INTRODUCTION

Seed dispersal, a mutualistic interaction resulting from fruit consumption and the deposition of intact seeds is tightly linked with macroevolutionary patterns (Burin et al. 2021) and is a major engine for trait diversification in both plants and frugivores (Gómez & Verdú 2012; Guimarães et al., 2011; Lengyel et al., 2010; Rojas et al., 2012, but see Scott, 2019). Mutual and reciprocal fitness benefits arise from fruit-frugivore interactions, as fruits are key resources for many vertebrates, and most tropical forest trees rely on fruit-eating vertebrates as seed dispersal vectors (Howe, 2014). Frugivory has evolved independently numerous times along the evolutionary history of vertebrates. This is also true for Neotropical primates (Hawes & Peres, 2014), a major Order of fruit-eating vertebrates (Fleming and Kress, 2013; Lim et al., 2021). Fruit intake is a condition for dispersal potential,
and all Platyrrhini primates routinely eat fruits (Hawes & Peres, 2014). The amount of fruit in the diet has significant consequences on the quality components of the seed dispersal effectiveness (Fuzessy et al., 2016, 2017). Yet, the extent to which variation in the degree of frugivory explains a possible shared evolutionary history between primates and Angiosperms remains largely unexplored (but see Gómez and Verdú 2012; Lim et al., 2021).

A persistent knowledge gap preventing a broader understanding of the role of fruit-frugivore interactions in shaping the evolution of primates and plants lies in the methodological challenges in addressing the consequences of seed dispersal to the concurrent diversification of both interacting clades (DeCasien et al 2017, Scott, 2019). Seed dispersal is a mutualistic interaction characterised by low degree of specialisation (Donatti et al., 2011; Stiles & Rosselli, 1993), and most primates have generalist feeding habits, consuming fruits from a wide range of plant species (Hawes & Peres, 2014; Heymann & Fuzessy, 2021; Lim et al., 2021). Thus, the potential of the mutual fitness benefits in promoting plant-primate co-diversification (correlation in speciation events) and/or coevolution (reciprocal evolutionary changes in traits) remains challenging to assess.

Considerable effort has been dedicated to uncover how Primates have diversified, and to what extent behavioural, ecological and morphological traits have contributed to both speciation and extinction rates (Gittleman & Purvis, 1998; Gómez & Verdú, 2012; Harcourt et al., 2002; Isaac et al., 2005; Matthews et al., 2011; Nunn et al., 2004). Primates are socially and ecologically complex organisms (Dunbar 2013; Lefebvre et al. 2004; Tran, 2014), but unequivocal evidence supporting the role of frugivory and seed dispersal in primate diversification remains scarce. The capacity to efficiently detect, consume fruits and disperse seeds is linked to larger geographical range sizes, higher diversification rates (Gómez & Verdú, 2012; Valenta et al., 2018) and cognitive complexity (DeCasien et al. 2017). However, among many traits, activity pattern, but not frugivory, was the only parameter correlated with primate diversification, with higher diversification rates observed in diurnal primates compared to nocturnal species (Scott, 2019). Considering that a diurnal habit is tightly related to primate colour vision, the emerging pattern may also reflect the strict relationship between foraging behaviour and the capacity to visually detect fruits (Kawamura, 2016). For example, the evolution of the trichromatic colour vision by most anthropoid primates has been linked to the efficient detection and selection of food, particularly ripe fruits among leaves in dappled light (Smith et al., 2003).

The evolutionary consequences of seed dispersal also remain unclear from the plant perspective. The intimate relation with certain groups of vertebrates is hypothesised to promote the evolution of dispersal syndromes, i.e. the non-random association of plant traits with specific disperser groups (Gautier-Hion et al., 1985). For instance, many studies attempted to distinguish bird-dispersed from primate-dispersed fruits based on fruit features such as colour, size and seed traits (Gautier-Hion et al., 1985; Janson, 1983; Lomáscolo et al., 2008; Voigt et al., 2004), but the specificity of such syndromes remains contentious (Valenta et al., 2018).

Theoretical studies on mutualistic networks suggest that the establishment of a link between two partners occurs when an interaction evolves successfully (Burin et al., 2021), and more species can be connected by evolutionary trait convergence (Guimarães et al., 2011). Although we are still unable to draw strong evidence of the potential of seed dispersal in promoting reciprocal selective responses between closely linked primates and plants (but see Guimarães et al., 2017), recent eco-phylogenetic tools allow us to assess both the shared evolutionary histories and the contribution of a particular clade to the coevolutionary dynamics (Blasco-Costa et al. 2021; Hutchinson et al., 2017). In particular, eco-phylogenetic analyses allow the detection of a non-random shared pattern or signal in the evolutionary trajectories of interacting species (e.g. frugivores and plants, hosts and parasites).

Estimation of the cophylogenetic signals (CS) in fruit-primates interactions allows quantifying the degree at which the topology and chronology of the phylogenies of interacting clades are congruent, and whether interaction between evolutionarily coupled taxa still occurs. Thus, CS underscores more precise inferences on how ecological interactions shape diversification patterns (Aizen et al., 2016; Balbuena et al., 2013; Hutchinson et al., 2017).

Here we tested for a phylogenetic congruence in primates and plants at both continental and regional scales in the Neotropics, the most biodiverse region of the planet (Raven et al., 2020). Under a strong CS, we expected a low overlap in fruit consumption among primates and other fruit-eating vertebrates, since strong associations may involve strong reciprocal and exclusive selection pressures (prediction I). We further assessed whether the magnitude of CS is driven by (1) the number of interactions performed (i.e. the generalism degree), (2) primate functional traits related to fruit consumption, such as feeding guilds and frugivory degree (the proportion of fruits in the diet relative to other items, Hawes & Peres, 2014), and (3) dispersal-relevant fruit and seed traits known to shape seed dispersal abilities (fruit length and seed diameter). The contribution of seed dispersal to primate diversification was mostly mediated by the broadening of mutualists’ geographic ranges (Gómez & Verdú, 2012). Therefore, we expect that primate species with a broad range of interaction partners belong to lineages with higher diversification and will contribute the most to the CS; plant species performing the lowest number of
interactions as a consequence of morphological constraints related to dispersal (large seeds and fruits) will contribute the most to the CS, since those species tend to be exclusively or primarily dispersed by primates (prediction II) (Jordano, 1995; Valenta et al., 2018). As a consequence, seed and fruit sizes are expected to be plant traits defining the CS strength (prediction III) (Jordano, 1995; Valenta et al., 2018). Finally, CS is particularly expected to arise in the most frugivorous lineages of primates, whereas less frugivorous feeding guilds and the lowest frugivory degrees will contribute less to define the past history shared among Neotropical primates and Angiosperms (prediction IV).

MATERIALS AND METHODS

Database

We performed a comprehensive literature search on plant species potentially dispersed by Neotropical primates, using both Web of Science and Google Scholar. Search details and the complete list of studies are available in Supplementary Material S1.

To evaluate the overlap in fruit consumption between primates with other vertebrates, based on the list for primates, we searched for other vertebrates also interacting with the fruits. We obtained records of frugivore-plant interactions whenever seeds were recorded to have been either swallowed and potentially defecated intact, or spat and/or spat out intact by animals away from the parent tree. We obtained data for birds, bats, small mammals and ungulates (Supplementary Material S2).

Phylogenetic reconstruction

We used ultrametric phylogenies and reconstructed both plant and primate phylogeny at the species-level. Detailed information on phylogenetic reconstruction methods is provided in Supplementary Material S3.

Cophylogenetic signal (CS)

To test for a CS between Angiosperms and primates, we implemented a Procrustean Approach to Cophylogeny (PACo; Balbuena et al., 2013). Under high CS values, diversification patterns of associated organisms over evolutionary time tend to be dependent, and congruences (i.e. topological similarity) between the phylogenies of interacting clades are expected to occur. The PACo approach addresses such phylogenetic congruence by optimising the topological fit of two phylogenies using interaction graphs of a given matrix (detailed explanation of the procedure is available in Supplementary Material S4).

Congruence expresses the extent to which each node in a given phylogenetic tree corresponds to a position in the phylogenetic tree of the interacting clade. A perfect congruence can be interpreted as a strong cospeciation evidence, which may or may not result from coevolutionary mechanisms. In nature, in addition to cospeciation, other types of evolutionary events can act concurrently, so such a perfect fit is rarely observed. Thus, the cophylogenetic signal is estimated by the squared residual distance, $r$, between two corresponding points in the phylogenetic trees of two interacting clades. PACo thus returns a quantification of the global fit of the topologies of the phylogenies based on observed interactions as the sum of squared residual distances ($R = \sum r^2$) between phylogenetic-interaction graphs (Balbuena et al., 2013). As in any regression analysis, the smaller the residual distances (Procrustean residuals), the better the fit of the two phylogenies to each other and the more support for a hypothesis of CS as reflected by the extant interactions (Hutchinson et al., 2017).

High CS illustrates a strong congruency of the phylogenies of the two partner groups. Procrustean residuals measure the variation in the topological fit that is not explained by the co-phylogenetic structure of the interaction matrix, so that interactions with small Procrustean residuals contribute the most to CS, whereas those deviating more from the expectation derived from the shared phylogenetic history contribute the less to CS. CS was considered to be significant when it was smaller than 95% of the values obtained from 1000 randomisations of the aggregated interaction dataset.

To ensure that the continental-level cophylogenetic pattern was not due to distinct geographical subsets of primate plants coexisting in different regions, we split our dataset into the major Neotropical forests (Morrone 2014): Atlantic, Amazon and Mesoamerican forests. We then filtered our dataset and obtained, for each region, specific phylogenies of plants and primates, and their respective seed dispersal interactions. We then measured the CS and tested for its significance following the same statistical approach as for the continent-wide assessment.

The influence of interaction degree and functional traits on CS

We firstly characterised each primate and plant species by its interaction degree, i.e. the total number of interactions performed in our dataset. To assess how the interaction degree of plants and primates affect the magnitude of the CS, we fitted two Generalised Linear Mixed Models (GLMMs) using the ‘lmerTest’ package (Kuznetsova et al., 2017), one for plants and one for primates. Abundant species tend to be generalists, so to account for the variation in geographic distribution area among primate species, we used primate geographic range as a proxy of their density, given the lack
of abundance data for each specific site. This approach is supported by strong associations between species distribution ranges and population sizes (Chiarello & Melo 2001; Estrada et al. 2017). Data on primate distribution range (in km²) were extracted from IUCN maps (IUCN 2021). In each model, we then used Procrustean residuals calculated by the PACo function as the response variable, the primate distribution area as a random effect, and the interaction degree of plants and primates as predictor variables.

Then, we obtained data on primate (body size, degree of frugivory and dietary guild) and plant functional traits (fruit length and seed diameter) from the literature. As primate functional traits, we included variables recognised to influence the potential of fruit consumption. We calculated primate frugivory degree as the percentage of the diet that corresponds to fruit with pulp, excluding seed predatory interactions (data from Hawes & Peres, 2014), and we divided primates into five dietary guilds based on the percentage of food items in the diet: frugivores, folivores, insectivores, omnivores and seed predators. For plant traits, we included variables recognised to define consumption by frugivores. Details on classification and data collection are provided in Supplementary Material S5. To test whether functional traits of primates and plants influence the magnitude of the CS, we fitted a GLMM and included the Procrustean residuals calculated by the PACo function as the response variable and functional traits as predictor variables. We tested the model for multicollinearity of the variables by using the variance inflation factor (VIF). Multicollinearity occurs when two or more predictors are correlated and provide redundant information about the response. VIF values for body size were found to be high (VIF = 10.83, tolerance = 0.09), which were also correlated with model intercept (0.9). Further, we obtained AIC values for both models, with and without body size, and found that the model without body size had the lowest value. Facing the likely confounding effects of body size added to the collinearity produced, and the AIC values, we removed this variable from our model (see Supplementary Material S6).

To account for the possible influence of sample size of interactions and primate geographic range, we also used the total time spent in the field and primate distribution area as random variables in the GLMM. Information on the total time spent in the field was extracted from the studies from which we obtained data on interactions (Supplementary Material S6).

The final GLMM model included then Procrustean residuals as a response variable, primate geographic range, plant species, sampling effort and primate distribution area as random effects, and functional traits (frugivory degree and dietary guilds for primates; fruit length and seed diameter for plants) as fixed effects.

**RESULTS**

We gathered information on 1632 interactions between 39 primate species, and 964 plant species distributed in 312 genera and 98 families across Neotropical rainforests. Primates were the exclusive seed dispersers of 627 (65%) species (in 233 genera) (Figure 1). Birds overlapped 147 plant species with primates (in 80 genera), bats 48 (in 32 genera), ungulates 17 (in 16 genera) and small mammals 9 (in nine genera). Finally, 112 plant species (in 68 genera) were found to be dispersed by primates and more than one vertebrate group. The most primate-exclusively dispersed family was Sapotaceae (70 species, or 5.5% of the total 1273 species), followed by Menispermaceae (22 species, or 5% of the 440 species) and Moraceae (44 species, or 3.7% of the 1180 species) (Figure S7).

**Continent-wide patterns**

We detected a coupled evolutionary history among primates and plants based on seed dispersal, as evidenced by a strong and significant cophylogenetic signal ($R_{Continental}^2 = 0.98, p < 0.001$). The residual contribution to the overall magnitude of the signal varied among interacting species. Specialisation or generalisation of each primate species (quantified by the number of interaction partners) predicted the CS of their interactions: the most interacting primates (those presenting highest degrees) had the shortest residual distances, thus contributing more to the CS ($t = -2.39, p = 0.02$). In turn, the number of interactions performed by plants had no influence on the CS ($t = 1.79, p = 0.07$, Figure S8).

The folivores (Alouatta) and the major frugivorous primates (Ateles, Lagothrix and Brachyteles) contributed more to the strength of the CS, followed by the omnivores (Cebus and Sapajus) and frugivore-insectivores (Leontopithecus, Leontocebus and Saginus). On the other hand, mostly seed predators (Cacajao, Chiropotes and Calliebus species) showed the largest Procrustean residual values, with the smallest contribution to the signal (Figure 2, Tables S9 and S10a).

Primate functional traits (degree of frugivory: $F_{Continental} = 3.18, p = 0.04$; feeding guild: $F_{Continental} = 574.56, p < 0.001$), but not plant traits (fruit length: $F_{Continental} = 2.53, p = 0.11$; seed diameter: $F_{Continental} = 0.89, p = 0.35$), determined the magnitude of the influence of the interaction on the overall CS.

**Regional-scale assessment**

The continental pattern was consistent when broken-down at the regional scale. We found a strong and significant CS on primate-fruit interactions in the Atlantic forest ($R_{Atlantic}^2 = 0.98, p < 0.001$), Amazon...
Feeding guild was the most important variable accounting for variation in the magnitude of the CS, regardless of biogeographic region (\(F_{\text{Atlantic}} = 373.7, p < 0.001\); \(F_{\text{Amazon}} = 323.7, p < 0.001\); \(F_{\text{Mesoamerican}} = 95.7, p = 0.01\), Tables S10). The contribution by each primate feeding guild in the Atlantic forest followed the continental pattern, where folivores contributed the most to the CS, followed by major frugivores, omnivores, insectivores and predators. In the Amazon, the major frugivores had the greatest contribution, whereas omnivores had the greatest contribution in Mesoamerican rainforests (Figure 3).

The only effect detected for plant functional traits was observed in the Amazon, where fruit length determined the contribution of the interaction to the CS (\(F_{\text{Amazon}} = 9.94, p = 0.002\)). We observed no other effect of plant traits (see Tables S10).

**DISCUSSION**

Understanding the eco-evolutionary dynamics of species interactions remains challenging due to intrinsic complexities involved in mutualisms, especially in species-rich communities (Hall et al., 2020). Further complexities arise from the multi-specific, mega-diversified nature of most mutualisms among free-living species, but also from difficulties to infer process from the simple co-phylogenetic patterns (Poisot, 2015). Seed dispersal has been hypothesised to drive coevolution between plants and frugivores, shaping the present-day patterns of interactions and the evolution of fruit traits (Eriksson, 2016; Guimarães et al., 2011, 2017; Lengyel et al., 2010; Rojas et al., 2012), but empirical evidence remains elusive both for pattern and process. Using an unprecedented database of fruit-frugivore interactions, we show a significant cophylogenetic signal (CS), which reveals that...
phylogenies of Neotropical primates and Angiosperms are strongly coupled, and provide support for the long-standing claims that interactions between Angiosperms and primates are shaped by coevolution (Eriksson, 2016; Sussman, 1991, 1995).

CS emerged independently in three tropical rainforests (Mesoamerica, Amazonia and Atlantic Forest), and the consistency between the continental and regional scales evidences spatial replication of the process and supports the idea of similar eco-evolutionary dynamics operating irrespective of species composition in regional meta-communities. The observed CS was mainly driven by generalist primates dispersing a diverse assemblage of angiosperms, partially in agreement with our prediction II. On the other hand, contrarily to prediction III, plant functional traits were not important to define CS either at continental or at local scales, except for fruit size in the Amazon. As for primates function, CS emerged primarily in primate lineages with the highest degrees of fruit consumption, whereas less frugivorous feeding guilds and/or the lowest frugivory degrees were less important to define the past history shared among Neotropical primates and Angiosperms regardless of spatial scales, as expected by our prediction IV.

Recent estimates suggest primate origins from 55.8 to 50.3 Mya, in the early Eocene (O’Leary et al., 2013), consistent with the rise and dominance of modern tropical rainforests lineages (Carvalho et al. 2021; Sussman, 2017). Primate diversification entailed changes in locomotion, reproductive biology, skull morphology, dentition and feeding niches, which likely arose as a coevolutionary response to fruiting plants (Sussman et al., 2013). In New World primates, the divergence times in the modern Platyrhini are estimated to be 20.1 Mya, during the Oligocene-Miocene boundary (Schrago, 2007). This was nearly the time of diversification of many extant lineages of fleshy-fruited Angiosperms. Even though many families seem to have conserved fruit traits ever since the Eocene (Eriksson et al., 2000), drupes and berries, the most consumed fleshy-fruits by primates, have evolved much more recently. For instance, Bolmgren and Eriksson (2005) found that almost half of fleshy-fruited clades are younger than 40 Mya, and many examples in dominant and keystone plant resources in Arecaceae, Rubiaceae, Solanaceae and in the Myrtales experienced a remarkable increase in diversification between 40 and 18 Mya (Berger et al. 2016; Eriksson, 2016). This timescale overlap in diversification of both Neotropical primates and fleshy-fruiting plants is consistent with the idea that seed dispersal played a role in their coupled evolutionary history.

Seed dispersal clearly drove the congruence in phylogenies, as evidenced from strong effects of the primate frugivory-related traits. Chronological evidence supports the origin of primate frugivory at the end of the first stage of Angiosperm diversification (Eriksson, 2016). As expected, the cophylogenetic relation among plants and primates was shaped mainly by the most frugivorous species, although specific plant traits were not pivotal in driving such a signal. Primarily frugivores and folivores belong to the Atelidae family, the largest among Neotropical primates. Atelids are the most important and legitimate seed dispersers in neotropics (Fuzessy et al., 2016) and contributed the most to the cophylogenetic signal. On the other hand, seed predators, belonging to the Pitheciidae family, had the least contribution. It suggests that similar mechanisms might be operating for dietary guilds within a same family. Seed predators are absent in Mesoamerican forests (Estrada et al. 2006), where frugivore-insectivores were found to contribute the least, whereas omnivores followed by frugivores and folivores were those with the highest contribution to the CS. Among-guild differences in contribution may be a consequence of the relative importance of guild species diversity in each region, added to their conservation status, and therefore the number of interactions performed. For instance, in the Atlantic Forest, the only frugivores are the species of *Brachyteles*, both threatened by extinction. On the

**FIGURE 2** Contribution to the overall co-phylogenetic signal (CS) by primate feeding guild at continental scale (Neotropic). Lower residual values represent higher contributions to the CS. Boxplots show the median, upper and lower quartiles, and minimum and maximum observed values. Scattered dots represent individual Procrustean residual values for each primate species.
other hand, in the Amazon, *Ateles* and the *Lagothrix* play the most important role as frugivores and are much better represented.

Recent studies on the visual adaptations of primates support intrinsic relationships between primate diversification and the capacity to detect plant resources (Valenta et al., 2018, Onstein et al., 2020, but see Heymann & Fuzessy, 2021). The evolution of modern primates, therefore, may be directly related to improved means of efficiently exploiting fleshy fruits (Sussman, 2017), although, in some cases, evidence is limited to particular clades and recent times (e.g. Onstein et al., 2020). Our analysis reveals tree-wide contributions to the CS, although some clades (e.g. Atelidae) had larger contributions. These findings, together with the evidence of an early frugivorous habit over primate diversification (Sussman et al., 2013), support the prominent phylogenetic congruence in Neotropical primate-fruit interactions driven by primate frugivore-related, functional traits. It also provides clues that diversification is an ongoing process, given the large contribution from recent Platyrrhini clades.

**FIGURE 3** Contribution to the overall co-phylogenetic signal (CS) by primate feeding guild at regional scale (a—Atlantic forest, b—Amazon forest, and c—Mesoamerican forest) considering plant phylogeny at species level. Lower residual values represent higher contributions to the CS. Boxplots show the median, upper and lower quartiles, and minimum and maximum observed values. Scattered dots represent individual Procrustean residual values for each primate species.
Reconstructing Angiosperm evolutionary history based on fruit traits as a response to selective pressures generated by interacting primates is still challenging. In this case, the phenotypic responses may be more strongly subjected to phylogenetic constraints, i.e. reflect inherited ancestral characteristics rather than traits adapted to an ecological niche (Jordan, 1995; Valenta & Nevo, 2020). Our results suggest that diaspor size, key morphological constraint to the establishment of mutualistic interactions (Dehling et al., 2014), was not an important driver of the CS. Most studies evaluating plant adaptive responses to mutualisms with primates based on current empirical data have found weak evidences of phylogenetic signals (Valenta et al., 2016, Valenta et al., 2018—fruit colour; Nevo et al., 2018, Nevo et al., 2020—fruit scent; Valenta et al., 2016—size, mass and hardness), a likely consequence of the low specificity found in extant interactions.

Frugivorous birds and mammals are the major seed dispersers in the tropics (Fleming and Kress 2013). Considering the primate-centered basis of our dataset, and despite the low overlap in fruit consumption among primates and other fruit-eating vertebrates, complying with our prediction I, our results reflect the trend of plants to share multiple dispersers. This is not only true for recent times, so it is important to consider that, throughout evolutionary history, primates were not the only interacting taxa with the radiating fruiting plants.

At the time of Angiosperm radiation, the diversification of the earliest modern-looking primates in parallel with that of other mammals and fruit-eating birds, propelled the beginning of a shared evolutionary relationship (Sussman, 2017). It helps us to explain how such low specificity prevents detection of strong phylogenetic patterns. However, the CS arose even in a supergeneralist system where other frugivorous groups may be also shaping plant diversification. We also observed a dissimilarity in terms of the number of interactions performed by each plant and primate species explaining the influence on the CS: primates dispersing the greater diversity of plants (i.e. generalists) contributed the most to CS strength, while no effect of plant degree was detected.

Besides fruits, Platyrrhini primates also include leaves, flowers, seeds, nuts, nectar, and animal prey as feeding resources (Hawes & Peres, 2014; Lim et al., 2021). Distinct amounts of each item vary across taxa (Hawes & Peres, 2014; Lim et al., 2021), thus less frugivorous primates and other coexisting frugivorous clades may act together favouring a process known as diffuse coevolution (Erikson 2016). It is unlikely that frugivores and plants share a very tight coevolutionary history (Valenta & Nevo, 2020), such as those observed in host-parasite interactions (Brooks, 1988; Gandon & Michalakis, 2002), or plant-pollinator interactions (Herrera, 2019). Instead, spatio-temporal asymmetries, disruptions in relationships between patterns, and shifts between periods of coevolution among coexisting clades should lead to reciprocal adaptive changes, ultimately resulting in a weak process (Erikson 2016), as suggested by our results. Primate evolution seems to have somehow “tracked” plant radiations, resulting in a coevolutionary history with asymmetric influences.

The multi-specific nature of the process delineating the CS does not mean that primates and plants have not coevolved or that coevolution has necessarily been the primary force fueling diversification (see, e.g. Althoff et al., 2014; Poisot, 2015). Instead, it seems to occur in a much more complex framework, including both direct and indirect effects underlined by three non-exclusive main processes (Guimarães et al., 2011, 2017). First, selection regimes imposed by generalised multiple-partner interactions, such as seed dispersal, are the outcome of a complex interplay among selection pressures operating through multiple pathways, leading to slow, but continuous, coevolution. Coevolution repeatedly reshapes selection regimes and species traits by speeding up the overall diversification rate in interacting clades. Second, coevolution results in higher trait complementarity among interacting partners (reduced mismatch and increased trait convergence), and the level of integration may provide a mechanism for the emergence of community-level trait patterns. Finally, convergence tends to be higher in the presence of super-generalists, here represented by the most frugivorous primates, which interact with a wide-range of plant species and thus establish the magnitude of the observed CS. Our results indicate a strong non-random pattern in the diversification of primarily primate-dispersed neotropical plants and their primate disperser partners, reinforced by the replicated consistency found in three major neotropical biomes. Although the processes involved in such high-level macroevolutionary patterns remain obscure, our approach highlights replicated consistency over large biogeographic extents and evidences the strong potential of highly diversified mutualisms among free-living species in macroevolution.

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AUTHORSHIP
All authors conceived the ideas; LFF and MV designed the methodology and analysed the data; LFF and FAOS led data collection; LFF led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.
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DATA AVAILABILITY STATEMENT
All published data sources are referenced in the manuscript (main text and/or supplementary material). At the time of publication, raw data will be available at https://github.com/LFuzessy (https://doi.org/10.5281/zenodo.5578594).

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