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

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-

he 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

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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 Naucleeae, 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 Naucleeae 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, ~88 Ma (37)]; and finally, (*iii*) several *Cephalanthus* fossils indicate the presence of the tribe Naucleeae 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 MR-CAs 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. 1111). 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 [$\approx 3-5^{\circ}S$ (1, 2, 45, 46, 50)], termed the "Western Andean Portal" (WAP, Fig. 2111) 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. 1. Combined chronogram and biogeographic analysis of Neotropical Rubiaceae. The tree is the 50% majority-rule consensus (with compatible groups added) from the Bayesian analysis, with branches proportional to absolute ages (in millions of years) calculated from mean branch lengths of 6,000 Bayesian trees. Green bars indicate 95% confidence intervals of node ages estimated from 1,000 trees randomly sampled from the Bayesian stationary distribution. Node charts show the relative probabilities of alternative ancestral distributions obtained by integrating dispersal-vicariance analysis (DIVA) optimizations over the 1,000 Bayesian trees; the first 4 areas with highest probability are colored according to their relative probability in the following order: white > red > blue > gray; any remaining areas (usually frequencies <0.01) are collectively given with black color. Stars indicate calibration points. Red arrows indicate clades with a posterior probability <0.90. Present ranges for each species are given after the species name. Brackets identify subfamilies and tribes: CHI, Chiococceae; CIN, Cinchoneae; GUE, Guettardeae; HAM, Hamelieae; HIL, Hillieae; ISE, Isertieae; NAU, Naucleeae; RON, Rondeletieae. Shaded boxes indicate approximate periods of Andean uplift phases. The biogeographic interpretation of events I–V is summarized in Fig. 2. (*Inset*) Areas used in the biogeographic analysis. A, Central America, B, West Indies; C, Northern Andes; D, Central Andes; E, Chocó; F, Amazonia; G, The Guiana Shield; H, Southeastern South America; I, Temperate North America; J, Africa; K, Australasia. Topographic map from the National Geophysical Data Center (www.ngdc.noaa.gov).



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 lead to the demise of Lake Pebas (27, 28) (Fig. 2V). Aquatic conditions, however, seem to have persisted in westerncentral 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

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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 Naucleeae) 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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Supporting Information

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SI Text

Expanded Materials and Methods. Phylogenetic analyses. We reconstructed phylogenetic relationships using DNA sequences from 5 different markers: the matK and rbcL genes, the ITS1-5.8S-ITS2 and trnL-F regions, and the rps16 intron. Most of these sequences were previously obtained by using the methodology published in Andersson and Antonelli (1), except for 6 additional species of Isertia (I. haenkeana, I. hypoleuca, I. parviflora, I. pittieri, I. rosea, and I. spiciformis) that were newly sequenced for this study to increase taxon sampling within tribe Isertieae. Accession numbers and source of all sequences included in this study are listed in Table S3. Following Andersson and Antonelli (1), we used representative genera of 4 families of Gentianales as outgroup for the analysis, based on evidence that the Rubiaceae are the sister group to the rest of Gentianales (e.g., ref. 2). The dataset also included representatives from the 2 other Rubiaceae subfamilies Rubioideae and Ixoroideae, as well as genera representing all tribes in subfamily Cinchonoideae (Fig. 1). In addition, an extra outgroup species (*Nicotiana tabacum*) was included in the phylogenetic analysis to determine the position of the root within the basal branch, as required by the molecular dating method described below, but this taxon was subsequently excluded from the results. The complete dataset was realigned by using MAFFT v. 5.64 (3) and adjusted manually. Gaps were coded as present/absent (0/1) following the principles outlined by Antonelli (4) and included in the phylogenetic analysis. The final aligned data matrix comprised 5,894 characters, of which 1,600 derived from matK, 1,398 from rbcL, 1,254 from trnL-F, 908 from rps16, 676 from the ITS region, and 58 from gap codings. Of 2,153 variable characters, 1,165 were parsimony informative.

Parsimony jackknife support values (5) were estimated in PAUP v. 4.0b10 (6) by running 10,000 replicates with 37% deletion, 10 random addition sequence replicates, using TBR branch swapping and saving up to 30 trees per replicate. Following recent works (7, 8), the Akaike Information Criterion, implemented in MrModelTest v. 2.2 (9), was used to choose the optimal model of sequence evolution for each DNA marker. Bayesian analyses were subsequently implemented in MrBayes v. 3.1.2 (10) using the GTR+G model for matK and rps16, and the GTR+I+G model for ITS, *rbcL*, and *trnL*-F. Gap-codings were analyzed as a separate partition under the Restriction Site (Binary) Model. Two simultaneous analyses with 8 Metropolis-Coupled MCMC chains each were run for 2.5 million generations, sampling every 500th generation. Each analysis was started from different, randomly sampled topologies and let to run until the average standard deviation of split frequencies became <0.01, indicating convergence of trees and model parameters across the runs (11). The burn-in value was set in all cases to 1,000,000 generations after visual inspection of the split (clade) frequencies using the software AWTY (12). Final results were based on the pooled values from the 2 independent analyses. The parsimony jackknife consensus tree was virtually congruent with the Bayesian tree (Fig. S3), and both were very similar to the tree of Andersson and Antonelli (1). Nearly 80% of all tree nodes were strongly supported (Bayesian posterior probability values, pp $\geq 95\%$ or jackknife $\geq 85\%$). Kerianthera was placed with strong support as the sister to Isertia, and within the latter there were 3 strongly supported clades of sister species (Fig. S3). The Bayesian and jackknife trees, as well as the aligned dataset used for the analyses, are available from TreeBase (www.treebase. org), accession numbers S2334 and M4437.

Molecular dating. Divergence times were estimated on the tree topology with the highest posterior probability from the Bayesian analysis, but using mean branch lengths calculated from 1,000 post-burn-in trees from the Bayesian sample (see below). A likelihood ratio test (13) strongly rejected the hypothesis of a molecular clock for the tree (P < 0.0001). The clock-independent algorithm Penalized Likelihood (14) implemented in the software r8s v. 1.70 (15) was then used for estimating divergence times in the tree.

Fossils have been assigned to some 10 genera of extant Rubiaceae in our phylogeny (ref. 17; the Paleobiology Database, http://paleodb.org). However, in the majority of cases, a careful evaluation of the original fossil descriptions made evident that almost none of these could be deemed fully reliable in terms of taxonomic placement and/or geologic age. Approximately more than half of these records are fragmentary traces of leaves, and the rest are pollen grains examined with light microscopy. Considering the great convergence of leaf and pollen forms among angiosperms, which render even fresh material very difficult to identify in the absence of various other vegetative and floral parts, we have therefore opted for not using any of these 2 types of fossils as calibration points.

An important exception among Rubiaceae fossils exists in Cephalanthus. The oldest fossil of the genus is Cephalanthus kireevskianus from the Late Eocene of Germany (18), which appears to be both correctly identified (E. M. Friis, personal communication) and dated (S. Schultka, personal communication). Additional findings indicate that this species was common and widespread from the Late Eocene onwards in Europe (19) and in Western Siberia, where the species was originally described (20, 21). Another fossil species of the genus, C. pusillus, was described from the Middle Miocene of Denmark (22). Fossil fruits of Cephalanthus possess several morphological features that make them taxonomically recognizable by means of overall similarity. These include fruit type, a schizocarp; mericarp obovoid, slightly dorsoventrally flattened; ventral face flat with a shallow median furrow; dorsal face convex, apically truncate; placentation apical; seed strophiolate, apotropous pendulous, obovoid, slightly dorsoventrally flattened; strophiole apical, sickle-shaped; hilum dorsal, marked by a narrow slit (ref. 22; Fig. S4A Inset).

The continuous fossil records in every single geologic epoch from the Late Eocene to the Pliocene in almost 20 fossil sites (Fig. S4 *B* and *C*), combined with the reliability of their taxonomic placement thanks to a high quality of preservation, make *Cephalanthus* an ideal calibration point for the Rubiaceae phylogeny. The oldest fossil finding in *Cephalanthus* was therefore used to place a minimum age constraint of 33.9 Ma on the first split in tribe Naucleeae, separating *Cephalanthus* and *Hallea* + *Nauclea*. The stratigraphical age of this fossil was converted to an absolute age by using the ending point of the geological epoch to which it was assigned, using the time scale of Gradstein et al. (23). In addition, a maximum age constraint of 78 Ma was independently enforced for the basal node in the tree based on the crown age of Gentianales, as estimated from a well-sampled analysis of the asterids calibrated with multiple fossils (16).

Although our approach of using a single fossil calibration may seem too conservative (it could be that some other Rubiaceae fossils are correctly identified and dated) most fossils described in the literature (including the *Posoqueria* and *Cosmibuena* fossils cited in ref. 17) are too young to produce any influence in the molecular dating results obtained by using the *Cephalan*- *thus* and root calibrations (Fig. S4). The only potential exceptions are the following 2 fossils, which may have been correctly dated, but that we do not consider reliable for the reasons specified:

1) Assigned name: Remijia tenuiflorafolia

Type of fossil: Leaf traces

Minimum age: 47.46

Country: Argentina

Primary reference: ref. 24. Comments: In our judgment, the leaf characters originally used for placing these fossils in *Remijia* (size 9.5×2.25 cm, secondary veins 10–13, camptodromous venation, equally acute apex and base), as well as the illustrations provided, indicate that these fossil leaves could equally well belong to several other species of Rubiaceae (e.g., *Agouticarpa curviflora*, *Kutchubaea surinamensis*, or *Alibertia bertierifolia*, in tribe Gardenieae) or even to other plant families. Moreover, this taxon was described from southwestern Argentina, which is some 3,000 km south of the southernmost border of the present-day range of the genus

(Fig. S1D).

2) Assigned name: Psychotria eogenica

Type of fossil: Seeds

Minimum age: 28.4

Country: Peru

Primary reference: ref. 25.

Comments: As traditionally circumscribed, *Psychotria* is one of the most species-rich genera among angiosperms, at most comprising some 1,950 species (26, 27). Phylogenetic analyses have shown, however, that "*Psychotria* is broadly paraphyletic and defined by lack of characters used to define other genera in the tribe" (26). It is thus not surprising that the placement of this fossil in *Psychotria* was originally done "with some hesitation for the reason that in all seeds of that genus that I have seen they are smaller than the fossils and the ribbing is more continuous and is also more pronounced, as is also true in the case of related genera such as *Phialanthus* Grisebach and *Ixora* Linné" (25).

All other fossils listed in the Paleobiology Database and the literature for the genera sampled in this study would have been far too recent to reach the lower bound of the age intervals calculated using the 2 calibration points we rely on (i.e., *Cephalanthus* and the root). In sum, we argue that it is a better approach to base our dating analysis on a single (but by all means reliable) fossil, than using several more or less dubious records.

Finally, to account for topological and branch length uncertainty in our age estimates, 1,000 randomly chosen trees from the Bayesian stationary sample were independently dated and results summarized to obtain the median value and 95% credibility intervals of node ages (Fig. S4 and Table S2), by using the softwares TreeAnnotator and FigTree v. 1.4 (28).

Biogeographic Analysis. *Distribution and altitudinal data.* Except for *Remijia*, all larger genera of Cinchoneae and Isertieae have been the subject of recent taxonomic revisions (29–32), and fairly detailed distribution data are available for all major taxa (29–39). Distribution data on the remaining Neotropical Rubiaceae and the paleotropical genera were obtained from Andersson (40), Bridson and Verdcourt (41), Mabberley (42), Puff (43), and Smith (44). In addition, we searched for recent collections retrieved from various herbaria at the Global Biodiversity Information Facility (www.gbif.org), but such datapoints were only used when they were deemed reliable (e.g., identifications by experts).

Operational areas used in the biogeographic analysis. The delimitation of areas for the biogeographic analysis was based on the extant distribution patterns of Rubiaceae taxa (i.e., congruent distributional ranges shared by 2 or more species) and on geological history, i.e., areas historically isolated from one another by dispersal barriers (45, 46). In South America, we also tried to maximize congruence with other biogeographic studies in the

region by selecting similarly defined areas (47–50). However, our altitudinal boundaries were usually lower than it is commonly adopted, reflecting ecological constraints in our taxon distributions (many Rubiaceae are premontane species). In all, 11 areas were defined:

A: Central America. From southern Mexico (Veracruz, Oaxaca, Tabasco, Campeche, Yucatán and Quintana Roo) south to Panama. Although this region has had a complex geologic history and its land and island connections to South America are still prone to discussion, there has been a long-term isolation from South America until the uplift of the Panama Isthmus at 3.5 Ma (45, 51).

B: West Indies. Excluding Trinidad and Tobago, which are geologically and biologically more closely related to South America than to the other Caribbean islands. Although the Greater and the Lesser Antilles have different geologic histories, they have been long isolated by water from other American landmasses.

C: Northern Andes (10° N–5° S). From Venezuela and Colombia south to northernmost Peru (Piura, Cajamarca, and Amazonas), from elevations >500 m. This area is roughly the same as the Páramo recognized in other biogeographic studies (e.g., refs. 48–50), except that occurrences in this area are arbitrarily coded beginning at altitudes somewhat inferior to the ones generally adopted (due to the distributional patterns shown by our study taxa). Our delimitation of the Northern Andes is slightly different from that of Taylor (45) ranging from 10°N to 3°S. The reason for this is that we wanted the boundary between the Northern and the Central Andes to coincide with the Western Andean Portal (Guayaquil Gap), following Hoorn (52) and Hurgerbühler et al. (53). This region seems to be a major biogeographic barrier for many groups of Neotropical plants (see ref. 54). Also, many Cinchoneae species endemic to area C extend their range down to 5-6°S.

D: Central Andes (5°5–18°5). From Peru (San Martin and La Libertad) southwards to the Tropic of Capricorn, from elevations higher than 500 m. Similarly to the operational area defined for the Northern Andes, this area approximately corresponds to the Puna or Altiplano commonly recognized (e.g., refs. 48–50) but is used here as if occurring from a lower altitudinal limit.

E: The Chocó area. Comprises areas west of the Andes and below 500-m altitude in Colombia (Chocó, El Valle, Cauca, and Nariño), Ecuador, and Peru (Tumbes, Piura). This area is usually recognized by bird biogeographers as a center of endemism (e.g., refs. 47 and 55).

F: Amazonia. Comprises the lowland (<500 m) vegetation in Colombia, Ecuador, Peru, Bolivia, Brazil, Venezuela, Guyana, Suriname, and French Guiana and includes the islands immediately off the South American coast.

G: The Guiana Shield. Includes the elevated (\geq 500 m) areas in northeastern South America, comprising parts of Venezuela, Guyana, Suriname, French Guiana, and Brazil. It corresponds to the Guianan Bedrock region.

H: Southeastern South America. Mostly comprised of the Brazilian Shield, but also including the lowlands in eastern Brazil and the Rio Paraná drainage. The area corresponds to the pre-Cambrian Brazilian Bedrock formation.

I: Temperate North America. From the Tropic of Cancer northwards. This area is ecologically separated from Central America by the current occurrence of arid and semiarid habitats in northern Mexico.

J: Africa. Separated from South America at about 110–100 Ma (56, 57).

K: Australasia. Includes also Madagascar and the Seychelles, which were separated from Africa together with India already at \approx 121 Ma (58).

Dispersal-vicariance analysis. Dispersal-vicariance analysis (DIVA, refs. 59 and 60) was used to infer ancestral distributions at

internal nodes in the phylogeny of Rubiaceae and to identify the biogeographic events involved in the history of the group. DIVA has the advantage over more cladistically oriented methods that it makes no a priori assumptions about the shape or existence of general biogeographic patterns, making it very useful in regions where area relationships have varied greatly through time such as the Neotropics (47). However, like most biogeographic inference methods (61, 62), DIVA requires completely resolved, fully bifurcated trees. This can be a problem because unresolved trees are a common feature of Andean radiations, presumably due to rapid diversification rates. To account for phylogenetic uncertainty in our biogeographic reconstructions, we used here a method that averages DIVA reconstructions over a Bayesian sample of trees (in this case n = 1,000) reflecting credibility values on each clade (63). Moreover, it has been observed that 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). DIVA analyses were run unconstrained, i.e., with no constraint in the maximum number of areas applied to ancestral area optimizations (59).

Altitudinal optimization. To infer ancestral altitudinal ranges, Maximum Parsimony (Fitch) optimizations were performed in the software Mesquite v. 2.0.1 (64). Independent optimizations were run on 5,000 Bayesian trees from a stationary tree sample and plotted on the 50% majority-rule consensus tree (compatible groups added) of the Bayesian analysis.

Expanded Results and Discussion. Diversity patterns in Neotropical Rubiaceae show that although it is cosmopolitan in distribution, the highest diversity in family Rubiaceae is distinctly confined to the tropics. Subfamily Rubioideae is pantropically distributed and comprises some highly diverse groups in the Neotropics (e.g., Palicoureeae and Spermacoceae) but is otherwise concentrated to the Old World where it probably originated (65, 66). The majority of species in the subfamily Ixoroideae (except for the "Condamineeae-Calycophylleae" alliance) are also found in the Paleotropics, whereas the subfamily Cinchonoideae, with the exception of tribe Naucleeae, is predominantly Neotropical. There are several tribes (Hamelieae, Hillieae, Chiococceae, Rondeletieae, and Guettardeae) distributed in Central America and the West Indies, where they inhabit wet lowlands and cloud forests, with some species extending their range to southern North America, and others (such as in the genus *Hillia*) confined to South America.

Cinchonoideae is represented in tropical South America by the sister tribes Cinchoneae and Isertieae (1). A list of all currently recognized genera is shown in Table S1 and their species listed in Table S4. Figs. S1–S2 show detailed distribution maps for the main genera in Cinchoneae and Isertieae still not depicted by earlier studies (drawn by Lennart Andersson).

The tribe Isertieae is more or less continuously distributed from Guatemala to central Bolivia and the Amazon mouth (Fig. 1*A*). The greatest species diversity (3–4 species per grid square) is found in eastern Guiana and adjacent parts of the Amazon basin, between 68°W and 74°W, but there is not a very pronounced center of diversity (the highest number of species per grid square is only 5). The range west of 72°W is occupied by *I. haenkeana*, *I. rosea*, and species of the subgenus *Cassupa*, which do not occur east of 66°W. The high species richness in the central Amazon basin is mainly due to the overlap of the ranges of widespread species, whereas the species richness in eastern Guiana and easternmost Amazonia depends mainly on the presence of narrowly restricted endemics.

The distribution range of Cinchoneae (Fig. S1B) comprises most of the Neotropical region. Within this range, two distinct centers of diversity can be found, one along the Andes and another in central Amazonia. The Andean center is formed mainly by species of *Cinchona, Joosia*, and *Ladenbergia*, whereas the eastern Amazonian center comprises a large number of endemic *Remijia* species. The genera *Ciliosemina, Cinchonopsis, Maguireocharis,* and *Pimentelia* have too few species to clearly show any center of diversity. Cinchoneae is also present in the Brazilian Shield through two species of *Ladenbergia* (*L. chapadensis* and *L. cujabensis*). There is one species of *Remijia* (*R. ferruginea*) isolated on the southeastern fringe of the Brazilian Shield as well as scattered occurrences of *Ladenbergia hexandra* in the Atlantic coast of southeastern Brazil.

The distribution range of *Ladenbergia* (Fig. S1*C*) covers most of the range of Cinchoneae, but species diversity is highest in the Andes. Most species have relatively restricted ranges (31), with only a few species extending further than 10° in direction north to south and no species further than a few degrees west to east except for *L. oblongifolia*. The majority of Andean species occur at relatively low altitudes, mainly <1,500 m (31). The wide range of *Ladenbergia* outside the Andean area is entirely attributable to a small cluster of species, which appeared to be closely related with each other according to morphology (31). The only exceptions are two species that are not related to these: *Ladenbergia oblongifolia*, which has its main range in the Andes, and also has a disjunct occurrence in the Serra da Neblina, and *L. hexandra*, which is endemic to the mountains of southeastern Brazil.

The distribution range of *Cinchona* (Fig. S2A) is Andean, except for *C. pubescens*, which extends its range into southern Central America and the coastal mountains of Venezuela. Like *Ladenbergia*, *Cinchona* has a predominantly premontane to montane range. All records outside this range are probably introductions and, occasionally, subsequent naturalization (32).

The distribution range of *Ciliosemina* (Fig. S2*B*) falls largely within the Northern Andes, but reaches marginal parts of the Amazon basin.

Maguireocharis neblinae is a montane species growing at considerable altitude in the Serra da Neblina (Fig. S2A). In the absence of sequence data, and because of its unrevealing morphology, its relationships are obscure (1).

The monotypic genus *Pimentelia* (Fig. S2*B*) occurs on the eastern slopes of the Andes in southern Peru and northern Bolivia, whereas *Stilpnophyllum* (Fig. S2*D*) occurs in the Eastern Cordillera of northern Peru and southern Ecuador, and in the Central Cordillera of northern Colombia. *Pimentelia* could not be sequenced, but morphology indicates that it is the sister group of *Stilpnophyllum* (1). These 6 related species are all confined to the Andes. *Stilpnophyllum* appears to have a wide distribution gap between southern Ecuador and Colombia. This is likely to be real, because montane forests of eastern Ecuador are fairly well explored. Montane forests of northern and central Peru, however, are poorly explored, so the number of localities shown in this region is probably an underestimate.

The range of *Joosia* (Fig. S2C) extends over most of the tropical Andean chain, and it is almost fully confined to it. One species, *J. umbellifera*, extends throughout the range of the genus and is the only species occurring north of 1°N. Except for the single locality of *J. multiflora* in southernmost Peru, *J. umbellifera* is also the only species occurring south of 11°S. The largest species diversity of *Joosia* occurs in Ecuador and northern Peru, although species richness is not great anywhere. The slight drop seen in northernmost Peru is almost certainly an artifact caused by poor exploration of this region. Most species occur in the premontane to lower montane life zones, but 3 have only been found in the lowlands, and only 1 in the montane zone (30).

The distribution range of *Cinchonopsis amazonica* (Fig. S1*E*), the only species of the genus, comprises most of the western half of the Amazon Basin, and adjacent parts of the Orinoco drainage. Scattered information on herbarium labels suggests that it grows mainly in upland forest on white sand, which also agrees with the single field observation made by one of us (A.A.).

The range of *Remijia* (Fig. S1D) comprises mainly the central parts of the Amazon basin (between ≈55°W and 74°W) and adjacent parts of the Guiana Shield. In addition, there are 2 apparently disjunct areas, 1 in the western Amazonia and 1 in the Serra do Espinhaço in southeastern Brazil. Each of these exclaves has but a single endemic species (R. chelomaphylla and R. ferruginea, respectively). Remijia has a very pronounced center of diversity in central Amazonia, mainly in the Rio Negro, Rio Branco, and Rio Madeira basins, but extending into the Guiana Highlands. This seems to be related to an ecological preference of Remijia species for savanna and Amazonian caatinga, biomes that are particularly common in these regions (e.g., ref. 67). However, this is not the only explanation, because some central Amazonian species seem to occur in the rain forest. The phylogeny of Andersson and Antonelli (1) showed 2 sister groups, R. chelomaphylla-R. macrocnemia and R. pacimonica-R. *ulei*. The first 2 species occur only west of the diversity center, R. chelomaphylla in the western exclave and R. macrocnemia in a relatively small range west and northwest of the Trapecio Amazónico, where it is the only species. The other 2 species occur within the region of high species richness and are morphologically very similar. The apparent east-west sister group relationship may be an artifact, however, because many Remijia species have not yet been sampled.

Effects of Taxon Sampling on Biogeographic Reconstruction and Age Estimates. *Biogeographic reconstruction*. Barber and Bellwood (68) showed that biogeographical optimization methods such as dispersal-vicariance analysis could be highly sensitive to incomplete taxon sampling, especially if the missing taxa occupy a basal position in the phylogeny and/or occur in areas that are underrepresented in the analysis.

Although taxon sampling may be a problem in a large family such as Rubiaceae, spanning nearly all continents, we have tried to overcome this problem by using very encompassing areas outside the Neotropics, where most of the Rubioideae and Ixoroideae are found, as well as the first diverging clade in Cinchonoideae, the tribe Naucleeae. Although the small tribes Hamelieae, Hillieae, and Chiococceae are poorly represented in number of taxa, the distribution of the species included here are representative of their tribes, which are all predominantly distributed in Central America and the Caribbean region. A possible source of "noise" in the optimizations may be caused by Guettarda speciosa, an Australasian species that alone represents the tribe Guettardeae. Except for that species, the tribe has a Neotropical distribution (mainly in Central America and the West Indies; ref. 69). To estimate the effect of this bias in ancestral area optimizations, we carried out a new analysis coding the *Guettarda* lineage as restricted to Central America, the West Indies, and both. Except for the ancestor of the clade comprising Guettarda and Rogiera, the new coding did not affect any other ancestral area optimizations.

For tribes Cinchoneae and Isertieae, which are the focus of this article and upon which our main biogeographic conclusions depend, the key areas are the Northern and Central Andes (areas C and D in Fig. 1 *Inset*). In special, the presence of Central Andean species in a basal position within these tribes could potentially alter the conclusions drawn based on the empirical sequences analyzed.

To test the sensitivity of DIVA reconstructions to incomplete taxon sampling, we first estimated the geographic bias in our sample for each genus included in tribes Isertieae and Cinchoneae. We compiled distribution data for all species, genera, and tribes (Table S4) and compared the representation of each area in the complete inventory against that of our molecular sample. Fig. S6 (left column) shows that the geographical distribution of species in our sample is fairly representative of the actual distribution of the specie. For tribes, area C is slightly underrepresented in Cinchoneae and area D in Isertieae. In contrast, area F (Amazonia) is fairly represented in both tribes. For genera, the geographical bias is largest in Joosia, Stilpnophyllum, and Cinchona, underrepresented in area D, and Remijia, which is missing all 8 species endemic to the Guiana Shield and the Brazilian Planalto (areas G and H). Some of these genera are also poorly represented in botanical collections: Most species of Stilpnophyllum, and many of Joosia, are known from the type specimen only (Fig. S6, right column). The paucity of specimens in botanical collections directly reflects the difficulty to find these species in the field, as evidenced by numerous botanical expeditions undertaken from our research institute in the last decades, as well as by the experience of many colleagues working locally. Although this paucity may sometimes be attributed to poor botanical exploration of some inaccessible regions, most species seem to be naturally rare. In addition, many species are only minimally distinct, which calls into question the species concept applied within each genus and suggests that some of the rarest species could in fact be variations of a more common species, or hybrids (e.g., ref. 32).

Based on this survey (Fig. S6), we then followed a similar approach to Barber and Bellwood (68). We performed a series of heuristic analyses in which a hypothetical missing taxon was added, one at a time, at certain points in the phylogeny. Instead of random addition, we kept the basic backbone of the phylogeny (all major nodes for Cinchonoideae genera) as we did for the temporal simulations (see below). The phylogenetic position of the missing taxa within each genus was chosen following morphological cladograms whenever available (e.g., for genera Joosia, Ladenbergia). When no morphological phylogeny was available (e.g., genus *Remijia*) or did not include any additional species (genus Isertia), the missing taxon was added, alternatively, to the most basal or the most distal node within the genus. The distribution of the missing (hypothetical) taxon was determined following the geographical bias estimated for each genus (see Fig. S6).

Fig. S7 shows the results of these simulations. Somewhat surprisingly, DIVA ancestral area reconstructions proved to be very stable to the addition of missing taxa. Adding hypothetical taxa from underrepresented areas to the phylogeny did not alter the original ancestral area reconstructions (with no taxon addition) for the key nodes in the Isertieae/Cinchoneae phylogeny. The only exception was node 51: The addition of a taxon from areas G or H to the base of the *Remijia* clade (J6, Fig. S7A) resulted in the ancestor of Remijia, Ciliosemina, and Ladenbergia being originally distributed in areas G and/or H (J6, Fig. S7B). However, this has no consequences for our biogeographic scenario, because ancestral area reconstructions for the rest of the nodes remained unaltered (Fig. S7B). That is, *Remijia* would have reached the Guiana Shield and Southeastern South America (areas G and H, respectively) earlier than we postulate in our scenario, but biogeographical inferences concerning the Western Andean Portal and Lakes Pebas would not change. Moreover, these changes in the ancestral distribution of Remijia would only occur if the missing taxon really is sister to the rest of the genus: A distal position among the remaining ≈ 40 species would instead produce the same outcome as in our original reconstruction (J7, Fig. S7).

The only scenario that could significantly alter our hypotheses is if the 2 species from area D (Central Andes) in genus *Joosia* (*J. multiflora* and *J. dichotoma*) and the 2 species from area D in genus *Isertia* (*I. krausei* and *I. reticulata*) came to occupy the 2 most basal positions within the phylogeny of each of these genera (simulations not shown). This would result in area D being inferred as the ancestral area of *Joosia* and of *Isertia*, and area CD as the ancestral area of tribe Cinchoneae. In other words, tribe Cinchoneae would have been originally present in the North and Central Andes instead of dispersing to the Central

Andes from the north after the uplift of the WAP as suggested by our original reconstruction. However, the probability that the 2 Central Andean species of Joosia would occupy the 2 most basal positions in the phylogeny of the genus is low. In Andersson's (30) morphological phylogeny, these 2 species are not phylogenetically close: Joosia multiflora occupies the most basal position in the cladogram, whereas J. dichotoma is "firmly nested within the genus" (p. 28 in ref. 30). Both Joosia multiflora and J. dichotoma are known from the type collection only, and we have been unable to find them in the field for sequencing. In the case of Isertia, Andersson's (70) morphological phylogeny of Isertieae did not include any Central Andean species. In the only published revision of the genus, Boom (29) included Isertia krausei and I. reticulata within section Cassupa, which is separated from section Isertia on the basis of the presence of fruits with fleshy endocarp. However, our molecular phylogeny does not support Boom's infrageneric classification: Isertia pittieri and I. laevis from section Cassupa appear nested within a clade composed exclusively by species from section Isertia. Moreover, Isertia krausei is known from the type collection only, whereas I. reticulata is "a seldom collected and imperfectly known species" that may occur also in Colombia (29).

Out of the eight species of *Cinchona* endemic to the Central Andean region, only one (*C. calisaya*) was possible to include in our analysis. Our molecular phylogeny places *Cinchona calisaya* as the sister group to the rest of the genus, and DIVA infers the Northern and Central Andes (area CD) as the geographic origin of *Cinchona* (node 46). Even if the other 7 Central Andean species in the genus come out as a basal grade in the phylogeny—an unlikely inference according to Andersson's (32) morphological cladogram, which places all Central Andean species in different clades—and D is inferred as the ancestral area of *Cinchona* (with later dispersal to C), this would not affect the ancestral reconstructions for the backbone nodes in Cinchoneae (nodes 44, 45, and 51), which still is inferred as having originated in the Northern Andes, area C.

All things considered, we believe that the possibility of missing taxa from area D to have a significant influence in our scenario is low and that the DIVA results presented here are stable to incomplete taxon sampling.

Age estimates. Linder et al. (71) demonstrated that limited taxon sampling may influence age estimates, generally resulting in younger ages for nodes than those obtained with a denser sample. To investigate these effects in the Rubiaceae, we followed a similar approach to Linder et al. (71): first, we identified 14 "core" nodes across the phylogeny corresponding to all genera in tribes Cinchoneae and Isertieae, main clades in Cinchonoideae including the Cephalanthus calibration point, subfamilies Rubioideae and Ixoroideae, and outgroup (nodes 2, 7, 13, 22, 25, 27, 32, 33, 41, 43, 46, 52, 55, and 56 in Fig. S4); second, we generated reduced matrices comprising 30%, 40%, 60%, and 80% of the original dataset by randomly deleting terminal taxa, while still keeping at least 2 species from each core node in order to keep the backbone of relationships in the phylogeny (the basic structure of the phylogeny); third, we perfomed Bayesian phylogenetic analyses and computed meanlength consensus phylograms for each of the reduced matrices, using the same settings as for the original dataset; and last, we estimated divergence times for each of those phylograms using the same methodology as previouly described.

The results of these analyses are summarized in Fig. S8. Although there is a slight tendency to underestimation of ages when taxon sampling decreases (Fig. S8A), for all core nodes in the phylogeny the ages obtained for the reduced matrices fall within the 95% confidence intervals estimated from complete dataset (Fig. S8B). Similarly, even a 2-fold increase in the number of taxa (trend lines in Fig. S8A) is not expected to result in node ages outside the confidence intervals of particular nodes.

The only exceptions are nodes 20, 31, and 33 in the dataset comprising only 30% of the original taxa: their ages are underestimated by 1.3, 0.7, and 0.53 Ma from the original confidence intervals, respectively. It seems that 30% represents a threshold level for those nodes, after which ages stabilize.

These results indicate that the addition of more taxa in Rubiaceae is not expected to significantly influence the empirical results presented, nor their temporal interpretation. Moreover, they support the finding by Linder et al. (71) that Penalized Likelihood is largely insensitive to taxon sampling, as compared with other dating methods such as nonparametric rate smoothing and Bayesian dating.

Generality of the Biogeographic Patterns Found. Western Andean Portal (WAP). Although it has never been properly analyzed, it is particularly noteworthy that several studies have demonstrated a biogeographic disjunction between the Northern and Central Andes, roughly corresponding to the latitude used in this study for separating these 2 operational areas (\approx 5°S). However, the dispersal barrier between these regions has received many different denominations in the literature, which seem to refer to approximately the same area: Western Portal, Andean Gate, Marañón Portal, Guayaquil Gap, Huancabamba Depression, Huancabamba Deflexion, Northern Peruvian Low, and Pirua Divide.

In plants, the pattern has been demonstrated in many families, such as Campanulaceae (72), Calceolariaceae (73), Tropaeolaceae (74), Loasaceae, Passifloraceae, Grossulariaceae (54, 75), and Alstroemeriaceae (76). In animals, Vuilleumier (77) showed such a disjunction for populations of the bird superspecies Asthenes flammulata (Furnariidae), Nores (78) defined the region north of this barrier (northwestern Andes) as a general area of bird endemism, and Cortés-Ortiz (79) defined this barrier as the southern limit of the monkey Alouatta palliata. Moreover, this region (the Río Marañon in Peru) has long been recognized as the turnover point between the Northern and Central Andean regions of bird endemism (80), further emphasizing the role of the WAP as a major biogeographic barrier. We have not attempted to compile an exhaustive account of taxa whose distribution is affected by the barrier, but anticipate that many other groups may show similar patterns.

Although our results demonstrate how and when the WAP influenced the distribution of a modern group of plants, they cannot provide evidence to its exact nature, geographic extension, and duration. The few paleontological studies in the area strongly support the existence a marine incursion that reached the western part of present-day Amazonia at least during the Late Eocene (81, 82) and Middle Miocene (53), but how long marine settings dominated the region before and after these epochs is a matter of debate. Moreover, contrary to several earlier studies (e.g., ref. 83), new marine geophysical data suggest that global sea levels may have been relatively stable from the Early Eocene onwards (ref. 84 and references therein). Therefore it seems plausible that a marine incursion dominated the WAP during most of the Eocene–Middle Miocene, ending with the uplift of the Eastern Cordillera.

The geographic extent of the WAP has also been under debate. From the Pacific coast, marine settings have been demonstrated to reach as far east as $\approx 75^{\circ}$ W in the Late Eocene (82) and $\approx 79^{\circ}$ W in the Middle Miocene (53). An aquatic connection between the Pacific and Amazonia has long been hypothesized (e.g., 53, 85, 86), but no conclusive evidence seems yet to support such a long connection (F. P. Wesselingh and C. Hoorn, personal communication). However, there is some evidence that intermittent connections existed between Lake Pebas in western Amazonia and the Pacific through the Ecuadorean Andes during the Middle Miocene (87). Even if the WAP would not have been filled by sea water during its entire existence and even if marine incursions did not reach as far east as Amazonia, it could still have acted as a dispersal barrier to montane organisms: either by being lowland corridors or by imposing ecological constraints (e.g., very dry climates were present in southern Ecuadorean Andes in the Miocene; ref. 87). We make here a plea for more studies that address these important issues.

Lake Pebas. The biogeographic scenario proposed here implies that all plant species adapted to dry land conditions (terra firme) in present-day western Amazonia have gained their current distribution after Lake Pebas drained (i.e., in the last ≈ 11 Ma; ref. 88). Similarly, in situ speciation in those terrestrial groups could not have taken place before that event.

These results may seem controversial given the fact that western Amazonia is characterized today by an outstanding number of endemic plant species, of which many belong to long-lived woody families not expected to speciate fast. In a recent study using large molecular chronograms from 5 angio-

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sperm families, Smith and Donoghue (89) showed that there is a strong correlation between growth mode and speciation rates: Herbs generally diversify much more rapidly than shrubs and trees. However, the few molecular dating studies that have dealt with a representative number of Amazonian trees have presented very contrasting results. Two of the largest genera of Neotropical trees have been inferred to have diversified recently: Inga (Fabaceae, 300 spp), which is estimated to have started its diversification either 9.8 Ma (based on cpDNA) or 1.6 Ma (based on nrDNA; ref. 90); and Guatteria (Annonaceae, 265 spp), where the crown age for all Neotropical taxa (including 2 large Amazonian subclades) is estimated at 7.4 \pm 1.4 Ma (91). Among animals, a recent study (92) suggested that the diversification of Amazonian poison frogs (Dendrobatidae) also occurred recently, with radiations in the last 10 Ma. These results fit well into the paleogeographic scenario for Amazonia proposed here and suggest that Amazonian organisms may have evolved at an unusually fast rate after the end of Lake Pebas.

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Fig. S1. Distribution patterns in Neotropial Rubiaceae: tribes Isertieae and Cinchoneae. (*A*) Geographical range and species richness of Rubiaceae tribe Isertieae s. str. (based on maps in ref. 29, specimen citations in ref. 38, and localities cited by ref. 36). (*B*) Geographical range and species richness of Rubiaceae tribe Cinchoneae. (*C*) Geographical range and species richness of the genus *Ladenbergia* (based on data from ref. 31). (*D*) Geographical range and species richness of the genus *Ladenbergia* (based on data from ref. 31). (*D*) Geographical range and species richness of the genus *Ladenbergia* (based on data from ref. 31). (*D*) Geographical range and species richness of the genus *Remijia* (based on Lennart Andersson's unpublished notes). (*E*) Recorded localities of *Cinchonopsis amazonica* (based on Lennart Andersson's unpublished notes). (*F*) Branch tip with inflorescences of *Ciliosemina pedunculata* and detail of a dissected flower (branch based on herbarium specimen Lleras et al. in Prance 16958, S and flower detail on St. John and Arcila 20627, GB. [Flower image in *F* reproduced with permission from ref. 1 (Copyright 2005, Olof Helje). Branch image in *F* reproduced with permission from ref. 35 (Copyright 1994, University of Gothenburg).]



Fig. S2. Distribution patterns in tribe Cinchoneae. (A) Geographical range and species richness of the genus Cinchona (based on data from ref. 32), and localities of Maguireocharis neblinae recorded by ref. 33. (B) Recorded localities of Ciliosemina and Pimentelia (based on Lennart Andersson's unpublished notes). (C) Recorded occurrences of the genus Joosia (based on data from ref. 30). (D) Geographical range and species richness of the genus Stilpnophyllum (based on specimen citations in ref. 35, and Lennart Andersson's unpublished notes).



Fig. S3. Summary of the Bayesian phylogenetic analysis (50% majority-rule consensus tree with compatible groups added). A thick-lined branch indicates that the branch was also present in the majority-rule consensus tree of the parsimony-jackknife analysis. Numbers above branches indicate the posterior probability of the clade. Numbers below branches show jackknife support values, whenever applicable. Brackets identify subfamilies and tribes (CHI, Chiococceae s. lat.; CIN, Cinchoneae s. str.; GUE, Guettardeae; HAM, Hamelieae; HIL, Hillieae; ISE, Isertieae s. str.; NAU, Naucleeae s. lat.; RON, Rondeletieae).



Fig. 54. Molecular dating analysis of the Rubiaceae. (A) Chronogram showing 95% confidence intervals of node ages (indicated by bars; n = 1,000). Tree topology is the 50% majority-rule Bayesian consensus tree with all compatible groups added. Node numbers refer to Table S2. Red stars indicate calibration points based on molecular and fossil evidence [Node 1: crown group age of Gentianales as determined in the large-scale analysis of Bremer et al. (16); node 22: fruits of *Cephalanthus kireevskianus* from the Late Eccene of Germany reported by Mai and Walther (18)]. (*Inset*) SEM pictures depicting fossil (I–II) and extant (III–IV) fruits of *Cephalanthus*. I, III: Apical part of mericarp with germination valve detached showing characteristic strophiole (ST) and seed (S). II, IV: seed. [Reproduced with permission from ref. 22 (Copyright 1985, The Royal Danish Academy of Sciences and Letters).] (*B*) Number of recovered *Cephalanthus* fruit fossils per geologic epoch. (C) Locations of all 19 records of *Cephalanthus* fossilized fruits.



Fig. S5. Altitudinal optimization on ancestral nodes. Pie charts represent relative frequencies of optimizations, based on 5,000 Bayesian trees from a stationary sample. Optimizations performed in the software Mesquite under the Maximum Parsimony (Fitch) criterion, and plotted on the 50% majority-rule consensus tree (compatible groups added) of Fig. 1. Only trees where the relevant node is present were considered in calculating the relative frequencies.



Fig. S6. Estimate of geographic bias on taxon sampling for tribes Isertieae and Cinchoneae. (*Left*) Geographic distribution of all species in the main genera and tribes of subfamily Cinchonoideae, compared with the distribution of species sampled in our phylogeny. The number of species for each chart is given immediately after it. Area codings as in Fig. 1. (*Right*) Number of collected specimens for each species in a genus/tribe. Note that these are minimal numbers, because only specimens verified in taxonomic revisions have been accounted for (additional specimens may have been collected recently and deposited at various herbaria).



Fig. 57. Effect of missing taxa on ancestral area reconstructions. (A) Phylogeny of tribes Isertieae and Cinchoneae, as estimated under the Bayesian analysis (50% majority-rule consensus tree with all compatible groups added). J_0-J_9 indicate branches where missing taxa might be expected to attach, as suggested by morphological evidence (e.g., morphology-based cladograms, accounts in taxonomic revisions); when no morphological evidence was available, the missing taxa neach to the most basal or most distal node of the genus. Node numbers as in Fig. 54. (*B*) Results from DIVA simulations after the addition of missing taxa at each intersection node (J_0-J_9) outlined above. The distribution of the missing (hypothetical) taxon added is given after each intersection node, according to the area codings in Fig. 1 (*Inset*) and based on the geographic bias estimated for each genus (Fig. S6). Ancestral area distributions differing from the original reconstruction (no taxon addition) are marked in yellow.



		Perc	entage	of		Ag	Fit of			
Node		taxa inc luded					(all ta xa included)			
No.	20	40	(0	90	100 Mean	Maan	95% Confide	ence Interval	D ²	
	30	40	00	00		Lower bound	Upper bound	ĸ		
20	46.5	48.9	49.0	50.3	51.3	51.3	47.8	54.6	0.92	
31	39.0	44.1	40.2	41.9	44.5	45.2	39.7	50.0	0.29	
32	20.1	22.6	22.7	24.3	25.1	25.0	19.9	31.4	0.93	
33	9.25	10.7	12.6	12.8	13.7	13.1	9.78	18.3	0.98	
40	24.3	26.7	25.3	22.3	28.0	28.6	22.9	35.1	0.02	
42	15.8	17.8	16.7	17.9	18.6	19.0	14.0	24.8	0.62	
44	14.3	15.7	14.8	15.8	16.8	17.2	14.0	24.8	0.60	
45	13.5	14.7	14.1	14.9	15.7	16.2	12.5	22.0	0.68	
46	6.45	6.95	6.92	7.16	9.08	9.00	6.05	12.9	0.57	
52	3.63	4.09	4.30	4.59	4.94	5.19	2.96	7.84	0.96	
55	12.7	13.8	13.1	13.9	13.8	14.4	10.5	19.7	0.47	
56	10.9	11.6	11.1	11.4	10.9	11.8	8.51	16.3	0.00	

Fig. S8. Effect of missing taxa on divergence time estimations. (*A*) Ages of key nodes estimated from randomly reduced datasets (including 30%, 40%, 60%, and 80% of the original number of species) and their expected development with the addition of missing taxa (projection curve calculated following a second-degree logarithmic function). Node numbers as in Fig. S4. (*B*) Ground data for the diagram above, compared with age node statistics calculated from the original dataset (all taxa included). Boxes marked in yellow fall outside the 95% confidence interval calculated from 1,000 Bayesian trees inferred from the whole dataset. The last column shows the statistical fit of the trend line to the data points in the diagram.

Table S1. Genera and number of species currently recognized in tribes Cinchoneae and Isertieae

Tribe	Genus	No. of species
Cinchoneae	Ciliosemina	2
	Cinchona	23
	Cinchonopsis	1
	Joosia	11
	Ladenbergia	34
	Maguireocharis	1
	Pimentelia	1
	Remijia	41
	Stilpnophyllum	4
Isertieae	Isertia	14
	Kerianthera	1

A complete list of the species in each genus is given in Table S4.

Table S2. Crown group ages of all nodes in the phylogeny

PNAS PNAS

Consensus phylogram

Estimates based on 1000 Bayesian trees

			95% credibility interval		
Node	Age	Median Age	Lower	Upper	
1	78.0	78.0	_	-	
2	68.6	68.8	63.8	74.5	
3	64.9	65.4	59.2	71.5	
4	43.8	44.0	37.6	51.1	
5	65.8	66.1	63.0	68.8	
6	64.8	65.1	61.5	67.8	
7	46.8	47.9	43.4	52.5	
8	44 9	46.0	41 3	50.6	
9	43.6	44.2	39.9	49 1	
10	30.8	31.8	27.4	36.7	
11	20.6	21.5	17.7	25.8	
17	20.0	21.5 62.7	F0 7	25.0	
12	02.7	02.7	59.7	00.Z	
13	48.1	48.1	43.8	52.4	
14	46.5	46.9	42.6	51.0	
15	40.4	41.0	35.7	45.5	
16	46.1	45.9	41.5	50.5	
17	43.7	43.7	38.9	48.2	
18	30.3	30.3	26.2	34.8	
19	15.7	15.8	12.5	19.5	
20	51.3	51.3	47.8	54.6	
21	36.4	36.3	34.8	38.1	
22	33.9	33.9	-	-	
23	28.2	28.3	23.1	32.5	
24	49.2	49.4	44.8	53.3	
25	35.2	35.3	29.9	40.8	
26	48.5	49.2	44.9	53.1	
27	46.4	47 3	42.5	52.1	
28	20.7	30.0	23.5	37.6	
20	19.9	19.2	16.1	22.2	
20	15.0	15.2	12.0	19.0	
30	15.5	15.0	12.0	10.9	
21	44.5	45.2	59.7 10.0	50.0	
32	25.1	25.0	19.9	31.4	
33	13.7	13.1	9.78	18.3	
34	6.34	5.76	2.90	10.2	
35	10.9	10.8	7.47	15.0	
36	8.56	8.80	4.71	14.0	
37	1.85	1.91	0.35	4.12	
38	10.0	10.3	6.52	14.0	
39	1.87	1.94	0.57	3.76	
40	28.0	28.6	22.9	35.1	
41	13.4	13.7	8.93	18.3	
42	18.6	19.0	14.0	24.8	
43	5.97	6.05	3.73	9.83	
44	16.8	17.2	14.0	24.8	
45	15.7	16.2	12.5	22.0	
46	9.08	9.00	6.05	12.9	
47	6.74	6.67	4.32	9.71	
48	6.02	5 98	3.88	8 60	
49	4 50	4 51	2 77	6.80	
50	3.40	3 42	1 87	5 39	
51	5. 4 0 15 1	5. 4 2 15 Ω	1.07	5.55 01 E	
57	13.1	E 10	2 04	21.2	
52	4.94	2.22	2.90	7.84	
22	3.08	3.27	1.05	5.43	
54	1.49	1.54	0.56	2.80	
55	13.8	14.4	10.5	19.7	
56	10.9	11.8	8.51	16.3	
57	8.46	9.07	6.31	13.0	
58	9.81	10.6	7.83	15.0	
59	8.66	9.42	6.69	13.1	
60	6.16	6.68	4.62	9.55	

Statistics based on 2 calculations: Consensus phylogram refers to ages calculated from mean branch lengths of trees sampled by a Bayesian MCMC analysis (n = 6,000); Estimates based on 1,000 Bayesian trees refer to median age estimates and 95% credibility intervals calculated by independently dating trees from a random subsample of the Bayesian MCMC analysis (n = 1,000). Node numbers refer to Fig. S4.

Table S3. Sequences used in the phylogenetic and biogeographic analyses, together with source and GenBank accession numbers

Taxon				GenBan	k accession	number	
Outgroup	Origin	Voucher	ITS	matK	rbcL	rps16	<i>trn</i> L–F
Gelsemium sempervirens (L.) J. StHil.	GenBank	_	_	Z70195	L14397	AF004092	AF159696
Geniostoma rupestre J.R. Foster & G. Foster	GenBank	—	—	Z70194	Z68828	—	_
Strychnos nux-vomica L.	GenBank	—	_	Z70193	L14410	AF004094	AF102484
Thevetia peruviana K. Schum.	GenBank	—	—	Z70188	X91773	—	_
Ingroup							
Cephalanthus occidentalis L.	GenBank	—	_	AY538377	X83629	AF004033	AF152692
Chiococca alba (L.) Hitchc.	GenBank	_		AY538378	L14394	AF004034	AF102400
Cinchona calisaya Wedd.	GenBank	—	AY538352	AY538379	AY538478	AF242927	AY538447
C. macrocalyx Pav. ex DC.	GenBank	—		A1228280	A1538479	AF538425	A 1 538448
C. officinalis I	GenBank	—	A 1 336333	AVE20201	AVE20400	A 1 336420	A 1 330449
C. nitavensis Wedd	GenBank	_	AV538355	AT 336361	AY538480	A1556427	A1556450
C nitavensis Wedd	GenBank	_		A1550502	A1330-01	ΔF242928	ΔF152684
C pubescens Vahl	GenBank		AY538356	_	_		AY538451
C. pubescens Vahl	GenBank	_	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Z70197	X83630	AF004035	
Cinchonopsis amazonica (Standl.) L. Andersson	GenBank	_	AY538357	AY538383	AY538482	AY538428	AY538452
Condaminea corvmbosa (Ruiz & Pay.) DC.	GenBank	_	_	AY538384	Y18713	AF004039	AF102406
Cosmibuena grandiflora (Ruiz & Pav.) Rusby	GenBank	_	_	AY538385	AY538483	AF242929	AF152686
Cubanola domingensis (Britton) Aiello	GenBank	_	_	AY538386	X83632	AF004044	AF152701
Exostema lineatum (Vahl) Roem. & Schult.	GenBank	_	_	AY538387	AY538484	AF242944	AF152698
Genipa americana L.	GenBank	_	_	AY538388	Z68839	AF200997	AF201045
Guettarda speciosa L.	GenBank	_	_	AY538389	AY538485	AF246924	AF152725
Hallea rubrostipulata (K. Schum.) JF. Leroy	GenBank	—	_	AY538390	AY538486	AF538429	AY538453
Hamelia papillosa Urb.	GenBank	—	_	AY538391	AY538487	AF004053	AF102439
Hymenodictyon floribundum Robinson	GenBank	—	—	AY538392	AY538488	AF004058	AY538454
Isertia coccinea (Aubl.) J.F. Gmel.	GenBank	—	AY538358	AY538393	AY538489	AY538430	AY538455
<i>I. laevis</i> (Triana) Boom	GenBank	—	AY538359	AY538394	AY538490	AY538431	AY538456
I. haenkeana DC.	Colombia	Alzate et al. 203 (GB),	DQ448607*	_	DQ448595*	DQ448601*	-
<i>I. hypoleuca</i> Benth.	Colombia	Andersson et al. 2173 (GB)	DQ448608*	—	DQ448596*	DQ448602*	_
I. parviflora Vahl	French Guiana	Andersson et al. 1969 (GB),	DQ448609*	—	DQ448597*	DQ448603*	_
<i>I. pittieri</i> (Standl.) Standl.	Colombia	Andersson et al. 2099 (GB)	DQ448610*	—	DQ448598*	DQ448604*	-
<i>I. rosea</i> Spruce ex K. Schum.	Ecuador	Hekker & Hekking 10.147 (GB)	DQ448611*	_	DQ448599*	DQ448605*	_
I. spiciformis DC.	French Guiana	Andersson et al. 1905 (GB	DQ448612*		DQ448600*	DQ448606*	—
Joosia aequatoria Steyerm.	GenBank	—	AY538360	AY538395	AY538491	AY538432	AY538457
J. umbellifera H. Karst.	GenBank	—	AY538361	AY538396	AY538492	AY538433	AY538458
Kerianthera preciara Kirkbr.	GenBank	—	AY538362	AY538397	AY538493	AF242970	AY538459
Ladenbergia amazonensis Ducke	GenBank	—	AY538363	AY538398	AY538494	A 1 538434	AY538460
L. Carua (Wedd.) Standt.	GenBank	—	A 1 336304	A1556599	A1556495	A 1 336433	A 1 3 3 6 4 0 1
L. macrocarpa (Vani) Notzsch	GenBank		AT538365	AV538400	AT538490	AF242971	AF132003
L. pavonii (Lamb.) Standl	GenBank		AY538367	AV538402	768801	AV538437	AV538463
/ sp (prob pova)	GenBank	_	ΔΥ538368	AY538403	AY538498	ΔΥ538438	ΔΥ538464
Lasianthus batangensis K. Schum	GenBank	_		AY538404	ΔΥ538499	ΔΥ538439	ΔΥ538465
Luculia grandiflora Ghose	GenBank	_	_	Z701999	X83648	AF242974	AF102453
Mitriostigma axillare Hochst.	GenBank	_	_	AY538405	X83650	AF201006	AF201054
Mussaenda raiateensis J.W. Moore	GenBank	_	_	AY538406	AY538500	AF242983	AY538466
Nauclea orientalis L.	GenBank	_	_	AY538407	AY538501	AY538440	AY538467
Ophiorrhiza mungos L.	GenBank	_	_	AY538408	X83656	AF004064	AF152610
Paederia foetida L.	GenBank	_	_	AY538409	AF332373	AF004065	AF152619
Pauridiantha sp.	GenBank	—	_	AY538410	AY538502	AF004068	AF102467
Phyllis nobla L.	GenBank	—	—	AY538411	Z68814	AF003613	AY538468
Posoqueria latifolia (Rudge) Roem. & Schult.	GenBank	—	—	AY538412	Z68850	AF242998	AF152680
Psychotria kirkii Hiern	GenBank	_	_	AY538413	X83663	AF410728	AY538469
Remijia chelomaphylla G.A. Sullivan	GenBank	—	AY538369	AY538414	AY538503	AY538441	AY538470
R. macrocnemia (Mart.) Wedd.	GenBank	—	AY538371	AY538415	AY538504	AY538442	AY538471
R. pacimonica Standl.	GenBank	—	AY538372	AY538416	AY538505	AY538443	AY538472
R. pedunculata (H. Karst.) Flueck.	GenBank	—	AY538373	AY538417	AY538506	AY538444	AY538473
R. ulei K. Krause	GenBank	—	AY538374	AY538418	AY538507	AY538445	AY538474
Rovaeanthus suffrutescens (Brandeg.) Borhidi	GenBank	—	_	AY538419	X83665	AF243003	AF152738
Sabicea aspera Aubl.	GenBank	—	_	AY538420	AY538508	AF004079	AY538475
Sipanea bitlora (L.t.) Cham. & Schltdl.	GenBank	—		AY538421	AY538509	AF004085	AF152675
Stilpnopnyllum granditollum L. Andersson	GenBank	_	AY538375	AY538422	AY538510	AY538446	AY538476
S. Gengaardii L. Andersson	GenBank	—	AY5383/6	AY538423	AY538511	AF243026	AY5384/7
vangueria madagascariensis J.F. Gmel.	GenBank	_	_	A1538424	79792	AF243033	Ar 152654

Data on origin and voucher are given only for sequences added in this study (indicated by an asterisk).

Table S4	. Checklist of	species and	genera in	the siste	r tribes	Cinchoneae	and Isertiea	e, their distribu	ition, and	number of	of collections
verified	in recent taxo	nomic revisi	ons (see <mark>S</mark>	I Text for	refere	nces)					

Species count in tribes	Species count in genera	Genus	Species	Distribution	No. of collections
1	1	Ciliosemina	pedunculata	CF	>6
2	2	Ciliosemina	purdieana	F	2
3	1	Cinchona	antioquiae	С	>6
4	2	Cinchona	asperifolia	D	2
5	3	Cinchona	barbacoensis	CE	>6
5	4	Cinchona	calisaya	D	>6
/ o	5	Cinchona	fruticoso	C	ס< ר
0 0	о 7	Cinchona	alandulifera		2 5 or 6
10	8	Cinchona	hirsuta	D	2
11	9	Cinchona	krauseana	D	5 or 6
12	10	Cinchona	lancifolia	C	>6
13	11	Cinchona	lucumifolia	С	>6
14	12	Cinchona	macrocalyx	CD	>6
15	13	Cinchona	micrantha	CD	>6
16	14	Cinchona	mutisii	с	>6
17	15	Cinchona	nitida	D	5
18	16	Cinchona	officinalis	С	>6
19	17	Cinchona	parabolica	CD	6
20	18	Cinchona	pitayensis	C	>6
21	19	Cinchona	pubescens	ACD	>6
22	20	Cinchona	pyrifolia	D	3
23	21	Cinchona	rugosa	C	>6
24	22	Cinchona	scrobiculata	C	1 (type only)
25	23	Cinchona	villosa	C F	2
20	1	Cinchonopsis	coccines	F	>0
28	1	Isertia	haenkeana	ABCEE	>6
29	2	Isertia	hypoleuca	ACEG	>6
30	4	Isertia	krausei	D	1 (type only)
31	5	Isertia	laevis	ACDF	>6
32	6	Isertia	longifolia	F	>6
33	7	Isertia	parviflora	FG	>6
34	8	Isertia	, pittieri	EC	>6
35	9	Isertia	reticulata	D	5
36	10	Isertia	rosea	F	>6
37	11	Isertia	scorpioides	А	5
38	12	Isertia	spiciformis	FG	>6
39	13	Isertia	verrucosa	F	5
40	14	Isertia	wilhelminensis	G	3
41	1	Joosia	aequatoria	С	2
42	2	Joosia	dichotoma	D	>6
43	3	Joosia	dielsiana	C	>6
44	4	Joosia	longisepala	C	2
45	5	Joosia	macrocalyx	C	1 (type only)
46	6	Joosia	multiflora	D	1 (type only)
4/	7	Joosia	obtusa	C	1 (type only)
48	8	Joosia	oligantha		i (type only)
49 50	10	Joosia	standlevana	CD C	20
50	11	Joosia	umbellifera	ACDE	5
57	1	Kerianthera	preclara	F	4
52	1	Ladenbergia	acutifolia	L L	4
54	2	Ladenbergia	amazonensis	DFG	>6
55	3	Ladenbergia	brenesii	A	>6
56	4	Ladenbergia	bullata	D	2
57	5	Ladenbergia	buntingii	С	3
58	6	Ladenbergia	carua	D	>6
59	7	Ladenbergia	chapadensis	Н	1 (type only)
60	8	Ladenbergia	cujabensis	н	>6
61	9	Ladenbergia	discolor	D	6
62	10	Ladenbergia	dwyeri	А	>6
63	11	Ladenbergia	epiphytica	С	1 (type only)
64	12	Ladenbergia	ferruginea	D	1 (type only)
65	13	Ladenbergia	graciliflora	DFH	>6
66	14	Ladenbergia	heterophylla	AC	>6
67	15	Ladenbergia	hexandra	Н	>6
68	16	Ladenbergia	klugii	D	1 (type only)
69	17	Ladenbergia	lambertiana	FG	>6
70	18	Ladenbergia	lauritolia	A	3

Species count in tribes	Species count in genera	Genus	Species	Distribution	No. of collections
71	19	Ladenbergia	lehmanniana	С	3
72	20	Ladenbergia	macrocarpa	С	>6
73	21	Ladenbergia	magdalenae	С	>6
74	22	Ladenbergia	moritziana	С	>6
75	23	Ladenbergia	muzonensis	CEF	>6
76	24	Ladenbergia	nubigena	А	5
77	25	Ladenbergia	oblongifolia	CDEFG	>6
78	26	Ladenbergia	obovata	C	3
79	27	Ladenbergia	paraensis	F	1 (type only)
80	28	Ladenbergia	pauciflora	C	1 (type only)
81	29	Ladenbergia	pavonii	CE CE	>0
02 02	21	Ladenbergia	rivereana	C	>0
84	37	Ladenbergia	rubiginosa	C CF	20
85	33	Ladenbergia	stenocarna	C C	2
86	34	Ladenbergia	undata	C	>6
87	1	Maguireocharis	neblinae	G	2
88	1	Pimentelia	glomerata	CD	>6
89	1	Remijia	amazonica	F	not revised
90	2	Remijia	amphithrix	F	not revised
91	3	Remijia	aracamuniensis	G	not revised
92	4	Remijia	argentea	FG	not revised
93	5	Remijia	asperula	F	not revised
94	6	Remijia	berryi	F	not revised
95	7	Remijia	bracteata	F	not revised
96	8	Remijia	chelomaphylla	CD	not revised
97	9	Remijia	delascioi	F or G	not revised
98	10	Remijia	densiflora	G	not revised
99	11	Remijia	duckei	F	not revised
100	12	Remijia	ferruginea	H	not revised
101	13	Remijia	tirmuia	FG	not revised
102	14	Remijia	globosa	G	not revised
103	15	Remijia	giomerala	r c	not revised
105	18	Remijia	involucrata	F	not revised
106	18	Remijia	leiocalyx	F	not revised
107	19	Remijia	longifolia	F	not revised
108	20	Remiiia	macrocnemia	F	not revised
109	21	Remijia	macrophylla	С	not revised
110	22	Remijia	maguirei	G	not revised
111	23	Remijia	marahuacensis	F	not revised
112	24	Remijia	megistocaula	F	not revised
113	25	Remijia	morilloi	F	not revised
114	26	Remijia	pacimonica	F	not revised
115	27	Remijia	paniculata	Н	not revised
116	28	Remijia	peruviana	F	not revised
117	29	Remijia	physophora	F	not revised
118	30	Remijia	pilosinervula	G	not revised
119	31	Remijia	prismatostylis	C	not revised
120	32	Remijia	reducta	F	not revised
121	33	Remijia	roraimae	FG	not revised
122	34	Remijia	sessilis	F	not revised
123	35	Remijia	sipapoensis	G	not revised
124	30 27	Remijia	tenuiflora	G E	not revised
126	38	Remijia	trianae	F	not revised
127	30	Remiiia	ulei	F	not revised
128	40	Remiiia	vaunesiana	, F	not revised
129	41	Remiiia	wurdackii	F	not revised
130	1	Stilpnophyllum	grandifolium	c	1 (type only)
131	2	Stilpnophvllum	lineatum	D	1 (type only)
132	- 3	Stilpnophyllum	oellgaardii	c	2
133	4	Stilpnophyllum	revolutum	С	1 (type only)

Species in bold type were included in the molecular phylogeny. Distribution codings are according to Fig. 1 Inset.