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Running head: PHYLOGENY, TIME, AND BIOGEOGRAPHY OF PLETHODONTIDS

Title: Enlarged Multilocus Dataset Provides Surprisingly Younger Time of Origin for the Plethodontidae, the Largest Family of Salamanders

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Abstract

Deep phylogenetic relationships of the largest salamander family Plethodontidae have been difficult to resolve, probably reflecting a rapid diversification early in their evolutionary history. Here, data from 50 independent nuclear markers (total 48,582 bp) are used to reconstruct the phylogeny and divergence times for plethodontid salamanders, using both concatenation and coalescence-based species-tree analyses. Our results robustly resolve the position of the enigmatic eastern North American Four-toed Salamander (Hemidactylium) as the sister taxon of Batrachoseps + Tribe Bolitoglossini, thus settling a longstanding question. Furthermore, we statistically reject sister taxon status of Karsenia and Hydromantes, the only plethodontids to occur outside the Americas, leading us to new biogeographic hypotheses. Contrary to previous longstanding arguments that plethodontid salamanders are an old lineage originating in the Cretaceous (more than 90 Ma), our analyses lead to the hypothesis that these salamanders are much younger, arising close to the K-T boundary (~66 Ma). These time estimates are highly stable using alternative calibration schemes and dating methods. Our data simulation highlights the potential risk of making strong arguments about phylogenetic timing based on inferences from a handful of nuclear genes, a common practice. Based on the newly obtained timetree and ancestral area reconstruction results, we argue that (i) the classic “Out of Appalachia” hypothesis of plethodontid origins is problematic; (ii) the common ancestor of extant plethodontids may have originated in northwestern North America in the early Paleocene; (iii) origins of Eurasian plethodontids likely result from two separate dispersal events from
western North America via Beringia in the late Eocene (~ 42 Ma) and the early Miocene (~ 23 Ma), respectively.

**Keywords:** phylogenomics, timetree, molecular dating, species tree, paleogeography, dispersal
Lungless salamanders (Plethodontidae) are the most successful radiation in the long evolutionary history of caudate amphibians, comprising more than two thirds of all living salamander species (AmphibiaWeb 2015). They occupy a surprisingly broad series of ecological niches, from temperate woodlands to deserts and tropical rainforests, and exhibit great diversity in life history, morphology, and ecology. Notably, plethodontids are the only salamanders with an extensive presence in tropical regions. Trait evolution in plethodontids is characterized by pervasive homoplasy (Wake 1966, 1991; Mueller et al. 2004), thus providing an important model system to study macroevolutionary patterns (Wake 2009). To understand evolutionary processes, a robust, reliable phylogeny is essential. However, despite recent molecular studies that have built an ever-growing database, a general consensus on the phylogenetic relationships among the major plethodontid lineages is lacking. For example, analyses based on a few nuclear genes (Min et al. 2005; Vieites et al. 2007), whole mtDNA genomes (Mueller et al. 2004) and combined datasets (Chippindale et al. 2004; Kozak et al. 2009; Pyron and Wiens 2011; Vieites et al. 2011) have yielded conflicting results in placing several key genera, especially *Hemidactylium, Aneides, Ensatina, Hydromantes* and *Karsenia*. This uncertainty about relationships has hindered our understanding of the macroevolutionary processes that underlie plethodontid diversity.

Plethodontids are distributed in a curious and unique, highly disjunct and asymmetric manner in the Holarctic region, with 98% of the species in America, a few in the central Mediterranean portion of Europe, and a sole species (*Karsenia koreana*).
on the Korean peninsula (Fig. 1). When and where plethodontids arose, and how and when they achieved their present distribution, remain compelling questions. For example, Appalachia was long thought to be the site of origin of plethodontids, until it was brought into question (Ruben and Boucot 1989; Mueller et al. 2004). Furthermore, the timing and route of colonization of Eurasia by plethodontids has long been a biogeographic enigma (reviewed by Wake 2013). Historically, North America has been connected to Eurasia by the relatively ephemeral Paleocene–Eocene North Atlantic land bridge (NALB) and the long-lasting Bering land bridge (Beringia), so Eurasian plethodontids might have used either route, (NALB: Lanza and Vanni 1981; Delfino et al. 2005. Beringia: Wake et al. 1978; Vieites et al. 2007; Wake 2013. Both: Lanza et al. 2005). However, because the NALB was completely broken after the Late Eocene (~39 Ma), Eurasian plethodontids could not use this route if they originated too late. Reliable estimation of relationships and splitting times are key elements in tackling such questions.

Because early plethodontid fossils are rare, our understanding of the timing of their diversification is based mainly on molecular data. However, recent dating analyses using molecular data have yielded conflicting results. For example, analyses based on whole mtDNA genome sequences (Mueller 2006; Zhang and Wake 2009) suggested that the most recent common ancestor of living plethodontids occurred 98–129 Ma; however, molecular inferences based on a few nuclear genes (Roelants et al. 2007; Vieites et al. 2007) supported a younger origin of 81–94 Ma. Recent empirical studies have consistently shown that slowly evolving markers such as
nuclear exons surpass mitochondrial data for dating a phylogeny covering a relatively wide time span (Zheng et al. 2011; Near et al. 2012). However, because nuclear exons are relatively conservative and contain fewer signal sites, more loci should be used to calculate reliable divergence times (Zheng et al. 2011).

In this study, we generated a large multilocus dataset of 50 unlinked nuclear protein-coding genes for 25 plethodontid species and 13 outgroup species, covering all major lineages of Plethodontidae. Our goal was to test previously proposed hypotheses on phylogenetic relationships, as well as to estimate divergence times of the main lineages in the family by large-scale data analyses. Our analyses generated new hypotheses of phylogenetic relationships and divergence timing, and provide new perspectives on the evolutionary history and historical biogeography of plethodontid salamanders.

**METHODS**

*Taxon and Gene Sampling, and Laboratory Protocols*

We sampled 25 plethodontid species representing all Holarctic plethodontid genera and all nine plethodontid tribes (Wake 2012). Currently Neotropical plethodontids are allocated to 12 genera (AmphibiaWeb 2014) and the monophyly of this group is unquestioned (Mueller et al. 2004; Pyron and Wiens, 2011; Vieites et al. 2007; Vieites et al. 2011). Our sampling of Neotropical plethodontids is limited to three genera (*Thorius*, *Bolitoglossa* and *Pseudoeurycea*), which span the entire tree of this group (see Pyron and Wiens 2011). Six additional salamanders, two frogs, and
four amniotes are included to provide a backbone phylogeny and calibration points for molecular dating analyses. African lungfish was used as an outgroup.

Total genomic DNA was extracted from ethanol-preserved tissues (liver or muscle) using a standard salt extraction protocol (Aljanabi and Martinez 1997). All extracted genomic DNA was quantified using the Nanodrop 2000 (Thermo Scientific) and diluted to a concentration of 50 ng /µL. PCR primers used to amplify the 50 nuclear genes were from a recently developed nuclear marker set (Shen et al. 2013). All of the 50 target regions (see Table 1) are nuclear protein-coding exons. Of the target 1900 sequences, 468 are available in public databases (NCBI and UCSC); 1432 sequences were generated de novo. Nuclear genes were amplified with a nested PCR strategy, as described by Shen et al. (2013) and bidirectionally sequenced by Sanger sequencing. All newly obtained sequences were examined by checking for the presence of premature stop codons (pseudogene) and frame-shifting mutations. Detailed information of species, nuclear markers, and GenBank accession numbers of newly generated sequences are given in Supplementary Table S1.

Sequence Alignment, Partition Strategy and Substitution Model Selection

Each nuclear gene was aligned using the program PRANK v.130410 (Löytynoja and Goldman 2008) according to their translated amino acid sequences (-translate). Ambiguous alignment regions were trimmed by using Gblocks 0.91b (Castresana 2000) with type of sequences set to codons (-t=c) and half gaps allowed (-b5=h); otherwise, default settings were assumed. It should be mentioned here that using
GBlocks (or any other alignment-filtering method) may worsen single-gene phylogenetic inference (Tan et al. 2015). However, this argument is still contentious and beyond the scope of our study. Therefore, we still used Gblocks for automated alignment refinement. All 50 refined alignments were combined into a concatenated dataset (48,582 bp) and we manually defined five different partitioning schemes: 1 partition (unpartitioned); 3 partitions (a separate partition for all first, second, and third codon positions); 50 partitions (one partition for each gene); 100 partitions (a separate partition for the first and second codon positions together in each gene and a partition for the third codon position in each gene); and 150 partitions (a separate partition for each codon position in each gene). Comparisons of the five partitioning strategies and selections of corresponding nucleotide substitution models were conducted under the Bayesian information criterion (BIC) implemented in PartitionFinder v1.0.1 (Lanfear et al. 2012). The 3-partition scheme was chosen as the best-fitting partitioning strategy, and all 3 partitions favored the GTR+Γ+I model. The data matrices and resulting trees were deposited in TreeBase (http://purl.org/phylo/treebase/phylows/study/TB2:S17263).

**Phylogenetic Analyses**

The concatenated dataset was separately analyzed with both maximum likelihood (ML) and Bayesian inference (BI) methods using 3 codon position partitioning. Partitioned ML analyses were implemented using RAxML version 7.2.8 (Stamatakis 2006), with the GTR+Γ+I model assigned to each partition. Supports for nodes were
assessed with 500 rapid bootstrapping replicates. The partitioned Bayesian inference was implemented in MrBayes 3.2 (Ronquist et al. 2012). All model parameters were unlinked. Two MCMC runs were performed with one cold chain and three heated chains (temperature set to 0.1) for 60 million generations and sampled every 1000 generations. The chain stationarity was checked by plotting $-\ln L$ against the generation number in Tracer version 1.4 (Rambaut and Drummond 2007). The effective sample sizes (ESS) are greater than 200 for all parameters after the first 20% of generations were discarded. Topologies and posterior probabilities were estimated from the remaining generations. Two runs for each analysis were compared for congruence.

Species tree analysis without gene concatenation was performed using the pseudo-maximum likelihood approach implemented in the program MP-EST v1.5 (Liu et al. 2010) under the coalescent model. First, for each nuclear gene, 500 bootstrapping gene trees were constructed using PHYML 3.0 (Guindon et al. 2010) under its best-fitting model, which is selected under the Bayesian information criterion (BIC) implemented in jModelTest v2.1.7 (Darriba et al. 2012). Second, the gene trees obtained for the 50 nuclear genes were rooted with the lungfish (Protopterus annectens) and supplied to the MP-EST to generate 500 bootstrapping species trees. The consensus and robustness of the species tree were evaluated with the 500 bootstrapping replicates.

Likelihood-based tests of alternative phylogenetic hypotheses were assessed based on the concatenated dataset. Site-wise log-likelihoods of all alternative
hypotheses (see Table 2) were first calculated with RAxML under the GTR+Γ+I model using the option (-f g). Then, the site log-likelihood file was supplied to the CONSEL program (Shimodaira and Hasegawa 2001) to estimate P-values for each alternative hypothesis using the approximately unbiased (AU) test, the Kishino-Hasegawa (KH) test, and the bootstrap probability (BP).

**Divergence Time Estimation**

No consensus prevails concerning which of many dating procedures using molecular data is the most accurate in time estimation. Accordingly, we used three popular Bayesian relaxed-clock dating programs, MultiDivTime (Thorne and Kishino 2002), MCMCTREE (dos Reis et al. 2014) and BEAST (Drummond and Rambaut 2007) to estimate divergence times. The reference topology was that derived from the ML analysis. All molecular dating analyses were performed with three codon partitions, as in the phylogenetic analyses.

Our analysis began by using three different calibration schemes used by previous studies (Mueller 2006; Vieites et al. 2007; Zheng et al. 2011) in the MultiDivTime analysis (see Supplementary Table S2 for detailed). Because alternative calibration schemes have little impact on divergence time estimation within living salamanders (Supplementary Fig. S1), we chose the calibration scheme with eleven time constraints used by Vieites et al. (2007) for our definitive molecular dating analyses. Three widely tested nodes were constrained with lower and upper bounds: the Amniota - Amphibia split (332–360 Ma; Benton and Donoghue 2007), the Mammalia
Aves split (312–330 Ma; Benton and Donoghue 2007), and the Aves - Crocodyla split (243–251 Ma; Benton and Donoghue 2007). Five less defined nodes were constrained with lower bounds only, based on known fossil records: the split between Aves and Squamata at least 252 Ma (*Protorosaurus speneri*; Evans and King 1993), the split between Anura and Caudata at least 230 Ma (*Triadobatrachus massinoti*; Rage and Rocek 1989), the split between Hynobiidae and Cryptobranchidae at least 155 Ma (*Chunerpeton tianyiensis*; Gao and Shubin 2003), the split between Amphiumidae and Plethodontidae at least 65.5 Ma (*Proamphiuma cretacea*; Gardner 2003), the split between eastern and western *Plethodon* clades at least 25 Ma (Tihen and Wake 1981).

For MultiDivTime analysis, lungfish (*Protopterus annectens*) was used as the outgroup and the Amniota - Amphibia split was regarded as the ingroup root. For each codon partition, the model parameters were calculated with an F84+Γ model using the BASEML program in PAML4.8 package (Yang 2007), and optimized branch lengths with their variance–covariance matrices of the DNA data set were estimated by the program Estbranches_dna, a component of MultiDivTime. The priors for the mean (rttm) and standard deviation (rttmsd) of the ingroup root age were set to 3.44 and 0.15 time units (one time unit = 100 Ma), respectively. The prior mean (rtrate) and standard deviation (rtratesd) for the gamma distribution describing the rate at the root node were both set to 0.089. These values were based on the median of the substitution path lengths between the root and each terminal, divided by rttm (as suggested by the authors). The autocorrelation parameter prior (brownmean) and its
SD (brownsd) were set to 0.58, such that brownmean multiplied by the rttm prior (3.44) equals 2.0. After an initial burn-in period of 100,000 cycles, MCMC chains were run for 3,000,000 cycles, with sampling intervals of every 100 cycles. Two independent runs were performed to examine whether similar results were observed.

For MCMCTREE analysis, we used the latest MCMCTREE in the PAML4.8 package which uses a new prior (gamma-Dirichlet prior) to describe substitution rates across multiple loci, thereby improving the accuracy of divergence time estimation (dos Reis et al. 2014). The approximate likelihood method was used for divergence time estimation (see MCMCTREE tutorial). The ML estimates of branch lengths for the three codon partitions were obtained using BASEML (in PAML) programs under the GTR + Γ model. For the gamma-Dirichlet prior for the overall substitution rate (rgene gamma), we used a quite diffuse (uninformative) prior $\alpha_{\mu} = 1$. Based on the mean estimate from three codon partitions using the strict molecular clock assuming 346 Ma constraint at the root, an average of the Amniota - Amphibia split (332–360 Ma; Benton and Donoghue 2007), the gamma-Dirichlet prior for the overall substitution rate (rgene gamma) was set at $G(1, 11.96, 1)$. The gamma-Dirichlet prior for the rate-drift parameter (sigma2 gamma) was set at $G(1, 4.5, 1)$. All eleven calibration constraints were not rigorously constrained (specified with 2.5% tail probabilities above or below their limits; this is a built-in function of MCMCTREE). The independent-rate and autocorrelated-rate models (clock=2 and clock=3 in MCMCTREE) were separately used to estimate divergence time. For each model, after a burn-in period of 2,000,000 cycles, the MCMC run was sampled every 400
cycles until a total of 10,000 samples was collected. Two separate MCMC runs were compared for convergence with two different random seeds and similar results were observed.

For BEAST analysis, we assigned a lognormal relaxed clock model (uncorrelated), a GTR+Γ+I site model, and a Yule tree prior for each of the three codon partitions. For calibration points with both minimum and maximum bounds, we used lognormal distributions to describe the priors of those calibration points with maximum boundaries, representing a “soft” calibration strategy. The means and standard deviations of the lognormal distribution for each calibration point were chosen so that 95% of the probability lies within the minimum bound and the maximum bound and the means are the arithmetical medians of the intervals. For the calibration points with only minimum bounds, the means and standard deviations were set so that the lognormal distributions have 2.5% of chance beyond the minimum bound. Two independent runs were performed for 300 million generations, with sampling every 6,000 generations. The convergence of the run was confirmed in Tracer version 1.4 (Rambaut and Drummond 2007) because effective sampling sizes (ESS) for all parameters are all >200. The two runs reached similar likelihood platforms and results of the two runs were summarized after discarding the first 25% generations.

**Historical biogeography reconstruction**

To test the likelihood of different ancestral distribution scenarios for plethodontid
salamanders, ML inference of the evolution of geographic ranges was explored with Lagrange v2.0 (Ree and Smith 2008). We used the plethodontid portion of the time tree generated by our molecular dating analyses as the input time-calibrated phylogeny, which included 25 in-group plethodontid species and two outgroups (Amphiumidae and Rhyacotritonidae). Based on the current distribution pattern of plethodontids (Fig. 1), we defined five possible ancestral distribution areas: western North America (WN), eastern North America (EN), eastern Eurasia (EE), western Eurasia (WE), and tropical region (TP). Although Amphiumidae is currently endemic to southeastern North America, its fossils are found across North America and its oldest fossils are all from western North America (reviewed by Bonett et al. 2014). Therefore, we set up two independent analyses that assume Amphiumidae has an ancestral distribution of western North America or an ancestral distribution across all of North America, respectively.

Paleogeographic connections between these areas were gathered from the literature. WN is continuously connected to EN except that an epicontinental seaway separated them during the Cretaceous (110-70 Ma) (Blakey 2011). TP was continuously connected to WN and EN but its connection to EN was blocked during the epicontinental seaway period (110-70 Ma) (Blakey 2011). The Beringian land bridge was opening and closing periodically during the Mesozoic and Cenozoic (Tiffney and Manchester 2001), so the connection between WN and EE was also continuous during these periods. The Turgai Sea separated EE from WE from the Middle Jurassic to the Oligocene (160-29 Ma) (Briggs 1995), beyond that, they
connected continuously. The opening of the North Atlantic began in the Late Cretaceous (90 Ma) but terrestrial connections between WE and EN persisted along various North Atlantic land bridges until at least the Late Eocene (39 Ma) (Tiffney 1985). Combining the above information, we set up the probability of dispersal success between two areas to one during the time period they connected, and to zero outside the time period.

For some nodes, except the most probable scenario, Lagrange also gave some alternative biogeographic scenarios with lower likelihood values. In this case, the statistical significance of likelihood differences between biogeographic scenarios was assessed using the conventional cutoff value of two log-likelihood units. Finally, we calculated the relative probability (fraction of the global likelihood) of a certain reported scenario.

**Investigating the Effect of Gene Number on Divergence Time Estimation**

To investigate the effect of gene number on plethodontid divergence time estimation, we generated data sets with different gene numbers by randomly sampling genes from the whole set of 50 nuclear genes. We generated 200 replicates for each of eight data points with gene number of 3, 6, 12, 24, 50, 100, 150, and 200, respectively (a total number of 1600 data sets). For each data set, divergence times were estimated by MCMCTREE using the independent rate model (clock=2) and the codon partitioning strategy. All other settings, including calibration choices, follow the MCMCTREE analysis procedure as described before. For the 200 replicates of each
data point, the mean divergence times for four representative nodes within the plethodontid timetree (the origin of Plethodontidae, the origin of Hemidactyliinae, the origin of Plethodontinae and the split between *Hemidactylium* and *Bolitoglossa*) were showed as boxplots.

**RESULTS**

*Data and Phylogenetic Analyses*

The supermatrix of 50 combined nuclear genes is 98.5% complete for the 38 taxa and 98.7% complete for the 25 plethodontid taxa (Supplementary Table S1). The aligned lengths of 50 nuclear genes ranged from 540 to 1500 bp (mean = 987). No significant compositional heterogeneity among sequences was detected for any genes among the 25 sampled plethodontid species (Table 1). The combined 50-gene dataset comprised 48,582 base pairs (bp) and displays no apparent substitution saturation within the 31 included salamander species (Supplementary Fig. S2).

Maximum Likelihood and Bayesian analyses of the concatenated data set and species tree inference without data concatenation produced similar topologies for the 25 plethodontid species (Fig. 2). The three analyses all find strong support for the view that plethodontids divided into two major clades (Plethodontinae and Hemidactyliinae), as recovered by most recent molecular studies (Chippindale et al. 2004; Mueller et al. 2004; Frost et al. 2006; Vieites et al. 2007; Kozak et al. 2009; Pyron and Wiens 2011; Vieites et al. 2011).

Within the subfamily Hemidactyliinae, the genus-level cladistic structure is well
resolved (bootstrap support > 95% or PP = 1.0; Fig. 2). The monotypic genus

*Hemidactylium* is robustly resolved as the sister taxon of *Batrachoseps +

*Bolitoglossini* (Fig. 2). Within the subfamily Plethodontinae, cladistic structure of the
six genera studied is not robustly resolved. However, an *Aneides + Desmognathus*
clade is strongly supported by ML and Bayesian analyses (Fig. 2A and 2B). Otherwise,
phylogenetic relationships among the remaining four genera are unclear (Fig. 2). The
positions of two monotypic genera, *Ensatina* and *Karsenia*, are peculiarly unstable
among the three analyses (Fig. 2). Excluding *Ensatina* and *Karsenia*, respectively,
from the analyses (Supplementary Fig. S3 and S4) did not provide further resolution
for placing these problematic genera. The internal branches separating these genera
are extremely short (close to zero), compared to branches elsewhere in the family,
suggesting that these ancient cladistic events occurred almost simultaneously.

**Divergence Times**

In this study, we used four well-known relaxed-clock dating algorithms,

MultiDivTime (correlated-rate model), BEAST (independent-rate model) and

MCMCTREE (both independent-rate and correlated-rate models), to calculate
divergence times for plethodontids. Because the average time deviation among the
four methods is small (~6.4%; see also Table 3), we choose to focus on the results
from the MCMCTREE (the independent-rate model; clock = 2) (its time estimates
center within the four results) as our preferred dating results. According to the
MCMCTREE time estimates, the two major clades of plethodontids began to split at
66.1 (57.8-74.9) Ma (Fig. 3A and Table 3), close to the KT-boundary (~ 65.5 Ma).

This time estimate is 20% - 50% younger than most previous estimates (at least 80
Ma; e.g., Mueller 2006; Roelants et al. 2007; Vieites et al. 2007; Zheng et al. 2011).
The initial diversification of Hemidactyliinae and Plethodontinae took place at 58.6
(51.2-66.6) Ma and 43.1 (37.5-48.7) Ma, respectively. The divergence among major
clades of Plethodontinae occurred almost simultaneously within a short time window
(~ 3 million years; Fig. 3A and Table 3).

Ancestral Area Reconstruction

By using the program Lagrange v2.0 (Ree and Smith 2008) and assuming
Amphiumidae has an ancestral distribution of western North America or an ancestral
distribution across North America, we calculated relative probabilities of possible
ancestral areas for Plethodontidae, Karsenia, all Hydromantes, and European
Hydromantes, respectively.

When assuming Amphiumidae has an ancestral distribution across North America,
eastern North America (EN) is the most probable ancestral area for Plethodontidae
(rel. prob. = 0.458; Fig. 3B), but western North America (WN) is the second probable
ancestral area for Plethodontidae (rel. prob. = 0.250; Fig. 3B). However, if assuming
Amphiumidae has an ancestral distribution of western North America, western North
America (WN) is the most probable ancestral area for Plethodontidae (rel. prob. =
0.557; Fig. 3B). Under this assumption, eastern North America (EN) becomes the
least probable ancestral area for Plethodontidae (rel. prob. = 0.110; Fig. 3B),
considerably less likely than the WN hypothesis.

For the Korean *Karsenia*, the most probable ancestral area is western North America (rel. prob. ~ 0.8; Fig. 3B) rather than eastern Eurasia (rel. prob. < 0.2; Fig. 3B). *Hydromantes* may have originated in western North America (rel. prob. > 0.9; Fig. 3B) and the ancestor of European *Hydromantes* most likely lived in eastern Eurasia (rel. prob. > 0.9; Fig. 3B).

**DISCUSSION**

**Phylogeny of Plethodontids**

The placement of the enigmatic four-toed salamanders of the monotypic genus *Hemidactylium* is one of the most compelling questions in plethodontid phylogenetics. It has been suggested to be the sister taxon to all other plethodontids (Macey 2005), to *Batrachocephos* (Mueller et al. 2004), to all other hemidactyliine plethodontid salamanders (Wake 1966; Chippindale et al. 2004; Kozak et al. 2009; Pyron and Wiens 2011), and to *Batrachocephos* + Bolitoglossini (Vieites et al. 2007). By assembling all available molecular data at that time (complete mitochondrial genomes and three nuclear genes), Vieites et al. (2011) placed *Hemidactylium* as the closest relatives of *Batrachocephos*, with an alternative but less likely placement as the sister taxon of *Batrachocephos* + Bolitoglossini. These contradictory results may be due to rapid evolution and high heterogeneity of mitochondrial genomes and the fact that mitochondrial genomes have a strong numerical advantage in characters over a few nuclear genes. Our concatenation and species-tree analyses based on 50 nuclear genes
strongly support *Hemidactylium* as the sister taxon to *Batrachoseps* + Bolitoglossini (MLBS = 100%, BPP = 1.00; Fig. 2). This result remained stable when mitochondrial genome data were added to the analyses (Supplementary Fig. S5). Importantly, all other possible hypotheses about the placement of *Hemidactylium* can be rejected by an approximately unbiased test (AU test) (Table 2).

Earlier molecular studies recovered the Asian genus *Karsenia* as the sister group of *Hydromantes*, although without strong support (Vieites et al. 2007; Kozak et al. 2009; Vieites et al. 2011). Contrary to this result, none of our analyses supported such a relationship. Moreover, the AU test statically rejects the grouping of *Karsenia* and *Hydromantes* (*P* = 0.001; Table 2), showing that these taxa have no direct genetic affinity. Although not fully resolved, the degree of resolution in our phylogeny among the rapidly diverging Plethodontinae is nevertheless encouraging, and holds promise that using increasingly large numbers of nuclear genes may lead to better phylogenetic resolution of these challenging deep nodes.

**Younger Time Estimation for Plethodontid Origin and Diversification**

Prior to this study, many attempts had been made to estimate times for the origin and diversification of plethodontids (Mueller 2006; Roelants et al. 2007; Vieites et al. 2007; Wiens 2007; Pyron 2010, 2011; Zheng et al. 2011). Analysis of complete mitochondrial genomes (Mueller 2006) provided estimates for the origin of plethodontids at about 129 (109-152) Ma. As part of a large scale divergence time study of all amphibians, Roelants et al. (2007) estimated the origin time for
plethodontids at 82 (62-103) Ma, based on four nuclear genes and one mtDNA fragment. Using data from three nuclear genes, Vieites et al. (2007) obtained an estimate of 94 (74-117) Ma. By using the three nuclear genes from Vieites et al. (2007) but calibration choices similar to those of Mueller (2006), Zheng et al. (2011) obtained a slightly younger time of 87 (72-100) Ma. Despite differences in time estimates, all four previous studies suggest that plethodontids originated at least 80 million years ago, in the Cretaceous. In addition, there are also some younger estimates for plethodontid origin: ~ 60 Ma (Wiens 2007), and ~ 70 Ma (Pyron 2010). Moreover, by using fossils as terminal taxa, Pyron (2011) proposed a much younger origin (~40 Ma) for plethodontids. However, these younger estimates are based only on a single nuclear locus (RAG1).

We estimate considerably younger times of divergence than the majority opinion (20% - 50% younger); the new estimates are to some degree close to previous young estimates. Furthermore, our time estimates (Fig. 3A and Table 3) coincide with all currently known plethodontid fossil records. For example, the oldest stratum that contains uncontroversial plethodontid fossils is the Cabbage Patch formation dated to ~25 Ma (Tihen and Wake 1981; Personal communication with Jonathan Calede).

Both western Aneides and western Plethodon were found in this layer, which means the western and eastern groups of both Aneides and Plethodon separated from each other before or near that date (that is, ca 25 Ma). Our estimates of divergence between eastern and western Plethodon are 40 Ma (node 24, Fig. 3A and Table 3), and 28.2 Ma (node 18, Fig. 3A and Table 3) between eastern and western Aneides, which fits
the fossil data. In addition, the oldest European *Hydromantes* fossil is known from the Middle Miocene of Slovakia (~14 Ma; Venczel and Sanchíz 2005), but previous time estimates suggested that European *Hydromantes* was at least 40 Ma (Mueller 2006; Vieites et al. 2007). Our results find that European *Hydromantes* arose no earlier than 24 Ma (node 23, Fig. 3A and Table 3), which is more congruent with the fossil record.

But could our young time estimates for plethodontids be a methodological artifact? Molecular time estimates can be affected by three major factors: data, calibration scheme, and dating method. Because the average time deviation among the four relaxed-clock methods used in this study is small (~6.4%), the choice of dating methods should have little effect on our time estimates. With respect to calibration schemes, our analyses show that, when using the same dating program and calibration scheme, our molecular time estimates using the 50-gene dataset largely agree with previous studies for the four older salamander nodes outside plethodontids, but are clearly younger for the four nodes within plethodontids (Fig. 4). This discrepancy remains when we successively applied three different calibration schemes used by previous studies to our dating analyses (Fig. 4). Such a phenomenon is likely because the nodes outside plethodontids are relatively tightly constrained by the working calibration points; however the nodes within plethodontids do not have such constraints. Meanwhile, further analyses suggest that using different calibration schemes has no obvious effect on both mean and 95% CI for all 24 nodes within plethodontids (Supplementary Fig. S1). These findings strongly suggest that our time estimates for plethodontids are robust against dating methods and calibration choices,
and the younger result is mainly caused by the time signals within our data.

**The effect of gene number on Plethodontid Divergence Time Estimation**

We have demonstrated that our young time estimates for plethodontids are mainly caused by the data, in other words, the number of genes used. Considering that we only used a modest number of nuclear genes (50 loci here), one may wonder what we will obtain if we use different numbers of genes. Can we get the same time results when we analyze 100 or even more nuclear genes? To answer these questions, we generated simulated data sets with different gene numbers by randomly sampling genes from the whole set of 50 nuclear genes and repeated our time estimation process (see Methods for details).

Our data simulation shows that when the number of genes goes up from three to two hundred, the median time estimates (from 200 replicates) for the four major plethodontid evolutionary events are all close to what we obtained by using 50 loci (Fig. 5). This result suggests that our dating results are robust against the number of genes used; similar time estimates for plethodontids might well be obtained using 100 or even more nuclear genes. However, the distribution of time estimates becomes considerably wider when fewer and fewer genes are used (Fig. 5). Notably, using only three nuclear genes, the time estimate for plethodontid origin can be from 48 to 106 Ma (Fig. 5). This observation coincides with previous empirical studies that used a few nuclear genes: 82 Ma, four nuclear genes (Roelants et al. 2007); 94 Ma, three nuclear genes (Vieites et al. 2007); 87 Ma, three nuclear genes (Zheng et al. 2011).
Overall, according to our simulation, it is possible to obtain accurate time estimates by using a few genes; however, this practice is prone to random errors because of limited gene sampling.

Our results support the recent argument that analyzing many nuclear loci improves the precision of posterior time estimation (Mulcahy et al. 2012; Zhu et al. 2015). These results not only have important implications for future molecular dating practices for non-model animals, but also raise questions for many molecular dating studies in the past. Previous studies have repeatedly shown that, in the absence of younger effective calibration points, mitochondrial genes tend to overestimate divergence times compared with results from more slowly evolving nuclear genes (Zheng et al. 2011; Near et al. 2012). Therefore, nuclear genes are normally the preferred data source in molecular dating, especially for relatively ancient evolutionary events. However, as indicated by our data simulation, using only a small number of nuclear genes (one to three, as in many recent studies) for molecular dating is risky because of random errors and is likely to produce inaccurate time estimates, especially when the ingroup has a relatively ancient root age but lacks younger calibration points (like the case of plethodontids). This phenomenon was also observed in the molecular dating practices for mammal evolution: the time estimate for *Mus-Rattus* divergence decreases from 23 Ma (3 nuclear genes; Adkins et al. 2001) to 16 Ma (19 nuclear genes; Springer et al. 2003), and eventually to 14 Ma (> 14000 nuclear genes; dos Reis et al. 2012). Because of the historical difficulty in generating large nuclear gene datasets for non-model animals, many molecular dating analyses
are still based on a handful of mitochondrial genes and one to three nuclear genes. These dating practices become highly questionable and many of the time estimates may be far too old. We suggest that any applications of those time estimates (e.g., biogeographic hypotheses) should be interpreted with caution until new analyses using more nuclear loci have confirmed them.

**Origin of Plethodontids and Historical Biogeography**

Plethodontids long were thought to have originated from Appalachia (eastern North America) because loss of lungs was thought to have evolved in mountain streams and this mountain system is old and stable. Furthermore, larvae were thought to be an ancestral state for the family and taxa with larvae are fully represented there. Appalachian plethodontids exhibit high present-day cladistic diversity, adaptive diversity and species richness (Wilder and Dunn 1920; Wake 1966; Beachy and Bruce 1992), further evidence to many that this area is not only the region of origin but also the area in which primary diversification occurred. This idea was further supported by analyses in the context of both geologic history (Beachy and Bruce 1992) and molecular phylogeny (Macey 2005), and thus is widely accepted. However, the “Out of Appalachia” hypothesis was challenged with the discovery of the basal split between two major clades: eastern American Hemidactyliinae and western American Plethodontinae. Mueller et al. (2004) proposed a transcontinental North American origin of plethodontids and Vieites et al. (2007) further inferred that the basal split of plethodontids was caused by a major vicariant event associated with the forming of a
midcontinental seaway in North America from 110 to 70 Ma. These biogeographic inferences are largely based on estimates of a Cretaceous divergence between the eastern Hemidactyliinae and the western Plethodontinae (Mueller 2006; Vieites et al. 2007). Based on our revised time estimate for the basal plethodontid split (~ 66 Ma), the midcontinental seaway may not have been the driving factor because the seaway, while still partially in existence, was closing rapidly at about 65 Ma (Blakey 2011).

The current distribution of plethodontids does not retain any clues on their early biogeographic history because the major clades, Plethodontinae and Hemidactyliinae, both having representatives in eastern and western North America (Fig. 1). Although high species and ecological diversity is observed in the Appalachian Mountains, this pattern can be misleading for inferring center of plethodontid origin if the distributions of the relevant fossils are not considered. In fact, Appalachian fossil plethodontids are no older before Mio-Pliocene (~ 7 Ma; Boardman and Schubert 2011), whereas the oldest fossil plethodontids (*Aneides* sp. and *Plethodon* sp.) were found in western North America (Montana) in late Oligocene (~ 25 Ma; Tihen and Wake 1981). For Amphiumidae (the closest extant relative of plethodontids), although it is currently endemic to southeastern North America, the oldest fossil record of this family was also found in western North America (*Proamphiuma cretacea*; Cretaceous of Montana, ~ 65 Ma). Notably, other American salamander families all have similar patterns: Sirenidae (oldest fossil *Habrosaurus prodilatus* from Cretaceous of Alberta, Canada), Dicamptodontidae (oldest fossil *Dicamptodon antiquus* from Paleocene of Alberta, Canada) and Ambystomatidae (oldest fossil *Ambystoma tiheni* from Eocene
of Saskatchewan, Canada). These fossil records suggest that the most likely ancestral area for these American salamander families (including Amphiumidae) is western North America. The apparent western bias on early fossil distributions of both plethodontids and relevant salamander families suggests that western North America could also be the ancestral area for the origin of plethodontids. In fact, when assuming that Amphiumidae has an ancestral distribution of western North America, our Lagrange analyses (Fig. 3B) did suggest that western North America (WN) is the most probable ancestral area for Plethodontidae (rel. prob. = 0.557; Fig. 3B), and both an eastern North American origin (EN) (rel. prob. = 0.110; Fig. 3B) and an origin across the whole North American (WN+EN) (rel. prob. = 0.269; Fig. 3B) have lower probabilities.

Multiple studies have suggested that salamanders originated in Asia with later expansion into America (Gao and Shubin 2001; Zhang et al. 2005). Given that a very well supported clade Rhyacotritonidae+(Amphiumidae, Plethodontidae) was consistently recovered by all recent studies (e.g., Roelants et al. 2007; Zhang and Wake 2009; Shen et al. 2013), the common ancestor of Rhacotritonidae, Plethodontidae and Amphiumidae most probably dispersed into western North America through Beringia (Fig. 6A). The exclusive western North America distribution for Rhacotritonidae adds more force to this hypothesis. The splits among the three salamander families are dated in the mid-Cretaceous (Table 3), a time when eastern and western North America were separated by a intracontinental seaway (Fig. 6A). Therefore, Rhyacotritonidae, Plethodontidae and Amphiumidae were likely
restricted to western North America until the seaway finally closed at the end of the Cretaceous. Bonett et al. (2013) proposed that amphiumids arose in western North America in association with the evolution of the Western Interior Coastal Plain, as the intracontinental seaway closed. It is likely that rhyacotritonids and plethodontids also arose in montane areas bordering this plain.

From the late Cretaceous to Paleocene, the climate of North America was ~ 10°C warmer than today with palm trees growing in Greenland (Scotese 2001). The climate in southern North America was generally subtropical to tropical, with high temperatures and rainfall, and with small seasonal differences between summer and winter. This warm phase had begun in the Cretaceous period, peaked in the early Eocene, and continued to the end of the Eocene (Zachos et al. 2001). Because salamanders are generally cool-adapted, the common ancestor of plethodontids most likely occurred well to the north, in northwestern North America (Fig. 6A).

Our general hypothesis for the dispersion of plethodontids postulates several steps. After the midcontinental seaway through North America finally closed at the end of the Cretaceous (Blakey 2011), plethodontids began to spread across the continent and adapted to any available ecological niches. This phase lasted from the Paleocene to the end of the Eocene (65 -33 Ma) and produced most major plethodontid lineages. Because southern North America was too hot for salamanders during this period, the major divergences of plethodontids most likely took place in the northern regions of North America (Fig. 6B). As the climate continued to cool in the Oligocene, plethodontids migrated to the south (Fig. 6C). Finally, the increasing
aridity of central North America during the Miocene interrupted the former
continuous distribution of plethodontids, which split into two major clades, the
plethodontines along the western coast area and the hemidactyliines in eastern North
America, centered in Appalachia (Fig. 6D). Previous studies argued that the high
species diversity of Plethodon and Desmognathus in Appalachia corresponds mainly
to recent radiations (Kozak et al. 2005; Wiens et al. 2006). We agree with this
argument and further suggest that the rapid species accumulation for these genera
occurred within the Miocene. Currently, the earliest fossils of plethodontids from
Appalachia, including Plethodon and Desmognathus, date back only to Mio-Pliocene
(Boardman and Schubert 2011).

**Origin of Eurasian Plethodontids and Dispersal Routes**

The current distribution of plethodontids is disjunctive and highly asymmetric,
with 98% of the species in Americas, a few in Europe and a sole species Karsenia
koreana in East Asia. More surprisingly, members of the genus Hydromantes are
distributed in western North America and in Europe and the American species and
European species of the genus are genetically more closely related to each other than
to other plethodontid taxa. How plethodontids attained their present distributions has
long been a biogeographic enigma, given their generally low dispersal capability.

Recent molecular studies suggested, although without strong support, that
Karsenia and Hydromantes are sister taxa and they together constitute the initial
branch of Plethodontinae (Vieites et al. 2007, 2011). If these relationships are real,
Karsenia and Hydromantes may have entered Eurasia separately from North America, or by a more likely biogeographic scenario as proposed by Vieites et al. (2007): their common ancestor entered Asia and underwent diversification there, which separated the two genera and left Karsenia in northeast Asia. Later Hydromantes migrated both into western Europe and back into western North America, and eventually went extinct in Asia. Our phylogenetic analyses based on much more data find no support for the hypothesis that Karsenia and Hydromantes are sister taxa; rather they are embedded in a near polytomy within the subfamily Plethodontinae, which possesses an otherwise entirely North American distribution. In addition, our ancestral distribution reconstruction also suggests that a western North American origin (rel. prob. ~ 0.8; Fig. 3B) is much more possible than an East Asian origin (rel. prob. ~ 0.2; Fig. 3B) for Karsenia. These results make the latter biogeographic scenario proposed by Vieites et al. (2007) highly unlikely, but suggest that the origin of Karsenia may have resulted from a dispersal event out of North America.

Although it is generally assumed that the genus Hydromantes dispersed into Europe from North America, there is no consensus on the timing and dispersal route for the event (Wake et al. 1978; Lanza et al. 1995; Delfino et al. 2005; Vieites et al. 2007; Carranza et al. 2008; Wake 2013; Pyron 2014). Geographically, Europe was connected with North America via the North Atlantic Land Bridge (NALB) but separated from Asia by the Turgai Strait from Paleocene to Eocene; with the breakup of NALB and the closure of the Turgai Strait during the Oligocene, the physical connection between Europe and North America was exclusively by way of the Bering
Land Bridge (Beringia) via the Asian continent (Jones 2011). Therefore, the time for separation between the American and European *Hydromantes* becomes the key: times no later than Eocene offer some support for the NALB as the dispersal route while times after Oligocene will favor Beringia. A single vertebra of *Hydromantes* (diagnosable as *H. (Speleomantes)* sp.) is known from the Miocene of Slovakia, dated 13.75 Ma (±1.25 Ma) (Venczel and Sanchiz 2005), suggesting that the separation between the American and European *Hydromantes* predated this time. Our time estimate for this split is about 23 Ma, congruent with the fossil record. This result also suggests that *Hydromantes* arrived in Europe from North America by crossing Beringia and the entire Asian continent in Miocene. Our biogeographic inferences support this idea because the ancestral distribution area of all *Hydromantes* is exclusively western North America (rel. prob. > 0.9; Fig. 3B), while the ancestral distribution area of European *Hydromantes* is exclusively eastern Eurasia (rel. prob. > 0.9; Fig. 3B).

**SUPPLEMENTARY MATERIALS**

Supplementary material, including data files and figures, can be found in the Dryad data repository (http://datadryad.org, doi:10.5061/dryad.h4qn5).

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**FIGURE CAPTIONS:**

Figure 1. Current distribution of extant plethodontid salamanders. The distribution area is roughly divided into five parts: western North America (blue), eastern North America (green), tropical (yellow), East Asia (purple), and central Europe (red). The locality for related fossil taxa is indicated as a black triangle.

Figure 2. Phylogenetic relationships of plethodontid salamanders inferred from 50 nuclear genes (48,582 bp). The trees were inferred by concatenation analyses using maximum likelihood (A) and Bayesian inference (B) and by species-tree analysis using the MP-EST program (C). Branch support values beside nodes indicate ML bootstrap (A), BI posterior probability (B) and MP-EST bootstrap (C), respectively. The placement of the root of each tree is indicated with an open circle. Outgroup taxa are not shown.

Figure 3. Timetree and ancestral area reconstruction results. (A) Evolutionary timetree inferred by MCMCTREE using an independent rate model (clock=2). Eleven time constraints used in the molecular dating are shown as black triangles. Gray horizontal bars represent 95% credibility intervals. At the top right, a color-coded square represents the five main regions considered: EE (purple), eastern Eurasia; WE (red), western Eurasia; WN (blue), western North America; EN (green), eastern North America; TP (yellow), tropical region. The same color-coding scheme was applied to species names. Detailed divergence time estimates for nodes with numbers are
provided in Table 3. (B) Inferences about the ancestral distribution area of four plethodontid groups calculated by Lagrange v2.0. Two probability values are given for every possible ancestral area, assuming that Amphiumidae has an ancestral distribution of whole North America (left) or an ancestral distribution of western North America (right), respectively.

Figure 4. Comparison of divergence time estimates for eight split events sharing across four studies. Each box represents a splitting event with its two branching lineages (if applicable) showing above and below the box. The circle within boxes represents the mean of the posterior estimate and the whiskers mark the upper and lower 95% highest posterior density of the age estimates. The comparison shows that (1) our age estimates remain stable when calibration schemes changed; (2) our new age estimates are largely congruent with previous studies for nodes outside plethodontids, but younger within plethodontids.

Figure 5. Box plot depicting the simulated time estimates for four major evolutionary events of plethodontids. Each box plot contains the time estimates resulted from 200 simulated data sets with different numbers of genes. Box plots show a thick line at the median time estimate, a surrounding box containing the middle 50% of the data, and whiskers extending to encompass the middle 95% of the data. Dots indicate outlier points.
Figure 6. Biogeographic hypothesis showing the origin and dispersal pattern of plethodontid salamanders. Landmasses are abbreviated as follows: western North America (WA), eastern North America (EA). (A) Ancestors of Amphiumidae and Plethodontidae dispersed into North America by way of Beringia and were restricted to the northern part of western North America. (B) Plethodontids began their initial diversification and dispersed eastward after the Central Continental Seaway quickly closed in Paleocene. (C) Plethodontids migrated southward when the northern North America continued to cool down during Oligocene. (D) Plethodontids were restricted to eastern and western North America as central North America became increasingly dryer in the Miocene.
TABLE 1. Summary information for the 50 NPCL amplified in 25 plethodontid salamanders.

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<td>0.0733</td>
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</table>

Note: Length, length of refined alignment; PI sites, parsimony informative sites. P distance value is estimated using MEGA5 (Tamura et al. 2011). Compositional homogeneity (P-value) is calculated with the X2 test implemented in PAUP* v4b10 (Swofford 2002).
### TABLE 2. Statistical confidence (P-values) for alternative branching hypotheses based on the 50-gene dataset

<table>
<thead>
<tr>
<th>Alternative topology tested</th>
<th>$\Delta \ln L$</th>
<th>P-value</th>
<th>Rejection</th>
</tr>
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<tr>
<td>Best tree</td>
<td>0</td>
<td>0.519</td>
<td>0.075</td>
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<td><em>Hemidactylum</em> sister to remaining genera</td>
<td>673.0532</td>
<td>3.00E-04</td>
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<tr>
<td><em>Hemidactylum</em> sister to (<em>Batrachocephus</em> + Bolitoglossini) + Spelerpini</td>
<td>328.8304</td>
<td>2.00E-07</td>
<td>0</td>
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<tr>
<td><em>Hemidactylum</em> sister to <em>Batrachocephus</em></td>
<td>102.8745</td>
<td>2.00E-42</td>
<td>0</td>
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<tr>
<td><em>Karsenia</em> sister to remaining subfamily Plethodontinae</td>
<td>7.767544</td>
<td>0.255</td>
<td>0.05</td>
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<tr>
<td><em>Karsenia</em> sister to <em>Plethodon</em></td>
<td>-1.6251</td>
<td>0.591</td>
<td>0.258</td>
</tr>
<tr>
<td><em>Karsenia</em> sister to <em>Hydromantes</em> + <em>Ensatina</em></td>
<td>47.80727</td>
<td>3.00E-30</td>
<td>0</td>
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<td><em>Karsenia</em> sister to <em>Hydromantes</em></td>
<td>58.33756</td>
<td>1.00E-142</td>
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<tr>
<td><em>Karsenia</em> sister to <em>Ensatina</em></td>
<td>65.97841</td>
<td>2.00E-129</td>
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<tr>
<td><em>Karsenia</em> sister to <em>Aneides</em> + <em>Desmognathus</em></td>
<td>-2.7957</td>
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<td><em>Karsenia</em> sister to <em>Desmognathus</em></td>
<td>17.31448</td>
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<td>0.017</td>
<td>0.003</td>
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<td><em>Ensatina</em> sister to remaining subfamily Plethodontinae</td>
<td>26.29407</td>
<td>0.121</td>
<td>0.026</td>
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<td><em>Ensatina</em> sister to <em>Plethodon</em></td>
<td>38.38214</td>
<td>0.039</td>
<td>0.002</td>
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<tr>
<td><em>Ensatina</em> sister to <em>Karsenia</em> + <em>Aneides</em> + <em>Desmognathus</em> + <em>Hydromantes</em></td>
<td>30.25031</td>
<td>0.036</td>
<td>0.002</td>
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<tr>
<td><em>Ensatina</em> sister to <em>Aneides</em> + <em>Desmognathus</em> + <em>Hydromantes</em></td>
<td>6.868215</td>
<td>0.355</td>
<td>0.051</td>
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<td><em>Ensatina</em> sister to <em>Aneides</em> + <em>Desmognathus</em></td>
<td>1.170115</td>
<td>0.511</td>
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<td><em>Ensatina</em> sister to <em>Desmognathus</em></td>
<td>28.15624</td>
<td>0.002</td>
<td>0.001</td>
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<td><em>Ensatina</em> sister to <em>Aneides</em></td>
<td>26.84005</td>
<td>0.018</td>
<td>0.002</td>
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<td><em>Hydromantes</em> sister to <em>Plethodon</em></td>
<td>58.38783</td>
<td>0.001</td>
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</tbody>
</table>

Note: Best tree refers to Figure 2A; $\ln L$, likelihood value; AU, Approximately Unbiased test; BP, Bootstrap Probability; KH, Kishino-Hasegawa test.
TABLE 3. Detailed results of Bayesian molecular dating based on the analyses of 50 NPCLs using MultiDivTime, MCMCTREE and BEAST. Node numbers refer to Figure 3A. Eight nodes are labeled with their corresponding age constraints. Values are mean estimates and 95% credible interval (within parentheses). Time unit: one million years.

<table>
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<tr>
<th>Nodes</th>
<th>MultiDivTime</th>
<th>MCMCTREE</th>
<th>BEAST</th>
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<td>1: Amniota - Amphibia</td>
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<tr>
<td>2: Mammalia - Squamata</td>
<td>354.0 (341.5-362.2)</td>
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<tr>
<td>3: Aves - Squamata</td>
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<td>4: Aves - Crocodylia</td>
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<tr>
<td>5: Anura - Caudata</td>
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<td>6: Bombina - Silurana</td>
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<td>7: Living salamanders</td>
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<td>8: Hydriphidae - Cryptobranchidae</td>
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<td>9: Salamandroidea</td>
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<td>10: Cynops - Ambystoma</td>
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<td>14: Plethodontinae</td>
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<tr>
<td>15: Karsenia - Aneides</td>
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<tr>
<td>16: Aneides - Hydromantes</td>
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<td>17: Aneides - Desmognathus</td>
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<td>18: A. aeneus - A. hardii</td>
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<td>19: A. hardii - A. lugubris</td>
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<td>23: <em>H. italicus</em> - <em>H. shastae</em></td>
<td>23.4 (17.9-29.8)</td>
<td>23.5 (18.3-28.8)</td>
<td>24.5 (18.4-30.9)</td>
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<td>24: Eastern - Western <em>Plethodon</em></td>
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<td>40.0 (34.2-45.8)</td>
<td>45.3 (35.2-55.4)</td>
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<td>25: <em>P. dunni</em> - <em>P. asupak</em></td>
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<td>29.7 (22.2-37.5)</td>
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<td>26: <em>P. jordani</em> - <em>P. cinereus</em></td>
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<td>17.1 (11.7-22.5)</td>
<td>21.3 (15.8-27.5)</td>
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<td>27: <em>Hemidactyliinae</em></td>
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<td>58.6 (51.2-66.6)</td>
<td>58.4 (46.4-71.6)</td>
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<td>28: <em>Eurycea</em> - <em>Stereochilus</em></td>
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<td>37.3 (30.4-44.1)</td>
<td>40.9 (31.6-50.6)</td>
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<td>29: <em>Stereochilus</em> - <em>Pseudotriton</em></td>
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<td>17.7 (12.2-23.2)</td>
<td>22.3 (16.5-28.8)</td>
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<td>30: <em>E. bilineata</em> - <em>E. quadridigitata</em></td>
<td>18.4 (13.7-24.1)</td>
<td>18.7 (13.6-24.1)</td>
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<td>31: <em>Hemidactylium</em> - <em>(Batrachoseps+ Bolitoglossini)</em></td>
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<td>32: <em>Batrachoseps</em> - <em>Bolitoglossini</em></td>
<td>47.6 (37.8-58.2)</td>
<td>51.2 (44.0-58.5)</td>
<td>50.0 (39.1-61.4)</td>
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<td>33: <em>Thorius</em> - <em>Bolitoglossa</em></td>
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<td>40.9 (34.1-47.8)</td>
<td>40.0 (31.1-49.4)</td>
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<tr>
<td>34: <em>Bolitoglossa</em> - <em>Pseudo</em></td>
<td>21.9 (16.8-27.9)</td>
<td>24.9 (19.0-31.1)</td>
<td>23.8 (18.2-30.3)</td>
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<td>35: <em>B. robustus</em> - <em>B. attenuatus</em></td>
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<td>39.6 (33.2-47.2)</td>
<td>39.9 (31.1-49.4)</td>
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<td>17.1 (12.3-22.1)</td>
<td>15.2 (11.0-19.9)</td>
</tr>
</tbody>
</table>
1. Plethodon
2. Aneides
3. Ensatina
4. Karsenia
5. Hydromantes
6. Batrachoseps
7. Desmognathus
8. Hemidactylium
9. Spelerpini
10. Bolitoglossini (tropical salamanders)

▲ Oldest fossil records (~25 Ma)
11. Plethodon sp.
12. Aneides sp.
Mean divergence time and 95% CI (Ma)

- mtDNA genome (Mueller 2006)
- 3 nuclear genes (Vieites et al. 2007)
- 3 nuclear genes (Zheng et al. 2011)
- 50 nuclear genes (this study; calibrations from Mueller 2006)
- 50 nuclear genes (this study; calibrations from Vieites et al. 2007)
- 50 nuclear genes (this study; calibrations from Zheng et al. 2011)

Taxa:
- Rhyacotritonidae
- Amphiumidae
- Hynobiidae
- Cryptobranchidae
- Salamandridae
- Ambystomatidae
- Plethodontidae
- Hemidactyliinae
- Plethodontinae
- Hemidactylium
- Bolitoglossa