

Consistent patterns of common species across tropical tree communities

<https://doi.org/10.1038/s41586-023-06820-z>

Received: 3 December 2022

Accepted: 1 November 2023

Published online: 10 January 2024

Open access

 Check for updates

Trees structure the Earth's most biodiverse ecosystem, tropical forests. The vast number of tree species presents a formidable challenge to understanding these forests, including their response to environmental change, as very little is known about most tropical tree species. A focus on the common species may circumvent this challenge. Here we investigate abundance patterns of common tree species using inventory data on 1,003,805 trees with trunk diameters of at least 10 cm across 1,568 locations^{1–6} in closed-canopy, structurally intact old-growth tropical forests in Africa, Amazonia and Southeast Asia. We estimate that 2.2%, 2.2% and 2.3% of species comprise 50% of the tropical trees in these regions, respectively. Extrapolating across all closed-canopy tropical forests, we estimate that just 1,053 species comprise half of Earth's 800 billion tropical trees with trunk diameters of at least 10 cm. Despite differing biogeographic, climatic and anthropogenic histories⁷, we find notably consistent patterns of common species and species abundance distributions across the continents. This suggests that fundamental mechanisms of tree community assembly may apply to all tropical forests. Resampling analyses show that the most common species are likely to belong to a manageable list of known species, enabling targeted efforts to understand their ecology. Although they do not detract from the importance of rare species, our results open new opportunities to understand the world's most diverse forests, including modelling their response to environmental change, by focusing on the common species that constitute the majority of their trees.

Tropical forests are a crucial component of the Earth system; they cover around 10% of the Earth's land surface⁸ but contribute approximately 33% of terrestrial net primary productivity⁹. They account for around 40% of the carbon stored in live vegetation¹⁰ and are globally important carbon sinks¹¹. Tropical forests are also extraordinarily biodiverse, harbouring two-thirds of all known species¹² and the majority of the world's biodiversity hotspots¹³. Of note, as many tree species can be found in a single hectare of tropical forest as in the entire native Western European tree flora¹⁴. Recent estimates suggest that there are approximately 37,900 named tropical tree species in the scientific literature¹⁵, with potentially thousands more yet to be identified by scientists¹⁶. This extraordinary diversity means that little is known about the biology of the vast majority of tropical tree species. Our understanding of tropical forest ecology, productivity and carbon storage and how they may respond to environmental change is hindered by this lack of knowledge. This limited understanding also curtails scientific input into land use, biodiversity, climate and other forest-related policy and management.

Our understanding of tropical forests may improve through a focus on the most common tree species. This is a promising avenue, given that species abundance distributions (SADs) showing a modest number of common species and much larger numbers of rare species have been documented across taxa globally^{17–19}. Indeed, analyses of tropical forest inventory data from Amazonia have shown that a relatively small number of common species comprise a majority of trees in the region^{6,20–24}. However, whether such patterns hold in other tropical forests is unknown, as there have been no comparable analyses for African or Southeast Asian tropical forests. Perhaps, given the substantial

differences in total tree species richness²⁵, forest structure¹, contemporary climate²⁶ and biogeographic and human-occupancy histories⁷ among continents, important contrasts in patterns of common species would be expected. Alternatively, if the same processes or mechanisms apply to all tropical forests²⁷, highly consistent patterns may be expected. Crucially, if a tractably modest number of common species do comprise the majority of tropical trees on Earth, this could open new ways of understanding tropical forests by investigating the ecology of the common species.

Cross-continental comparisons of common species patterns are complicated by unresolved differences in the results from published Amazon forest studies^{6,20,22}. Estimates of hyperdominance—describing the minimum number of species required to account for 50% of all trees in a sample—range from 1.4% to 8.2% of the total number of species found in each of the Amazon forest datasets analysed (corresponding to 224 and 1,312 hyperdominant species respectively, assuming 16,000 Amazon tree species). Therefore, here we: (1) investigate sample-related biases and standardize our sampling to enable meaningful comparisons among datasets; (2) test whether patterns of hyperdominance differ across Amazonia, Africa and Southeast Asia; (3) extrapolate our results to assess how many species comprise half of all Earth's tropical trees; (4) assess species abundance patterns, with differing classifications of 'common species' beyond hyperdominance; and (5) use resampling techniques to assess which sampled species are likely to be hyperdominant.

We analyse species abundance data from networks of inventory plots across three continents. We limit our analysis to closed canopy

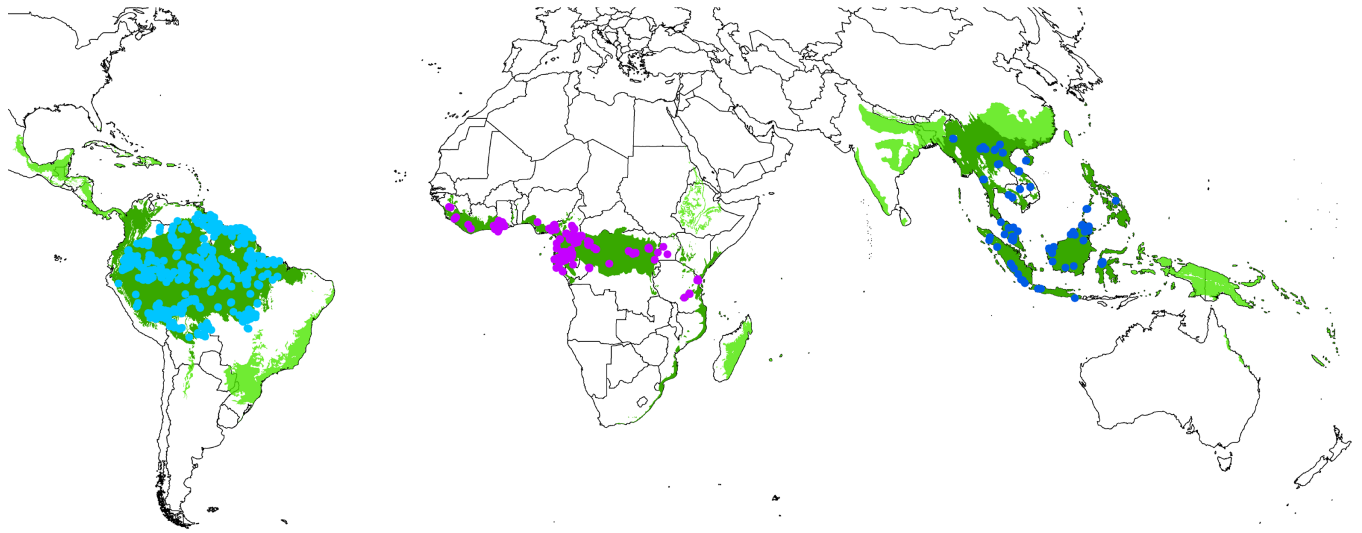


Fig. 1 | Location of the 1,568 plots, tropical forest regions, and tropical forest biome extent used in the study. Dots show the location of the plots analysed, coloured by continental region. Dark green shows the Amazonia,

Africa and Southeast Asia regions that we extrapolate to. Light green shows 'tropical and subtropical moist broadleaf forests'⁶⁰, which we extrapolate to as the closed canopy tropical forest biome.

structurally intact old-growth tropical forests. For Amazonia, defined as the lowland Amazon Basin and Guiana Shield, we use the Amazon Tree Diversity Network and RAINFOR datasets ($n = 1,097$ plots). For Africa, encompassing West, central and East Africa, we use the African Tropical Rainforest Observatory Network (AfriTRON)¹, Central African Plot Network, and two smaller networks^{2,3} ($n = 368$ plots). For Southeast Asia, defined as extending from Myanmar in the West to Sulawesi in the East, we use a tree diversity⁴ and a carbon monitoring⁵ network ($n = 103$ plots). We limit our analysis to trees with trunk diameter of at least 10 cm at breast height (1.3 m along the stem or above any buttresses or deformities), the widely used minimum size for inventorying tropical trees. The combined dataset includes 1,003,805 trees, of which 93.3% are identified to species (Fig. 1 and Extended Data Table 1).

Consistent patterns of commonness

The Africa, Amazonia and Southeast Asia datasets differ in the number and size of plots sampled and the number of trees sampled (Extended Data Table 1). We therefore excluded small plots (below 0.9 ha; Extended Data Fig. 1 and Methods) and used rarefaction—that is, repeated random subsampling of plots to comparable numbers of trees—to standardize sampling across the three datasets (Fig. 2).

Rarefying to a common sample size of 77,587 stems, the size of the Asia dataset (equivalent to 150, 116 and 103 plots in Africa, Amazonia and Southeast Asia respectively), we find that 77 species (95% confidence interval: 62–92) in Africa comprise 50% of individual trees, compared with 174 species (95% confidence interval: 134–215) in Amazonia and 172 species (95% confidence interval: 125–217) in Southeast Asia (Table 1 and Fig. 2). However, the substantially lower number of hyperdominant species in Africa compared with Amazonia and Southeast Asia scales with the substantially lower number of total species. We find just 1,132 species in our standardized 77,587 tree sample in Africa, compared with 2,565 and 2,585 species in Amazonia and Southeast Asia, respectively for the same sample size. Consequently, percentage hyperdominance is statistically indistinguishable among the continents at 6.79% (95% confidence interval: 5.39%–8.20%), 6.80% (95% confidence interval: 5.24%–8.36%) and 6.65% (95% confidence interval: 4.59%–8.71%) in Africa, Amazonia and Southeast Asia, respectively (Table 1). This consistency is not affected by the aggregated spatial distribution of plots within each region (Extended Data Fig. 2) and holds true for analyses based

solely on 1-ha plots (Methods). Thus, once sampling is standardized, there is marked pan-tropical consistency in the proportion of the total number of tree species accounted for by the most common species.

The consistency of commonness is not limited to defining common species as those that account for 50% of all individual trees in a dataset. The proportions of the total number of species required to account for thresholds between 10% and 90% of individual trees are also highly consistent across the rarefied data for the three continents (Fig. 3 and Extended Data Table 3). Thus, the data from the three continents appear to result from the same underlying statistical distribution.

Our rarefaction analysis shows that the number of hyperdominants, the total number of species and the percentage hyperdominance are dependent on sample size. This is because as plots—and therefore trees—are added to the sample, increasing numbers of rare species start to appear. Meanwhile, most common species have, by definition, already appeared, but their abundances increase. Thus, with increasing sample size, the number of hyperdominants increases, but at an ever-decreasing rate that tends towards saturation (Fig. 2 and Extended Data Fig. 3). The total number of species increases at a decreasing rate with increasing sample size, without apparent saturation. Therefore, as sample sizes increase, the percentage hyperdominance decreases gradually, but does not appear to saturate (Fig. 2 and Extended Data Fig. 3). This sample size dependence is likely to explain the published differences in percentage hyperdominance in Amazonian forests, which follow expectations given the sample size in each study^{6,20,22}.

Amazonia and Southeast Asia show remarkably similar patterns of commonness and diversity. The rarefaction curves of the number of species accounting for 50% of all trees (Fig. 2a), total number of species (Fig. 2b), percentage hyperdominance (Fig. 2c) and Fisher's α —the parameter of the log series distribution shown to best describe tropical tree species abundance distributions²¹ (Fig. 2d)—are almost identical between the two datasets. Furthermore, the numbers of species required to account for any threshold between 10% and 90% of trees in the respective rarefied samples of 77,587 trees are statistically indistinguishable (Table 1 and Extended Data Tables 2 and 3). This equivalence in overall tropical forest diversity patterns between these similarly species-rich regions is particularly striking given their very different biogeographic, climatic and anthropogenic histories, and the fact that Amazonia is one large contiguous region, whereas Southeast Asia is a series of islands and island-like regions.

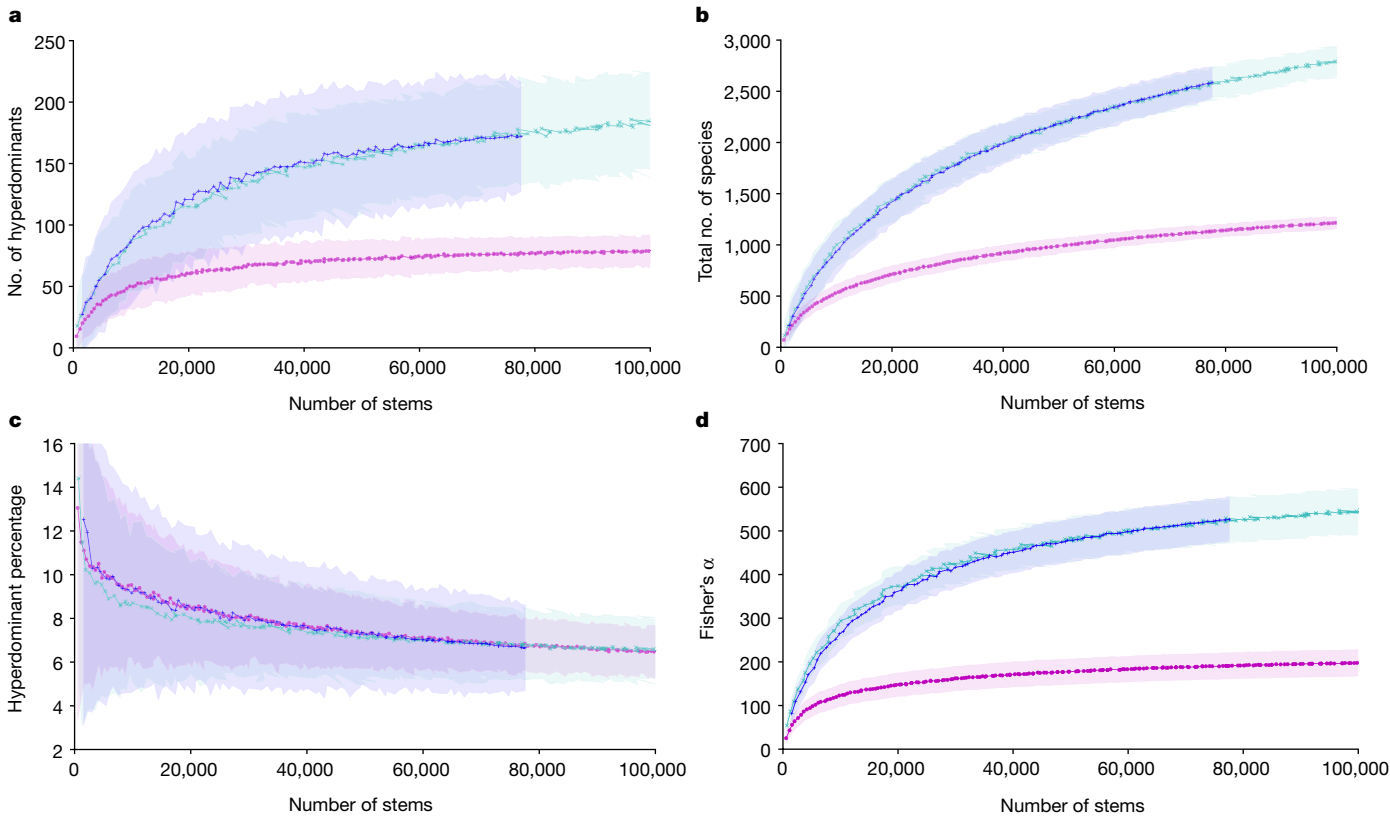


Fig. 2 | Rarefaction curves showing the effect of increasing sample size on the number of hyperdominants, total species, hyperdominant percentage and fitted values of Fisher's α in tropical tree communities. a–d, The effect of increasing sample size on the number of hyperdominants (a), total species (b), hyperdominant percentage (c) and fitted values of Fisher's α (d) in tropical Africa (magenta), Amazonia (cyan), Southeast Asia (blue). Rarefied data (mean

values across iterations of subsamples) are shown as points joined by lines for clarity, shaded areas represent 95% confidence intervals (derived via the s.d. across iterations of subsamples taken with replacement at each sampling point). Note that resampling for rarefaction was by subsampling of plots, but curves are re-plotted on an x axis of number of stems.

In contrast to the similarity between Amazonia and Southeast Asia, our results provide sample size-corrected validation of the 'odd-one-out' observation^{28,29} of much lower tree species richness in Africa compared with Amazonia and Southeast Asia. Here we add a similar odd-one-out observation of a much lower number of common species in Africa than in Amazonia and Southeast Asia. However, in combination these two results lead to an almost identical percentage hyperdominance in the African, Amazonian and Southeast Asian rarefied data. This consistency extends to the proportion of species required to account for all thresholds between 10% and 90% of trees in the rarefied data (Fig. 3 and Extended Data Table 3). This pan-tropical invariance recasts the tropical forests of Africa from 'odd' in terms of

species richness to statistically indistinguishable from those in Amazonia and Southeast Asia in terms of proportional patterns of abundance. Overall, using standardization by rarefaction, we find consistent patterns of species abundance across Africa, Amazonia and Southeast Asia.

Scaling to the study region

Next, we estimate commonness patterns in each of our three study regions: Africa, Amazonia and Southeast Asia. We extrapolate log series fits to the empirical Africa, Amazonia and Southeast Asia datasets (Extended Data Fig. 4), including a correction to account for the clumped spatial occurrence of species, to the total number of trees with trunk diameter of at least 10 cm in each study region. We estimate that just 104 species (95% confidence interval: 101–107) account for 50% of the 113 billion trees in Africa's closed canopy tropical forests (Table 2). We also estimate that just 299 species (95% confidence interval: 295–304) account for 50% of the 344 billion trees in Amazonia's closed canopy tropical forest, and 278 (95% confidence interval: 268–289) account for 50% of the 129 billion trees in Southeast Asia's closed canopy tropical forests (Table 2). Our results from Amazonia match those derived using a different extrapolation approach³⁰.

Our extrapolations again outline consistent percentage hyperdominance: just 2.2% of African, 2.2% Amazonian and 2.3% of Southeast Asian species account for 50% of all trees with trunk diameters of at least 10 cm in each region (Table 2). The dominant proportions of total species required to account for 10% to 90% of trees are also very similar across continents (Fig. 3 and Extended Data Table 5). The lower percentage dominance values from the extrapolated data compared

Table 1 | Tree species hyperdominance results for African, Amazonian and Southeast Asian tropical forests, resampled to the common sample size of 77,587 trees

	Number of hyperdominants	Total species	Hyperdominant percentage	Fisher's α
Africa	77 [62, 92]	1,132 [1,069, 1,194]	6.79 [5.39, 8.20]	191 [161, 220]
Amazonia	174 [134, 215]	2,565 [2,419, 2,711]	6.80 [5.24, 8.36]	525 [475, 575]
Southeast Asia	172 [125, 219]	2,585 [2,440, 2,730]	6.65 [4.59, 8.71]	526 [476, 577]

Numbers in brackets are confidence intervals derived from the s.d. across iterations of subsamples taken with replacement at the sample size of the Asia dataset. Resampling done by plot; 77,587 is the size of the Southeast Asia dataset.

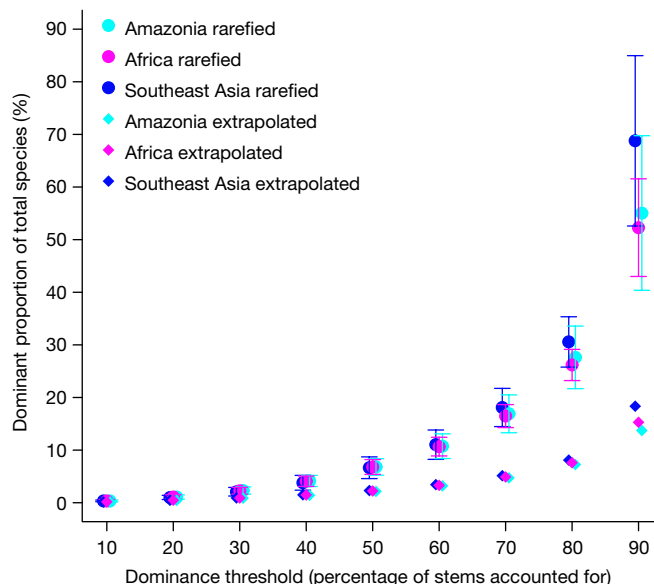


Fig. 3 | The minimum percentage of total species required to account for given dominance thresholds of the total number of stems when this varies from 10% to 90%. Circles show results as rarefied to the size of the Southeast Asia dataset (mean values across iterations of subsamples with 77,587 stems). Diamonds show the extrapolated results at the scale of the regions. Estimated rarefaction confidence intervals are derived from the s.d. across iterations of subsamples taken with replacement at 77,587 stems.

with those from the rarefied data are consistent with the pattern, described above, of many more rare species being added as the number of trees increases while many fewer common species are added (Fig. 2). Overall, the extrapolated results show that there are a tractable number of common species in tropical forests in Africa, Amazonia and Southeast Asia.

Scaling to the tropics

We next estimate the number of common tropical tree species on Earth by multiplying the pan-tropical proportion of common species by the total number of tropical tree species on Earth. Our results suggest a pan-tropical hyperdominant percentage of 2.24% (Table 2). However, our extrapolations cannot provide an estimate of the total number of tropical tree species because we do not—for this study—have data from all tropical regions, including a lack of data from Central America, New Guinea and Micronesia. Furthermore, there is no consensus estimate of the total number of tropical tree species on Earth.

A compilation of lists of species known to science suggests a total of 60,065 tree species globally¹⁵. Tropical forest biomes likely comprise 63% of this list (E. Beech, personal communication, 2021), implying that there are around 37,900 known tropical tree species. This minimum estimate does not account for species that are yet to be identified and described by scientists. An alternative extrapolation method estimated that there are 46,900 species for the closed canopy tropical forest biome²⁵ (range 40,500–53,300 species), implying that there are 9,000 yet-to-be-identified species. This is in agreement with a recent global study suggesting that there are around 9,200 tree species remaining yet to be formally named, almost all in the tropics¹⁶. Thus, together, these studies suggest there are likely to be approximately 47,000 tropical tree species in the world's closed canopy tropical forests.

Our best estimate is that 1,053 tree species (2.24% of 47,000 species) account for half of Earth's 800 billion trees with trunk diameters of at least 10 cm found in the closed-canopy tropical forest biome. Although the true number may be lower or higher, the conclusion that a tractable

Table 2 | Extrapolated tree species hyperdominance results for African, Amazonian, Southeast Asian tropical forests at the regional scale

	Number of hyperdominants	Total species	Hyperdominant percentage
Africa	104 [101, 107]	4,638 [4,511, 4,764]	2.23
Amazonia	299 [295, 304]	13,826 [13,615, 14,036]	2.16
Southeast Asia	278 [268, 289]	11,963 [11,451, 12,475]	2.32
Total^a	681 [664, 700]	30,427 [29,577, 31,275]	2.24

^aCalculated as the sum of the number of hyperdominants and total species across the three major tropical forest regions with hyperdominance percentage derived therefrom. Prediction intervals (in brackets) combine uncertainty from the standard error of predicted means and the residual s.d. of the regression of the bias correction fit.

number of species dominate tropical forests is clear. Some of these species are likely to be extraordinarily common: our best estimate is that just 61 species account for 80 billion individual trees (0.13% of 47,000 species). At the other end of the spectrum, we estimate that the rarest approximately 39,500 species account for just 80 billion trees, or 10% of individuals. Meanwhile, the other 90% of all trees are estimated to belong to just 7,487 species (15.93% of 47,000 species). Thus, these results open the possibility of focusing efforts on understanding the biology of a tractable number of species in tropical forests to approximate the whole stand.

Identifying the most common species

Our analyses showing that 104, 299 and 278 common species account for 50% of the trees in our African, Amazonian and Southeast Asian study regions, respectively, do not yield a list of named species. To assess which named species are likely to be hyperdominant, we use a subsampling procedure similar to the rarefaction methodology above. We randomly subsample from approximately 10,000 trees per subsample (drawn by plot) and increase the size of the subsample in 10,000-tree increments until the size of each regional dataset is reached, and repeat this process 100 times. For each sampled increment of 10,000 trees we then calculate the proportion of random subsamples in which each species qualifies as hyperdominant (Supplementary Table 1). We then assign the species to one of four groups:

- (1) Both hyperdominant in the full data and hyperdominant in the majority of subsamples even at very small sample sizes. These 50, 95 and 105 species in our Africa, Amazonia and Southeast Asia datasets, respectively, represent 3.5%, 2.1% and 4.1% of sampled species in each dataset. These species are likely to be geographically widespread and abundant.
- (2) Both hyperdominant in the full data and hyperdominant in the majority of subsamples, but at the smallest sample sizes only occasionally hyperdominant. These 32, 129 and 67 species in our Africa, Amazonia and Southeast Asia datasets, respectively, represent 2.3%, 2.9% and 2.6% of sampled species in each dataset. These species are likely to be geographically widespread but not always abundant.
- (3) Not quite hyperdominant in the full data, but hyperdominant in a substantial proportion of subsamples. These 102, 339 and 200 species in our Africa, Amazonia and Southeast Asia datasets, respectively, represent 7.2%, 7.5% and 7.7% of sampled species in each dataset. These species are probably locally abundant but not necessarily geographically widespread.
- (4) Not hyperdominant in the full data and almost never hyperdominant in the subsamples. These 1,232, 3,929 and 2,213 species in our Africa, Amazonia and Southeast Asia datasets, respectively, represent 87%, 87.5% and 85.6% of sampled species in each dataset. These species are probably neither geographically widespread nor abundant.

We suggest that if all trees in a region were sampled, the hyperdominant species would be drawn from the first three groups, which are listed in Supplementary Table 2. This candidate list of 1,119 hyperdominant species contains 184 species in Africa, 563 species in Amazonia and 372 species in Southeast Asia, with no species appearing on more than one region's list. Thus, the list of species that are likely candidates for hyperdominance is manageably small.

There is uncertainty in our candidate hyperdominant list owing to the limitations of the underlying samples of plots across the landscape. Specifically, some species that always have low local abundance but are geographically widespread and lack habitat restrictions may require larger sample sizes for their hyperdominance to become clear. Similarly, species that combine low local abundance and habitat specificity pose challenges. If the distribution and extent of specialist habitat is great enough to result in hyperdominance of specialists but is not sufficiently captured in our sampling, such species might not appear in our candidate list. By contrast, some species in our candidate hyperdominant list will not be true hyperdominants. Of particular note, some apparently common species may actually comprise a group of cryptic species, with none of these cryptic species being hyperdominant by itself^{31–33}. However, the striking similarity in species abundance patterns across the Africa, Amazonia and Southeast Asia datasets, despite differing sampling intensity on each continent, suggests that these potential limitations do not substantially affect the overall patterns found. We therefore expect a high overlap between our list of candidate hyperdominant species and eventual elucidation of the actual hyperdominants of these three regions and the pan-tropics.

Our list of 1,119 candidate hyperdominant species represents a tractable number of species on which to prioritize autecological research. Indeed, given their commonness, ecological data already exists for many of these species: 95% have some autecological data recorded in a large global database³⁴; 83% have at least 10 different types of measurement, typically including their growth form, maximum height, wood density and aspects of leaf chemistry. This indicates that these species are already relatively well known. Therefore, only limited additional data may be required to open new approaches to better understanding tropical forests through their most common tree species, including how they may react to today's era of rapid global environmental change.

Discussion

Charles Darwin wrote in *The Origin of Species* that “rarity is the attribute of a vast number of species of all classes and in all countries”³⁵. If this is the case, then common species are themselves rare. Our results concur: despite their formidable diversity, the trees in tropical forests fit the ‘rare is common, common is rare’ pattern³⁶ which has been documented in many other taxa^{17–19,36,37}. Beyond this, our analyses reveal highly consistent patterns of commonness across three major tropical forest regions. Notably, despite substantial inter-continental variation in biogeographic history, contemporary environment, forest structure and species composition, we have found an emergent property of the tropical forest system. For the trees that structure tropical forests, a consistent ~2.2% of the total species pool accounts for 50% of all individual trees in Africa, Amazonia and Southeast Asia. This consistency is all the more notable given relatively lower tree species richness of African tropical forests compared with Amazonian and Southeast Asian forests, probably owing to higher extinction rates in African forests, with evidence of major losses of African species at the Oligocene–Miocene boundary³⁸, and contractions of rainforest area due to drier conditions during repeated glacial–interglacial cycles over the past 2.6 million years³⁹.

We find common diversity patterns despite the very different histories of human occupancy in Amazonian, African and Southeast Asian tropical forests⁴⁰. The relatively recent arrival of humans in Amazonia approximately 20,000 years ago has been linked to greater Pleistocene extinctions, in contrast to much longer human occupancy in the

tropical forests of Africa and Southeast Asia⁴¹. Some have also suggested that Amazonian forest composition was altered by humans through the incipient domestication of tree species, increasing the abundance of a small number of favoured species⁴². Others have reported large areas of deforestation associated with the African Iron Age⁴³. How can such different human histories result in near-identical patterns of tree species dominance? The most parsimonious explanation is that the system tends to return to a state with a similar species abundance pattern.

Nevertheless, consistent patterns of commonness do not necessarily imply the same causal mechanisms. The ubiquity of the broad ‘rare is common, common is rare’ pattern in ecology, which is also found in non-biological complex systems⁴⁴, means inferences as to the cause of this broad pattern are challenging^{27,45}. Although combinatoric methods⁴⁵ and models that maximize the entropy of information^{46,47} both produce the ubiquitous ‘reverse lazy-J’ pattern, empirical observations show fewer common species and more rare species than expected by statistical controls alone⁴⁵. Similarly, neutral models produce the same broad pattern, but produce too few individuals of the most common Amazonian tree species⁴⁸. This suggests that biological mechanisms influence tree community assembly to produce a consistent proportion of common species across continents.

Recent analyses have revealed that the same few families contribute most of the species richness in Africa and Amazonia⁴⁹, which when combined with analyses showing that more diverse families have more common species⁵⁰, may indicate a role for deep evolutionary mechanisms driving the patterns we find. Yet, considering the substantially smaller regional species pool in Africa compared with Amazonia and Southeast Asia, one might expect differing continental patterns of species dominance if evolutionary drivers were the primary mechanism, not the highly consistent patterns that we find. Similarly, if environmental filtering were a key mechanism, the different contemporary environments, with Africa much drier on average than the other two continents²⁶, and Southeast Asia consisting of scattered island-like areas of forest compared with the contiguous forested region of Amazonia, would also imply differing continental patterns of species dominance, not the near-identical patterns that we find. These constraints limit the potential mechanisms that could apply across our three-continent context.

One potential cross-continental mechanism is dispersal limitation, where the dispersal capabilities of species result in some suitable habitat patches remaining unoccupied. Another mechanism is density- or distance-dependent mortality, which appears widespread across tropical forests⁵¹. Here, specialist species-specific natural enemies such as pathogens and herbivores reduce seed or juvenile conspecific survival rates near conspecific adults or in areas of high juvenile conspecific density, thereby reducing competitive exclusion and contributing to the maintenance of high tree species richness in tropical forests⁵¹. It is possible that common species have largely evaded density- and/or distance-dependent mortality. Analyses showing that species abundance can be either high or low within given genera⁵² support this hypothesis. Further progress on putative mechanisms can be made, for example, by exploring whether ecological or functional traits differ between common and rare species, and assessing the consistency of any differences among tropical continents⁵³. Although deducing mechanisms is complex, the identification of a tractable number of common species in tropical forests will facilitate progress in understanding of tropical forests beyond species abundance distributions.

Refining our results, particularly the naming of common species, requires improved sampling of tropical forests, both in terms of geographic scope and taxonomic identification of trees within plots. Expanding sampling to include Central America, New Guinea, Micronesia and other regions would improve the generality of our results. Better identifying trees in existing plots would increase the utility of available samples: in our Southeast Asia region we excluded 142 plots

(approximately 120,000 stems) because they did not have more than 80% of trees identified to species. Furthermore, additional taxonomic research on even the most common species is needed given that some of the most common Amazonian³³ and African^{54,55} tree species have been found to be complexes of several distinct species that are difficult to distinguish in the field. However, the similarity of our results across the three continental regions suggests that the occurrence of such species complexes may also be similar across the continental regions, again implying the operation of fundamental processes in differing forests. Overall, our work underscores the need for investment in taxonomy, particularly given the thousands of rare species we and others¹⁸ document, but also when considering the most common species.

Our best estimate, using extrapolation, that for the tropics as a whole just 1,053 species account for half of Earth's 800 billion tropical trees has potentially profound implications. Rather than attempting to understand tens of thousands of species of tropical trees, a focus on just a few hundred of the most common species can provide a simplified characterization of these otherwise complex forests. Our analyses indicate that the most common of these species are reliably named and relatively well known. Our list of candidate hyperdominants can therefore readily serve new research, including in facilitating targeted autecological data collection to understand their role in providing ecological functions and services. Practically, this species-specific information could enhance tropical forest modelling by focusing on common species instead of relying on functional types or traits, thereby potentially improving predictions of future forest change.

In the future, analyses should be extended to investigate forest carbon stocks and hyperdominant species and their role in the provision of ecosystem services. In Amazonia, even fewer tree species were found to account for 50% of aboveground carbon stocks than the minimum number required to account for 50% of trees²². More generally, the set of common species is likely to include foundation species that define broader community assemblages, the environmental sensitivity of which will probably drive tropical forest responses to environmental change⁵⁶. Of course, striving to understand and protect rare and non-hyperdominant species remains crucial, particularly as they face greater extinction risk and probably also contribute to the functioning of ecosystems, particularly when more functions⁵⁷, longer timescales⁵⁸ and imposed environmental changes⁵⁹ are considered, and given that the hyperdominants of the future may be rarer today. Nonetheless, with a complementary grasp of the most common species, mapping, understanding and modelling of the world's tropical forests will be a much more tractable proposition.

Online content

Any methods, additional references, NaturePortfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-023-06820-z>.

- Lewis, S. L. et al. Above-ground biomass and structure of 260 African tropical forests. *Phil. Trans. R. Soc. B* **368**, 20120295 (2013).
- Rovero, F. & Ahumada, J. The Tropical Ecology, Assessment and Monitoring (TEAM) Network: an early warning system for tropical rain forests. *Sci. Total Environ.* **574**, 914–923 (2017).
- Anderson-Teixeira, K. J. et al. CTFs—Forest GEO: a worldwide network monitoring forests in an era of global change. *Glob. Change Biol.* **21**, 528–549 (2015).
- Slik, J. W. F. et al. Phylogenetic classification of the world's tropical forests. *Proc. Natl Acad. Sci. USA* **115**, 1837–1842 (2018).
- Qie, L. et al. Long-term carbon sink in Borneo's forests halted by drought and vulnerable to edge effects. *Nat. Commun.* **8**, 1966 (2017).
- ter Steege, H. et al. Hyperdominance in the Amazonian tree flora. *Science* **342**, 1243092 (2013).
- Corlett, R. T. & Primack, R. B. *Tropical Rain Forests: An Ecological and Biogeographical Comparison* (John Wiley & Sons, 2011).
- IPCC Climate change 2022. *Impacts, Adaptation and Vulnerability* (eds Pörtner, H.-O. et al.) (Cambridge Univ. Press., 2022).
- Gough, C. Terrestrial primary production: fuel for life. *Nat. Educ. Knowl.* **3**, 28 (2011).
- Erb, K.-H. et al. Unexpectedly large impact of forest management and grazing on global vegetation biomass. *Nature* **553**, 73–76 (2018).
- Hubau, W. et al. Asynchronous carbon sink saturation in African and Amazonian tropical forests. *Nature* **579**, 80–87 (2020).
- Dirzo, R. & Raven, P. H. Global state of biodiversity and loss. *Annu. Rev. Env. Res.* **28**, 137–167 (2003).
- Mittermeier, R. A., Turner, W. R., Larsen, F. W., Brooks, T. M. & Gascon, C. in *Biodiversity Hotspots* (eds Zachos, F. & Habel, J.) 3–22 (Springer, 2011).
- Valencia, R., Balslev, H. & Paz Y Miño, C. G. High tree alpha-diversity in Amazonian Ecuador. *Biodivers. Conserv.* **3**, 21–28 (1994).
- Beech, E., Rivers, M., Oldfield, S. & Smith, P. P. GlobalTreeSearch: the first complete global database of tree species and country distributions. *J. Sustain. For.* **36**, 454–489 (2017).
- Cazzolla Gatti, R. et al. The number of tree species on Earth. *Proc. Natl Acad. Sci. USA* **119**, e2115329119 (2022).
- McGill, B. J. et al. Species abundance distributions: moving beyond single prediction theories to integration within an ecological framework. *Ecol. Lett.* **10**, 995–1015 (2007).
- Enquist, B. J. et al. The commonness of rarity: global and future distribution of rarity across land plants. *Sci. Adv.* **5**, eaaz0414 (2019).
- Baldrige, E., Harris, D. J., Xiao, X. & White, E. P. An extensive comparison of species-abundance distribution models. *PeerJ* **4**, e2823 (2016).
- Draper, F. C. et al. Amazon tree dominance across forest strata. *Nat. Ecol. Evol.* **5**, 757–767 (2021).
- ter Steege, H. et al. Biased-corrected richness estimates for the Amazonian tree flora. *Sci. Rep.* **10**, 10130 (2020).
- Fauset, S. et al. Hyperdominance in Amazonian forest carbon cycling. *Nat. Commun.* **6**, 6857 (2015).
- Pitman, N. C. A., Silman, M. R. & Terborgh, J. W. Oligarchies in Amazonian tree communities: a ten-year review. *Ecography* **36**, 114–123 (2013).
- Pitman, N. C. A. et al. Dominance and distribution of tree species in upper Amazonian terra firme forests. *Ecology* **82**, 2101–2117 (2001).
- Slik, J. W. et al. An estimate of the number of tropical tree species. *Proc. Natl Acad. Sci. USA* **112**, 7472–7477 (2015).
- Parmentier, I. et al. The odd man out? Might climate explain the lower tree α -diversity of African rain forests relative to Amazonian rain forests? *J. Ecol.* **95**, 1058–1071 (2007).
- McGill, B. J. & Nekola, J. C. Mechanisms in macroecology: AWOL or purloined letter? Towards a pragmatic view of mechanism. *Oikos* **119**, 591–603 (2010).
- Richards, P. W. in *Tropical Forest Ecosystems of Africa and South America: A Comparative Review* (eds Meggers, B. J., Ayensu, E. S. & Duckworth, W. D.) 21–26 (Smithsonian Institution Press, 1973).
- Couvreux, T. L. Odd man out: why are there fewer plant species in African rain forests? *Plant Syst. Evol.* **301**, 1299–1313 (2015).
- Tovo, A. et al. Upscaling species richness and abundances in tropical forests. *Sci. Adv.* **3**, e1701438 (2017).
- Cardoso, D. et al. Amazon plant diversity revealed by a taxonomically verified species list. *Proc. Natl Acad. Sci. USA* **114**, 10695–10700 (2017).
- ter Steege, H. et al. Towards a dynamic list of Amazonian tree species. *Sci. Rep.* **9**, 3501 (2019).
- Damasco, G. et al. Revisiting the hyperdominance of Neotropical tree species under a taxonomic, functional and evolutionary perspective. *Sci. Rep.* **11**, 9585 (2021).
- Kattge, J. et al. TRY plant trait database—enhanced coverage and open access. *Glob. Change Biol.* **26**, 119–188 (2020).
- Darwin, C. *On The Origin of Species by Means of Natural Selection: Or, the Preservation of Favored Races in the Struggle for Life* (J. Murray, 1859).
- McGill, B. J. in *Biological Diversity: Frontiers in Measurement and Assessment* (ed. Magurran, A. E. & McGill, B. J.) 105–122 (2021).
- Henderson, P. A. & Magurran, A. E. Linking species abundance distributions in numerical abundance and biomass through simple assumptions about community structure. *Proc. R. Soc. B* **277**, 1561–1570 (2010).
- Currano, E., Jacobs, B. & Pan, A. Is Africa really an “odd man out”? Evidence for diversity decline across the Oligocene–Miocene boundary. *Int. J. Plant Sci.* **182**, 551–563 (2021).
- Morley, R. J. *Origin and Evolution of Tropical Rain Forests* (John Wiley & Sons, 2000).
- Scerri, E. M. L., Roberts, P., Maizumi, S. Y. & Malhi, Y. Tropical forests in the deep human past. *Phil. Trans. R. Soc. B* **377**, 20200500 (2022).
- Sandom, C., Faurby, S., Sandel, B. & Svenning, J.-C. Global late Quaternary megafauna extinctions linked to humans, not climate change. *Proc. R. Soc. B* **281**, 20133254 (2014).
- Levis, C. et al. Persistent effects of pre-Columbian plant domestication on Amazonian forest composition. *Science* **355**, 925–931 (2017).
- Garcin, Y. et al. Early anthropogenic impact on Western Central African rainforests 2,600 y ago. *Proc. Natl Acad. Sci. USA* **115**, 3261–3266 (2018).
- Nekola, J. C. & Brown, J. H. The wealth of species: ecological communities, complex systems and the legacy of Frank Preston. *Ecol. Lett.* **10**, 188–196 (2007).
- Diaz, R. M., Ye, H. & Ernest, S. K. M. Empirical abundance distributions are more uneven than expected given their statistical baseline. *Ecol. Lett.* **24**, 1739–2039 (2021).
- Harte, J. & Newman, E. A. Maximum information entropy: a foundation for ecological theory. *Trends Ecol. Evol.* **29**, 384–389 (2014).
- Harte, J., Brush, M., Newman, E. A. & Umemura, K. An equation of state unifies diversity, productivity, abundance and biomass. *Commun. Biol.* **5**, 874 (2022).
- Pos, E. et al. Scaling issues of neutral theory reveal violations of ecological equivalence for dominant Amazonian tree species. *Ecol. Lett.* **22**, 1072–1082 (2019).
- Silva de Miranda, P. L. et al. Dissecting the difference in tree species richness between Africa and South America. *Proc. Natl Acad. Sci. USA* **119**, e2112336119 (2022).
- Webb, C. O. & Pitman, N. C. Phylogenetic balance and ecological evenness. *Syst. Biol.* **51**, 898–907 (2002).

51. Comita, L. S. et al. Testing predictions of the Janzen–Connell hypothesis: a meta-analysis of experimental evidence for distance- and density-dependent seed and seedling survival. *J. Ecol.* **102**, 845–856 (2014).
52. Ricklefs, R. E. & Renner, S. S. Global correlations in tropical tree species richness and abundance reject neutrality. *Science* **335**, 464–467 (2012).
53. Koffel, T., Umemura, K., Litchman, E. & Klausmeier, C. A. A general framework for species-abundance distributions: linking traits and dispersal to explain commonness and rarity. *Ecol. Lett.* **25**, 2359–2371 (2022).
54. Ikabanga, D. U. et al. Combining morphology and population genetic analysis uncover species delimitation in the widespread African tree genus *Santiria* (Burseraceae). *Phytotaxa* **321**, 166 (2017).
55. Koffi, K. G. et al. A combined analysis of morphological traits, chloroplast and nuclear DNA sequences within *Santiria trimera* (Burseraceae) suggests several species following the Biological Species Concept. *Plant Ecol. Evol.* **143**, 160–169 (2010).
56. Ellison, A. M. et al. Loss of foundation species: consequences for the structure and dynamics of forested ecosystems. *Front. Ecol. Environ.* **3**, 479–486 (2005).
57. Lefcheck, J. S. et al. Biodiversity enhances ecosystem multifunctionality across trophic levels and habitats. *Nat. Commun.* **6**, 6936 (2015).
58. Isbell, F. et al. Quantifying effects of biodiversity on ecosystem functioning across times and places. *Ecol. Lett.* **21**, 763–778 (2018).
59. Isbell, F. et al. High plant diversity is needed to maintain ecosystem services. *Nature* **477**, 199–202 (2011).
60. Olson, D. M. et al. Terrestrial Ecoregions of the World: a new map of life on Earth: a new global map of terrestrial ecoregions provides an innovative tool for conserving biodiversity. *Bioscience* **51**, 933–938 (2001).

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2024

Declan L. M. Cooper^{1,2}, Simon L. Lewis^{1,3}, Martin J. P. Sullivan^{3,4}, Paulo I. Prado⁵, Hans ter Steege^{6,7}, Nicolas Barbier^{8,9}, Ferry Slik¹⁰, Bonaventure Sonké^{9,11}, Corneille E. N. Ewango¹², Stephen Adu-Bredu¹³, Kofi Affum-Baffoe¹⁴, Daniel P. P. de Aguiar^{15,16}, Manuel Augusto Ahuite Reategui¹⁷, Shin-Ichiro Aiba¹⁸, Bianca Weiss Albuquerque¹⁹, Francisca Dionizia de Almeida Matos²⁰, Alfonso Alonso²¹, Christian A. Amani^{22,23}, Dário Dantas do Amaral²⁴, Iêda Leão do Amaral²⁰, Ana Andrade²⁵, Ires Paula de Andrade Miranda²⁰, Ilondea B. Angoboy²⁶, Alejandro Araujo-Murakami²⁷, Nicolás Castaño Arboleda²⁸, Luzmila Arroyo²⁷, Peter Ashton²⁹, Gerardo A. Aymard³⁰, Cláudia Baide^{31,32}, Timothy R. Baker³³, Michael Phillippe Bessike Balinga³³, Henrik Balslev³⁴, Lindsay F. Banin³⁵, Olaf S. Bánki⁶, Chris Baraloto³⁶, Edelcio Marques Barbosa³⁰, Flávia Rodrigues Barbosa³⁷, Jos Barlow³⁸, Jean-Francois Bastin³⁹, Hans Beekman⁴⁰, Serge Begne^{3,9,11}, Natacha Nssi Bengone⁴¹, Erika Berenguer^{38,42}, Nicholas Berry⁴³, Robert Bitarinho⁴⁴, Pascal Boeckx⁴⁵, Jan Bogaert⁴⁶, Bernard Bonyoma⁴⁷, Patrick Boundja^{22,48}, Nils Bourland^{40,49,50,51}, Faustín Boyemba Bosela⁵², Fabian Brambach⁵³, Roel Brien³, David F. R. P. Burslem⁵⁴, José Luis Camargo⁵⁰, Wegliane Campelo⁵⁵, Angela Cano^{56,57}, Sasha Cárdenas⁵⁶, Dairon Cárdenas López²⁸, Rainiellen de Sá Carpanedo³⁷, Yrma Andreina Carrero Márquez⁵⁸, Fernanda Antunes Carvalho^{59,60}, Luísa Fernanda Casas⁵⁶, Hernán Castellanos⁶¹, Carolina V. Castilho⁶², Carlos Cerón⁶³, Colin A. Chapman^{64,65,66}, Jerome Chave⁶⁷, Phourin Chhang⁶⁸, Wanlop Chutipong⁶⁹, George B. Chuyong⁷⁰, Bruno Barçante Ladvocat Cintra⁷¹, Connie J. Clark⁷², Fernanda Coelho de Souza^{59,73,74}, James A. Comiskey^{75,76}, David A. Coomes⁷⁷, Fernando Cornejo Valverde⁷⁸, Diego F. Correa^{66,79}, Flávia R. C. Costa⁵⁹, Janaina Barbosa Pedrosa Costa⁸⁰, Pierre Couteron^{8,9}, Heike Culmsee⁸¹, Aida Cuni-Sanchez^{92,83}, Francisco Dallmeier²¹, Gabriel Damasco⁸⁴, Gilles Dauby⁸⁵, Nállarett Dávila⁸⁵, Hilda Paulette Dávila Doza⁸⁶, Jose Don T. De Alban^{97,88}, Rafael L. de Assis⁸⁹, Charles De Caniere⁹⁰, Thales De Haulleville⁴⁰, Marcelo de Jesus Veiga Carim⁹¹, Layon O. Demarchi¹⁹, Kyle G. Dexter^{92,93}, Anthony Di Fiore^{94,95}, Hazimah Haji Mohammad Din⁹⁶, Mathias I. Disney⁹⁷, Brice Yannick Djiofack^{40,97,98}, Marie-Noël K. Djuikouo^{9,70}, Tran Van Do⁹⁹, Jean-Louis Doucet¹⁰⁰, Freddie C. Drape¹⁰¹, Vincent Droissart⁸¹, Joost F. Duivenvoorden¹⁰², Julien Engel^{81,103}, Vittoria Estienne⁴⁸, William Farfan-Rios^{104,105}, Sophie Fauset¹⁰⁶, Kenneth J. Feeley^{107,108}, Yuri Oliveira Feitosa¹⁰⁹, Ted R. Feldpausch^{73,110}, Cid Ferreira^{20,293}, Joice Ferreira¹¹¹, Leandro Valle Ferreira²⁴, Christine D. Fletcher¹¹², Bernardo Monteiro Flores¹¹³, Alusine Fofanah¹¹⁴, Ernest G. Foli¹³, Émile Fonty^{115,116}, Gabriella M. Fredriksson¹¹⁷, Alfredo Fuentes^{105,118}, David Galbraith⁷, George Pepe Gallardo Gonzales⁸⁶, Karina Garcia-Cabrera¹¹⁹, Roosevelt Garcia-Villacorta^{120,121}, Vitor H. F. Gomes^{22,122}, Ricardo Zárate Gómez¹²⁴, Therany Gonzales¹²⁵, Rogério Gribel²⁰, Marcelino Carneiro Guedes⁸⁰, Juan Ernesto Guevara^{126,127}, Khalid Rehman Hakeem¹²⁸, Jefferson S. Hall¹²⁹, Keith C. Hamer¹³⁰, Alan C. Hamilton¹³¹, David J. Harris⁹³, Rhett D. Harrison¹³², Terese B. Hart^{133,134}, Andy Hector¹³⁵, Terry W. Henkel¹³⁶, John Herbohn¹³⁷, Mireille B. N. Hockemba⁴⁸, Bruce Hoffman¹³⁸, Milena Holmgren¹³⁹, Euridice N. Honorio Coronado^{140,141}, Isau Huamantupa-Chuquimaco¹⁴², Wannes Hubau^{3,40,143}, Nobuo Imai¹⁴⁴, Mariana Victória Irumé²⁰, Patrick A. Jansen^{145,146}, Kathryn J. Jeffery¹⁴⁷, Eliana M. Jimenez⁴⁸, Tommaso Jucker¹⁴⁹, André Braga Junqueira¹⁵⁰, Michelle Kalamandeen¹⁵¹, Narcisse G. Kamdem^{9,11}, Kuswata Kartawinata¹⁵², Emmanuel Kasongo Yakusu^{40,98,153}, John M. Katembo⁵², Elizabeth Kearsley¹⁵⁴, David Kenfack¹²⁹, Michael Kessler¹⁵⁵, Thiri Toe Khaing^{156,157}, Timothy J. Killeen¹⁵⁸, Kanehiro Kitayama¹⁵⁹, Bente Klitgaard¹⁶⁰, Nicolas Labrière⁶⁷, Yves Laumonier¹⁶¹, Susan G. W. Laurance¹⁶², William F. Laurance¹⁶², Félix Laurent^{40,97,98}, Tinh Cong Le¹⁶³, Trai Truong Le¹⁶³, Miguel E. Leal¹⁶⁴, Evtlyn Márcia Leão de Moraes Novo¹⁶⁵, Aurora Levesley³, Moses B. Libalah^{9,11,166}, Juan Carlos Licona¹⁶⁷, Diógenes de Andrade Lima Filho²⁰, Jeremy A. Lindsell^{168,169}, Aline Lopes¹⁷⁰, Maria Aparecida Lopes¹⁷⁷, Jon C. Lovett¹⁷², Richard Lowe⁷³, José Rafael Lozada¹⁷⁴, Xinghui Lu¹⁷⁵, Nestor K. Luambua^{40,97,176,177}, Bruno Garcia Luize⁸⁵, Paul Maas⁶, José Leonardo Lima Magalhães^{178,179}, William E. Magnusson⁵⁹, Ni Putu Diana Mahayani¹⁸⁰, Jean-Remy Makana¹⁸¹, Yadvinder Malhi⁴², Lorena Maniguate Rincón²⁰, Asyraf Mansor^{182,183}, Angelo Gilberto Manzatto¹⁸⁴, Beatriz S. Marimon¹⁸⁵, Ben Hur Marimon-Junior¹⁸⁵, Andrew R Marshall^{82,186,187}, Maria Pires Martins²⁰, Faustín M. Mbaya¹⁸⁸, Marcelo Brilhante de Medeiros¹⁸⁹, Italo Mesones¹⁹⁰, Faizah Metalni¹⁹¹, Vianet Mihindou^{192,193}, Jerome Millet¹⁹⁴, William Milliken¹⁹⁵, Hugo F. Mogollón¹⁹⁶, Jean-François Molino^{8,9}, Mohd. Nizam Mohd. Said¹⁹⁷, Abel Monteagudo Mendoza^{198,199}, Juan Carlos Montero^{16,167}, Sam Moore⁴², Bonifacio Mostacedo¹⁹⁹, Linder Felipe Mozombite Pinto⁸⁶, Sharif Ahmed Mukul^{137,200}, Pantaleo K. T. Munishi²⁰¹, Hidetoshi Nagamasu²⁰², Henrique Eduardo Mendonça Nascimento²⁰, Marcelo Trindade Nascimento²⁰³, David Neill²⁰⁴, Reuben Nilus²⁰⁵, Janaina Costa Noronha³⁷, Laurent Nsenga⁴⁰, Percy Núñez Vargas¹⁸⁸, Lucas Ojo²⁰⁶, Alexandre A. Oliveira³, Edmar Almeida de Oliveira¹⁸⁵, Fidèle Evouna Ondo¹⁹², Walter Palacios Cuenca²⁰⁷, Susamar Pansin²⁰⁸, Marcelo Petratti Pansonato^{20,32}, Marcos Rios Paredes⁸⁶, Ekananda Paudel²⁰⁹, Daniela Pauletto²¹⁰, Richard G. Pearson², José Luis Marcelo Pena²¹¹, R. Toby Pennington^{93,110}, Carlos A. Peres²¹², Andrea Permana²¹³, Pascal Petronelli²¹⁴, Maria Cristina Peñañuela Mora²¹⁵, Juan Fernando Phillips²¹⁶, Oliver L. Phillips³, Georgia Pickavance³, Maria Teresa Fernandez Piedade¹⁹, Nigel C. A. Pitman²¹⁷, Pierre Ploton^{8,9}, Andreas Popelier^{40,98,153}, John R. Poulsen^{72,218}, Adriana Prieto²¹⁹, Richard B. Primack²²⁰, Hari Priyadi²²¹, Lan Qie^{3,222}, Adriano Costa Quaresma^{19,223}, Helder Lima de Queiroz²²⁴, Hirma Ramirez-Angulo²²⁵, José Ferreira Ramos²⁰, Neidiane Farias Costa Reis²⁰⁸, Jan Reitsma²²⁶, Juan David Cardenas Revilla²⁰, Terhi Riutta^{42,227}, Gonzalo Rivas-Torres^{98,228}, Iyan Robiansyah^{229,230}, Maira Rocha¹⁹, Domingos de Jesus Rodrigues³⁷, M. Elizabeth Rodriguez-Rondero^{97,231}, Francesco Rovero^{232,233}, Andes H. Rozak²³⁴, Agustín Rudas²³⁹, Ervan Rutishauser²³⁵, Daniel Sabatier^{8,9}, Le Bienfaiteur Sagang^{9,11,236}, Adeliza Felipe Sampaio²⁰⁸, Ismayadi Samsodin²³⁷, Manichanh Satdichanh²³⁹, Juliana Schieth²²⁰, Jochen Schöngart¹⁹, Veridiana Vizoni Scudeller²³⁸, Naret Seauturien²³⁹, Douglas Sheil²⁴⁰, Rodrigo Sierra²⁴¹, Miles R. Silman¹⁹, Thiago Sanna Freire Silva²⁴², José Renan da Silva Guimarães²⁴³, Murielle Simo-Droissart^{9,31}, Marcelo Fragomeni Simon¹⁸⁹, Plinio Sist²⁴⁴, Thainae R. Sousa²⁴⁵, Emanuelle de Sousa Farias^{246,247}, Luiz de Souza Coelho²⁰, Dominick V. Spracklen²⁴⁸, Suzanne M. Stas²⁴⁸, Robert Steinmetz²³⁹, Pablo R. Stevenson³⁶, Juliana Stropp²⁴⁹, Rahayu S. Sukri¹⁹⁶, Terry C. H. Sunderland^{22,250}, Eizi Suzuki²⁵¹, Michael D. Swaine²⁵², Jianwei Tang²⁵³, James Taplin²⁵⁴, David M. Taylor²³¹, J. Sebastián Tello²⁵⁵, John Terborgh^{256,257}, Nicolas Texier²⁵⁸, Ida Theilade²⁵⁹, Duncan W. Thomas²⁶⁰, Raquel Thomas²⁶¹, Sean C. Thomas²⁶², Milton Tirado²⁴¹, Benjamin Toirambe^{40,263}, José Julio de Toledo⁵⁵,

Kyle W. Tomlinson^{156,264}, Armando Torres-Lezama²²⁵, Hieu Dang Tran¹⁶³, John Tshibamba Mukendi^{40,153,265}, Roven D. Tumaneng^{88,266}, Maria Natalia Umaña²⁶⁷, Peter M. Umuay^{268,269}, Ligia Estela Urrego Giraldo²⁷⁰, Elvis H. Valderrama Sandoval^{271,272}, Luis Valenzuela Gamarra¹⁸⁸, Tinde R. Van Andel^{6,273}, Martin van de Bult²⁷⁴, Jaqueline van de Pol²⁷⁵, Geertje van der Heijden²⁷⁶, Rodolfo Vasquez¹⁸⁸, César I. A. Vela²⁷⁷, Eduardo Martins Venticinque²⁷⁸, Hans Verbeek²⁷⁹, Rizza Karen A. Veridiano^{28,280}, Alberto Vicentin¹⁵⁹, Ima Célia Guimarães Vieira²⁴, Emilio Vilanova Torre^{225,268}, Daniel Villarreal^{272,281}, Boris Eduardo Villa Zegarra²⁸², Jason Vleminckx^{36,283}, Patricio von Hildebrand²⁸⁴, Vincent Antoine Vos²⁸⁵, Corine Vriesendorp²¹⁷, Edward L. Webb^{286,287}, Lee J. T. White^{41,147,288}, Serge Wich²⁸⁹, Florian Wittmann^{16,223}, Roderick Zagt²⁹⁰, Runguo Zang²⁹¹, Charles Eugene Zartman²⁰, Lise Zemagho^{9,11}, Eglee L. Zent²⁹² & Stanford Zent²⁹²

¹Department of Geography, University College London, London, UK. ²Centre for Biodiversity and Environment Research, Department of Genetics, Evolution and Environment, University College London, London, UK. ³School of Geography, University of Leeds, Leeds, UK. ⁴Department of Natural Sciences, Manchester Metropolitan University, Manchester, UK. ⁵Instituto de Biociências, Departamento de Ecologia, Universidade de São Paulo (USP), São Paulo, Brazil. ⁶Naturalis Biodiversity Center, Leiden, The Netherlands. ⁷Quantitative Biodiversity Dynamics, Department of Biology, Utrecht University, Utrecht, The Netherlands. ⁸AMAP, Université de Montpellier, IRD, Cirad, CNRS, INRAE, Montpellier, France. ⁹International Joint Laboratory DYCOFAC, IRD-UYI-IRGM, Yaoundé, Cameroon. ¹⁰Environmental and Life Sciences, Faculty of Science, Universiti Brunei Darussalam, Gadong, Brunei Darussalam. ¹¹Plant Systematics and Ecology Laboratory, Higher Teachers' Training College, University of Yaoundé I, Yaoundé, Cameroon. ¹²Faculty of Renewable Natural Resources Management and Faculty of Sciences, University of Kisangani, Kisangani, Democratic Republic of the Congo. ¹³Forestry Research Institute of Ghana (FORIG), Kumasi, Ghana. ¹⁴Mensuration Unit, Forestry Commission of Ghana, Kumasi, Ghana. ¹⁵Procuradoria-Geral de Justiça, Ministério Público do Estado do Amazonas, Manaus, Brazil. ¹⁶Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ¹⁷Medio Ambiente, PLUSPRETOL, Iquitos, Peru. ¹⁸Faculty of Environmental Earth Science, Hokkaido University, Sapporo, Japan. ¹⁹Ecology, Monitoring and Sustainable Use of Wetlands (MAUA), Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ²⁰Coordenação de Biodiversidade, Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ²¹Center for Conservation and Sustainability, Smithsonian Conservation Biology Institute, Washington, DC, USA. ²²Center for International Forestry Research (CIFOR), Bogor, Indonesia. ²³Université Officielle de Bukavu, Bukavu, Democratic Republic of the Congo. ²⁴Coordenação de Botânica, Museu Paraense Emílio Goeldi, Belém, Brazil. ²⁵Projeto Dinâmica Biológica de Fragmentos Florestais, Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ²⁶Institut National pour l'Etude et la Recherche Agronomiques, Bukavu, Democratic Republic of the Congo. ²⁷Museo de Historia Natural Noel Kempff Mercado, Universidad Autónoma Gabriel Rene Moreno, Santa Cruz, Santa Cruz, Bolivia. ²⁸Herbario Amazónico Colombiano, Instituto SINCHI, Bogotá, Colombia. ²⁹Bullard Emeritus Professor of Forestry, Harvard University, Cambridge, MA, USA. ³⁰Programa de Ciencias del Agro y el Mar, Herbario Universitario (PORT), UNELLEZ-Guanare, Guanare, Venezuela. ³¹The Mauritius Herbarium, Agricultural Services, Ministry of Agro-Industry and Food Security, Reduit, Mauritius. ³²Instituto de Biociências, Departamento de Ecologia, Universidade de São Paulo (USP), São Paulo, Brazil. ³³Tetra Tech ARD, Accra, Ghana. ³⁴Department of Biology, Aarhus University, Aarhus C, Aarhus, Denmark. ³⁵UK Centre for Ecology and Hydrology, Penicuik, UK. ³⁶International Center for Tropical Botany, Department of Biological Sciences, Florida International University, Miami, FL, USA. ³⁷ICNHS, Federal University of Mato Grosso, Sinop, Brazil. ³⁸Lancaster Environment Centre, Lancaster University, Lancaster, UK. ³⁹TERRA Teaching and Research Centre, Gembloux Agro-Bio Tech, University of Liege, Gembloux, Belgium. ⁴⁰Service of Wood Biology, Royal Museum for Central Africa, Tervuren, Belgium. ⁴¹Ministry of Forests, Seas, Environment and Climate, Libreville, Gabon. ⁴²Environmental Change Institute, School of Geography and the Environment, University of Oxford, Oxford, UK. ⁴³The Landscapes and Livelihoods Group, Edinburgh, UK. ⁴⁴Institute of Tropical Forest Conservation, Mbarara University of Science and Technology (MUST), Mbarara, Uganda. ⁴⁵Isotope Bioscience Laboratory (ISOFYS), Ghent University, Ghent, Belgium. ⁴⁶Biodiversity and Landscape Unit, Gembloux Agro-Bio Tech, Université de Liege, Liège, Belgium. ⁴⁷Section de la Foresterie, Institut National pour l'Etude et la Recherche Agronomique Yangambi, Yangambi, Democratic Republic of the Congo. ⁴⁸Congo Programme, Wildlife Conservation Society, Brazzaville, Republic of Congo. ⁴⁹CIFOR, Bogor, Indonesia. ⁵⁰Forest Resources Management, Gembloux Agro-Bio Tech, University of Liège, Liège, Belgium. ⁵¹Resources and Synergies Development, Singapore, Singapore. ⁵²Laboratory of Ecology and Forest Management, Faculty of Sciences, University of Kisangani, Kisangani, Democratic Republic of the Congo. ⁵³Biodiversity, Macroecology and Biogeography, University of Göttingen, Göttingen, Germany. ⁵⁴School of Biological Sciences, University of Aberdeen, Aberdeen, UK. ⁵⁵Universidade Federal do Amapá, Ciências Ambientais, Macapá, Brazil. ⁵⁶Laboratório de Ecologia de Bosques Tropicales y Primatología, Universidad de los Andes, Bogotá, Colombia. ⁵⁷Cambridge University Botanic Garden, Cambridge, UK. ⁵⁸Programa de Maestria de Manejo de Bosques, Universidad de los Andes, Mérida, Mérida, Venezuela. ⁵⁹Coordenação de Pesquisas em Ecologia, Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ⁶⁰Departamento de Genética, Ecologia e Evolução, Instituto de Ciências Biológicas, Belo Horizonte, Brazil. ⁶¹Centro de Investigaciones Ecológicas de Guayana, Universidad Nacional Experimental de Guayana, Puerto Ordaz, Venezuela. ⁶²Centro de Pesquisa Agroflorestal de Roraima, Embrapa Roraima, Boa Vista, Brazil. ⁶³Escuela de Biología Herbario Alfredo Paredes, Universidad Central, Quito, Ecuador. ⁶⁴Biology Department, Vancouver Island University, Nanaimo, British Columbia, Canada. ⁶⁵Shaanxi Key Laboratory for Animal Conservation,

Article

Northwest University, Xi'an, China. ⁶⁶School of Life Sciences, University of KwaZulu-Natal, Scottsville, South Africa. ⁶⁷Laboratoire Évolution et Diversité Biologique, CNRS and Université Paul Sabatier, Toulouse, France. ⁶⁸Institute of Forest and Wildlife Research and Development (IRD), Phnom Penh, Cambodia. ⁶⁹Conservation Ecology Program, King Mongkut's University of Technology Thonburi, Bangkok, Thailand. ⁷⁰Faculty of Science, Department of Plant Science, University of Buea, Buea, Cameroon. ⁷¹Instituto de Biociências, Departamento Botanica, Universidade de Sao Paulo (USP), São Paulo, Brazil. ⁷²Nicholas School of the Environment, Duke University, Durham, NC, USA. ⁷³University of Leeds, Leeds, UK. ⁷⁴BeZero, London, UK. ⁷⁵Inventory and Monitoring Program, National Park Service, Fredericksburg, VA, USA. ⁷⁶Smithsonian Conservation Biology Institute, Washington, DC, USA. ⁷⁷Department of Plant Sciences and Conservation Research Institute, University of Cambridge, Cambridge, UK. ⁷⁸Andes to Amazon Biodiversity Program, Madre de Dios, Madre de Dios, Peru. ⁷⁹The University of Queensland, Brisbane, Queensland, Australia. ⁸⁰Empresa Brasileira de Pesquisa Agropecuária, Embrapa Amapá, Macapá, Brazil. ⁸¹State Agency for Environment, Nature Conservation and Geology, Güstrow, Germany. ⁸²Department of Environment and Geography, University of York, York, UK. ⁸³Department of International Environmental and Development Studies (NORAGRIC), Norwegian University of Life Sciences, Ås, Norway. ⁸⁴Gothenburg Global Biodiversity Centre, University of Gothenburg, Gothenburg, Sweden. ⁸⁵Departamento de Biologia Vegetal, Instituto de Biologia, Universidade Estadual de Campinas (UNICAMP), Campinas, Brazil. ⁸⁶Servicios de Biodiversidad EIRL, Iquitos, Peru. ⁸⁷Centre for Nature-Based Climate Solutions, Department of Biological Sciences, National University of Singapore, Singapore, Singapore. ⁸⁸Phillipines Programme, Fauna and Flora International, Cambridge, UK. ⁸⁹Biodiversity and Ecosystem Services, Instituto Tecnológico Vale, Belém, Brazil. ⁹⁰Landscape Ecology and Vegetal Production Systems Unit, Universite Libre de Bruxelles, Brussels, Belgium. ⁹¹Departamento de Botânica, Instituto de Pesquisas Científicas e Tecnológicas do Amapá (IEPA), Macapá, Brazil. ⁹²School of Geosciences, University of Edinburgh, Edinburgh, UK. ⁹³Royal Botanic Garden Edinburgh, Edinburgh, UK. ⁹⁴Department of Anthropology, University of Texas at Austin, Austin, TX, USA. ⁹⁵Estación de Biodiversidad Tiputini, Colegio de Ciencias Biológicas y Ambientales, Universidad San Francisco de Quito (USFQ), Quito, Ecuador. ⁹⁶Institute for Biodiversity and Environmental Research, Universiti Brunei Darussalam, Bandar Seri Begawan, Brunei Darussalam. ⁹⁷Institut National pour l'Etude et la Recherche Agronomiques (INERA), Wood Laboratory of Yangambi, Yangambi, Democratic Republic of the Congo. ⁹⁸UGent-Woodlab, Laboratory of Wood Technology, Department of Environment, Faculty of Bioscience Engineering, Ghent University, Ghent, Belgium. ⁹⁹Silviculture Research Institute, Vietnamese Academy of Forest Sciences, Hanoi, Vietnam. ¹⁰⁰Forest Is Life, TERRA, Gembloux Agro-Bio Tech, Liège University, Liège, Belgium. ¹⁰¹Department of Geography and Planning, University of Liverpool, Liverpool, UK. ¹⁰²Institute of Biodiversity and Ecosystem Dynamics, University of Amsterdam, Amsterdam, The Netherlands. ¹⁰³Florida International University, Miami, FL, USA. ¹⁰⁴Living Earth Collaborative, Washington University in Saint Louis, St Louis, MO, USA. ¹⁰⁵Missouri Botanical Garden, St Louis, MO, USA. ¹⁰⁶School of Geography, Earth and Environmental Sciences, University of Plymouth, Plymouth, UK. ¹⁰⁷Department of Biology, University of Miami, Coral Gables, FL, USA. ¹⁰⁸Fairchild Tropical Botanic Garden, Coral Gables, FL, USA. ¹⁰⁹Programa de Pós-Graduação em Biologia (Botânica), Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ¹¹⁰Department of Geography, College of Life and Environmental Sciences, University of Exeter, Exeter, UK. ¹¹¹Empresa Brasileira de Pesquisa Agropecuária, Embrapa Amazônia Oriental, Belém, Brazil. ¹¹²Forest Research Institute Malaysia, Kepong, Malaysia. ¹¹³Postgraduate Program in Ecology, Federal University of Santa Catarina, Florianópolis, Brazil. ¹¹⁴The Gola Rainforest National Park, Kenema, Sierra Leone. ¹¹⁵Direction Régionale de la Guyane, Office National des Forêts, Cayenne, French Guiana. ¹¹⁶Université de Montpellier, Montpellier, France. ¹¹⁷Pro Natura Foundation, Balikpapan, Indonesia. ¹¹⁸Herbario Nacional de Bolivia, Instituto de Ecología, Carrera de Biología, Universidad Mayor de San Andrés, La Paz, Bolivia. ¹¹⁹Biology Department and Center for Energy, Environment and Sustainability, Wake Forest University, Winston Salem, NC, USA. ¹²⁰Programa Restauración de Ecosistemas (PRE), Centro de Innovación Científica Amazónica (CINCIA), Tambopata, Peru. ¹²¹Peruvian Center for Biodiversity and Conservation (PCBC), Iquitos, Peru. ¹²²Escola de Negócios Tecnologia e Inovação, Centro Universitário do Pará, Belém, Brazil. ¹²³Universidade Federal do Pará, Belém, Brazil. ¹²⁴PROTERRA, Instituto de Investigaciones de la Amazonia Peruana (IIAP), Iquitos, Peru. ¹²⁵ACEER Foundation, Puerto Maldonado, Peru. ¹²⁶Grupo de Investigación en Biodiversidad, Medio Ambiente y Salud-BIOMAS, Universidad de las Américas, Quito, Ecuador. ¹²⁷The Field Museum, Chicago, IL, USA. ¹²⁸Department of Biological Sciences, Faculty of Science, King Abdulaziz University, Jeddah, Saudi Arabia. ¹²⁹Forest Global Earth Observatory (ForestGEO), Smithsonian Tropical Research Institute, Washington, DC, USA. ¹³⁰School of Biology, University of Leeds, Leeds, UK. ¹³¹Honorary Professor, Kunming Institute of Botany, Chinese Academy of Science, Kunming, China. ¹³²World Agroforestry, Lusaka, Zambia. ¹³³Lukuru Wildlife Research Foundation, Kinshasa, Democratic Republic of the Congo. ¹³⁴Division of Vertebrate Zoology, Yale Peabody Museum of Natural History, New Haven, CT, USA. ¹³⁵Department of Plant Sciences, University of Oxford, Oxford, UK. ¹³⁶Department of Biological Sciences, California State Polytechnic University, Humboldt, Arcata, CA, USA. ¹³⁷Tropical Forests and People Research Centre, University of the Sunshine Coast, Maroochydore DC, Queensland, Australia. ¹³⁸Amazon Conservation Team, Arlington, USA. ¹³⁹Resource Ecology Group, Wageningen University and Research, Wageningen, The Netherlands. ¹⁴⁰Instituto de Investigaciones de la Amazonia Peruana (IIAP), Iquitos, Peru. ¹⁴¹University of St Andrews, St Andrews, UK. ¹⁴²Herbario HAG, Universidad Nacional Amazónica de Madre de Dios (UNAMAD), Puerto Maldonado, Peru. ¹⁴³Department of Environment, Laboratory of Wood Technology (Woodlab), Ghent University, Ghent, Belgium. ¹⁴⁴Department of Forest Science, Tokyo University of Agriculture, Tokyo, Japan. ¹⁴⁵Smithsonian Tropical Research Institute, Ancon, Panama. ¹⁴⁶Department of

Environmental Sciences, Wageningen University and Research, Wageningen, The Netherlands. ¹⁴⁷Department of Biological and Environmental Sciences, University of Stirling, Stirling, UK. ¹⁴⁸Grupo de Ecología y Conservación de Fauna y Flora Silvestre, Instituto Amazónico de Investigaciones Imani, Universidad Nacional de Colombia sede Amazonia, Leticia, Colombia. ¹⁴⁹School of Biological Sciences, University of Bristol, Bristol, UK. ¹⁵⁰Institut de Ciència i Tecnologia Ambientals, Universitat Autònoma de Barcelona, Barcelona, Spain. ¹⁵¹School of Earth, Environment and Society, McMaster University, Hamilton, Ontario, Canada. ¹⁵²Integrative Research Center, The Field Museum of Natural History, Chicago, IL, USA. ¹⁵³Faculté de Gestion de Ressources Naturelles Renouvelables, Université de Kisangani, Kisangani, Democratic Republic of the Congo. ¹⁵⁴Computational and Applied Vegetation Ecology (CAVElab), Department of Environment, Faculty of Bioscience Engineering, Ghent University, Ghent, Belgium. ¹⁵⁵Department of Systematic and Evolutionary Botany, University of Zurich, Zurich, Switzerland. ¹⁵⁶Center for Integrative Conservation, Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, Menglun, Mengla, China. ¹⁵⁷University of the Chinese Academy of Sciences, Beijing, China. ¹⁵⁸Agteca—Amazonia, Santa Cruz, Bolivia. ¹⁵⁹Graduate School of Agriculture, Kyoto University, Kyoto, Japan. ¹⁶⁰Department for Accelerated Taxonomy, Royal Botanic Gardens, Richmond, UK. ¹⁶¹Forest and Environment Program, Center for International Forestry Research (CIFOR), Bogor, Indonesia. ¹⁶²Centre for Tropical Environmental and Sustainability Science and College of Science and Engineering, James Cook University, Cairns, Queensland, Australia. ¹⁶³Viet Nature Conservation Centre, Hanoi, Viet Nam. ¹⁶⁴Uganda Programme, Wildlife Conservation Society, Kampala, Uganda. ¹⁶⁵Divisão de Sensoriamento Remoto (DSR), Instituto Nacional de Pesquisas Espaciais (INPE), São José dos Campos, Brazil. ¹⁶⁶Department of Plant Biology, Faculty of Science, University of Yaoundé I, Yaoundé, Cameroon. ¹⁶⁷Instituto Boliviano de Investigacion Forestal, Santa Cruz, Santa Cruz, Bolivia. ¹⁶⁸The RSPB, Sandy, UK. ¹⁶⁹A Rocha International, Cambridge, UK. ¹⁷⁰Department of Ecology, Institute of Biological Sciences, University of Brasília, Brasília, Brazil. ¹⁷¹Instituto de Ciências Biológicas, Universidade Federal do Pará, Belém, Brazil. ¹⁷²Herbarium, Royal Botanic Gardens Kew, Richmond, UK. ¹⁷³Botany Department, University of Ibadan, Ibadan, Nigeria. ¹⁷⁴Facultad de Ciencias Forestales y Ambientales, Instituto de Investigaciones para el Desarrollo Forestal, Universidad de los Andes, Mérida, Mérida, Venezuela. ¹⁷⁵Institute of Forest Ecology, Environment and Protection, Chinese Academy of Forestry, Beijing, China. ¹⁷⁶Faculty of Renewable Natural Resources Management, University of Kisangani, Kisangani, Democratic Republic of the Congo. ¹⁷⁷Faculté des sciences Agronomiques, Université Officielle de Mbujimayi, Mbujimayi, Democratic Republic of the Congo. ¹⁷⁸Programa de Pós-Graduação em Ecologia, Universidade Federal do Pará, Belém, Brazil. ¹⁷⁹Embrapa Amazônia Oriental, Belém, Brazil. ¹⁸⁰Faculty of Forestry, Universitas Gadjah Mada, Yogyakarta, Indonesia. ¹⁸¹Faculté des Sciences, Laboratoire d'Écologie et Aménagement Forestier, Université de Kisangani, Kisangani, Democratic Republic of the Congo. ¹⁸²School of Biological Sciences, Universiti Sains Malaysia, George Town, Malaysia. ¹⁸³Centre for Marine and Coastal Studies, Universiti Sains Malaysia, George Town, Malaysia. ¹⁸⁴Departamento de Biologia, Universidade Federal de Rondônia, Unir, Porto Velho, Brazil. ¹⁸⁵Programa de Pós-Graduação em Ecologia e Conservação, Universidade do Estado de Mato Grosso, Nova Xavantina, Brazil. ¹⁸⁶Flamingo Land, Kirby Misperton, UK. ¹⁸⁷Forest Research Institute, University of the Sunshine Coast, Sippy Downs, Queensland, Australia. ¹⁸⁸Jardín Botánico de Missouri, Oxapampa, Peru. ¹⁸⁹Embrapa Recursos Genéticos e Biotecnologia, Brasília, Brazil. ¹⁹⁰Department of Integrative Biology, University of California, Berkeley, CA, USA. ¹⁹¹Environmental and Life Sciences Programme, Faculty of Science, Universiti Brunei Darussalam, Bandar Seri Begawan, Brunei Darussalam. ¹⁹²Agence Nationale des Parcs Nationaux, Libreville, Gabon. ¹⁹³Ministère de la Forêt, de la Mer, de l'Environnement, Chargé du Plan Climat, Libreville, Gabon. ¹⁹⁴Office français de la biodiversité, Vincennes, France. ¹⁹⁵Department for Ecosystem Stewardship, Royal Botanic Gardens, Richmond, UK. ¹⁹⁶Endangered Species Coalition, Silver Spring, MD, USA. ¹⁹⁷Institute of Climate Change, Universiti Kebangsaan Malaysia, Bangi, Malaysia. ¹⁹⁸Herbario Vargas, Universidad Nacional de San Antonio Abad del Cusco, Cuzco, Peru. ¹⁹⁹Facultad de Ciencias Agrícolas, Universidad Autónoma Gabriel René Moreno, Santa Cruz, Santa Cruz, Bolivia. ²⁰⁰Department of Environment and Development Studies, United International University, Dhaka, Bangladesh. ²⁰¹Department of Ecosystems and Conservation, Sokoine University of Agriculture, Morogoro, Tanzania. ²⁰²The Kyoto University Museum, Kyoto University, Kyoto, Japan. ²⁰³Laboratório de Ciências Ambientais, Universidade Estadual do Norte Fluminense, Campos dos Goytacazes, Brazil. ²⁰⁴Universidad Estatal Amazónica, Puyo, Ecuador. ²⁰⁵Forest Research Centre, Sandakan, Malaysia. ²⁰⁶University of Abeokuta, Abeokuta, Nigeria. ²⁰⁷Herbario Nacional del Ecuador, Universidad Técnica del Norte, Quito, Ecuador. ²⁰⁸Programa de Pós-Graduação em Biodiversidade e Biotecnologia PPG-Bionorte, Universidade Federal de Rondônia, Porto Velho, Brazil. ²⁰⁹Centre for Mountain Ecosystem Studies, Kunming Institute of Botany, Chinese Academy of Sciences, Kunming, China. ²¹⁰Instituto de Biodiversidade e Florestas, Universidade Federal do Oeste do Pará, Santarém, Brazil. ²¹¹Universidad Nacional de Jaén, Cajamarca, Peru. ²¹²School of Environmental Sciences, University of East Anglia, Norwich, UK. ²¹³University of Warwick, Warwick, UK. ²¹⁴Cirad UMR Ecofog, AgrosParisTech, CNRS, INRAE, Université Guyane, Kourou Cedex, France. ²¹⁵Universidad Regional Amazónica IKIAM, Tena, Ecuador. ²¹⁶Fundación Puerto Rastrojo, Bogotá, Colombia. ²¹⁷Science and Education, The Field Museum, Chicago, IL, USA. ²¹⁸The Nature Conservancy, Boulder, CO, USA. ²¹⁹Instituto de Ciencias Naturales, Universidad Nacional de Colombia, Bogotá, Colombia. ²²⁰Biology Department, Boston University, Boston, MA, USA. ²²¹Department of Resource and Environmental Economics (ESL), IPB University, Bogor, Indonesia. ²²²School of Life Sciences, University of Lincoln, Lincoln, UK. ²²³Wetland Department, Institute of Geography and Geoecology, Karlsruhe Institute of Technology (KIT), Rastatt, Germany. ²²⁴Diretoria Técnico-Científica, Instituto de Desenvolvimento Sustentável Mamirauá, Tefé, Brazil. ²²⁵Instituto de Investigaciones para el Desarrollo Forestal (INDEFOR), Universidad de los Andes, Mérida, Mérida, Venezuela.

²²⁶Waardenburg Ecology, Culemborg, The Netherlands. ²²⁷College of Life Sciences, University of Exeter, Exeter, UK. ²²⁸University of Florida, Gainesville, FL, USA. ²²⁹Department of Biological Sciences, King Abdulaziz University, Jeddah, Kingdom of Saudi Arabia. ²³⁰Center for Plant Conservation Bogor Botanic Gardens, Indonesian Institute of Science, Bogor, Indonesia. ²³¹Department of Geography, National University of Singapore, Singapore, Singapore. ²³²Department of Biology, University of Florence, Sesto Fiorentino, Italy. ²³³Tropical Biodiversity Section, Museo delle Scienze (MUSE), Trento, Italy. ²³⁴Research Center for Plant Conservation, Botanic Gardens and Forestry, National Research and Innovation Agency (BRIN), Bogor, Indonesia. ²³⁵InfoFlora, Botanical Garden of Geneva, Geneva, Switzerland. ²³⁶Institute of the Environment and Sustainability, University of California, Los Angeles, CA, USA. ²³⁷Forest Research and Development Center, Research, Development and Innovation Agency, Ministry of Environment and Forestry, Bogor, Indonesia. ²³⁸Departamento de Biologia, Universidade Federal do Amazonas (UFAM)–Instituto de Ciências Biológicas (ICB1), Manaus, Brazil. ²³⁹World Wildlife Fund Thailand, Bangkok, Thailand. ²⁴⁰Forest Ecology and Forest Management Group, Wageningen University and Research, Wageningen, The Netherlands. ²⁴¹GeoS, Quito, Ecuador. ²⁴²Biological and Environmental Sciences, University of Stirling, Stirling, UK. ²⁴³Amcel Amapá Florestal e Celulose SA, Santana, Brazil. ²⁴⁴Cirad-ES, Campus International de Baillarguet, TA C-105/D, Montpellier, France. ²⁴⁵Programa de Pós-Graduação em Ecologia, Instituto Nacional de Pesquisas da Amazônia (INPA), Manaus, Brazil. ²⁴⁶Laboratório de Ecologia de Doenças Transmissíveis da Amazônia (EDTA), Instituto Leônidas e Maria Deane, Fiocruz, Manaus, Brazil. ²⁴⁷Instituto Oswaldo Cruz (IOC/FIOCRUZ), Rio de Janeiro, Brazil. ²⁴⁸School of Earth and Environment, University of Leeds, Leeds, UK. ²⁴⁹Biogeography Department, Trier University, Trier, Germany. ²⁵⁰Faculty of Forestry, University of British Columbia, Vancouver, British Columbia, Canada. ²⁵¹Research Center for the Pacific Islands, Kagoshima University, Kagoshima, Japan. ²⁵²Department of Plant and Soil Science, School of Biological Sciences, University of Aberdeen, Aberdeen, UK. ²⁵³Key Laboratory of Tropical Plant Resources and Sustainable Use, Xishuangbanna Tropical Botanical Garden, Chinese Academy of Sciences, Mengla, China. ²⁵⁴UK Research and Innovation, Innovate UK, London, UK. ²⁵⁵Center for Conservation and Sustainable Development, Missouri Botanical Garden, St Louis, MO, USA. ²⁵⁶Department of Biology and Florida Museum of Natural History, University of Florida, Gainesville, FL, USA. ²⁵⁷James Cook University, Cairns, Queensland, Australia. ²⁵⁸Université Libre de Bruxelles, Brussels, Belgium. ²⁵⁹Department of Food and Resource Economics, University of Copenhagen, Copenhagen, Denmark. ²⁶⁰School of Biological Sciences, Washington State University, Vancouver, WA, USA. ²⁶¹Iwokrama

International Centre for Rain Forest Conservation and Development, Georgetown, Guyana. ²⁶²Institute of Forestry and Conservation, University of Toronto, Toronto, Ontario, Canada. ²⁶³Ministère de l'Environnement et Développement Durable, Kinshasa, Democratic Republic of the Congo. ²⁶⁴Center of Conservation Biology, Core Botanical Gardens, Chinese Academy of Sciences, Menglun, China. ²⁶⁵Faculté des Sciences Appliquées, Université de Mbujimayi, Mbujimayi, Democratic Republic of the Congo. ²⁶⁶Emerging Technology Development Division, Department of Science and Technology Philippine Council for Industry, Energy and Emerging Technology Research and Development (DOST-PCIEERD), Taguig City, Philippines. ²⁶⁷Department of Ecology and Evolutionary Biology, University of Michigan, Ann Arbor, MI, USA. ²⁶⁸Wildlife Conservation Society, New York, NY, USA. ²⁶⁹Yale School of Forestry and Environmental Studies, Yale University, New Haven, CT, USA. ²⁷⁰Departamento de Ciencias Forestales, Universidad Nacional de Colombia, Medellín, Colombia. ²⁷¹Department of Biology, University of Missouri, St Louis, MO, USA. ²⁷²Universidad Nacional de la Amazonia Peruana, Iquitos, Peru. ²⁷³Wageningen University, Wageningen, The Netherlands. ²⁷⁴Doi Tung Development Project, Social Development Department, Chiang Rai, Thailand. ²⁷⁵Compagnie des Bois du Gabon, Port Gentil, Gabon. ²⁷⁶University of Nottingham, Nottingham, UK. ²⁷⁷Escuela Profesional de Ingeniería Forestal, Universidad Nacional de San Antonio Abad del Cusco, Puerto Maldonado, Peru. ²⁷⁸Centro de Biociências, Departamento de Ecologia, Universidade Federal do Rio Grande do Norte, Natal, Brazil. ²⁷⁹CAVELab—Computational and Applied Vegetation Ecology, Department of Environment, Ghent University, Ghent, Belgium. ²⁸⁰FORLIANCE, Bonn, Germany. ²⁸¹Fundación Amigos de la Naturaleza (FAN), Santa Cruz, Bolivia. ²⁸²Dirección de Evaluación Forestal y de Fauna Silvestre, Magdalena del Mar, Peru. ²⁸³Faculté des Sciences, Service d'Évolution Biologique et Écologie, Université Libre de Bruxelles, Brussels, Belgium. ²⁸⁴Fundación Estación de Biología, Bogotá, Colombia. ²⁸⁵Instituto de Investigaciones Forestales de la Amazonía, Universidad Autónoma del Beni José Ballivián, Riberalta, Beni, Bolivia. ²⁸⁶Viikki Tropical Resources Institute, Department of Forest Sciences, University of Helsinki, Helsinki, Finland. ²⁸⁷Helsinki Institute of Sustainability Science (HELSUS), Helsinki, Finland. ²⁸⁸Institut de Recherche en Écologie Tropicale, Libreville, Gabon. ²⁸⁹School of Biological and Environmental Sciences, Liverpool John Moores University, Liverpool, UK. ²⁹⁰Tropenbos International, Ede, The Netherlands. ²⁹¹Key Laboratory of Forest Ecology and Environment of State Forestry Administration, Institute of Forest Ecology, Environment and Protection, Chinese Academy of Forestry, Beijing, China. ²⁹²Laboratory of Human Ecology, Instituto Venezolano de Investigaciones Científicas (IVIC), Caracas, Venezuela. ²⁹³Deceased: Cid Ferreira.

Methods

Data compilation and pre-processing

We collated data from forest inventory plots ≥ 0.2 ha in size, situated in structurally intact (no detectable past logging or fire), closed canopy (not dry forest or savanna) tropical forest, with enumeration of all stems ≥ 10 cm diameter, in which $\geq 80\%$ of stems are identified to the species level. Following Sullivan et al.⁶¹, small (≤ 0.5 ha) plots within 1 km of each other were grouped for analysis to minimize the effect of stochastic tree fall events in smaller areas⁶². These criteria allow direct comparisons to be made with hyperdominance results from Amazonia^{6,21}. The data from each continent comprise the following:

Africa: 483 plots, covering a total of 504 ha (mean plot area 1.04 ha, median 1 ha, range 0.2–10 ha). These data are from four sources: 299 plots from the African Tropical Rainforest Observatory Network^{1,63} (AfriTRON: www.afritron.org, accessed 1 March 2020), curated at <http://www.ForestPlots.net>⁶⁴; 127 plots from the Central African Plot Network (<https://central-african-plot-network.netlify.app>); 52 plots from the TEAM network²; and 5 \times 1 ha plots from 5 different soil types, extracted from one 50-ha plot in Korup, Cameroon from the SIGEO/CTFS network³.

Amazonia: 1,417 plots, covering a total of 1,591 ha (mean plot area 1.12 ha, median 1 ha, range 0.1–78.8 ha) from the Amazon Tree Diversity Network (ATDN: <http://atdn.myspecies.info/>, includes plots from the RAINFOR network), accessed 8 January 2020.

Southeast Asia: 230 plots, covering a total of 202 ha (mean plot area 0.88 ha, median 0.49 ha, range 0.21–4.5 ha). These data are from two sources: 143 plots from Slik et al.^{4,25}—a decrease from the published Indo-Pacific dataset in Slik et al.^{4,25} due to our $\geq 80\%$ species identification criterion and our Southeast Asia study region excluding Australia, India, and Papua New Guinea; and 87 plots from the T-Forces network⁶⁴ curated at <http://www.ForestPlots.net>, accessed 03/02/2021.

Species names were checked for orthography and standardized (synonyms identified from the reference databases corrected to their accepted names) using the African Flowering Plants Database (<https://www.ville-ge.ch/musinfo/bd/cjb/africa>), Taxonomic Name Resolution Service⁶⁵, and Asian Plant Synonym Lookup (F. Slik, personal communication), for Africa, Amazonia and Southeast Asia, respectively. Trees not identified to species level (7.3%, 6.3% and 8.4% of stems in the Africa, Southeast Asia datasets respectively) were classed as ‘indeterminate’ (Indet). Indet stems contributed to plot-level and dataset-wide stem abundance totals but are necessarily absent from species totals.

For the purposes of our study we delimited tropical forests according to the ‘tropical and subtropical moist broadleaf forests’ biome delineation from the World Wildlife Fund ecoregion map⁶⁰. The total number of tropical trees ≥ 10 cm trunk diameter in each of our regions was then estimated by summing tree abundances in countries in which we have at least one sampled plot from the ‘map of Global Tree Density’⁶⁶ (derived from 429,775 ground-based estimates of tree density) and masking according to the ‘tropical and subtropical moist broadleaf forests’ borders using ArcGIS v3.10.1⁶⁷. Thus, we estimate that there are ~92 billion, ~331 billion trees, and ~217 billion trees in our Africa, Amazonia, and Southeast Asia regions, respectively, totalling 640 billion trees. Including abundance from countries in the ‘tropical and subtropical moist broadleaf forests’ biome in which we have no sampled plots, we estimate ~799 billion total trees across all of Earth’s moist tropical forests.

Data format, commonness and diversity parameters

The species abundance distribution (SAD), defined as a vector of abundances (number of individuals observed) of all species encountered in a community¹⁷, formed the basis for our analyses of the three tropical forest datasets. For each dataset, we tallied the number of trees of each species in each plot to give plot-level SADs and combined these SADs across all plots to get regional-level abundance matrices with rows representing plots, columns representing species, and entries

representing the abundance of each species in each plot. To capture patterns of commonness and species composition we calculated the number of hyperdominants (H#), defined as the minimum number of species required to account for 50% of the population of an assemblage⁶, hyperdominant species identities, total number of species (TS), hyperdominant percentage of total species ($H\% = H\#/TS$) and Fisher’s α (ref. 68). To investigate the sensitivity of results to the ‘hyperdominant’ definition of the most common species, we looked beyond the 50% threshold used for hyperdominance, at the minimum number of species required to account for 10%, 20%, 30%, ..., 90% of the population, here termed ‘dominants’.

Sampling standardization, subsampling and comparison of continental data

We identified variations in the number of plots, stems, and species, and the size and spatial clustering of plots as potential confounding factors liable to skew dominance and diversity results from our regional datasets and impede rigorous comparisons between them. We used sample-based rarefaction to quantify and account for the effect of differences in sample size (number of plots and stems) on our diversity measures of interest; namely species richness, number, ranking and identity of hyperdominants, hyperdominant percentage of total species, and Fisher’s α . To quantify the effect of plot size, which is smaller in Southeast Asia data (mean 0.88 ha, median 0.49 ha) than in Amazonia and Africa data (both mean ~1 ha, median 1 ha) we compared results from the full data to those from plots > 0.9 ha. We found that small plots (< 1 ha) inflate per-plot species totals relative to larger plots (because the rate of encountering new species is higher the smaller the plot size; Extended Data Fig. 1), so we limited our analyses to plots > 0.9 ha to enable like-for-like comparison.

For Africa, we retained 368 plots covering 450 ha (mean plot area 1.22 ha, median 1 ha, range 0.92–10 ha; 2% of plots 0.9–0.99 ha, 88% of plots 1 ha, 8% of plots 1.01–5 ha, 1% of plots > 5 ha) with mean temperature of 24.3 °C (range 16.2–27.6 °C), mean annual precipitation 1,802 mm yr⁻¹, (range 1,066–2,747 mm yr⁻¹), and mean elevation of 511 m above sea level (range 41–2,070 m) per WorldClim⁶⁹. For Amazonia we retained 1,097 plots covering 1,434 ha (mean plot area 1.31 ha, median 1 ha, range 0.9–78.8 ha; 2% of plots 0.9–0.99 ha, 90% of plots 1 ha, 7% of plots 1.01–5 ha, 1% of plots > 5 ha) with mean temperature of 26.0 °C (range 20.9–27.6 °C), mean annual precipitation 2,397 mm yr⁻¹ (range 1,119–4,284 mm yr⁻¹), and mean elevation of 154 m (range 0–1,142 m). For Southeast Asia we retained 103 plots covering 164 ha (mean plot area 1.59 ha, median 1 ha, range 0.96–4.5 ha; 1% of plots 0.9–0.99 ha, 48% of plots 1 ha, 52% of plots 1.01–5 ha, 0% of plots > 5 ha) with mean temperature of 25.7 °C (range 20.1–27.5 °C), mean precipitation 2,680 mm yr⁻¹ (range 1,466–3,941 mm yr⁻¹), and mean elevation of 288 m (range 10–934 m). We assessed if the remaining differences in plot size affected the results, using only the 1 ha plots from Africa ($n = 323$) and Amazonia ($n = 988$), rarefied to the size of the Asia dataset, again finding near-identical per cent hyperdominance on the two continents (Africa: 7.30%, 95% confidence interval: 6.56–8.04; Amazonia: 7.35%, 95% confidence interval: 6.61–8.10).

To quantify the effect of the spatial clustering of plots, we compared results from the full Amazonia data, as the largest dataset, to those from subsets of the Amazonia data in which 1, 2, 3, ..., 10 plots were sampled from each spatial cluster. We found that spatial clustering had a negligible and not statistically significant effect on hyperdominant percentage and fitted values of Fisher’s α (Extended Data Fig. 2). Therefore, we retain all plots for our analyses to maximize sample sizes. Computation of percentage hyperdominance and dominance accounts for the effects of variations in species richness on the number of hyperdominants and dominants.

For sample-based rarefaction, 200 subsamples of 1, 2, ..., N_p plots were drawn, without replacement, from the N_p total number of plots in the p th dataset, the stems contained in each subsample were pooled,

and the mean total species, number of hyperdominants, hyperdominance percentage, and Fisher's α were calculated across the subsamples. Similarly, we tallied the number of subsamples in which each species in the dataset qualified as hyperdominant at each level of subsampling and compared results between datasets at subsample sizes equating to a mean 10,000, 20,000, ..., I_p individual trees, where I_p is the total number of trees in the p th dataset. Confidence intervals were calculated as confidence interval = $\mu \pm 1.96 \times \sigma$, where μ values are the means of the diversity metrics calculated across the 200 iterations of subsamples taken without replacement, and σ values are the s.d. of the mean of diversity metrics calculated across the 200 iterations of subsamples taken with replacement (to reduce the degree to which confidence intervals were conditional on the sample). For point estimates, all datasets were compared at the common sample size of the Southeast Asia dataset (77,587 stems equivalent to 150, 116 and 103 plots in Africa, Amazonia and Southeast Asia, respectively).

Extrapolation and bias correction of log series fits to the empirical data

We extrapolated our empirical SADs to SADs at the scale of the entire Amazonian, African, and Southeast Asian regional level via analytical expansion and bias correction of Fisher's log series fits following the methodology of ter Steege et al.²¹ developed using the ATDN data that comprise our Amazonia dataset.

Ter Steege²¹ et al. found that simulations of sampling of plots with conspecific aggregation from log series-modelled SADs provide extremely good approximations of the processes that generate tropical forest inventory data—that is, non-random sampling of plots containing species with limited dispersal and/or ecological preferences. They further found that estimates of species richness derived from samples taken with conspecific aggregation from the simulated SADs substantially underestimated the true species richness of the simulated SADs, but that a linear relationship with low variance existed between the true and sample-derived values. Thus, although conspecific aggregation in the empirical data introduces bias in the log series-modelled SADs extrapolated therefrom, quantification and correction of the effects of this bias on regional estimates of species richness is possible. Therefore, to estimate species richness at the regional level, they fitted Fisher's log series to empirical species abundance data, quantified the effect of conspecific aggregation on these estimates via simulation, and applied quantified corrections to give more accurate estimates of regional species richness taking into conspecific aggregation. Thus, this approach corrects for species-specific aggregation at the plot scale depending on species density.

To estimate regional numbers and proportions of dominants and hyperdominants as well as species richness, we extended the methodology of ter Steege et al.²¹ to log series-derived estimates of regional numbers and proportions of dominants and hyperdominants. Initially, values of Fisher's α were fitted to the empirical species abundance vectors from each region using maximum likelihood and numerical optimization in the 'sads' R package⁷⁰ and fits visualized with Preston plots⁷¹ and rank abundance distributions (RAD)³⁶ (Extended Data Fig. 4). Regional species totals S , not accounting for bias introduced by conspecific aggregation, were then estimated⁶⁸ via $S = \alpha \times \ln\left(1 + \frac{N}{\alpha}\right)$ with total number of trees ≥ 10 cm trunk diameter at the continental level (N) from the Global Tree Density map of Crowther et al.⁶⁶ with each tropical region delineated within the 'tropical and subtropical moist broadleaf forests' biome of Olson et al.⁶⁰. An inverse quantile function from the sads R package⁷⁰ was then applied to generate (uncorrected) continental-scale SADs for each region using the above fitted α , estimated S and N .

For the quantification of bias and computation of corrections, we first simulated 250 log series SADs with known values of total species, S_k , randomly drawn from the range of plausible regional species totals (10,000–25,000 in Amazonia and Southeast Asia; 2,000–10,000 in

Africa) and N , the number of trees in each region ≥ 10 cm trunk diameter from Crowther et al.⁶⁶. We calculated known values of numbers of hyperdominants, H_k , and percentage hyperdominance, P_k , from each of these simulated distributions. Using a negative binomial distribution to simulate conspecific aggregation per ter Steege et al.²¹, we then simulated J random samples of 1-ha plots from each of the 250 simulated SADs, with J equal to the number of plots in the empirical data, and the expected abundance of each species in each plot equal to its mean regional density (total abundance/regional area). We then estimated (uncorrected) species richness, S_u , from each of the samples by fitting Fisher's α to the sampled data and applying the formula $S_u = \alpha \times \ln\left(1 + \frac{N}{\alpha}\right)$. From each of the samples we also derived continental-scale uncorrected SADs (see above), from which the number of hyperdominants, H_u , and percentage hyperdominance, P_u , could be directly calculated, via analytical expansion of the log series using the fitted values of α and corresponding values of S_u . We then regressed the known values of S_k , H_k and P_k from the simulated SADs against the estimated (uncorrected) values S_u , H_u and P_u from the samples drawn with conspecific aggregation across all 250 simulations—that is, fit linear models of the form $A_k = m \times A_u + c$ for $A = S, H, P$. This same procedure was also applied to the number and proportion of dominants.

Across all three regional datasets, the above procedure outlined a linear relationship with low variance between known values of species richness, number of dominants and hyperdominants, and percentage hyperdominance and dominance, and values thereof estimated from sampling with conspecific aggregation (Extended Data Fig. 5). Thus, constant terms with low variance were readily applicable to correct for bias in the point estimates of species richness, number of dominants/hyperdominants, and percentage hyperdominance/dominance, derived from the empirical Africa, Amazonia, and Southeast Asia data. To capture uncertainty around each bias-corrected point estimate, prediction intervals (PI) were derived as $PI = \mu \pm 1.96 \times \sigma_{PI}$, where μ is the predicted mean value of the point estimate according to the linear regression, and σ_{PI} is the PI standard error, calculated as $\sigma_{PI} = \sqrt{\sigma^2 + \sigma_R^2}$, where σ is the standard error of predicted means and σ_R is the residual s.d. (and 1.96 is the 0.05 quantile of a t -distribution).

Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

The species abundance data that support the findings of this study are available from <https://doi.org/10.6084/m9.figshare.21670883> (formatting notes: a column for each species, rows for each plot, entries are the number of trees ≥ 10 cm diameter of each species in each plot). WorldClim⁶⁹ bioclimatic data are available from <https://www.worldclim.org/data/bioclim.html>.

Code availability

R code (version 4.3.1) to run the analyses and produce the figures and tables is available from <https://github.com/declancooper/CommonSpecies2022.git>.

- Sullivan, M. J. P. et al. Long-term thermal sensitivity of Earth's tropical forests. *Science* **368**, 869–874 (2020).
- Clark, D. B. & Clark, D. A. Landscape-scale variation in forest structure and biomass in a tropical rain forest. *For. Ecol. Manag.* **137**, 185–198 (2000).
- ForestPlots.net. Taking the pulse of Earth's tropical forests using networks of highly distributed plots. *Biol. Conserv.* **260**, 108849 (2021).
- Lopez-Gonzalez, G., Lewis, S. L., Burkitt, M. & Phillips, O. L. ForestPlots.net: a web application and research tool to manage and analyse tropical forest plot data. *J. Veg. Sci.* **22**, 610–613 (2011).
- Boyle, B. et al. The taxonomic name resolution service: an online tool for automated standardization of plant names. *BMC Bioinformatics* **14**, 16 (2013).

66. Crowther, T. W. et al. Mapping tree density at a global scale. *Nature* **525**, 201–205 (2015).
67. Esri, A. D. *ArcGIS Release 10. Documentation Manual* (Environmental Systems Research Institute, 2011).
68. Fisher, R. A., Corbet, A. S. & Williams, C. B. The relation between the number of species and the number of individuals in a random sample of an animal population. *J. Anim. Ecol.* **12**, 42–58 (1943).
69. Fick, S. E. & Hijmans, R. J. WorldClim 2: new 1-km spatial resolution climate surfaces for global land areas. *Int. J. Climatol.* **37**, 4302–4315 (2017).
70. Prado, P. I., Miranda, M. D., Chalom, A., Prado, M. P. I. & Imports, M. sads: maximum likelihood models for species abundance distributions. R package version 0.4.2 (2018).
71. Preston, F. W. The commonness, and rarity, of species. *Ecology* **29**, 254–283 (1948).

Acknowledgements D.L.M.C. was supported by the London Natural Environmental Research Council Doctoral Training Partnership grant (grant no. NE/L002485/1). This paper developed from analysing data from the African Tropical Rainforest Observatory Network (AfriTRON), curated at ForestPlots.net. AfriTRON has been supported by numerous people and grants since its inception. We sincerely thank the people of the many villages and local communities who welcomed our field teams and without whose support this work would not have been possible. Grants that have funded the AfriTRON network, including data in this paper, are a European Research Council Advanced Grant (T-FORCES; 291585; Tropical Forests in the Changing Earth System), a NERC standard grant (NER/A/S/2000/01002), a Royal Society University Research Fellowship to S.L.L., a NERC New Investigators Grant to S.L.L., a Philip Leverhulme Award to S.L.L., a European Union FP7 grant (GEOCARBON; 283080), Leverhulme Program grant (Valuing the Arc); a NERC Consortium Grant (TROBIT; NE/D005590/), NERC Large Grant (CongoPeat; NE/R016860/1) the Gordon and Betty Moore Foundation the David and Lucile Packard Foundation, the Centre for International Forestry Research (CIFOR), and Gabon's National Parks Agency (ANPN). This paper was supported by ForestPlots.net approved Research Project 81, 'Comparative Ecology of African Tropical Forests'. The development of ForestPlots.net and data curation has been funded by several grants, including NE/B503384/1, NE/N012542/1, ERC Advanced Grant 291585—'T-FORCES', NE/F005806/1, NERC New Investigators Awards, the Gordon and Betty Moore Foundation, a Royal Society University Research Fellowship and a Leverhulme Trust Research Fellowship. Fieldwork in the Democratic Republic of the Congo (Yangambi and Yoko sites) was funded by the Belgian Science Policy Office BELSPO (SD/AR/01A/COBIMFO, BR/132/A1/AFRIFORD, BR/143/A3/HERBAXYLAREDD, FED-TWIN2019-prf-075/CongoFORCE, EF/211/TREE4FLUX); by the Flemish Interuniversity Council VLIR-UOS (CD2018TEA459A103, FORMONCO II); by L'Académie de recherche et d'enseignement supérieur ARES (AFORCO project) and by the European Union through the

FORETS project (Formation, Recherche, Environnement dans la TShopo) supported by the Xlth European Development Fund. EMV was supported by fellowship from the CNPq (Grant 308543/2021-1). RAPELD plots in Brazil were supported by the Program for Biodiversity Research (PPBio) and the National Institute for Amazonian Biodiversity (INCT-CENBAM). BGL post-doc grant no. 2019/03379-4, São Paulo Research Foundation (FAPESP). D.A.C. was supported by the CCI Collaborative fund. Plots in Mato Grosso, Brazil, were supported by the National Council for Scientific and Technological Development (CNPq), PELD-TRAN 441244/2016-5 and 441572/2020-0, and Mato Grosso State Research Support Foundation (FAPEMAT)—0346321/2021. We thank E. Chezeaux, R. Condit, W. J. Eggeling, R. M. Ewers, O. J. Hardy, P. Jeanmart, K. L. Khoon, J. L. Lloyd, A. Marjokorpi, W. Marthy, H. Ntahobavuka, D. Paget, J. T. A. Proctor, R. P. Salomão, P. Saner, S. Tan, C. O. Webb, H. Woell and N. Zweifel for contributing forest inventory data. We thank numerous field assistants for their invaluable contributions to the collection of forest inventory data, including A. Nkwasiwbe, ITFC field assistant.

Author contributions D.L.M.C. and S.L.L. conceived and developed the study. D.L.M.C. performed the analysis with M.J.P.S. and P.I.P. and input from S.L.L. D.L.M.C., P.I.P., G.C.P., A.L. and M.J.P.S. developed tools to support the analysis. D.L.M.C. and S.L.L. wrote the manuscript with significant input from M.J.P.S., R.G.P. and M.I.D. S.L.L., B.S. and C.E.N.E. curated the AfriTRON forest plot data. N.B., P.P. and G.D. curated the Central African Plot Network forest plot data. H.T.S. curated the ATDN forest plot data. F.S. curated the Slik et al.^{4,25} Southeast Asia forest plot data. S.L.L. and O.L.P. curated the T-FORCES Southeast Asia carbon monitoring network. O.L.P., S.L.L., T.R.B., B.S. B.S.M., C.E., E.H., L.Q., A.L. and G.P. provided ForestPlots.net pan-tropical data management. All co-authors contributed data, reviewed, approved and had the opportunity to comment on the manuscript.

Competing interests The authors declare no competing interests.

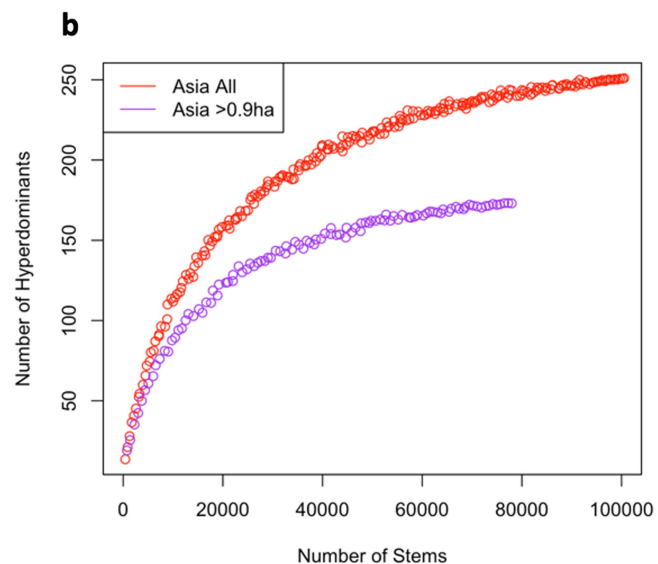
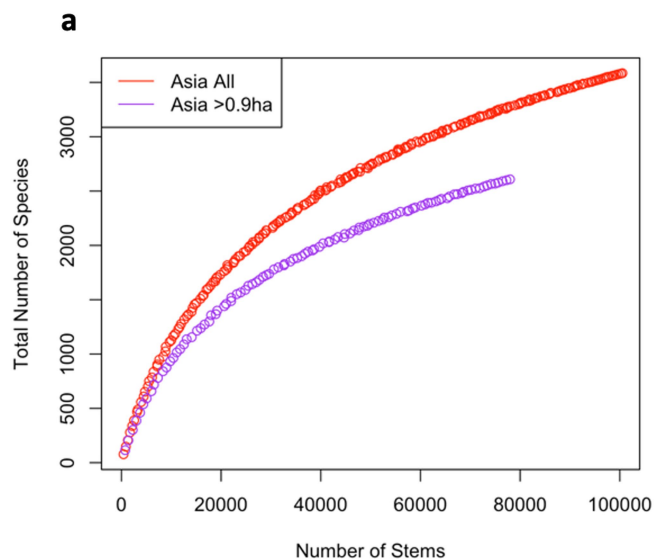
Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41586-023-06820-z>.

Correspondence and requests for materials should be addressed to Declan L. M. Cooper or Simon L. Lewis.

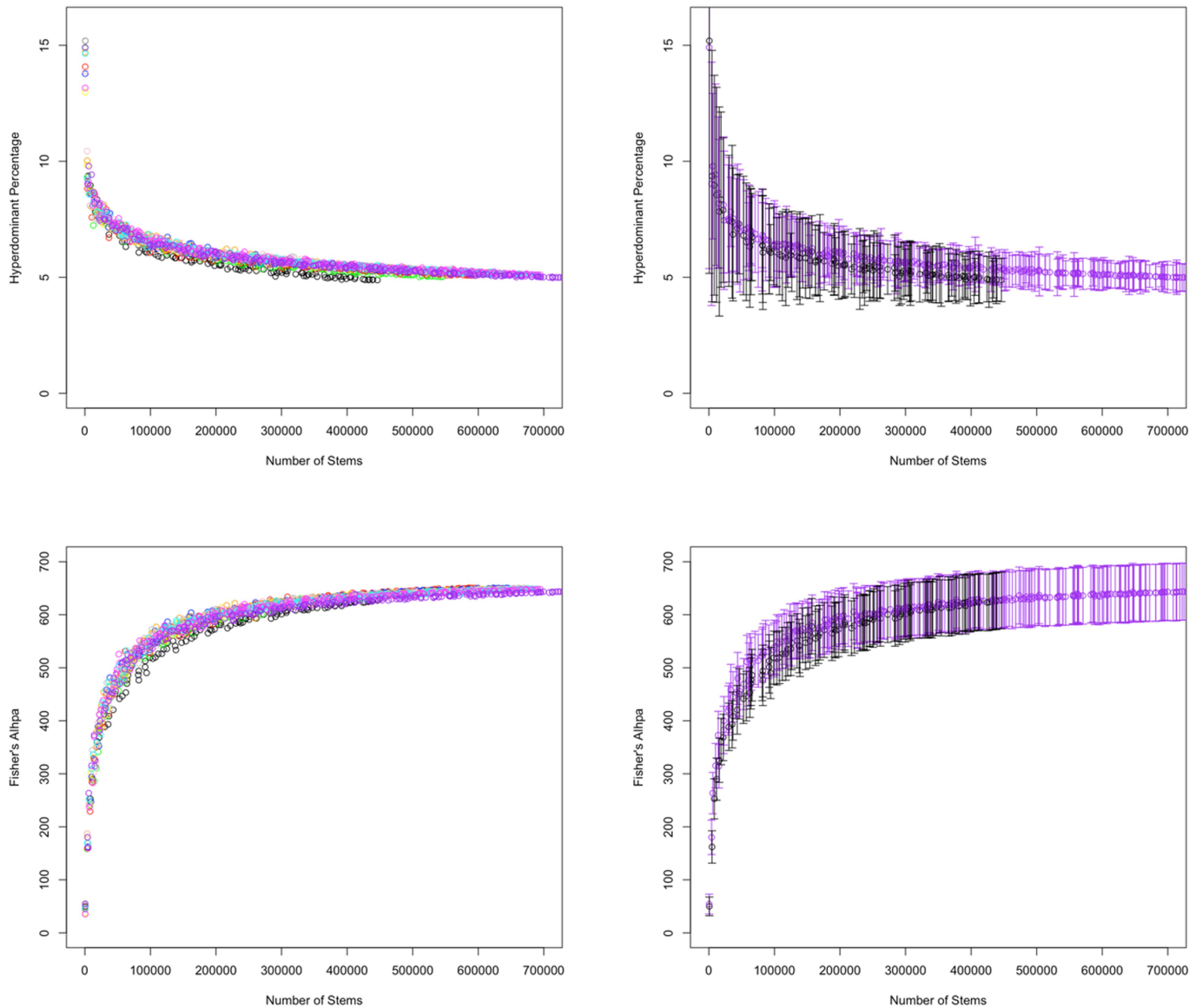
Peer review information *Nature* thanks Miguel Martinez-Ramos, Thorsten Wiegand and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Peer review reports are available.

Reprints and permissions information is available at <http://www.nature.com/reprints>.



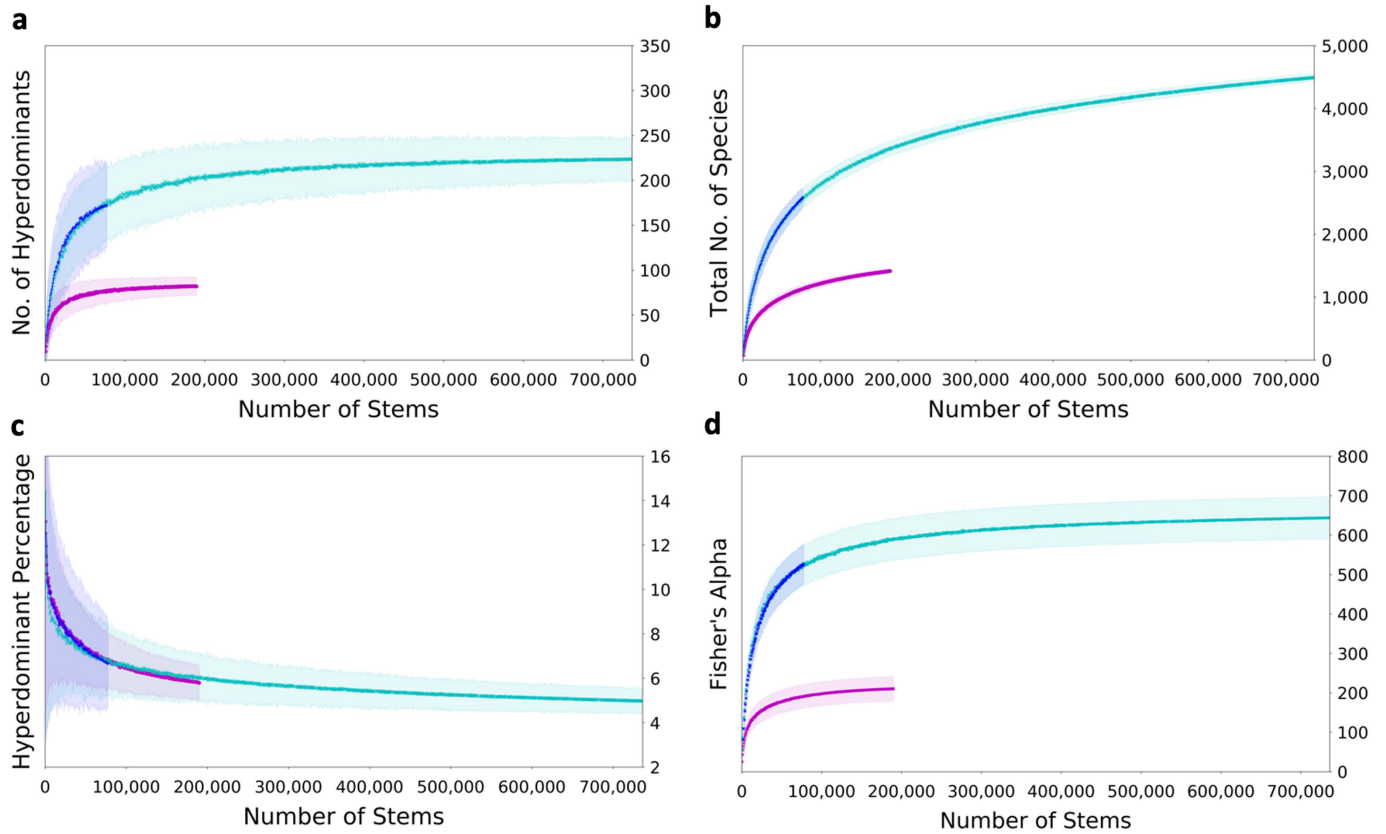
Extended Data Fig. 1 | Impact of plot size on rarefaction curves of total species (a) and number of hyperdominants (b) in the Asia data. Red points represent the full Southeast Asia data (mean values across iterations of

subsamples), including all plot sizes (mean plot size: 0.877 ha, median plot size: 0.5 ha); Purple points represent the Southeast Asia data restricted to plots ≥ 0.9 ha (mean plot size: 1.59 ha, median plot size: 1 ha).



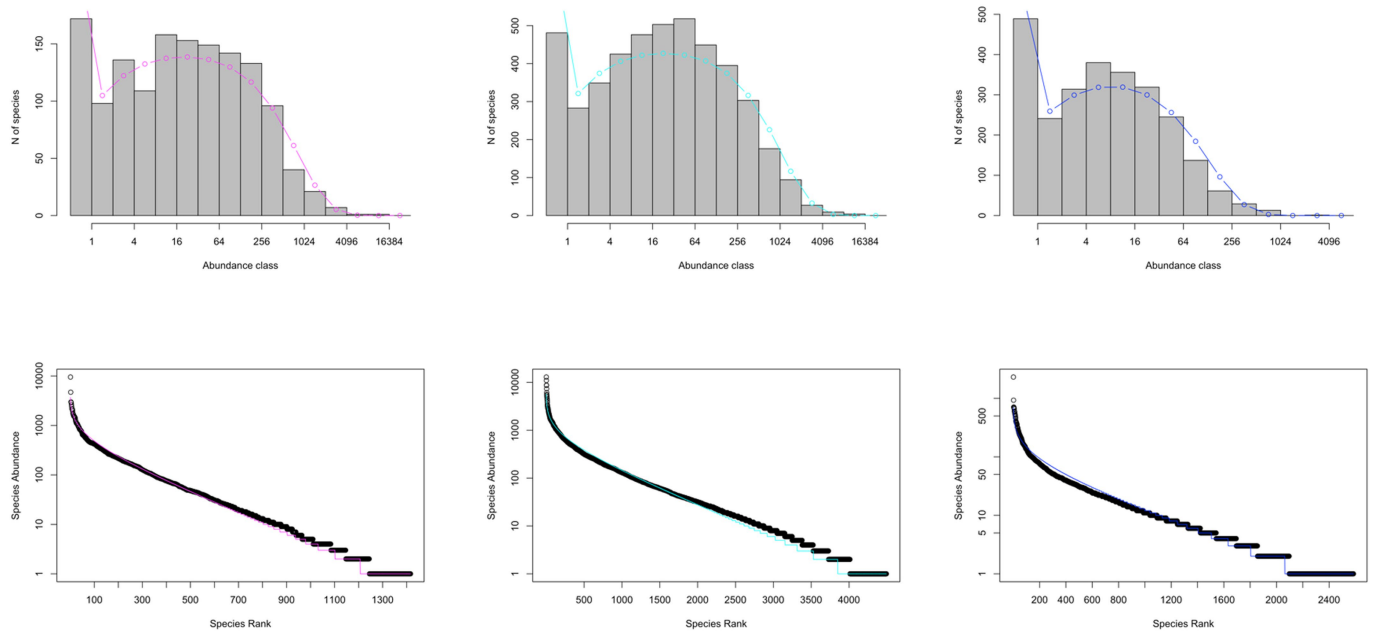
Extended Data Fig. 2 | Impact of spatial clustering of plots on rarefaction curves of hyperdominant percentage (first row) and Fisher's Alpha (second row) in the Amazonia data. Purple points and confidence intervals represent the full data; black points and confidence intervals represent a subset of the data in which one plot is sampled from each spatial cluster of plots; other coloured points represent subsets of the data in which 2, 3, 4, ..., 10 plots (or the

total number of plots in the cluster) are sampled from each spatial cluster of plots. Points give the mean values across iterations of subsamples. Confidence intervals are derived via the standard deviation across iterations of subsamples taken with replacement at each sampling point. Note that although resampling for rarefaction was done by subsampling tree inventory plots, the curves are re-plotted with an x-axis of number of stems.



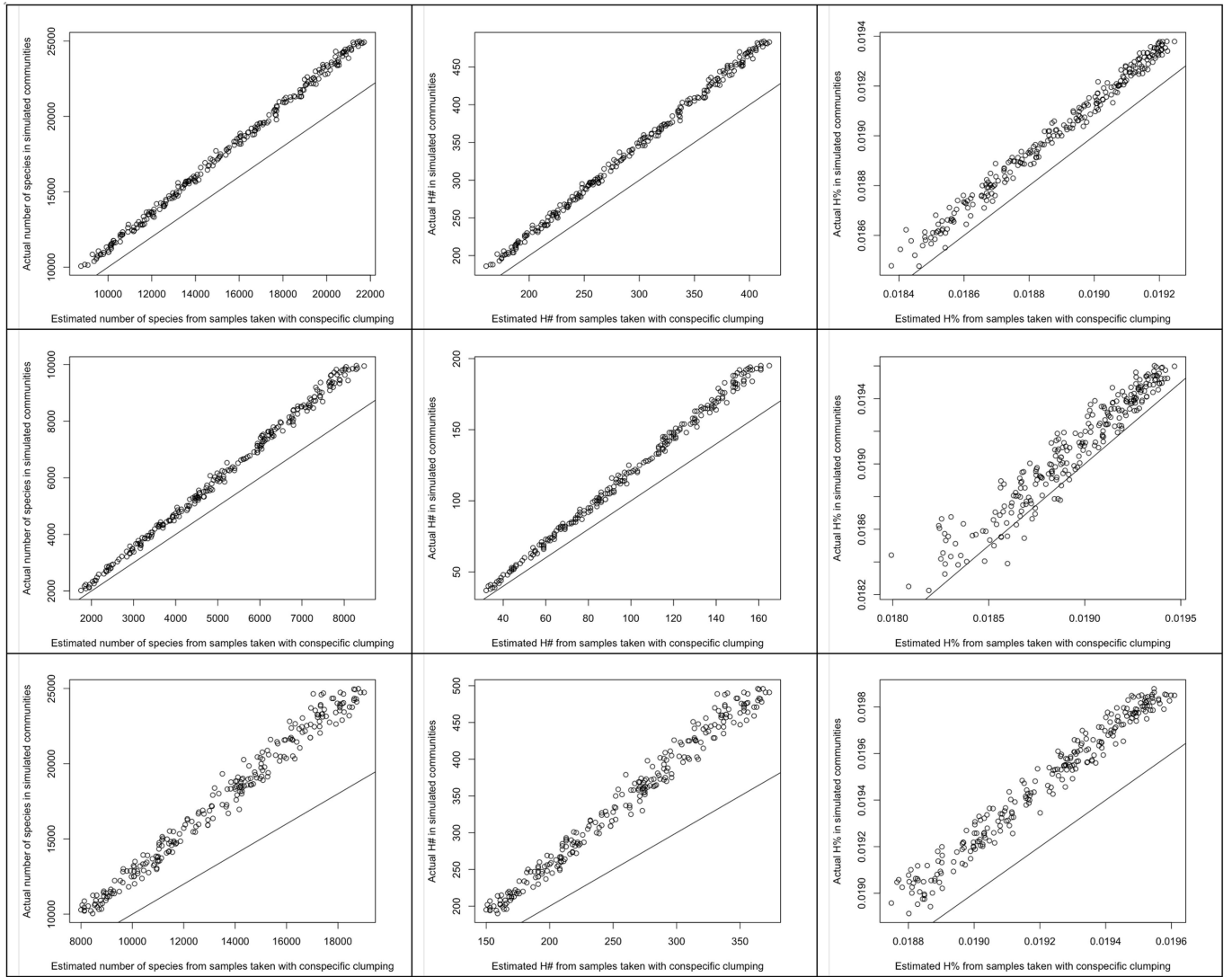
Extended Data Fig. 3 | Complete rarefaction curves showing the effect of increasing sampling on the number of hyperdominants (a), total species (b), hyperdominant percentage (c), and fitted values of Fisher's α (d). In tropical Africa (magenta), Amazonia (cyan), Southeast Asia (blue). Markers represent rarefied points (mean values across iterations of subsamples); shaded areas

represent confidence intervals (CIs). Confidence intervals are derived via the standard deviation across iterations of subsamples taken with replacement at each sampling point. Note that although resampling for rarefaction was done by subsampling tree inventory plots, the curves are re-plotted with an x-axis of number of stems.



Extended Data Fig. 4 | Preston plots (top row) and rank abundance distributions (bottom row) showing the empirical species abundance distributions for Africa (left) Amazonia (middle) and Southeast Asia (right) with log series fits overlaid. Histogram bars display the empirical species abundance distributions as Preston plots (top row); black markers show the

empirical species abundance distributions as rank abundance distributions (bottom row); overlaid points and lines show log series fits to empirical species abundance distributions in Africa (magenta), Amazonia (cyan), and Southeast Asia (blue).



Extended Data Fig. 5 | Bias correction of estimates of species richness (first column), number of hyperdominants (second column), percentage hyperdominance (third column) for the Amazonia (first row), Africa (second row) and Southeast Asia (third row) datasets. X-axes show estimated values derived from samples of the simulated communities taken with conspecific aggregation, Y-axes show true values of the simulated communities. Points show estimated true values for each of the 250 simulated communities.

1:1 equivalence shown by straight line in each plot. For number of hyperdominants and total species plots, simulated communities containing 100 to 25,000 species in Amazonia and Southeast Asia, 100 to 10,000 species in Africa are shown. For percentage hyperdominance, simulated communities containing 10,000 to 25,000 species in Amazonia and Southeast Asia, 2,000 to 10,000 species in Africa are shown.

Extended Data Table 1 | Empirical summary statistics and hyperdominance results for tree species data in Africa, Amazonia, and Southeast Asia

	Plots	Stems	%ID	H#	TS	H%	α
Africa	368	189,948	92.7	82	1,416	5.79	210
Amazonia	1,097	736,270	93.7	224	4,492	4.99	644
Southeast Asia	103	77,587	91.6	172	2,585	6.65	526

#H = Number of hyperdominants, TS = Total Species, H% = Percentage hyperdominance, α = Fisher's α , Stems = Total number of stems, Plots = Total number of plots, %ID = Percentage of stems identified to the species level.

Extended Data Table 2 | Rarefied minimum number of species required to account for 10%, 20%, ..., 90% of trees in the Africa, Amazonia, and Southeast Asia data, resampled to the common sample size of the Asia dataset (77,587 stems)

	10%	20%	30%	40%	50%	60%	70%	80%	90%
Africa	4 [2,7]	13 [8,17]	26 [19,33]	46 [36,57]	77 [62,92]	121 [102,139]	186 [163,209]	296 [264,328]	592 [494,689]
Amazonia	8 [4,13]	28 [17,38]	60 [42,77]	106 [79,133]	174 [134,215]	276 [214,337]	434 [339,528]	709 [553,864]	1413 [1029,1797]
Southeast Asia	9 [5,13]	26 [17,35]	54 [36,72]	98 [66,130]	172 [125,219]	285 [220,350]	468 [380,556]	790 [670,910]	1778 [1427,2129]

Percentage headings represent the different dominance thresholds. Confidence intervals are derived from the standard deviation across iterations of subsamples taken with replacement at the sample size of the Asia dataset.

Extended Data Table 3 | Rarefied proportion of total species required to account for 10%, 20%, ..., 90% of trees in the Africa, Amazonia, and Southeast Asia data, resampled to the common sample size of the Asia dataset (77,587 stems)

	10%	20%	30%	40%	50%	60%	70%	80%	90%
Africa	0.36 [0.12,0.61]	1.11 [0.71,1.51]	2.28 [1.64,2.93]	4.09 [3.1,5.07]	6.79 [5.39,8.20]	10.67 [8.89,12.44]	16.47 [14.28,18.65]	26.18 [23.22,29.15]	52.28 [43.00,61.56]
Amazonia	0.33 [0.15,0.51]	1.09 [0.67,1.50]	2.32 [1.63,3.01]	4.14 [3.09,5.19]	6.80 [5.24,8.36]	10.74 [8.40,13.09]	16.90 [13.30,20.50]	27.63 [21.68,33.58]	55.06 [40.37,69.76]
Southeast Asia	0.35 [0.19,0.51]	1.01 [0.61,1.41]	2.09 [1.27,2.91]	3.79 [2.38,5.20]	6.65 [4.59,8.71]	11.03 [8.24,13.82]	18.10 [14.47,21.73]	30.56 [25.78,35.34]	68.78 [52.50,84.97]

Percentage headings represent the different dominance thresholds. Confidence intervals are derived from the standard deviation across iterations of subsamples taken with replacement at the sample size of the Asia dataset.

Extended Data Table 4 | Extrapolated minimum number of species required to account for 10%, 20%, ..., 90% of trees in Africa, Amazonia, Southeast Asia, and the cross-regional total at the regional scale

	10%	20%	30%	40%	50%	60%	70%	80%	90%
Africa	7 [4,10]	22 [19,25]	41 [38,44]	69 [66,71]	104 [101,107]	154 [151,157]	228 [225,231]	354 [351,357]	713 [709,717]
Amazonia	20 [16,25]	62 [58,66]	120 [116,124]	196 [192,201]	299 [295,304]	443 [438,447]	651 [647,656]	1000 [995,1005]	1892 [1886,1899]
Southeast Asia	12 [1,23]	51 [40,62]	106 [95,116]	179 [168,189]	278 [268,289]	417 [406,427]	625 [614,636]	988 [977,1000]	2243 [2225,2262]
Total	39 [21,58]	135 [117,153]	267 [249,284]	444 [426,461]	681 [664,700]	1014 [995,1031]	1504 [1486,1523]	2342 [2323,2362]	4848 [4820,4878]

Percentage headings represent the different dominance thresholds. Prediction intervals combine uncertainty from the standard error of predicted means and the residual standard deviation of the regression of the bias correction fit. 'Total' minimum number of species required to account for 10%–90% of trees across all of the regions are calculated as the sum of the number of hyperdominants across the three major tropical forest regions.

Extended Data Table 5 | Extrapolated proportion of total species required to account for 10%, 20%, 30%, ..., 90% of trees in Africa, Amazonia, Southeast Asia, and the cross-regional total at the regional scale

	10%	20%	30%	40%	50%	60%	70%	80%	90%
Africa	0.16	0.48	0.90	1.48	2.23	3.31	4.90	7.60	15.29
Amazonia	0.14	0.44	0.86	1.41	2.16	3.21	4.72	7.25	13.73
Southeast Asia	0.14	0.46	0.91	1.5	2.32	3.45	5.14	8.10	18.34
Total	0.13	0.44	0.88	1.46	2.24	3.33	4.94	7.70	15.93

Percentage headings represent the different dominance thresholds. 'Total' minimum proportion of total species required to account for 10%–90% of trees across all of the regions are calculated as the proportion between the sum of the number of hyperdominants and the sum of total species across the three major tropical forest regions.

Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our [Editorial Policies](#) and the [Editorial Policy Checklist](#).

Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

n/a Confirmed

- | | | |
|-------------------------------------|-------------------------------------|--|
| <input type="checkbox"/> | <input checked="" type="checkbox"/> | The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> | A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | The statistical test(s) used AND whether they are one- or two-sided
<i>Only common tests should be described solely by name; describe more complex techniques in the Methods section.</i> |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> | A description of all covariates tested |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> | A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> | A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals) |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | For null hypothesis testing, the test statistic (e.g. F , t , r) with confidence intervals, effect sizes, degrees of freedom and P value noted
<i>Give P values as exact values whenever suitable.</i> |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated |

Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection No software was used for data collection.

Data analysis The custom R code (version 4.3.1) used to run the analyses and produce the figures is available on GitHub with the identifier <https://github.com/declancooper/CommonSpecies2022.git>

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](#) for further information.

Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our [policy](#)

The species abundance data that support the findings of this study are available from <https://doi.org/10.6084/m9.figshare.21670883>. WorldClim bioclimatic data are available from <https://www.worldclim.org/data/bioclim.html>.

Human research participants

Policy information about [studies involving human research participants and Sex and Gender in Research](#).

Reporting on sex and gender	NA
Population characteristics	NA
Recruitment	NA
Ethics oversight	NA

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☒ Life sciences ☐ Behavioural & social sciences ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	No sample size calculation was performed. We selected all available plots meeting the following criteria: forest inventory plots ≥ 0.2 ha in size, situated in structurally intact (no logging or fire), closed canopy (not dry forest or savanna) tropical forest, with enumeration of all stems ≥ 10 cm diameter, in which $\geq 80\%$ of stems are identified to the species level. These criteria allow direct comparisons to be made with results from previous studies investigating common species abundances in Amazonia. The major tropical forest regions of Amazonia, Africa, and Southeast Asia are adequately represented. This is the largest dataset of repeatedly measured plots ever used to calculate long-term trends in African forest carbon dynamics.
Data exclusions	Plots not meeting the following criteria were excluded from analysis: forest inventory plots ≥ 0.2 ha in size, situated in structurally intact (no logging or fire), closed canopy (not dry forest or savanna) tropical forest, with enumeration of all stems ≥ 10 cm diameter, in which $\geq 80\%$ of stems are identified to the species level. These conditions were pre-established in line with previous studies.
Replication	Replication of repeated random subsampling 200 times ensured that derived results were stable and reproducible.
Randomization	Forest inventory data were partitioned into the the Amazonia, Africa, and Southeast Asia study regions by location. In all of the analyses, sampling and sub-sampling by plots and by trees within plots was done completely randomly within regions.
Blinding	Blinding is not relevant to this study.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> Antibodies
<input checked="" type="checkbox"/>	<input type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology
<input checked="" type="checkbox"/>	<input type="checkbox"/> Animals and other organisms
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data
<input checked="" type="checkbox"/>	<input type="checkbox"/> Dual use research of concern

Methods

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging