PDss: $r_{\text{Latitude}} = -0.18$, $P < 0.01$; ses.MNTD: $r_{\text{Latitude}} = -0.17$, $P < 0.01$), ses.MPD shows a strong longitudinal gradient decreasing from west to east ($r_{\text{Longitude}} = -0.52$, $P < 0.001$). PDss was greatest in communities on young and intermediate-aged geological formations (Fig. 3e), whereas ses.PDss and ses.MNTD were greatest in communities on young and old geological formations (Fig. 3f,h). ses.MPD was greatest in young geological formations (Fig. 3g). These spatial patterns are conserved across the different forest types within the moist forest biome (e.g. terra firme, floodplain) (see Table 1).

Among the moist forest communities, montane and white-sand forests have the lowest PDss values and high values for ses.PDss, ses.MPD and ses.MNTD, at least in western Amazonia. In contrast to this, flooded and terra firme forests in central Amazonia have high PDss and the lowest values for ses.PDss, ses.MPD and ses.MNTD (Table 1). For all metrics, PD values of savannas were similar to moist forest communities, whereas SDTF consistently showed lower PD (Fig. 3e–h). Overall, PDss and ses.MPD were significantly

lower in potentially more stressful habitats, i.e. savanna, SDTF and white-sand forest (mean \pm 95% confidence interval: 2379 ± 305 Ma and -1.75 ± 0.40 respectively), than potentially less stressful habitats (3702 ± 118 Ma and -0.44 ± 0.14 ; $t_{\text{PDss}} = 8.28$, d.f. = 34, $P < 0.001$ and $t_{\text{ses.MPD}} = 6.27$, d.f. = 32, $P < 0.001$), whereas ses.PDss and ses.MNTD were not significantly different between the two (ses.PDss: -1.39 ± 0.50 vs. -1.47 ± 0.17 , $t_{\text{ses.PDss}} = -0.28$, d.f. = 31, $P = 0.78$; and ses.MNTD: -1.02 ± 0.35 vs. -1.11 ± 0.13 , $t_{\text{ses.MNTD}} = -0.51$, d.f. = 33, $P = 0.61$).

PDss ($r^2 = 0.002$, $P = 0.20$), ses.MPD ($r^2 = 0.004$, $P = 0.15$), and ses.MNTD ($r^2 = 0.007$, $P = 0.08$) showed no relationship with the percentage of unidentified individuals excluded per plot, whereas ses.PDss ($r^2 = 0.013$, $P < 0.05$) shows a very weak relationship (see Fig. S4). In addition, for plots with sufficient sample size to assess, we found a strong 1 : 1 relationship between phylogenetic diversity metrics (PDss, ses.PDss, ses.MPD, and ses.MNTD) calculated with rarefactions of 500 vs. 249 individuals (see Fig. S5). We also found that the patterns of PD metrics across Amazonia were qualitatively identical when using the sequenced-based genus-level phylogeny vs. the PHYLOCOM phylogeny (see Figs S6 & S7).

DISCUSSION

Our study has revealed a highly non-random spatial and environmental distribution of phylogenetic diversity (PD) across tree communities of Amazonia, by whichever metric it is evaluated, with some areas and environments holding significantly more, or less, phylogenetic diversity than others (Fig. 3). Phylogenetic diversity *sensu stricto* (PDss) and the mean nearest taxon distance (MNTD) in the Amazon correlate strongly with species richness (SR; Fig. 2a,c) following positive and negative trends, respectively. Therefore, diverse communities on young- and intermediate-aged substrates

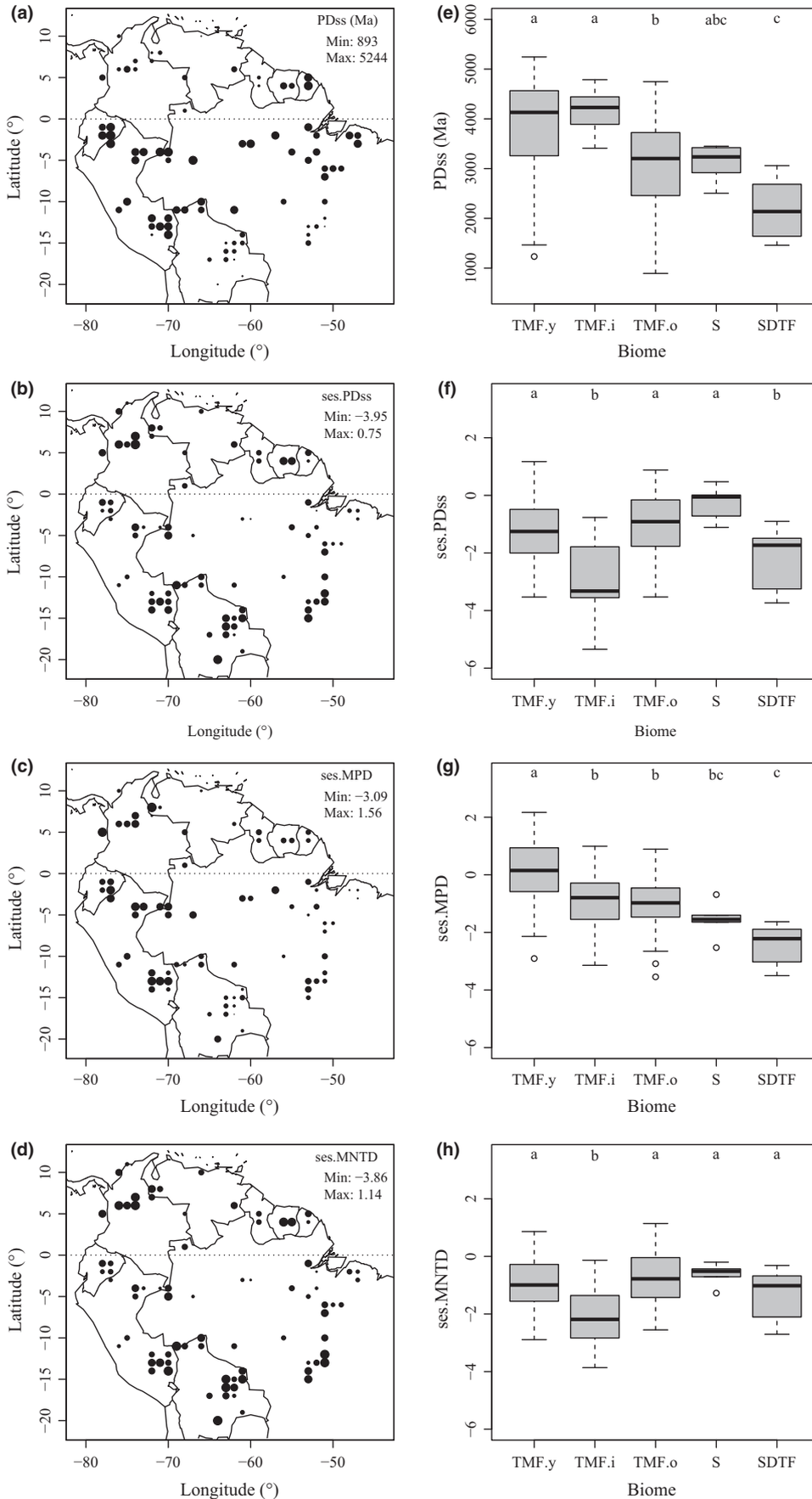


Figure 3 Variation in phylogenetic diversity, as evaluated by several metrics, across Amazonia. The results for phylogenetic diversity *sensu stricto* (PDss), its equivalent standardized for variation in species richness (ses.PDss), and the standardized measures of mean pairwise phylogenetic distance between species (ses.MPD) and mean nearest taxon distance (ses.MNTD) are shown in different rows. (a–d) The maps show the spatial distribution of values for each metric, with the size of circles corresponding to their values. If there were multiple plots in a given one-degree grid, the mean value is shown. (e–h) The tropical moist forest biome is classified based on maximum age of geological formations (TMF.y: < 20 Ma; TMF.i: 20–100 Ma, TMF.o: > 500 Ma), whereas savanna and seasonally dry tropical forest are indicated as S and SDTF respectively. Letters in boxplots indicate significant difference among mean values (Tukey's HSD; $P < 0.05$).

(western and central Amazonia respectively) have the highest PDss and the lowest MNTD values. Once variation in SR is controlled for, we found that the youngest and oldest substrates (the latter on the Brazilian and Guiana Shields) have the highest ses.PDss and ses.MNTD. The lowest values of ses.PDss and ses.MNTD were found in potentially more stressful environments, in particular white-sand forest and SDTF.

We also found that the mean pairwise phylogenetic distance between species (MPD) and its standardized equivalent, ses.MPD, depend primarily on how evenly taxa are distributed among the three major angiosperm clades (Magnoliids, Monocots and Eudicots), which is shown by the strong positive correlation between their values and the proportion of taxa in plots that are Magnoliids and Monocots (the two rarer clades; Fig. 2b). Thus, communities in western Amazonia, that have many Magnoliids and Monocots present, have the greatest MPD and ses.MPD values. While it is important to have a measure of how evenly distributed taxa are across the major clades of a phylogeny, MPD and ses.MPD do not seem to reflect lineage diversity *per se*. Moreover, ses.PDss and ses.MNTD were strongly positively correlated, giving similar patterns across geological substrates environments. We therefore focus below primarily on patterns with respect to PDss and ses.PDss.

Has the greatest phylogenetic diversity been accumulated in communities overlaying old geological formations?

Communities on old geological substrates in the Brazilian and Guianan Shields showed lower PDss than communities on young or intermediately aged geological substrates (Fig. 3e), which is unsurprising given their lower species richness. The communities on old geological substrates did show a higher median ses.PDss (Fig. 3f), but the distribution of ses.PDss values overlapped broadly with those for communities on the youngest substrate. The same pattern was found for ses.MNTD. Thus, our prediction that PD would be positively correlated with substrate age was falsified. However, we suggest that different processes may explain the high ses.PDss values observed in different communities across Amazonia. The high ses.PDss and ses.MNTD found in the Guiana and Brazilian Shields may very well be explained by their long-term geological history and the accumulation of lineages over many millions of years.

To understand the rejection of the hypothesis that geologically older substrates show the greatest PD, we need to consider why tree communities of western Amazonia show such high ses.PDss and ses.MNTD. That communities of western Amazonia show high PDss is unsurprising, as PDss is strongly correlated with SR, and SR is substantially higher in the western Amazon (ter Steege *et al.*, 2003). However, much of this species diversity is due to recently radiated species-rich genera (Gentry, 1982) such as *Inga* (Richardson *et al.*, 2001) and *Guatteria* (Erkens *et al.*, 2007), and short phylogenetic branches such as those within these genera do not greatly increase PD (Swenson, 2009). Moreover, low MNTD would be explained by the presence of short phylogenetic branches separating the nearest taxa in these diverse communities. However, another exceptional aspect of western Amazonian tree communities is that they are occupied by lineages from the entirety of the angiosperm phylogeny, which leads these communities to have high ses.PDss, and apparently also high ses.MNTD. One explanation might be related to the potentially high phylogenetic diversity found in the adjacent Andes, which provides a proximate resource to 'invade' western Amazonia (see also Chave *et al.*, 2007). Another explanation might be related to the particular environmental and ecological conditions (relatively fertile and aseasonal environments) in the west, which may be easier to invade by multiple lineages with diverse evolutionary backgrounds. Moreover, the ability of diverse lineages to establish in the western and southern Amazon may also be related to the high rates of disturbance and turnover in the region (Quesada *et al.*, 2012; Marimon *et al.*, 2013; Baker *et al.*, 2014). Thus, in the same way that more fertile, dynamic, and disturbed tropical forests have more open nutrient-cycles on ecological time-scales (Vitousek & Sanford, 1986), they also appear to be more open to repeated establishment of plant lineages on evolutionary time-scales.

Do environments with more potential ecophysiological barriers to growth show the lowest PD in their tree communities?

We expected that environments with potentially more stressful ecological conditions, namely marked seasonality of precipitation and/or low soil fertility, would have the lowest phylogenetic diversity, because these may represent ecophysiological barriers that are difficult for many lineages to surmount evolutionarily (Anacker & Harrison, 2012; Miller *et al.*, 2013; Qian *et al.*, 2013). Both savannas and SDTF

have a pronounced dry season, but they show contrasting patterns of PD. While PD metrics of savannas were similar to those of nearby communities in tropical moist forest, SDTF generally has low PD (Fig. 3e–h). Savannas and tropical moist forest communities may share similar lineages across the angiosperm phylogeny, a pattern which supports previous studies that suggested that Brazilian savannas are formed by the numerous independent colonizations of lineages from nearby biomes around 4–10 Ma (Simon *et al.*, 2009; Simon & Pennington, 2012). Conversely, the low PD values shown for SDTF communities suggest that fewer clades have succeeded in colonizing SDTF, and that consequently, SDTF is occupied by closer relatives. However, our conclusions must be taken as preliminary given the low sample size and limited geographical extent of our savanna and SDTF plots.

Previous studies have indicated strong habitat specialization in white-sand communities as indicated by the high number of individuals that represent white-sand specialist species (Fine *et al.*, 2010), and by the distinct ecophysiology and defences against herbivores that these species have evolved in order to live on such poor soils (Fine *et al.*, 2004). Therefore, we also expected that white-sand forests would have a high frequency of closely related species and low phylogenetic diversity. But while our results showed that both white-sand communities of the Guiana Shield and the western Amazon have low PDss, only those communities in the Guyana Shield have low ses.PDss values compared to neighbouring terra firme or flooded forest. We found higher values of ses.PDss in the small patches of white-sand forests of western Amazonia than in the Guiana Shield, suggesting a greater influence of the regional pool (i.e. species present in the surrounding phylogenetically diverse terra firme forest entering white-sand patches) than in the larger, more contiguous white-sand patches of the Guiana Shield.

Conservation priorities

Conservation planning based upon species richness (SR) gives the same value to communities with equal SR regardless of the total phylogenetic diversity of the species that they contain (e.g. Forest *et al.*, 2007). But if we are to preserve the full spectrum of lineage diversity and the evolutionary processes that led to the exceptional biodiversity of Amazonian communities, regional conservation planning must incorporate phylogenetic information.

In this study, we showed that while PDss is strongly correlated with SR (see also Forest *et al.*, 2007; Cadotte *et al.*, 2012), communities can vary greatly in their deviation from expected PD given SR, as measured by ses.PDss. While communities in the central and western Amazon have the greatest tree species richness in the basin (ter Steege *et al.*, 2003), the central Amazon shows much lower phylogenetic diversity than expected given its species richness (ses.PDss

compared to the western Amazon (Fig. 3e), thus suggesting that the western Amazon basin may hold a higher value for conservation of lineage diversity.

In addition, we found that the mean pairwise phylogenetic distance between species (MPD) is not strongly correlated with species richness, which could suggest that it is a better metric of phylogenetic diversity than PDss. However, we found that MPD and its standardized equivalent (ses.MPD) are strongly dependent on how evenly divided the species in a tree community are among the three major angiosperm clades (Magnoliids, Monocots and Eudicots; Fig. 2b). While this division is certainly interesting from an ecological and evolutionary perspective, we suggest that MPD and ses.MPD may not be the most useful metrics of phylogenetic diversity for conservation prioritization. Meanwhile, MNTD shows a strong inverse relationship with SR, and ses.MNTD essentially conveys the same information as ses.PDss (i.e. they are strongly positively correlated). Thus, we suggest that, PDss and ses.PDss may provide the most straightforward, interpretable means to evaluate lineage diversity in communities. While PDss is strongly correlated with SR and could perhaps be inferred from it, a phylogeny is clearly necessary to calculate ses.PDss and determine whether communities show more or less lineage diversity than expected given their species richness. An urgent priority for conservation should be to develop bigger community phylogenies that include all lineages, greater numbers of species within lineages, and greater phylogenetic resolution. Such phylogenies would allow evolutionary information to be properly incorporated into conservation decisions.

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REFERENCES

- Anacker, B.L. & Harrison, S.P. (2012) Historical and ecological controls on phylogenetic diversity in Californian plant communities. *The American Naturalist*, **180**, 257–269.
- Baker, T.R., Pennington, R.T., Magallon, S. *et al.* (2014) Fast demographic traits promote high diversification rates of Amazonian trees. *Ecology Letters*, **17**, 527–536.
- Bremer, B., Bremer, K., Chase, M.W., Fay, M.F., Reveal, J.L., Soltis, D.E., Soltis, P.S., Stevens, P.F., Anderberg, A.A., Moore, M.J., Olmstead, R.G., Rudall, P.J., Sytsma, K.J., Tank, D.C., Wurdack, K., Xiang, J.Q.Y., Zmarzty, S. & Angiosperm Phylogeny, G. (2009) An update of the Angiosperm Phylogeny Group classification for the orders and families of flowering plants: APG III. *Botanical Journal of the Linnean Society*, **161**, 105–121.
- Cadotte, M.W., Jonathan Davies, T., Regetz, J., Kembel, S.W., Cleland, E. & Oakley, T.H. (2010) Phylogenetic diversity metrics for ecological communities: integrating species richness, abundance and evolutionary history. *Ecology Letters*, **13**, 96–105.
- Cadotte, M.W., Dinnage, R. & Tilman, D. (2012) Phylogenetic diversity promotes ecosystem stability. *Ecology*, **93**, S223–S233.
- Chave, J., Chust, G. & Thebaud, C. (2007) The importance of phylogenetic structure in biodiversity studies. *Scaling biodiversity* (ed. by D. Storch, P.L. Marquet and J.H. Brown), pp. 150–167. Cambridge University Press, Cambridge, UK.
- Erkens, R.H.J., Chatrou, L.W., Maas, J.W., van der Niet, T. & Savolainen, V. (2007) A rapid diversification of rainforest trees (*Guatteria*; Annonaceae) following dispersal from Central into South America. *Molecular Phylogenetics and Evolution*, **44**, 399–411.
- Faith, D.P. (1992) Conservation evaluation and phylogenetic diversity. *Biological Conservation*, **61**, 1–10.
- Faith, D.P., Reid, C.A.M. & Hunter, J. (2004) Integrating phylogenetic diversity, complementarity, and endemism for conservation assessment. *Conservation Biology*, **18**, 255–261.
- Fine, P.V.A. & Kembel, S.W. (2011) Phylogenetic community structure and phylogenetic turnover across space and edaphic gradients in western Amazonian tree communities. *Ecography*, **34**, 552–565.
- Fine, P.V.A., Mesones, I. & Coley, P.D. (2004) Herbivores promote habitat specialization by trees in Amazonian forests. *Science*, **305**, 663–665.
- Fine, P.V.A., García-Villacorta, R., Pitman, N.C.A., Mesones, I. & Kembel, S.W. (2010) A floristic study of the white-sand forests of Peru. *Annals of the Missouri Botanical Garden*, **97**, 283–305.
- Forest, F., Grenyer, R., Rouget, M., Davies, T.J., Cowling, R.M., Faith, D.P., Balmford, A., Manning, J.C., Proches, S., van der Bank, M., Reeves, G., Hedderson, T.A.J. & Savolainen, V. (2007) Preserving the evolutionary potential of floras in biodiversity hotspots. *Nature*, **445**, 757–760.
- Gastauer, M. & Meira-Neto, J.A.A. (2013) Avoiding inaccuracies in tree calibration and phylogenetic community analysis using Phylocom 4.2. *Ecological Informatics*, **15**, 85–90.
- Gentry, A.H. (1982) Neotropical floristic diversity: phytogeographical connections between Central and South-America, Pleistocene climatic fluctuations, or an accident of the Andean orogeny. *Annals of the Missouri Botanical Garden*, **69**, 557–593.
- Gentry, A.H. (1988) Tree species richness of upper Amazonian forests. *Proceedings of the National Academy of Sciences USA*, **85**, 156–159.
- Gonzalez, M.A., Roger, A., Courtois, E.A., Jabot, F., Norden, N., Paine, C.E.T., Baraloto, C., Thébaud, C. & Chave, J. (2010) Shifts in species and phylogenetic diversity between sapling and tree communities indicate negative density dependence in a lowland rain forest. *Journal of Ecology*, **98**, 137–146.
- Helmus, M.R., Bland, T.J., Williams, C.K. & Ives, A.R. (2007) Phylogenetic measures of biodiversity. *The American Naturalist*, **169**, E68–E83.
- Hoorn, C., Wesselingh, F.P., ter Steege, H., Bermudez, M.A., Mora, A., Sevink, J., Sanmartín, I., Sanchez-Meseguer, A., Anderson, C.L., Figueiredo, J.P., Jaramillo, C., Riff, D., Negri, F.R., Hooghiemstra, H., Lundberg, J., Stadler, T., Särkinen, T. & Antonelli, A. (2010) Amazonia through time: Andean uplift, climate change, landscape evolution, and biodiversity. *Science*, **330**, 927–931.
- Kembel, S.W. & Hubbell, S.P. (2006) The phylogenetic structure of a Neotropical forest tree community. *Ecology*, **87**, S86–S99.
- Kembel, S.W., Cowan, P.D., Helmus, M.R., Cornwell, W.K., Morlon, H., Ackerly, D.D., Blomberg, S.P. & Webb, C.O. (2010) Picante: R tools for integrating phylogenies and ecology. *Bioinformatics*, **26**, 1463–1464.
- Malhi, Y., Phillips, O.L., Lloyd, J. *et al.* (2002) An international network to monitor the structure, composition and dynamics of Amazonian forests (RAINFOR). *Journal of Vegetation Science*, **13**, 439–450.
- Marimon, B.S., Marimon-Junior, B.H., Feldpausch, T.R., Oliveira-Santos, C., Mews, H.A., Lopez-Gonzalez, G., Lloyd, J., Franczak, D.D., de Oliveira, E.A., Maracahipes, L., Miguel, A., Lenza, E. & Phillips, O.L. (2013) Disequilibrium and hyperdynamic tree turnover at the forest–cerrado transition zone in southern Amazonia. *Plant Ecology & Diversity*, **7**, 281–292.
- Miller, E.T., Zanne, A.E. & Ricklefs, R.E. (2013) Niche conservatism constrains Australian honeyeater assemblages in stressful environments. *Ecology Letters*, **16**, 1186–1194.
- Myers, N., Mittermeier, R.A., Mittermeier, C.G., da Fonseca, G.A.B. & Kent, J. (2000) Biodiversity hotspots for conservation priorities. *Nature*, **403**, 853–858.

- Phillips, O. & Miller, J. (2002) *Global patterns of plant diversity: Alwyn H. Gentry's forest transect data set*. Missouri Botanical Garden Press, St. Louis, MO.
- Polasky, S., Csuti, B., Vossler, C.A. & Meyers, S.M. (2001) A comparison of taxonomic distinctness versus richness as criteria for setting conservation priorities for North American birds. *Biological Conservation*, **97**, 99–105.
- Qian, H., Zhang, Y., Zhang, J. & Wang, X. (2013) Latitudinal gradients in phylogenetic relatedness of angiosperm trees in North America. *Global Ecology and Biogeography*, **22**, 1183–1191.
- Quesada, C.A., Lloyd, J., Anderson, L.O., Fyllas, N.M., Schwarz, M. & Czimczik, C.I. (2011) Soils of Amazonia with particular reference to the RAINFOR sites. *Biogeosciences*, **8**, 1415–1440.
- Quesada, C.A., Phillips, O.L., Schwarz, M. *et al.* (2012) Basin-wide variations in Amazon forest structure and function are mediated by both soils and climate. *Biogeosciences*, **9**, 2203–2246.
- Richardson, J.E., Pennington, R.T., Pennington, T.D. & Hollingsworth, P.M. (2001) Rapid diversification of a species-rich genus of Neotropical rain forest trees. *Science*, **293**, 2242–2245.
- Rodrigues, A.S.L. & Gaston, K.J. (2002) Maximising phylogenetic diversity in the selection of networks of conservation areas. *Biological Conservation*, **105**, 103–111.
- Sechrest, W., Brooks, T.M., da Fonseca, G.A.B., Konstant, W.R., Mittermeier, R.A., Purvis, A., Rylands, A.B. & Gittleman, J.L. (2002) Hotspots and the conservation of evolutionary history. *Proceedings of the National Academy of Sciences USA*, **99**, 2067–2071.
- Simon, M.F. & Pennington, R.T. (2012) Evidence for adaptation to fire regimes in the tropical savannas of the Brazilian cerrado. *International Journal of Plant Sciences*, **173**, 711–723.
- Simon, M.F., Grether, R., de Queiroz, L.P., Skema, C., Pennington, R.T. & Hughes, C.E. (2009) Recent assembly of the Cerrado, a neotropical plant diversity hotspot, by *in situ* evolution of adaptations to fire. *Proceedings of the National Academy of Sciences USA*, **106**, 20359–20364.
- Sombroek, W. (2001) Spatial and temporal patterns of Amazon rainfall: consequences for the planning of agricultural occupation and the protection of primary forests. *Ambio*, **30**, 388–396.
- ter Steege, H., Pitman, N., Sabatier, D. *et al.* (2003) A spatial model of tree α -diversity and tree density for the Amazon. *Biodiversity and Conservation*, **12**, 2255–2277.
- ter Steege, H., Pitman, N.C.A., Sabatier, D. *et al.* (2013) Hyperdominance in the Amazonian tree flora. *Science*, **342**, 1243092.
- Swenson, N.G. (2009) Phylogenetic resolution and quantifying the phylogenetic diversity and dispersion of communities. *PLoS ONE*, **4**, e4390.
- Vane-Wright, R.I., Humphries, C.J. & Williams, P.H. (1991) What to protect?—Systematics and the agony of choice. *Biological Conservation*, **55**, 235–254.
- Vitousek, P.M. & Sanford, R. (1986) Nutrient cycling in moist tropical forest. *Annual Review of Ecology and Systematics*, **17**, 137–167.
- Webb, C.O. (2000) Exploring the phylogenetic structure of ecological communities: an example for rain forest trees. *The American Naturalist*, **156**, 145–155.
- Webb, C.O. & Donoghue, M.J. (2005) Phylomatic: tree assembly for applied phylogenetics. *Molecular Ecology Notes*, **5**, 181–183.
- Webb, C.O., Ackerly, D.D., McPeck, M.A. & Donoghue, M.J. (2002) Phylogenies and community ecology. *Annual Review of Ecology and Systematics*, **33**, 475–505.
- Webb, C.O., Ackerly, D.D. & Kembel, S.W. (2008) Phylocom: software for the analysis of phylogenetic community structure and trait evolution. *Bioinformatics*, **24**, 2098–2100.
- Wikström, N., Savolainen, V. & Chase, M.W. (2001) Evolution of the angiosperms: calibrating the family tree. *Proceedings of the Royal Society B: Biological Sciences*, **268**, 2211–2220.
- Winter, M., Devictor, V. & Schweiger, O. (2013) Phylogenetic diversity and nature conservation: where are we? *Trends in Ecology and Evolution*, **28**, 199–204.

SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Table S1 Floristic tree inventories compiled from RAINFOR forest plot network.

Table S2 Fossil-based calibrations used in sequenced-based genus-level phylogeny.

Figure S1 Phylogenetic tree for the whole species pool for 283 floristic inventories compiled from RAINFOR dataset.

Figure S2 Relationships and correlation values among phylogenetic diversity metrics, species richness and the proportion of species of Magnoliids and Monocots.

Figure S3 Distributions of species across the three major angiosperm clades for tree communities with increasing values of MPD.

Figure S4 Relationships between phylogenetic diversity metrics and the percentage of unidentified individuals excluded for each plot.

Figure S5 Sensitivity analysis of phylogenetic diversity metrics for the tropical moist forest biome calculated using different number of individuals per plot.

Figure S6 Variation in phylogenetic diversity metrics across Amazonia, calculated using phylogenies resolved to genus and family level.

Figure S7 Sensitivity analysis of phylogenetic diversity metrics calculated using genus- and family-level phylogenies.

BIOSKETCH

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