Diversification of rhacophorid frogs provides evidence for accelerated faunal exchange between India and Eurasia during the Oligocene

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Contributed by David M. Hillis, January 15, 2013 (sent for review November 19, 2012)

The accretion of the Indian subcontinent to Eurasia triggered a massive faunal and floral exchange, with Gondwanan taxa entering into Asia and vice versa. The traditional view on the Indian–Asian collision assumes contact of the continental plates during the Early Eocene. Many biogeographic studies rely on this assumption. However, the exact mode and timing of this geological event is still under debate. Here we address, based on an extensive phylogenetic analysis of rhacophorid tree frogs, if there was already a Paleogene biogeographic link between Southeast Asia and India; in which direction faunal exchange occurred between India and Eurasia within the Rhacophoridae; and if the timing of the faunal exchange correlates with one of the recently suggested geological models. Rhacophorid tree frogs showed an early dispersal from India to Asia between 46 and 57 Ma, as reconstructed from the fossil record. During the Middle Eocene, however, faunal exchange ceased, followed by increase of rhacophorid dispersal events between Asia and the Indian subcontinent during the Oligocene that continued until the Middle Miocene. This corroborates recent geological models that argue for a much later final collision between the continental plates. We predict that the Oligocene faunal exchange between the Indian subcontinent and Asia, as shown here for rhacophorid frogs, also applies for other nonvolant organisms with an Indian–Asian distribution, and suggest that previous studies that deal with this faunal interchange should be carefully reinvestigated.

phylogeography | molecular dating | tectonics

The tectonic collision between India and Asia induced a major biotic exchange from the former Gondwanan continent to tropical Asia during the Early Cenozoic (1). Recent molecular studies have confirmed that Late Cretaceous faunal and floral elements entered Asia from Africa and Madagascar via the northward drifting Indian subcontinent (2–8). Thus, this tectonic event can be considered a major driver of biotic diversification, comparable to the “Great American Interchange” between North and South America after the closure of the Isthmus of Panama (9). However, the timing of the Indian–Asian collision and, of equal importance, the position of the Indian plate on its move northwards relative to the adjacent continents, is still under debate (10–17). A set of recent hypotheses is given in Table 1. Early studies suggested that the India–Asia collision occurred during the Paleocene or Early Eocene at 66–56 Ma (18), and many biogeographic studies relied on this date (1, 3, 4). More recent geological and tectonic plate modeling analyses suggest that the final collision occurred much later, either around 35 Ma (10, 12) or between 25 and 20 Ma (16), and was just predated by the collision of either Greater India with an intraoceanic volcanic island arc (10, 12) or the collision of a Tibetan–Himalayan microcontinent with Asia around 52 Ma (16). Here we ask if these revised geological dates are in accordance with biogeographic data for a group of amphibians that occurs throughout this area and is restricted by the marine environment.

Tectonic collisions do not necessarily produce terrestrial connections. Thus, biogeographic research is needed to understand the biological implications of plate movements and collisions (19). In the case of northward drifting India, this applies especially for potential connections between India and Southeast Asia, as recently suggested, based on freshwater crab biogeography (19). Amphibians are ideal organisms to infer geological and environmental history due to their poor marine dispersal capabilities (3, 20–24). Frogs of the family Rhacophoridae (Old World tree frogs or shrub frogs) are a particularly appropriate animal group to test the timing of the Indian–Asian collision as their range comprises both India and East/Southeast Asia (21, 25) (Fig. S1 and Table S1). Previous studies identified the Madagascan Mantellidae as the sister group of the Rhacophoridae and concluded that the latter reached Eurasia via the northward drifting Indian subcontinent (3, 21, 26, 27), whereas three rhacophorid species that currently inhabit tropical Africa were shown to be the result of more recent dispersal from Asia (28, 29).

Here, we integrate phylogenetic, biogeographic, and molecular dating methods to reconstruct the comprehensive molecular phylogeny of rhacophorid frogs. In this time-calibrated phylogenetic framework, we reconstruct ancestral geographical areas to test (i) if there was already a Paleogene biogeographic link between Southeast Asia and India; (ii) in which direction faunal exchange occurred between India and Eurasia within the Rhacophoridae; and (iii) if the timing of the faunal exchange correlates with one of the recently suggested geological hypotheses (Table 1), thus providing evidence for the paleoposition of the Indian subcontinent as it moved northward.

**Results**

**Sequence Characteristics.** The aligned mtDNA gene fragments consisted of 2,041 sites (excluding positions of unstable alignment). The mitochondrial data correspond to sites 726–2,666 of the Polypedates megacephalus mitochondrial genome (AY458598). Transitions and transversions within the five nDNA partitions accumulated in a linear way and gave no indication of saturation. The concatenated nuclear dataset comprised 3,225 base pairs (bp), consisting of five


The authors declare no conflict of interest.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database. For a list of accession numbers, see Table S1.

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This article contains supporting information online at www.pnas.org/cgi/suppl/doi:10.1073/pnas.1300881110/-/DCSupplemental.
gene fragments: brain-derived neurotrophic factor (BDNF) (614 bp), proopiomelanocortin (POMC) (601 bp), recombination activating gene 1 (RAG-1) (1164 bp), rhodopsin exon 1 (RHOD) (315 bp), and tyrosinase exon 1 (TYR) (531 bp).

Phylogenetic Inference Identifies Rhacophoridae Dispersal Between India and Southeast Asia. The phylogeny based on the combined mitochondrial and nuclear data are largely consistent with the results of the nuclear data alone (Fig. 1 and Figs. S2 and S3); only the position of the genus Chiromantis differs in the two analyses (neither topology with strong support). Also the estimated divergence times for our combined dataset and the nuclear data were largely equally consistent (Fig. 1 and Figs. S2 and S3).

Our combined phylogeny confirms the sister-group relationship of Rhacophoridae and Madagascan Mantellidae that separated around 59 Ma (53–68 Ma highest posterior density (HPD), i.e., 95% HPD interval; see node 1 in Fig. 1). The dichotomy between the East Asian Buergeriinae (only including Buergeria) and Rhacophorinae (comprising the remaining rhacophorids) is strongly supported and likely occurred during the Early Eocene around 50 Ma (46–57 Ma, 95% HPD; node 2 in Fig. 1).

Within the subfamily Rhacophorinae the genus Liixalus (Southeast–East Asia) is recovered as the sister group of the other rhacophorines, separating around 44 Ma (39–50 Ma, 95% HPD; node 3 in Fig. 1). The genera Thelodema and Nyctixalus (East Asia–Insular Southeast Asia) form a clade that also diverged early (36–47 Ma, 95% HPD; node 4 in Fig. 1). Within the remaining Racomorphae that share a common ancestor around 40 Ma (35–46 Ma, 95% HPD; node 5 in Fig. 1), several clades with robust support can be identified: a clade of Karixalus, Raorchestes, and Pseudophilautus (“group 2,” time to the most recent common ancestor (tMRCA) 31–42 Ma, 95% HPD; node 8 in Fig. 1); the genus Chiromantis (diverging at 33–43 Ma, 95% HPD; node 7 in Fig. 1), comprising African representatives that separated between 19 and 28 Ma from their Asian relatives; and a clade consisting of the genera Polypedates, Taruga, Ghatixalus, and Feihyla (“group 1,” tMRCA 30–39 Ma, 95% HPD; node 10 in Fig. 1) that is sister to the genus Rhacophorus (tMRCA 31–41 Ma, 95% HPD; node 9 in Fig. 1).

The results of the biogeographic inference (Fig. 2 and Table S2) indicated that the most recent common ancestor (MRCA) of the extant Rhacophoridae inhabited East/Southeast Asia. The earliest splits within Rhacophoridae most likely occurred in East Asia, resulting in the genera Buergeria and Liixalus (Fig. 1), which currently inhabit South China and the islands of Taiwan, Hainan, and Japan. Hypothesis testing in the program Lagrange indicated that dispersal between India and Asia started after 35 Ma but before 25 Ma (Table S3). We could identify several dispersal events from East/Southeast Asia and Insular Southeast Asia to the Indian subcontinent (Figs. 1 and 2 and Table S2): Four events from Insular Southeast Asia at 28–38 Ma, 23–32 Ma, 15–22 Ma, and 14–23 Ma (three events within “group 1” and Rhacophorus malabaricus); and two events from East/Southeast Asia (estimated between 13 and 30 Ma). However, more frequent dispersal events occurred from East/Southeast Asia to Insular Southeast Asia (five events between 6 and 50 Ma), and from Insular Southeast Asia to East/Southeast Asia (five events between 7 and 37 Ma). Within the genus Polypedates rhacophorids dispersed from India to Insular Southeast Asia between 10 and 28 Ma.

Discussion

Origin of Rhacophoridae. Our phylogenetic analyses support the sister relationship of Rhacophoridae and Mantellidae and is consistent with previous studies inferring the interrelationships of these two families with Ranixalidae and Dicroglossidae (3, 26, 27). As the Mantellidae currently occur only on Madagascar and Mayotte Island, and Ranoidea as a whole supposedly evolved in Africa (3), it has been suggested that Rhacophoridae and Mantellidae have a Gondwanan origin, and the northward drifting Indian plate mediated the dispersal of rhacophorids to Asia (3). This dispersal pathway is generally proposed for Late Cretaceous Gondwanan fauna and flora (2–8), including several other groups of anurans (3, 7). However, there is evidence for a post-Cretaceous biogeographic connection between Madagascar and India and/or the Seychelles (30–32). This dispersal path could also apply
for the rhacophorid ancestors, as the MRCA of Rhacophoridae and Mantellidae is estimated at 53–68 Ma, and thus after the tectonic split of India and Madagascar (30, 31).

The oldest phylogenetic splits within the Rhacophoridae occurred in Asia, with all known extant Indian rhacophorids originating from later dispersal events to India (Figs. 1 and 2). Thus,
extant rhacophorids clearly have an Asian origin. Nevertheless, the occurrence of Eocene fossil Rhacophoridae on the Indian subcontinent (33), which we have used to calibrate the MRCA of the family, argues for an Indian origin of Rhacophoridae, followed by dispersal to Asia that must have occurred before 46 and 57 Ma (Fig. 1). Previous studies indicated that the Indian subcontinent experienced dramatic latitudinal and climatic changes that caused massive extinctions in its biota in the early Cenozoic (1, 34), and this might have also affected the rhacophorids.

How Rhacophorids Reached Asia Before the Final Collision Between India and Asia. Long-distance dispersal over the Tethys Ocean has to be considered unlikely for rhacophorid frogs, given their sensitivity to salinity and their arboreal habitat. Thus, a plausible geological model of the Indian–Asian collision should be able to explain (i) the early dispersal of rhacophorids from India to Asia between 46 and 57 Ma (Fig. 1); (ii) the cessation of faunal exchange during the Middle Eocene (39–46 Ma); and (iii) the increase of rhacophorid dispersal events between the Indian subcontinent and Asia starting in the Late Eocene and continuing until the Middle Miocene (Figs. 1 and 2). Although the collision between India and Asia is currently the largest active orogeny (10), there exists much uncertainty regarding the position of India on its move northwards relative to the adjacent continents and the timing of the initial collision (19, 35), which are both highly controversial (11, 14–17).

Multiple studies have suggested that the Indian–Asian collision occurred during the Paleocene/Early Eocene at 50–55 Ma (36, 37). Most previous biogeographic studies rely on this date (38, 39). Also the estimated time frame when the rhacophorid ancestor dispersed from India to Asia fits into this scenario. However, recent geological models draw a more complex picture of the Indian–Asian collision (10, 16), proposing that the final collision that led to the uplift of the Himalaya was preceded around 50 Ma by the collision of the Indian subcontinent with an intraoceanic island arc (10), followed by the continuing northward drift of the Indian subcontinent along Southeast Asia that might have allowed terrestrial connections (40). Alternatively, the Early Eocene collision event was caused by the accretion of a Tibetan–Himalayan microcontinent at 49–55 Ma that broke off the Indian subcontinent before (16). Both models are therefore able to explain an Early Eocene faunal exchange, followed by another period of continental isolation. The existence of a microcontinent that acted as a “ferry” for the rhacophorid ancestor that is dated between 46 and 57 Ma (Fig. 1) is compelling. As megathermal rainforests were widespread across India and Asia at the Paleocene–Eocene boundary (41) (Fig. 2), rhacophorids are likely to have expanded their range onto the Asian mainland very quickly. This could explain why the clades that diverged earliest within Rhacophoridae (Buergeria and Liuxialus) currently only occur in East Asia, far from the Indian subcontinent.
No Faunal Exchange Between Southeast Asia and India During the Middle Eocene. During the Eocene (52–34 Ma) rhacophorids dispersed between East/Southeast Asia and Insular Southeast Asia (Fig. 2), as evidenced by the presence of the genera Nyctixalus and Theloderma in Southeast Asia (areas III and IV, Fig. 1). There was, however, no exchange of species between Asia and India. The most straightforward interpretation is that there were no terrestrial connections with India at that time. The “traditional” geological view on the Indian–Asian collision (assuming a final contact of plates during the early Eocene) thus fails to explain why dispersal between India and Asia ceased during the following 10 million years, although climatic conditions supposedly would have been favorable for tropical tree frogs during this period (41). The more recent geological models that argue for a later final collision seem to fit the present biogeographic data better. Van Hinsbergen et al. (14–17) suggested that there was no continent–continent contact from 52 Ma (the accretion of the Tibetan–Himalayan microcontinent) to 25–20 Ma (the final collision of the Indian subcontinent). In contrast, Atchison and Ali (10–13) inferred that the final continent–continent collision began around the Eocene/Oligocene boundary (34 Ma).

Faunal Exchange Between Asia and India Since the Oligocene. According to the rhacophorid data, frequent faunal exchange between Southeast Asia and the Indian subcontinent started during the Early Oligocene (Fig. 2 and Table S2, node 10). This is consistent with the model of Ali and Atchison (10) for the timing of the final continental collision. In contrast, a final collision at 25–20 Ma (16) appears to be much later than our time estimates of Neogene rhacophorid dispersal from Asia to India. Faunal exchange between the Indian subcontinent and Asia ceased again at the transition between Middle and Late Miocene around 12 Ma. We propose that this correlates with the rapid uplift of the Himalayan mountains that continued until 10 Ma (42), leading to the establishment of the Indian and East Asian monsoon (43), and thus of more seasonal climates. Whereas in northern India extant rhacophorids are absent due to the monsoon climate, the humid and warm conditions of East and Southeast Asia from the Miocene onwards (41) likely resulted in the rapid diversification of Rhacophoridae (e.g., the genera Rhacophorus, Polyedates, and Kuriulus).

The biotic interchange between the Indian subcontinent and Asia significantly contributed to the vast biodiversity of present-day East and Southeast Asia. This is consistent with inferences from mammalian fossils and paleobotanical data (44–46). We predict that the Oligocene faunal exchange between the Indian subcontinent and Asia, as shown here for rhacophorid frogs, also applies for other terrestrial nonvolant organisms with an Indian–Asian distribution (8, 19), and suggest that previous studies that deal with this faunal interchange should be carefully reinvestigated.

Materials and Methods

Taxon Sampling and Data Collection. Our sampling covered the entire distribution of Rhacophoridae from tropical Africa and Asia to temperate China and Japan (Table S1). Whenever possible, samples from the type localities of nominal taxa were included and multiple samples were used to represent widespread species. Currently, 15 genera with 321 species of rhacophorids are recognized (25, 47). The mitochondrial DNA (mtDNA) dataset included 317 samples from all 15 rhacophorid genera, representing 114 species. For the analyses of the nuclear DNA (nDNA) markers, at least three species were sampled from each genus, totaling 64 taxa. We used two species of the family Dicroglossidae, 1 from the Ranidae, and 10 from the Mantellidae as outgroup taxa. Both the nDNA dataset and the combined mtDNA and nDNA data were used to perform phylogenetic analyses, molecular dating, and biogeographic reconstruction.

DNA Extraction, Amplification, and Sequencing. Genomic DNA was extracted from toe clips, muscle, or liver tissue preserved in 95% or 100% ethanol. Tissue samples were digested using proteinase K and then followed a standard three-step phenol/chloroform extraction procedure (48, 49). The mtDNA sequences were amplified using partial 12S and 16S rRNA together with the complete tRNA for valine using primers from previous studies (50–52). PCR was conducted as described previously (50, 51). Forward and reverse strands of purified PCR products were sequenced with an ABI 3730 automated DNA sequencer controlled via BLAST searching (53) against GenBank to ensure that the target sequences had been amplified. The nDNA data originate from a previous study (51) and include five gene fragments: BDNF, POMC, RAG-1, RHOD, and TYR. GenBank accession numbers for both the new and previously deposited data are given in Table S1.

Phylogenetic and Divergence Time Analyses. Sequences were aligned using ClustalX 1.81 (54) with default parameters, and subsequently minor adjustments were made to improve alignments following visual inspection. Positions that could not be confidently aligned were excluded to remove potentially misleading data. Gaps resulting from the alignment were treated as missing data. Because all mtDNA gene sequences are effectively inherited as one locus, they were concatenated into a single contiguous alignment for the analyses. For the mtDNA and each nDNA gene, possible saturation of substitution types was checked by plotting the number of transitions and transversions versus F84 distance using DAMBE (55). We simultaneously inferred phylogeny and divergence time estimates using a Bayesian relaxed molecular clock approach on the combined mtDNA and nDNA datasets using BEAST 1.6.3 (56). We incorporated partitioned strategies (57) and treated each gene as one part separately. After a preliminary series of test runs, we chose the following models to fit best to the data: the uncorrelated lognormal relaxed molecular clock model, a Yule tree prior, and GTR+I+Γ for the substitution model for each gene dataset. We performed a total of eight BEAST runs. The Markov chain Monte Carlo (MCMC) was run for 30 million generations and sampled every 1,000 generations. Convergence of MCMC chains was checked in Tracer 1.4.1 (58) and subsequently verified using the cumulative and compare functions in AWTY (59).

We calibrated the age of the most recent common ancestor of the Rhacophoridae with the age of fossil Indorastra prasadil from Early Eocene sediments of India (33). The prior distribution for a fossil-based age constraint was modeled with a 1°-distribution (shape value = 4; scale factor = 2) with the fossil minimum age (45 Ma) as the lower boundary and 48 Ma as the split between Boophis tephraemystax, an inhabitant of the Comoros island of Mayotte, and its sister species (Boophis doulioti) (30, 60) as an age constraint for the second calibrated time. Mayotte is a volcanic, oceanic island with no known previous connections to a larger landmass; thus the B. tephraemystax lineage could not have split from its sister lineage before 15 Ma, the maximum age of the island (30, 60). Therefore, we modeled the calibration age constraint using a broad uniform prior distribution from 15 Ma to present. We show the nuclear Bayesian and combined mitochondrial and nuclear data trees using the fossil and biogeographic calibrations (combined mitochondrial and nuclear data, Fig. 1; nuclear phylogeny Fig. S2).

Area Delimitation and Biogeographical Reconstruction. We compiled data of rhacophorid species from the literature (25) and assigned the individual reports to the respective areas. To answer questions concerning the direction of faunal exchange between the Indian subcontinent and Eurasia, we divided the rhacophorid range into four areas (Fig. 1): Africa (area I); the Indian subcontinent (area II); Insular Southeast Asia including the Malay Peninsula (area III); and East/Southeast Asia (area IV). Each species was assigned to its associated area according to its contemporary distribution range. Widespread species were assigned to more than one area (Table S1).

Biogeographical inferences were obtained by applying statistical dispersal–vicariance analysis (S-DIVA) and Bayesian binary MCMC (BBM) analysis implemented in RASp v2.0 using the default settings (61). Random 1,000 BEAST output trees were used and the maximum number of individual units was set to two. In addition to these event-based biogeographic methods, we also applied a parametric approach based on dispersal, extinction, and cladogenesis as implemented in the software Lagrange v. 20120508 (62). The analysis used the maximum clad credibility tree of the Bayesian phylogenetic analysis and the same area coding as for the event-based methods. The maximum size of ancestral ranges was set to two; adjacency of areas was not constrained. To test if the faunal exchange between the Indian subcontinent and Eurasia fits better to an earlier final collision of the two landmasses around 35 Ma (10) or rather a later one around 25 Ma (16), we compared different temporal dispersal models: an unconstrained model, a model that only allows faunal exchange after 35 Ma and one allowing dispersal only after 25 Ma. Based on the global likelihood of the models we ranked them according to their Akaikes Information Criterion values and Akaike weights (Table S3).