Phylogenomic assessment of microhylid frogs reveals widespread taxonomic confusion in the Asterophryinae and establishes the timing of diversification in Australia

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*Highlights:*

* A phylogenomic assessment of the globally distributed frog family Microhylidae
* The subfamily Asterophryinae radiated explosively in New Guinea
* Australian species represent two distinct clades
* Assignments of species to genera by morphological means are often unreliable

# Abstract

Microhylid frogs are a hyper-diverse family thought to have radiated explosively around the Cretaceous-Paleogene boundary. Roughly half of microhylid species richness is concentrated into a single subfamily, the Asterophryinae, which is centered in New Guinea and surrounds, and has been a rich source for species discovery over the past 50 years. However, resolving Asteroprhyinae phylogenetics has remained a challenge, with frequent taxonomic reshuffling. To address this instability, we generated a sequence-capture molecular dataset to investigate the phylogenetics of the group. This included 71 species of Asteroprhyinae, across 13 of 17 recognized genera representing extensive sampling of the New Guinea radiation and full sampling of Australian microhylid species. Our dated species tree supports an explosive diversification of microhylids in New Guinea near the start of the Miocene, approximately 20 million years ago. Asterophryinae expansion into northern Australia occurred much later (~10 ma) and is marked by well supported clades of *Austrochaperina* and *Cophixalus* that show temporally consistent splits from their New Guinea ancestors. Our phylogeny allows us to identify several instances of polyphyly which are at odds with our current understanding of intergeneric relationships within the Asterophryinae. We suggest this confusion is a result of rapid radiation and morphological variability across some poorly defined genera. This work establishes a reliable phylogenetic framework that can form a foundation for a more stable taxonomy of the Asterophryinae.

# Introduction

Narrow-mouthed frogs, Family Microhylidae, are one of the most species-rich groups of frogs worldwide. These tropical and subtropical frogs comprise nearly 800 species with a wide array of ecologies and morphologies (Frost, 2025). Their diversity is so broad that it defies easy summary but includes everything from round-bodied fossorial to slender toe-padded arboreal species, desert and rainforest specialists, and a variety of larval strategies (aquatic, direct-developing, foam-nest living, non-feeding). They are also geographically widespread, ranging from North, Central and South America, Africa, Madagascar, India, and southeast Asia, to New Guinea and northern Australia (Fig.1). Microhylids are allocated to 12 subfamilies and the relationships within and between them have been the subject of many molecular phylogenetic assessments (van der Meijden et al. 2007; Kurabayashi et al. 2011; da Sá et al. 2012; Peloso et al. 2016; Tu et al. 2018; Streicher et al. 2020) (see Peloso et al. 2016 Fig.1). The increasing size of molecular datasets has brought consensus to most intrafamilial relationships, despite using different marker types (Feng et al. 2017; Streicher et al. 2020; Hime et al. 2021). This level of agreement, however, has not extended to relationships within the most species-rich subfamily—the Asterophryinae.

The Asterophryinae is the largest subfamily of microhylids with over 370 species and 17 genera. Asterophryines are distributed from mainland southeast Asia (*Siamophryne*, *Vietnamophryne*), to Borneo (*Gastrophrynoides*), the Philippines (*Aphantophryne*), New Guinea (14 genera), and northern Australia (*Austrochaperina, Cophixalus*) (Frost, 2025). Richness peaks in New Guinea (Fig.1), with over 250 species and likely many more to be described (Ferreira et al. 2024; Ferreira et al. 2025). In contrast, microhylid diversity in Australia is limited to ~25 species, with just two genera (*Austrochaperina* and *Cophixalus*) on the mainland (Fig.1; Zweifel 1985; Hoskin 2004; Hoskin 2013), and two recently described species (*Callulops* and *Choerophryne*) on a far northern island of the Torres Strait. These two newly described species are endemic to Dauan Island, politically part of Australia, but geographically adjacent to New Guinea (Hoskin 2025). The vast majority of microhylid diversity in Australia comprises *Cophixalus* species in the Wet Tropics rainforests, with smaller numbers of species in the drier Cape York region and a single species of *Austrochaperina* in the far north of the Northern Territory (Fig.1). The evolutionary history of microhylid frogs in Australia is currently unresolved, including whether the two mainland genera represent clades relative to New Guinean congeners, or are the result of multiple dispersals from New Guinea.

Phylogenetic relationships within and among genera of the Asterophryinae have been examined thoroughly, but with little consensus, hindered by frequent findings of para- and polyphyly (Kohler and Gunther, 2008; Rivera et al. 2017; Hill et al. 2022, Hill et al. 2023). This is likely due to rapid radiation of the group after crossing Wallace’s Line ~20 million years ago, resulting in bursts in speciation and ecomorphological diversification. As a result of highly variable morphologies, and because it is rare for diagnoses of Asterophryinae species to include molecular evidence, generic assignments in this subfamily have changed frequently (Frost, 2025). The frustrations of Asterophryinae generic assignments have been so extreme that Dubois et al. (2021) proposed that all 360+ species (with exception of *Gastrophrynoides, Siamophryne,* and *Vietnamophryne*) be lumped under a single genus, *Asterophrys*. This suggestion, however, has not been adopted by researchers in the field (Frost, 2025).

Here we present a phylogenomic perspective on the diversification of Asterophryinae microhylids, particularly among New Guinean and Australian taxa. We started by generating a sequence-capture dataset to investigate the topology and timing of Sahulian asterophryine diversification, with the goal of providing a reliable backbone of intergeneric relationships. We assess the timing of diversification in New Guinea and Australia, and we resolve how the Australian species fit into the evolutionary history of this group. While we know a great deal about many aspects of Australian frog biology (Tyler 1998; Anstis 2017, Brennan et al. 2023), comparatively little is known about the phylogenetics and history of the ~25 species of microhylids (Zweifel 1985; Hoskin 2004). We also evaluated commonly used morphological information to quantify the distribution and utility of traits in generic assignments. Our work aims to provide insight into the phylogenetics of a taxonomically volatile group and expand our understanding of the biotic interchange between Australia and New Guinea.

# Materials and Methods

*Phylogenomics*

We assembled a sequence-capture dataset comprising 149 frog samples across 107 species that span nearly all microhylid subfamilies (10 of 12 recognized). Exceptions are limited to Hoplophryninae and Melanobatrachinae frogs of East Africa and India. Sampling focused on the Asterophryinae and represents 71 species from 13 of 17 recognized genera (with exceptions *Gastrophrynoides*, *Paedophryne*, *Siamophryne*, *Vietnamophryne*) (Table S1). We include near-complete sampling of the Australian Asterophryinae species (5 *Austrochaperina* spp.; 19 *Cophixalus* spp.) with the exception of *Cophixalus peninsularis* which is known only from two specimens collected in the 1980s and which is likely to be conspecific to *C. crepitans* (Hoskin, 2004).

We generated new Anchored Hybrid Enrichment (AHE—Lemmon et al. 2012) data for 96 samples and combined these with outgroup samples from Hime et al.’s (2021) amphibian phylogenomic dataset. We initiated this process by blasting AHE loci against the *Xenopus tropicalis* genome using *metablastr* (*blast\_best\_reciprocal\_hit*) (Benoit & Drost 2021) and renaming loci according to their orthologs in *Xenopus*. We similarly carried out this process on anuran samples from Hime et al. (2021) to harmonize target sequences across datasets. Samples across different AHE projects were combined using the *pipesnake* workflow (Brennan et al. 2024) to align and trim sequence data, and estimate locus and species trees. Briefly, sequences were aligned with *mafft* (Katoh et al. 2013), trimmed for gappy sites using *clipkit* (Steenwyk et al. 2020), then locus trees (n=450) were estimated under maximum-likelihood in IQTREE2 (Minh et al. 2020), allowing the program to assign the best fitting model of nucleotide substitution using ModelFinder (Kalyaanamoorthy et al. 2017) and then perform 1,000 ultrafast bootstraps (Minh et al. 2013), before being passed to hybrid weighted-ASTRAL (Zhang et al. 2018) to estimate a species tree.

To verify the identity of newly sequenced samples we assembled off-target reads of the mitochondrial loci CYTB and ND4 and combined these data with the alignments of Hill et al. (2023). We started by loosely mapping raw sequence reads to the *Microhyla pulchra* mitochondrial genome using *BBMAP* (Bushnell, 2014), then assembled mapped reads using *SPAdes* (Prjibelski et al. 2020). We added new sequences to the existing alignments with *mafft*, concatenated the alignments, and estimated a single mitochondrial topology using IQTREE2.

*Divergence Dating*

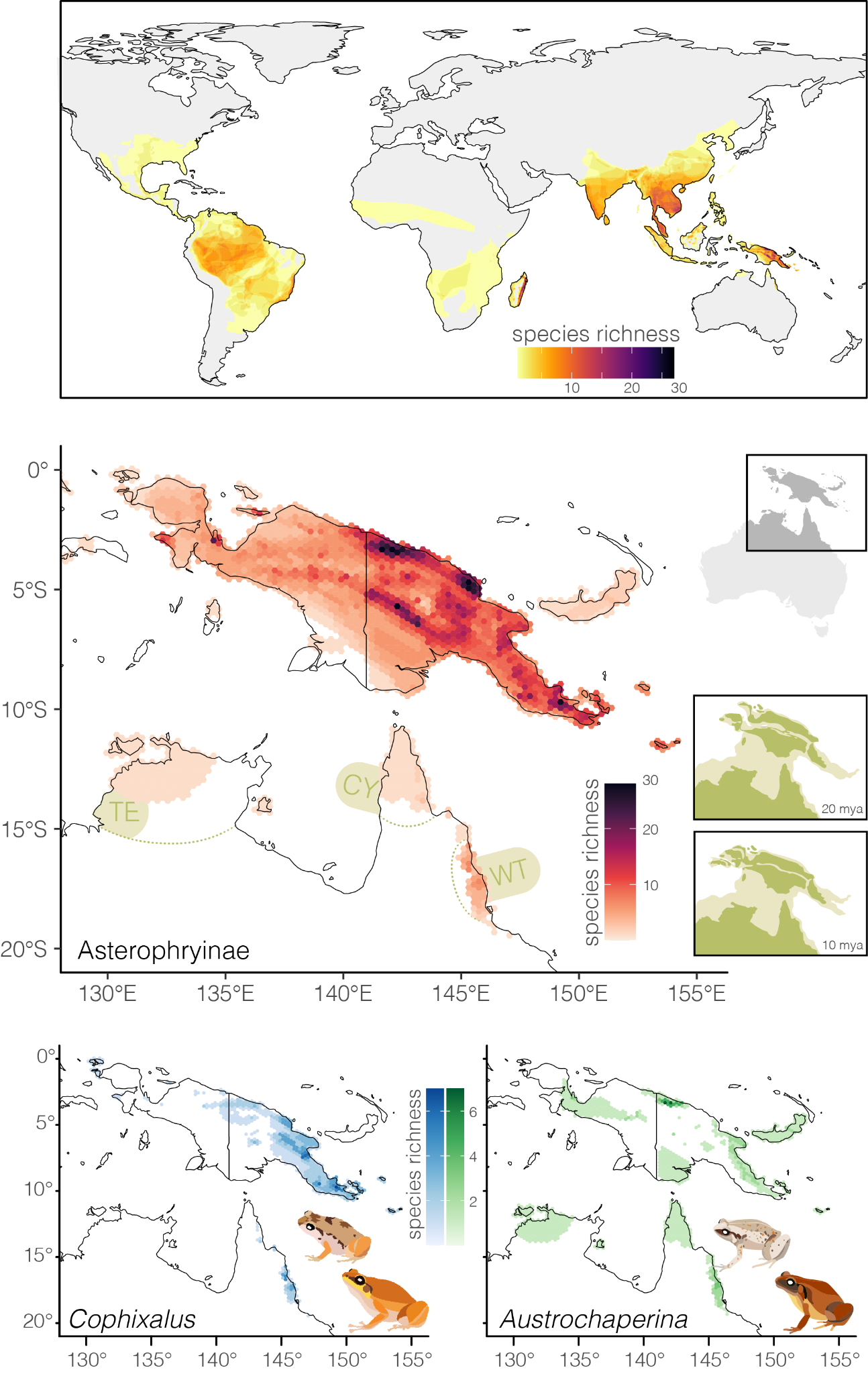
To estimate divergence times among taxa on the ASTRAL species tree we applied a series of fossil calibrations, first compiled by Feng et al. (2017) (Table S2), and used the Bayesian divergence time software MCMCtree (Rannala & Young 2007). We started by downloading orthologous coding sequences from the Orthologous Matrix (Altenhoff et al. 2024) for *Xenopus* and *Bufo* and aligned the exonic loci via MACSE (Ranwez et al. 2018). We then added our ingroup microhylid sequences to these codon-aligned sequences via *mafft* (*--add*, *--keeplength*), concatenated them, and partitioned first and second codon positions together following the strategy of dos Reis et al. (2018). Complex partitioning strategies such as filtering by evolutionary rate are possible but less influential than the absolute number of partitions (dos Reis et al. 2012). Additional data partitions ultimately incur substantial computational costs for modest increases in dating precision, and so we opted instead for a more conservative approach. We then used *baseml* to estimate approximate likelihoods (dos Reis & Yang 2011) and branch lengths before running *mcmctree* on the gradient and Hessian (in.BV file) for four replicate analyses. We inspected mcmc files for stationarity and compared for convergence, then combined them using logCombiner, and used this combined mcmc file to summarize divergence times on our tree (*print = -1* in .ctl file). Sample, alignment, and gene trees are available alongside all other materials on Dryad (doi:) and GitHub (<https://github.com/IanGBrennan/Asterophryinae>).

*Taxonomy*

For consistency, we adopt the taxonomy of Amphibian Species of the World v6.2 (Frost, 2025), which shows minor differences in the number of recognized species, recognized genera, and generic assignments from other amphibian authorities such as AmphibiaWeb. Notably, Amphibian Species of the World has incorporated the taxonomic suggestions of Rivera et al. (2017) regarding assignment of Asterophryinae genera and has not implemented the recommendation of Dubois et al. (2021) to lump most asterophryines into *Asterophrys*.

*Morphology*

We quantified commonly collected morphological traits used in Asterophryinae species descriptions, to evaluate a potential explanation for why generic assignments have been so unstable. Asterophryinae genera diagnoses often rely on internal anatomical traits such as presence/absence of clavicles, procoracoids, and the shape of the jaw. However, these traits are not always assessed for new species. We started by collecting eight morphological measurements from species descriptions in the microhylid literature (40 publications): snout-urostyle length (SUL), tibia length (TL), head width (HW), internasal distance (IN), eye-nasal distance (EN), eye diameter (EYE), third finger disc diameter (F3D), and fourth toe disc diameter (T4D). To remove the effect of size on individual traits (allometry) we calculated the geometric mean of all traits by individual and used this to transform measurements into log-shape ratios. We retained the geometric mean as a ninth trait (SIZE). We then used current generic assignments as a discrete character and carried out a RandomForest (Liaw & Wiener, 2022) analysis on the morphological traits to determine rates of mischaracterization based on gross phenotype. For comparison with other commonly used methods, we also applied Linear Discriminant (LDA) and Flexible Discriminant (FDA) analyses. To visualize the partitioning of morphological space we used dimensionality reduction techniques (PCA, LDA, FDA) and plotted the first two axes of variation.

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**Figure 1**. The Microhylidae are a global frog radiation with multiple hotspots of species richness. (Top) Mapping the global distribution of microhylid species highlights high richness on the east coast of Madagascar, across southeast Asia, and particularly on New Guinea. (Middle) Across New Guinea, richness is dominated by the subfamily Asterophryinae, and surrounding islands and landmasses show comparatively low diversity. Australian regions mentioned in the text are delineated by dotted lines: Top End (TE), Cape York (CY), Wet Tropics (WT). Inset maps at right indicate the proximity and connectivity of Australia and the various geological blocks that ultimately formed New Guinea at two different time periods: 20 million years ago during the rapid diversification of Asterophryinae microhylids in this region, and 10 million years ago when *Austrochaperina* and *Cophixalus* are hypothesized to have dispersed onto the Australian continent. Geological reconstructions are derived from Gold et al. 2020. (Bottom) The contemporary distribution of microhylids in Australia is concentrated in the Wet Tropics (WT) of North Queensland. Maps at indicate the spatial dynamics of species richness in *Cophixalus* (top) and *Austrochaperina* (bottom). One Australian species *Austrochaperina gracilipes* is also found in southern Papua New Guinea.

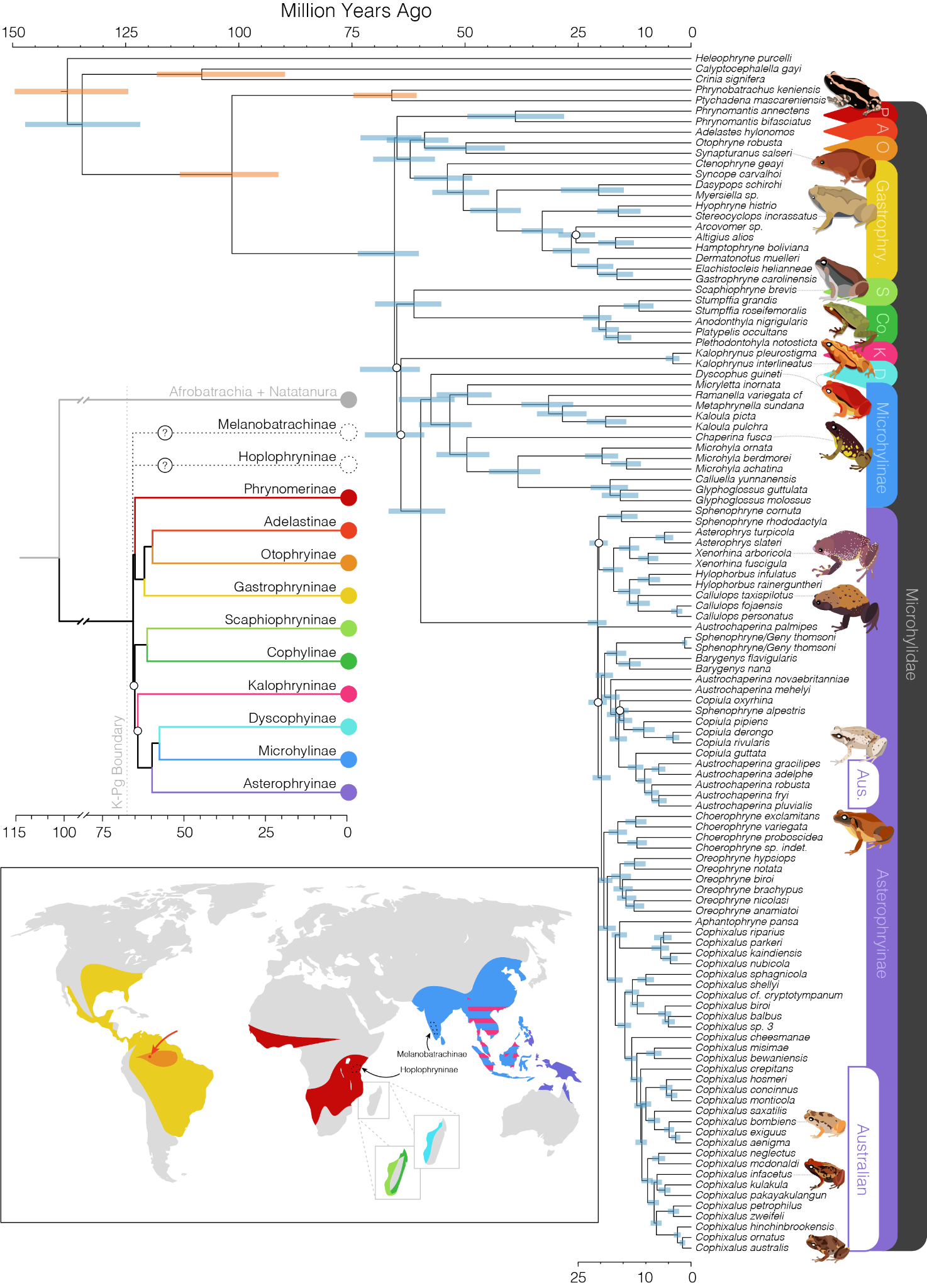
Results

*Phylogenomics and Divergence Dating*

We present a phylogenetic hypothesis of relationships for 107 species of microhylid frogs based on new anchored hybrid enrichment (AHE) data for 95 new samples. Sampling includes most recognized Asterophryinae genera (13 of 17) and, for the first time, all mainland Australian species of Asterophryinae. We combine this new molecular dataset with existing AHE data from Hime et al. (2021) to estimate relationships among 10 of the 12 microhylid subfamilies (Fig.2; S1).

We captured 356 loci with a mean alignment length of 1564 bp (max. = 5535, min. = 377) and sample occupancy of 120 individuals (max. = 148, min. = 34). Species tree analyses with ASTRAL provided strong support (>90 local posterior probability) for most branches of the tree (Fig.2; S1), with some exceptions among subfamilial splits (e.g. position of Kalophryninae and Scaphiophryninae/Cophylinae relative to the South American and Asian clades) and among some Asterophryinae clades. The subfamily topology is entirely consistent with Hime et al. (2021) and differs only in a small number of changes from Feng et al. (2017) (placement of Phrynomerinae) and Streicher et al. (2020) (position of Kalophryninae and Phrynomerinae). The microhylid stem branch is very long (>35 ma) and leads to an explosive radiation into major microhylid clades (subfamilies) roughly coincident with the Cretaceous-Paleogene boundary. The four earliest splits in the family all occur within a 1.5 million year window at this time (Fig.2; S2).

Our tree topology for the Asterophryinae differs considerably from recent investigations by Rivera et al. (2017) and Hill et al. (2022), and in some ways bears greater similarity to Tu et al. (2018). To verify the identity of new species we successfully recovered off-target mitochondrial data for 84 of 95 new samples (Fig.S3). While our species sampling (~70 spp.) represents roughly a third of the species included in those works, there are notable differences in relationships among major clades, which we will focus on. Importantly, we estimate *Oreophryne* is a deeply nested clade sister to *Cophixalus*, and not an early branch of the tree as seen in Hill et al. (2022) (‘*Oreophryne* A’). We place *Aphantophryne pansa* among *Cophixalus* and not *Oreophryne* (‘*Oreophryne* B*’*)*.* We do not find *Barygenys* embedded within *Austrochaperina*; rather, we identify a sister relationship between *Barygenys* and *Sphenophryne (Genyophryne) thomsoni*. Some topological differences cannot be addressed with our smaller taxon sampling, such as the monophyly of *Oreophryne.* There are, however, important similarities which lend support to an emerging consensus in the intergeneric relationships of Asterophryinae. *Asterophrys*, *Callulops*, *Mantophryne*—not sampled in our data, *Hylophorbus*, *and Xenorhina*, and are recovered consistently as a clade. *Austrochaperina* is clearly paraphyletic, appearing as four lineages/clades in our tree. *Austrochaperina palmipes* is a highly divergent lineage not closely related to other *Austrochaperina* or any other single group. *Sphenophryne* is also clearly paraphyletic, appearing in three positions in the tree. *Cophixalus* (here including *Aphantophryne pansa*) forms a well defined New Guinean and Australian clade.



**Figure 2.** Time calibrated species tree of microhylid frogs highlights the global radiation of the family around the K-Pg boundary (~66 mya) and subsequent explosive diversification of the subfamily Asterophryinae in the early Miocene (from ~20 mya). Inset tree shows the relationships and divergence times among microhylid subfamilies, and color codings match to the geographic distributions of those clades on the map below. Note (1) the small distribution of Adelastinae indicated by an arrow in northern South America, (2) the overlapping distribution of Microhylinae and Kalophryninae in southeast Asia indicated by pink stripes, (3) the presence of three subfamilies on Madagascar, and (4) the uncertain phylogenetic position of the Hoplophryninae (east African) and Melanobatrachinae (south Indian). Primary tree at right shows species level relationships and divergence times, unlabelled nodes have posterior probabilities >0.9 in the weighted-ASTRAL species tree, white circles at nodes indicate branches with local posterior probabilities <0.9. 95% Confidence intervals of divergence times are shown as shaded rectangles at nodes, with orange CIs indicating calibrated nodes.

The timing of Asterophryinae diversification is rapid and resembles the rapid pace of diversification among microhylid subfamilies. A long stem branch that spans ~40 ma was followed by explosive diversification around 20 million years ago and was quickly followed by eight early splits which occurred within a 1.5 million year window. Asterophryinae expansion into northern Australia occurred much later (~10 mya) and is marked by well supported clades of *Austrochaperina* and *Cophixalus* that show contemporaneous splits from their New Guinea sister lineages. The internal topologies of these clades are broadly consistent with their last phylogenetic assessment two decades ago (Hoskin 2004).

Our morphological dataset covered ~519 individual asterophryine frogs representing 120 species (~30% of species diversity) and all 17 recognized genera. Visualizations of dimensionality reduced morphological data (PCA, LDA, FDA) identified large regions of morphospace shared among genera, with some highly distinct forms (*Barygenys*, *Paedophryne*, *Xenorhina*) (Fig. 3). Some groups (*Choerophryne*, *Cophixalus*, *Sphenophryne*) show large or discontinuous distributions that indicate variable morphologies (Fig.3). Classification error for generic assignments using RandomForests were bimodal, though typically low (<10%) (Fig.S4). However, five genera (*Copiula, Choerophryne, Oreophryne, Mantophryne, Siamophryne*) show moderate (10<x<30%) rates of error and three others (*Aphantophryne, Sphenophryne, Vietnamophryne)* show high rates (30–100%). This suggests that morphological characters often used to describe genera do not differentiate them cleanly, even in combination.

# Discussion

Microhylids comprise the third largest family of living amphibians, with almost 800 species distributed across the tropics. Phylogenetics of this group have been a popular topic over the last two decades, greatly improving our understanding of the patterns of diversification and taxonomy of the group (van der Meijden et al. 2007; Kurabayashi et al. 2011; da Sá et al. 2012; Peloso et al. 2016; Tu et al. 2018; Streicher et al. 2020). However, much still remains to be discovered, resolved, and described (Ferreira et al. 2025). Recent global amphibian phylogenomics initiatives (Feng et al. 2016; Hime et al. 2020) have helped establish the evolutionary context in which microhylids have succeeded, and have built substantial data sets to test earlier hypotheses that were based on fewer loci. Our microhylid phylogenomics study presented here helps to clarify our understanding of this globally distributed frog group and provides substantial new information on the diverse Asterophryinae, especially in New Guinea and Australia.

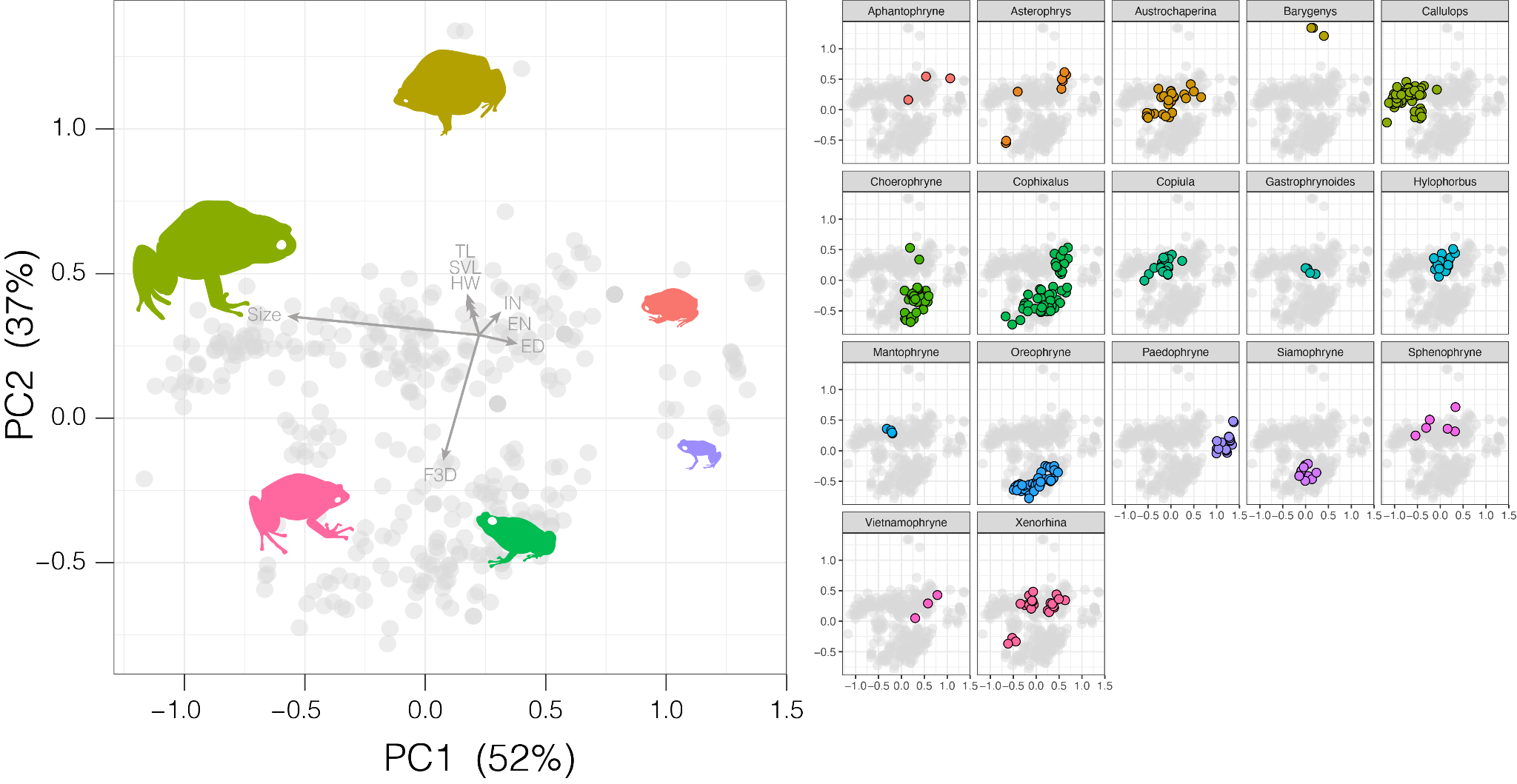
Phylogenetic relationships and divergence times among microhylid subfamilies estimated here are largely consistent with other phylogenomic investigations (Feng et al. 2017; Streicher et al. 2020; Hime et al. 2021). We confirm an explosive radiation at the base of the microhylid tree, rapidly separating the subfamilies. This event is coincident with, or closely follows, the K-Pg turnover. The timing of microhylid diversification is important because they split from the rest of Ranoidea nearly 100 million years ago. In light of this, the long stem branch leading to the Microhylidae (>35 ma) is likely indicative of elevated Cretaceous extinction and a dramatic rebound in the Paleogene. Importantly, the rapid diversification of the group in the wake of the K-Pg turnover—including multiple splits in the first 1.5 ma and establishment of all subfamilies in <10 ma—has the unfortunate effect of blurring the true branching order of the tree. One consequence is ambiguity in the position of Kalophryninae and Cophylinae/Scaphiophryninae. Frequent successive speciation events can be difficult for molecular phylogenetic methods to resolve because high levels of incomplete lineage sorting and gene tree incongruence can obscure the topology (Linkem et al. 2016).

Our topology and divergence dating offers an opportunity to assess the biogeographic history of an old, diverse group, with a wide distribution. Looking across our tree, we see among-subfamily relationships that might best reflect a Gondwanan history. For example, North and South American subfamilies (Adelastinae, Otophryninae, Gastrophryninae) that are sister to an African group (Phrynomerinae), and Madagascan and East Asian clades as sisters. However, the timing of microhylid diversification is at odds with a Gondwanan diversification-by-vicariance scenario. The rapid splitting of subfamilies from one another occurred around the Cretaceous-Paleogene boundary roughly 66 million years ago, long after the separation of Africa, Madagascar and India from Gondwana. Thus, ruling out vicariance as the primary driver of contemporary distributional patterns. Instead, the history of this group has likely been driven by significant long-distance dispersal events followed by subsequent diversification. The number and nature of these events seem almost implausible. For example, van der Meijden et al. (2007) suggested that better taxon sampling might resolve the relationships among the Microhylinae (Southeast Asia), Asterophryinae (eastern Malesia & New Guinea), and Dyscophinae (Madagascar). We interpret their comment to suggest that increased phylogenetic resolution might cluster the Microhylinae and Asterophryinae and provide a more parsimonious biogeographic story. Instead, more and better data presented here only confirms the topology they first presented (Asterophryinae as sister to Dyscophinae and Microhylinae), emphasizing the convoluted dispersal history of this group. As a result, microhylids either undertook two separate dispersals from Madagascar to Asia and Australia, or a single dispersal followed by a return of the Dyscophinae to Madagascar. Like van der Meijden et al. nearly two decades ago, we also find ourselves wishing for more molecular data and more comprehensive taxon sampling to pair with powerful contemporary biogeographic models to “unambiguously resolve the biogeographic history of the Microhylidae.”

Among the Microhylidae, nearly half of all species richness belongs in the Asterophryinae. The majority of this richness is concentrated in and around New Guinea, however there is also an early branching clade (not included in our study) spread across Vietnam and Thailand (*Siamophryne, Vietnamophryne*), peninsular Malaysia (*Gastrophrynoides*), and on Borneo (*Gastrophrynoides*) (Kurabayashi et al. 2011; Suwannapoom et al. 2018; Poyarkov et al. 2018). Given that the closest relatives of the Asterophryinae are likely the Microhylinae of Asia (and inexplicably the Dyscophinae of Madagascar), it seems plausible that Asterophryines dispersed via a stepping stone procession from southeast Asia, crossing Wallace’s Line just once (see Fig.1 of Poyarkov et al. 2018). Continued dispersal among island groups within Malesia and expansion across New Guinea have likely continued to drive species richness. As it currently stands, there are more than 250 described species found on New Guinea, across a diversity of habitats. This naturally has led to the idea that ecological opportunity has strongly contributed to asterophryine ecomorphological diversification (Rivera et al. 2017; Hill et al. 2022). Ecological and resulting morphological transitions are observable at deep timescales among genera (e.g. arboreal *Oreophryne,* fossorial *Xenorhina*) but also at shallow timescales (e.g., the arboreal *Xenorhina arboricola*, in an otherwise terrestrial genus *Xenorhina*). This process is ongoing, and species discovery rates in the Asterophryinae remain remarkably high (Ferreira et al. 2024; Ferreira et al. 2025). Here, we show that this incredible diversity has arisen within the last 20 million years.

The explosive diversification of New Guinean microhylids has resulted in relationships among genera that are poorly resolved, and further undermined by unreliable generic diagnoses. The vast majority of Asterophryinae species have been described by morphological investigation, and without a molecular understanding. Typically, species are assigned to genera based on suites of anatomical features, including osteological characters. However, these too may be more variable than anticipated. This is highlighted by the discovery by Günther et al. (2023) that two *Cophixalus* species possess procoracoids, lack of these elements was previously considered diagnostic of the genus. Here we show (Fig. 3) that standard linear measurements—which contribute to generic and species diagnoses and help to assign species to genera—often fail to accurately characterize genera. It is clear that some genera fill highly divergent and identifiable morphospaces, such as *Barygenys*, *Callulops*, and *Paedophryne*. Many others overlap considerably, complicating easy generic assignments. This is particularly the case for morphologically variable groups, such as *Cophixalus,* *Oreophryne*, and *Sphenophryne*. Others like *Austrochaperina* and *Copiula* are morphologically conservative but are not easily differentiated from one another. The result is a perfect storm where taxonomic uncertainties persist, muddled further by variable morphologies, and limited molecular phylogenetic information (sampling of both species and genetic loci).

While our data support prior studies in identifying a clade comprising *Asterophrys*, *Callulops*, *Hylophorbus*, *Mantophryne*, and *Xenorhina*, the number and position of remaining genera are hard to determine. Our phylogenomic perspective nests *Aphantophryne* among *Cophixalus*, and together confidently associates them with *Choerophryne* and *Oreophryne* (but with limited species sampling in these two genera). Other assessments have identified more than one major clade of *Oreophryne* (Rivera et al. 2017; Hill et al. 2022), however our sampling lacks a representative of the *Oreophryne* ‘B’ clade. The assignment of species to *Austrochaperina*, *Copiula*, *Sphenophryne* and associated genera, however, remain inconsistent and clearly unresolved. While our topology is mostly well supported, it is important to note that our sampling is incomplete, leaving a thorough taxonomic reassessment out of reach. It is tempting to resolve the situation by lumping these genera all into *Asterophrys* following Dubois et al. (2020), however, we agree with Hill et al. (2022) and Frost (2023) in that this is not the preferred solution. Further phylogenomic sampling paired with a careful morphological assessment will be necessary to resolve outstanding Asterophryinae taxonomic issues.



**Figure 3**. Asterophryinae morphospace shows considerable overlap among genera, with a small number of highly divergent groups (*Barygenys*, *Callulops*, *Paedophryne*). This morphospace, approximated by linear measurements, does not allow for a neat assignment of species to genera through traditional discriminant means (machine learning, linear or flexible discrimination). We suggest this underlies why some genera remain difficult to diagnose and species assignments are unstable. (Left) PCA of first two principal components which together explain ~90% of the variation. Diversity is primarily described by size and the diameter of the toe pad on the third finger, but there are obvious other morphological axes of diversity as seen by the species silhouettes. (Right) the same PCA, but highlighting the distribution of genera within this space.

The southwest Pacific is a region of complex geological and biogeographic history (Wallace, 1869; Hall, 1997). This area sits at the confluence of several continental plates that have been undergoing a complicated dance of movement and compression over millions of years. As a result, thousands of islands have emerged and subsided, including one of the world’s largest, New Guinea. The island of New Guinea itself is a composite formed by the amalgamation of several geologic terranes which accreted roughly around the same time as the explosive diversification of the Asterophryinae, ~20 million years ago (Davies 2012; Gold et al. 2020). This suggests that the interplay of New Guinea’s foundational geologic units were influential in the radiation of asterophryine microhylids (Hill et al. 2023). We stress, however, that complex biogeographic scenarios, such as presented in Hill et al. (2023), must be confirmed in the presence of new phylogenetic evidence that may impact the evolutionary narrative.

Regardless of the order of movements among New Guinea’s biogeographic regions, it is evident that microhylids are capable dispersers, and our Australian-focused sampling adds a new layer to their history. We find evidence of two independent dispersals into Australia, one each for the genera *Austrochaperina* and *Cophixalus*. One taxon, *A. gracilipes* occurs on both landmasses but sits in the Australian clade of species, suggesting a dispersal back to New Guinea (possibly during known land connections in the Pleistocene; Lewis et al. 2013). While the southern portion of New Guinea (the Australian Craton) has long been attached to the Australian continental plate, deeper time land connections (i.e., over millions of years) are poorly understood. Temporally consistent dispersal events from New Guinea to Australia by both *Austrochaperina* and *Cophixalus* suggest an elevated dispersal likelihood ~10 million years ago. This was possibly the result of substantially lower sea levels which facilitated expansion not only into Cape York, but also the Top End of northern Australia (Fig. 1) (Miller et al. 2020). Our dating for microhylids is broadly consistent with dating of faunal interchange between other vertebrate groups between the two regions (e.g., marsupials, Mitchell et al. 2014) Once present in Australia, subsequent diversification of both *Austrochaperina* and *Cophixalus* has been centered in the Wet Tropics of Far North Queensland (Fig. 1). This wet and topographically complex landscape has long been a source of endemic diversification, likely a result of persistent rainforest habitats and sufficient refugia during periods of climatic volatility (Martin 2006; VanDerWal et al. 2009).

Our phylogenomic perspective on the Asterophryinae provides much needed resolution to many parts of the evolutionary tree, such as the timing and position of Australian species. Despite our progress, many questions remain for this species-rich and enigmatic group. Further molecular sampling will undoubtedly help us to explore the complex biogeographic history of microhylids and unravel the amazing diversification of Asterophryinae on New Guinea.

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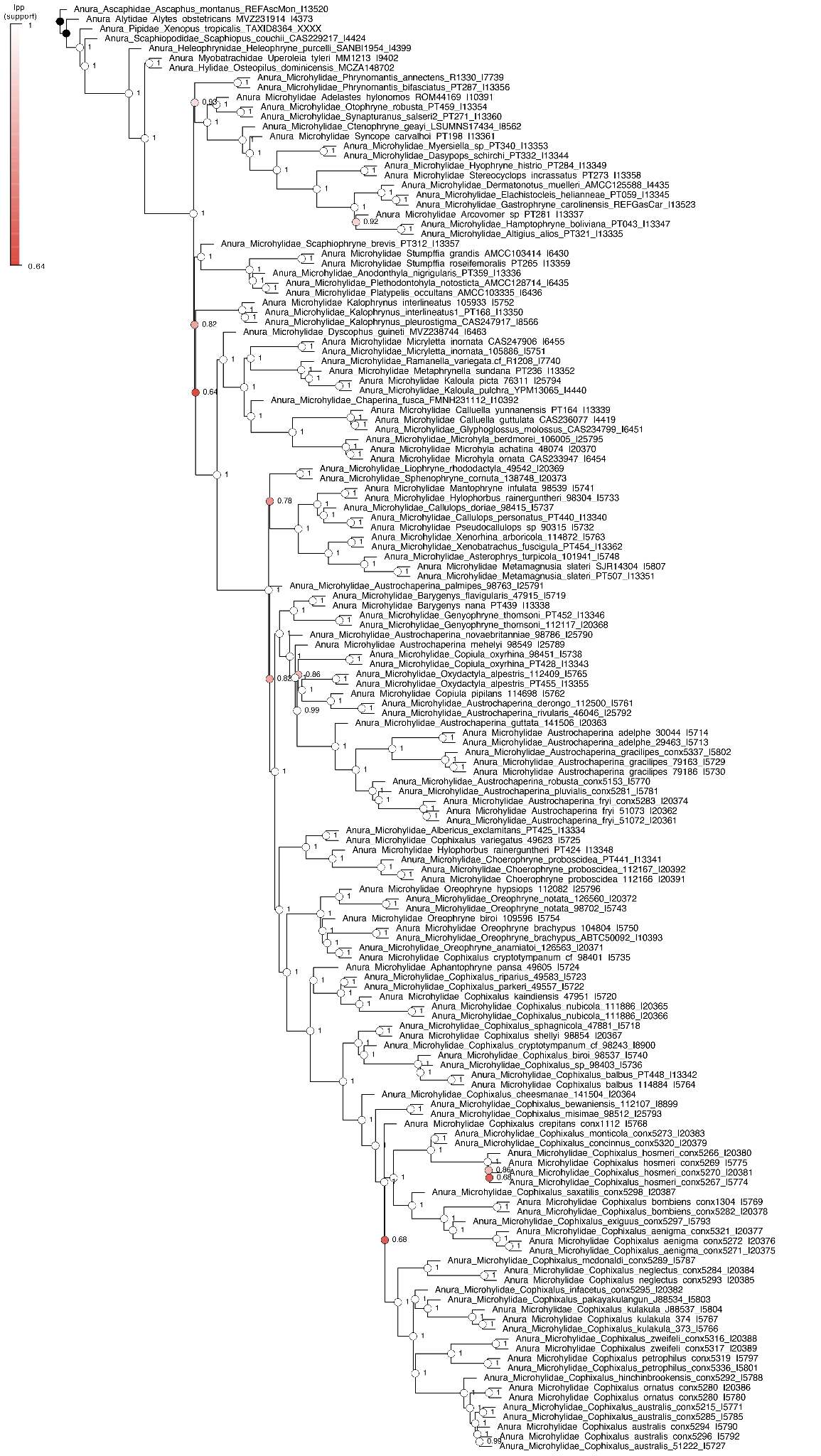
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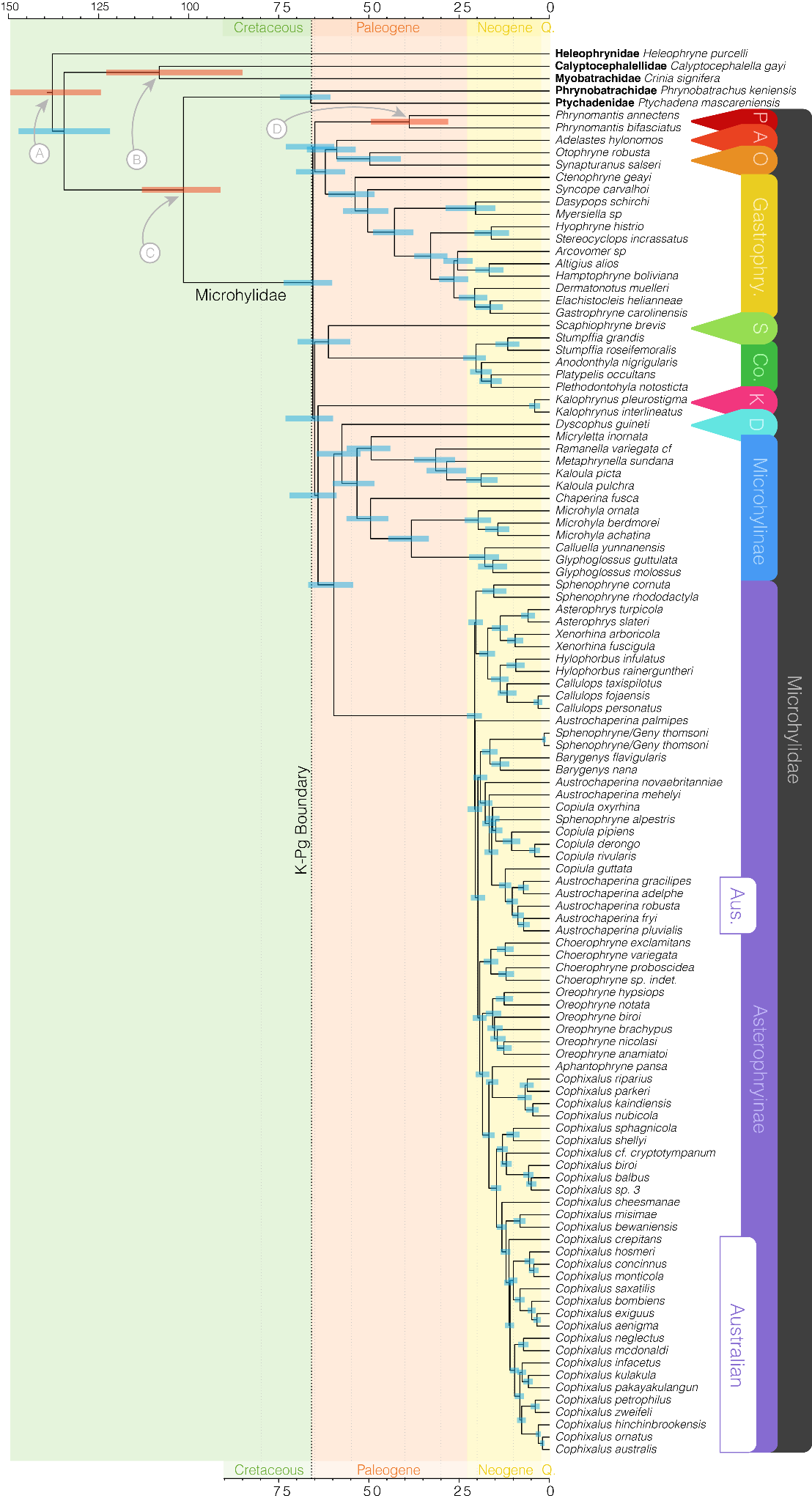
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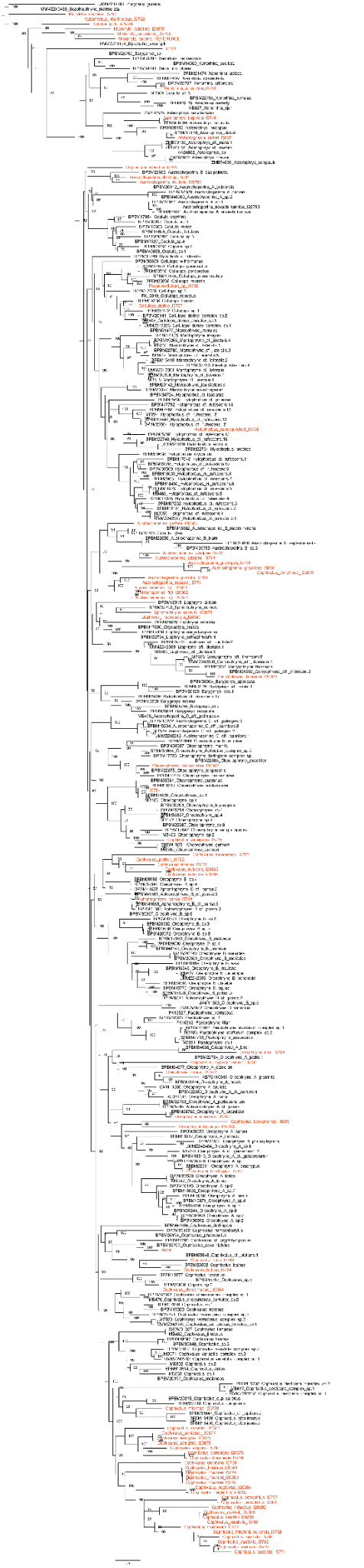
# Supplement



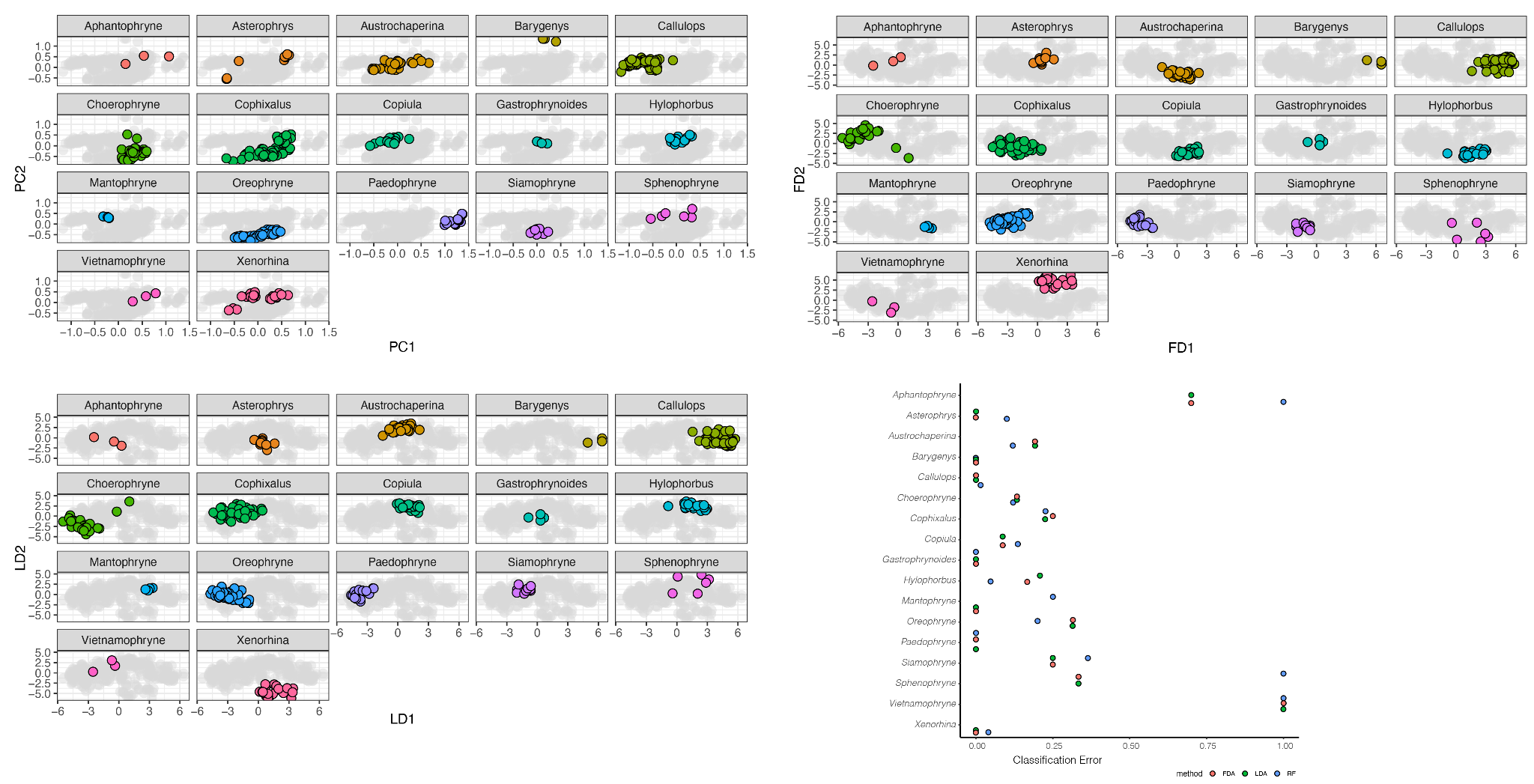
**Figure S1.** Fully sampled sequence capture tree, estimated by weighted hybrid ASTRAL using IQTREE genetree inputs. Branch support values are indicated at nodes and colored according to value (white = 1; red <= 0.9).



**Figure S2.** Time calibrated species tree of microhylid frogs highlights with confidence intervals indicated at nodes. Orange colored bars annotated with a circle indicate nodes calibrated by fossil evidence. These correspond to (A) *Beelzebufo ampinga* as a 66 million year minimum on the crown divergence of Neobatrachia; (B) *Calyptocephalella pichileufensis* as a 47.5 million year minimum on the split between Calyptocephalellidae and Myobatrachoidea; (C) *Thamatosaurus gezei* as a 33.9 million year minimum on the crown of Ranoidea; and (D) Ptychadenidae fossil as a 25 million year minimum on the split of Ptychadenidae and Phrynobatrachidae.



**Figure S3.** IQTREE gene tree of concatented mitochondrial loci (CYTB, ND4). Newly assembled and placed samples are indicated by orange text. Incorporating new samples into the alignments of Hill et al. (2023) allows for a shared understanding of Asterophryinae taxonomy between mitochondrial and nuclear datasets.



**Figure S4.** Visualizations of morphological dimensionality reduction and classification error under three discrimination techniques (RF, LDA, FDA) are largely consistent.