

Tracing the impact of the Andean uplift on Neotropical plant evolution

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Recent phylogenetic studies have revealed the major role played by the uplift of the Andes in the extraordinary diversification of the Neotropical flora. These studies, however, have typically considered the Andean uplift as a single, time-limited event fostering the evolution of highland elements. This contrasts with geological reconstructions indicating that the uplift occurred in discrete periods from west to east and that it affected different regions at different times. We introduce an approach for integrating Andean tectonics with biogeographic reconstructions of Neotropical plants, using the coffee family (Rubiaceae) as a model group. The distribution of this family spans highland and montane habitats as well as tropical lowlands of Central and South America, thus offering a unique opportunity to study the influence of the Andean uplift on the entire Neotropical flora. Our results suggest that the Rubiaceae originated in the Paleotropics and used the boreotropical connection to reach South America. The biogeographic patterns found corroborate the existence of a long-lasting dispersal barrier between the Northern and Central Andes, the “Western Andean Portal.” The uplift of the Eastern Cordillera ended this barrier, allowing dispersal of boreotropical lineages to the South, but gave rise to a huge wetland system (“Lake Pebas”) in western Amazonia that prevented *in situ* speciation and floristic dispersal between the Andes and Amazonia for at least 6 million years. Here, we provide evidence of these events in plants.

biogeography | Neotropical biodiversity | Rubiaceae

The uplift of the tropical Andes in the Neogene had a profound impact on the history of the South American continent. It changed the course of the Amazon system from flowing northwestwards to the modern system that flows to the Atlantic side (1, 2) and affected the climate of the region by forming the only barrier to atmospheric circulation in the Southern Hemisphere (3). Recent phylogenetic studies have shown that the Andean orogeny had also a major role in the evolution of the Neotropical flora. The Neotropics hold the highest plant species diversity in the world (4). This richness has traditionally been explained in terms of environmental factors (5), but lately, more integrative explanations have been advanced that emphasize the role of historical and evolutionary factors in the shaping of Neotropical diversity (6, 7). The “tropical conservatism hypothesis,” for example, argues that there are more plant species in the Neotropics simply because more lineages originated and diversified there, owing to the long-term climatic stability of the region and the tendency of species to retain their climatic niches over evolutionary time (7, 8). It is now also clear that part of this richness has been gained by the migration of lineages from other biogeographic regions (6). For instance, pantropically distributed plant families such as Malpighiaceae, Fabaceae, and Annonaceae (6, 9, 10, 11) originated at temperate latitudes as part of the former “boreotropical flora” (12–14) and subsequently entered the Neotropics via the mountain ranges of Central America and the newly formed Northern Andes. One point in common to these hypotheses is the key role that the formation of the tropical Andes would have played in the

historical diversification of the Neotropical flora (15). Recent phylogenetic studies have shown that the Andean uplift acted both as a dispersal route for boreotropical lineages (16, 17) and as a driver in promoting rapid diversification, via allopatric speciation and ecological displacement, in highland (16–19) and montane (11) habitats.

Fewer studies, however, have documented the impact of the Andean uplift on the lowland Amazonian flora. Clearly, the uplift must have affected these taxa by forming a new biotic barrier and profoundly changing the hydrology and climate of the region (20). Furthermore, previous biogeographic studies on Andean radiations have typically considered the Andean orogeny as a single, time-limited event, usually in connection with the final (Miocene to Pleistocene) uplift of the Andes (11, 19). This contrasts with geological reconstructions indicating that the uplift took place in discrete periods, progressing from south to north and from west to east and affecting different regions at different times (2, 3, 21, 22). Episodic marine incursions, related to global sea level rises during the extensional tectonic phases that followed periods of major uplift, had a dramatic impact in the drainage patterns of the region, as evidenced by paleogeographic and paleontological evidence (1, 2, 23–28). These marine incursions have been discussed in relation to their role as a pathway in the evolutionary transition from marine to freshwater habitats of Neotropical fishes (24, 29), but they could also have acted as barriers to dispersal or as vicariance events fragmenting the ranges of terrestrial animals and plants. It seems surprising that, despite increasingly detailed geological reconstructions (2, 24, 26–28), thus far no study has attempted to document the effect of these events on the evolution of the Neotropical flora. Generally, detailed reconstructions have been hampered by the lack of resolution in many Andean species-rich clades (19). Current biogeographic methods require well-resolved phylogenies, and uncertainty in phylogenetic relationships makes it difficult to reconstruct the specific sequence of geological vicariance and speciation events.

Here, we use an integration of phylogenetic, biogeographic, and molecular dating methods to reconstruct the evolutionary history of tribes Cinchoneae and Isertieae, which together form one of the major clades of Neotropical Rubiaceae. The distri-

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bution of this clade spans highland and montane habitats (the Andes, the Guiana, and the Brazilian Shields), as well as lowland tropical forests (the Amazonia and Chocó). It thus offers a unique opportunity to disentangle the evolutionary processes underlying botanical evolution in the region. Our results reveal an extraordinary level of congruence between the evolution of the Neotropical Rubiaceae and the progressive west-to-east Andean uplift, which brought about a series of marine incursions and lacustrine systems that blocked the dispersal of plants and shaped the distribution of the modern flora.

Study Group

The coffee family (Rubiaceae) is the fourth largest family of flowering plants, with some 13,100 species in 611 genera and 3 subfamilies (30, 31). Although cosmopolitan in distribution, its highest diversity is distinctly confined to the tropics. Subfamily Rubioideae is pantropically distributed and comprises some highly diverse groups in the Neotropics (e.g., Palicoureeae and Spermacocae), but it is otherwise concentrated to the Old World where it probably originated (32, 33). Subfamily Ixoroideae shows a similar pattern, because it comprises a species-rich Neotropical clade (the “Condaminae–Calycophylleae” alliance) but is otherwise concentrated in the Paleotropics. Contrastingly, except for tribe Naucleaeae, the large subfamily Cinchonoideae is predominantly Neotropical. In tropical South America, Cinchonoideae is represented by sister tribes Cinchoneae and Isertieae, which have been shown to build a strongly supported clade (34) and is sometimes treated as a single tribe (30). Comprising some 130 species of small trees and shrubs divided into 11 genera [see [supporting information \(SI\) Table S1](#)], the Cinchoneae and Isertieae are important ecological components of a wide array of habitats. Some species are also economically important as a source of quinine. The distribution of Isertieae is concentrated in the lowlands of the Amazon basin and eastern Guiana, whereas Cinchoneae species are mainly confined to the highland and montane habitats of the Northern and Central Andes, reaching up to 3,300 m ([Figs. S1 and S2](#)).

Results and Discussion

Gentry (35), following Raven and Axelrod (36), listed the Neotropical Rubiaceae as a Gondwana-derived group, evolving in isolation since the separation of South America from Africa. This hypothesized origin predicts that (i) the group has a minimal age of 100 Ma (37), and (ii) Old and New World lineages are reciprocally monophyletic (6). In contrast, the competing hypothesis of boreotropical origin predicts that (i) South American groups are derived from northern relatives with Old World taxa as sister groups, (ii) the divergence between Old and New World groups occurred between 40 and 50 Ma, i.e., around the Eocene climatic optimum, which favored the exchange of tropical floristic elements between these land masses (38), and (iii) Early Tertiary fossils have been found in North America, Europe, or Asia (6).

Our biogeographic reconstruction ([Fig. 1 I and II](#)) corroborates a boreotropical origin in all 3 predictions: (i) our phylogeny ([Fig. S3](#)) shows the Neotropical sister tribes Cinchoneae and Isertieae nested within a clade of mainly Central American and Antillean tribes, together sister to the essentially African tribe Naucleaeae and the Paleotropical subfamily Ixoroideae; (ii) our divergence time estimates place the most recent common ancestor (MRCA) of the Rubiaceae in the Early Paleocene (66.1 Ma, 63.0–68.8; see [Table S2](#)), whereas the minimum age of Cinchonoideae is estimated as only 51.3 (47.8–54.6) Ma, well after the last known island chain between Africa and South America possibly existed [Late Cretaceous, \approx 88 Ma (37)]; and finally, (iii) several *Cephalanthus* fossils indicate the presence of the tribe Naucleaeae in Europe from the Late Eocene (39) (see [SI Text](#)). Until the Late Eocene or Early Oligocene, a continuous

belt of boreotropical vegetation covered much of southern North America, southern Eurasia, and northwestern Africa (40). At that time, plant migration through direct land connection or across limited water gaps (6) could have been possible through the North Atlantic “Thulean” land bridge (13, 41) or through the Early–Mid Tertiary “Beringian land bridge” (42). Although climates during the Early Eocene were warmer than today (38), Beringia was in a considerably higher paleolatitude than the Thulean bridge. Dispersal of boreotropical elements is therefore considered more likely across the North Atlantic during this period (13, 42), which is also supported by the fact that the oldest *Cephalanthus* fossils have been found in Europe (39). Together, these lines of evidence strongly suggest that the Rubiaceae used the corridors provided by boreotropical vegetation and the North Atlantic land bridge as a pathway to reach North America in the Late Paleocene/Early Eocene ([Fig. 2I](#)).

From North America, dispersals to South America ([Fig. 2II](#)) may have been facilitated by the proto-Greater Antilles in the Early Eocene and later by the Greater Antilles and the Avies Ridge around the Eocene/Oligocene boundary [GAARlandia, 33–35 Ma (43)]. Our divergence time estimates suggest that the ancestors of Cinchonoideae arrived in northwestern South America around the Early/Middle Eocene (49.2 Ma, 44.9–53.1; node 26 in [Fig. S4](#)). At that time, sea levels some 50 m above today’s (44) created a marine incursion from the Caribbean that limited land dispersal eastwards (1, 24) and another incursion from the Pacific that blocked dispersal to the south (24, 45, 46) ([Fig. 2III](#)). Lowland areas were covered by closed-canopy tropical rainforests (47). Most of the Andes had not yet been formed, except for some low mountains in the regions now corresponding to the Central and Southern Andes (3, 21). Interestingly, the MRCA of tribes Isertieae and Cinchoneae is reconstructed as being lowland-adapted (see [Fig. S5](#)). But starting in the Middle Eocene, the Andean orogeny went through a major phase of mountain building, sometimes referred to as the Incaic II (21, 48). This phase was longitudinally widespread, and in the northern region it caused uplift of the Central Cordillera (21, 49). The newly formed montane habitats must have acted as an ecological barrier to lowland taxa, and this seems to explain the geographic disjunction (Andes vs. Amazonia) between the MRCA of tribes Cinchoneae and Isertieae ([Fig. 1III and Fig. S5](#)).

In the case of Isertieae, their MRCA is most likely reconstructed as being confined to lowland Amazonia ([Fig. 2III](#)), where it first radiated in the Late Oligocene, giving rise to the genera *Kerianthera* and *Isertia* ([Fig. 1III and Fig. S5](#)). Diversification in Isertieae occurred mainly in the Middle and Late Miocene, which is strikingly coincident with the uplift of the Eastern Cordillera in the Northern Andes [[Fig. 1IV](#); (2, 21)]. For tribe Cinchoneae, ecological adaptation to higher altitudes seems to have been the key to its diversification, with most speciation events confined to montane habitats in the Northern and Central Andes ([Fig. 1 III and IV and Fig. S5](#)).

Western Andean Portal. Most optimizations indicate that the MRCA of Cinchoneae was confined to the Northern Andes, presumably in the new habitats created by the Central Cordillera ([Fig. 1III](#)). From the Eocene to the Middle Miocene, some studies have proposed that marine incursions from the Pacific invaded a lowland corridor between the Northern and Central Andes at the latitude of southern Ecuador/northern Peru [\approx 3–5°S (1, 2, 45, 46, 50)], termed the “Western Andean Portal” (WAP, [Fig. 2III](#)) or “Guayaquil Gap” (23). Indication for these incursions comes from fossil occurrences of marine organisms and palynomorphs (23, 46) and paleosedimentary evidence (1, 2, 50). The WAP is suggested to have been uplifted, and marine incursions ended, in connection with the uplift of the Eastern Cordilleras of the Central and Northern Andes from the Middle Miocene onwards (13–11 Ma) (2, 21, 50).

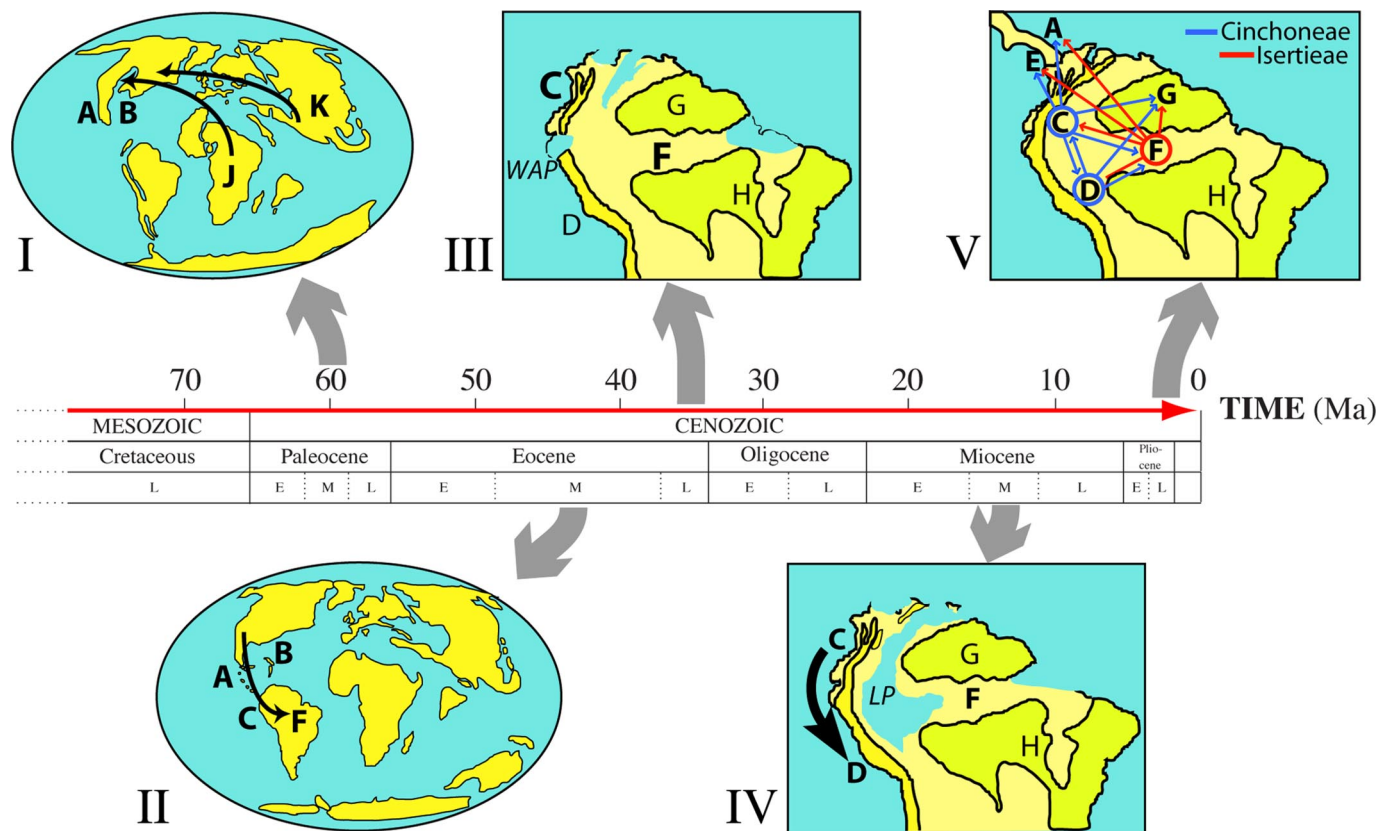


Fig. 2. Spatiotemporal evolution of the Neotropical Rubiaceae. (I) Paleocene: Rubiaceae ancestors use the boreotropical route to reach North America from the Paleotropics. (II) Early Eocene: Dispersal into South America, presumably facilitated by occasional island chains. (III) Late Eocene: North Andean and Amazonian lineages become isolated by marine incursions such as the Western Andean Portal (WAP). (IV) Middle Miocene: The gradual uplift of the Eastern Cordillera creates a huge watershed, Lake Pebas (LP). It also closes the WAP, enabling dispersal of plant lineages from the Northern to the Central Andes. (V) The Pebas system drains, promoting land dispersal of several lineages and rapid speciation of terrestrial plants in western Amazonia. Area codings as in Fig. 1. (Maps I–II are based on C. R. Scotese’s PALEOMAP project (www.scotese.com); maps III–V modified from refs. 2 and 28).

Our biogeographic reconstruction of subfamily Cinchonoideae (Fig. 1III) corroborates the existence of a dispersal barrier between the Northern and the Central Andes coincident with the WAP, and provides indication for the persistence of this barrier until the Middle Miocene. Throughout the Oligocene and Early Miocene, all ancestral area reconstructions in tribe Cinchoneae occur—partly or exclusively—in the Northern Andes (Fig. 1III). The uplift of the WAP is then observed as at least 5 independent migrations from the Northern to the Central Andes within the genera *Cinchona* and *Ladenbergia* (Fig. 1IV). All 5 events are dated as occurring around the Middle/Late Miocene (12–10 Ma, Fig. S4 and Table S2), almost simultaneously with the suggested end of marine incursions in the WAP (1, 2, 21, 50).

Several questions remain unanswered concerning the geographic extent and duration of marine settings in the WAP (24, 28). What is clear, however, is that it constituted an important and long-lasting biogeographic barrier, as evidenced by the fact that many montane taxa exhibit endemism centers in either side of the WAP (see *SI Text* for an expanded discussion), suggesting that species in those groups were not able to cross the WAP during their time of radiation. In plants, this pattern has been recognized in many families, such as Campanulaceae, Calceolariaceae, Tropaeolaceae, Loasaceae, Passifloraceae, Alstroemeriaceae, and Grossulariaceae (see *SI Text* for references). In birds, this area has long been recognized as a turnover point between the Northern and Central Andean biogeographic regions of endemism (51).

Lake Pebas. Until the end of the Oligocene (≈ 24 Ma), a fluvial system referred to as the paleo-Orinoco dominated the drainage of northwestern Amazonia and the foreland Andean basins toward Lake Maracaibo, on the Caribbean coast. Then, in the Early Miocene (≈ 23 Ma) geotectonic changes in the Amazon Basin associated with the ongoing uplift of the Eastern Cordillera in the Central Andes (28) caused western Amazonia to gradually become submerged, from south to north and from west to east. This process created a huge (>1 million km^2) system of long-lived lakes and wetlands from at least 17 to 11 Ma, known as “Lake Pebas” or the “Pebas Sea” (24–28). Whether it was a purely fluvio-lacustrine (fresh water) system or whether marine settings were also present is still a matter of debate (24, 27, 28). However, most authors agree that western Amazonia was completely flooded by some kind of wetland system from at least the Middle to the Late Miocene and that this system was connected to the Caribbean marine incursion in the north (Fig. 2IV). From the Late Miocene onwards (11–7 Ma), there was a new period of rapid mountain uplift, affecting mainly the Eastern Andean Cordilleras [sometimes termed Quechua phases II and III (21, 49)]. This presumably caused the western margin of the Guiana Shield to emerge, which closed the Caribbean connection of the paleo-Orinoco, shifted the drainage of the Amazon Basin eastwards, and led to the demise of Lake Pebas (27, 28) (Fig. 2V). Aquatic conditions, however, seem to have persisted in western-central Amazonia until at least 7 Ma, when the modern Amazon system came into place (28).

Previous studies have provided indications to the potential role played by Lake Pebas as a pathway for marine organisms to

disperse into fresh water biotopes, based on fossil occurrences of mollusks and DNA phylogenies of Neotropical fishes (23, 24, 26). On the other hand, if this wetland system was as large and interconnected as suggested, it should also have acted as a biotic barrier to the dispersal of terrestrial organisms between the Andes and the eastern Amazonian and Guiana regions.

Indeed, the existence of a dispersal barrier between the Andes and lowland Amazonia during the Middle Miocene is corroborated by our biogeographic reconstructions (Fig. 1IV). Until the Early/Middle Miocene boundary, several ancestors in tribe Cinchoneae are inferred to have been widespread in both of those areas. But after that—coinciding with the time Lake Pebas is proposed to have existed—all ancestral lineages in Cinchoneae and Isertieae appear to have occupied either the Andes or Amazonia, but not both. Most endemic species of Isertieae are currently found in Guiana and eastern Amazonian lowlands (Fig. S1A), whereas western-central Amazonian distributions are mainly represented by widespread species, suggesting that these distributions are the result of recent range expansion, probably after the drying of Lake Pebas. Similarly, several ancestral nodes in *Remijia* are reconstructed as occurring on both sides of the barrier (Fig. 1V), but these nodes are dated after the Miocene/Pliocene boundary (≈ 5.3 Ma, Fig. S4) and thus postdate the closing of the Pebas wetland system. The final reestablishment of land connections between the Andes and Amazonia is evidenced by at least 7 independent colonization events in Cinchoneae and Isertieae from the Late Miocene onwards (Fig. 1V). Eventually, the emergence of new lands in Central America after the closing of the Panama Isthmus (3.5 Ma) (21, 52) provided suitable areas for northwards dispersal of South American lineages. Species such as *Joosia umbellifera* and *Isertia haenkeana*, which are now widespread in Central America, probably dispersed soon after the establishment of that land connection (Fig. 2V).

Materials and Methods

Phylogenetic analyses were performed under parsimony and Bayesian methods as implemented in PAUP* (53) and MrBayes3 (54). We used representative genera of 4 families of Gentianales as outgroup, based on evidence that the Rubiaceae are the sister group to the rest of Gentianales (e.g., ref. 55). The ingroup included representatives of all Rubiaceae subfamilies, with focus on subfamily Cinchonoideae in which all tribes were represented (Fig. 1). The final dataset comprised 62 species and 5,894 characters, derived from matK, rbcL, ITS1–5.8S–ITS2, trnL-F, and rps16 (Table S3). Thus, our dataset comprised some 3–3.5 times more characters than similar studies (e.g., refs. 9 and 19), which resulted in a robust phylogeny where 80% of all tree nodes were strongly supported (Bayesian posterior probability values, $pp \geq 0.95$ or jackknife $\geq 85\%$). Dataset and trees are available from TreeBase (www.treebase.org) accession nos. S2334 and M4437. Evolutionary rates were estimated with the Penalized Likelihood algorithm implemented in r8s (56). Accurate fossil calibration has been suggested as a key factor in age estimation (57). Here, absolute ages were estimated by using fossil fruits and seeds of *Cephalanthus*, a genus with an exceptionally rich and reliable fossil record from the Late Eocene onwards (Fig. S4). Additionally, a maximum age constraint of 78 Ma was independently enforced on the basal node of the tree

based on the crown age of Gentianales (58). Finally, to incorporate topological and branch length uncertainty in our age estimates, 1,000 trees randomly sampled from the Bayesian stationary distribution were independently dated and results summarized to obtain median values and 95% credibility intervals of node ages (Fig. S4 and Table S2). See *SI Text* for a detailed description of the phylogenetic and dating methods used.

Eleven areas (Fig. 1 *Inset*) were defined for the biogeographic analysis based on the extant distribution patterns of Rubiaceae and on geological history (3, 21). Whenever possible, we tried to maximize congruence with other biogeographic studies in South America (59, 60). Dispersal-vicariance analysis [DIVA (61, 62)] was used to infer ancestral distributions and historical events involved in the biogeographic history of Rubiaceae. To overcome the uncertainty associated with phylogenetic estimation, we use an approach that averages DIVA biogeographic and temporal reconstructions over a Bayesian sample of highly probable trees (in this case $n = 1,000$), generating credibility support values for alternative phylogenetic relationships (63). Integrating over the posterior distribution of trees often reveals preference for a single or more restricted set of solutions, thus reducing the uncertainty in DIVA optimizations (63).

Incomplete taxon sampling may cause problems for historical inference methods because it reduces the accuracy of ancestral state reconstructions (see ref. 64 for a different view). This is particularly problematic in a large family such as Rubiaceae, spanning nearly all continents. To overcome this problem, we used encompassing areas outside the Neotropics, where most of Ixoroideae (as well as the first diverging clade in Cinchonoideae, the tribe Naucleaeae) are found. We also estimated the geographic bias in our sample for the Neotropical tribes Cinchoneae and Isertieae, the main focus of this article. Results showed that our taxon sampling is fairly representative of the actual distribution of the species within each genus and tribe (see Table S4, Fig. S6, and *SI Text* for a detailed discussion of distribution patterns in these tribes). In addition, to test the sensitivity of our DIVA reconstructions to missing taxa, we performed a series of heuristic simulations in which we added hypothetical taxa from underrepresented areas to key nodes in the phylogeny of these 2 tribes. DIVA ancestral reconstructions proved to be very stable to the addition of missing taxa, at least for the nodes involved in the WAP and Lake Pebas scenario (Fig. S7). We also examined the effect of missing taxa on divergence time estimations by randomly deleting taxa from the original sample and calculating divergence times on these reduced datasets. As Linder et al. (65) suggested, PL proved to be largely insensitive to taxon sampling, and for most nodes the estimated ages were within the 95% confidence interval from the complete dataset (Fig. S8; see *SI Text* for more details on simulations). This gives us confidence that the biogeographic scenario presented here (Fig. 2) would not be significantly altered by the addition of missing taxa.

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