

Title: Molecular phylogenetics of sub-Saharan African natricine snakes, and the biogeographic origins of the Seychelles endemic *Lycognathophis seychellensis*

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Abstract:

Phylogenetic relationships of sub-Saharan African natricine snakes are understudied and poorly understood, which in turn has precluded analyses of the historical biogeography of the Seychelles endemic *Lycognathophis seychellensis*. We inferred the phylogenetic relationships of Seychelles and mainland sub-Saharan natricines by analysing a multilocus DNA sequence dataset for three mitochondrial (mt) and four nuclear (nu) genes. The mainland sub-Saharan natricines and *L. seychellensis* comprise a well-supported clade. Two maximally supported sets of relationships within this clade are (*Limnophis*, *Natriciteres*) and (*Afronatrix*, (*Hydraethiops*, *Helophis*)). The relationships of *L. seychellensis* with respect to these two lineages are not clearly resolved by analysing concatenated mt and nu data. Analysed separately, nu data best support a sister relationship of *L. seychellensis* with (*Afronatrix*, (*Hydraethiops*, *Helophis*)) and mt data best support a sister relationship with all mainland sub-Saharan natricines. Methods designed to cope with incomplete lineage sorting strongly favour the former hypothesis. Genetic variation among up to 33 *L. seychellensis* from five Seychelles islands is low. Fossil calibrated divergence time estimates support an overseas dispersal of the *L. seychellensis* lineage to the Seychelles from mainland Africa ca. 43–25 Ma, rather than this taxon being a Gondwanan relic.

Key Words: biogeography, Gondwana, Natricinae, Natricidae, overseas dispersal, systematics

1. Introduction

The Seychelles archipelago is an unusual island system in that it has both a remote, oceanic setting and continental origins as a fragment of Gondwana. Thus, its extant biota is a mosaic of ancient, relictual lineages and more recent overwater arrivals (Ali, 2017). This provides an attractive system for studying evolutionary processes, though exploiting this system for evolutionary research depends on improved knowledge of whether each resident lineage is ancient or relatively recent, as well as the status of populations on multiple Seychelles islands. Recently, some of this information has been generated for several lineages of Seychelles amphibians and reptiles (e.g., Brandley et al., 2005; Rocha et al., 2010, 2011, 2016a, b; Tolley et al., 2013; Maddock et al., 2014), but not yet for any of the native snakes (see Williams et al., 2020 for the latest assessment of the non-native status of Seychelles *Indotyphlops braminus*).

The snake *Lycognathophis seychellensis* (Schlegel, 1837) is endemic to the granitic islands of the Seychelles (Nussbaum, 1984). Several aspects of the systematics and historical biogeography of *L. seychellensis* remain unclear. DNA sequence data (for two partial nuclear genes, *cmos* and *rag2*) have been published for only one or two individuals of *L. seychellensis* (Vidal et al., 2008). This species has been recorded from six of the inner, granitic Seychelles islands (Fig. 1; Gerlach and Ineich, 2006) and exhibits notable colour polymorphism (Nussbaum, 1984). Other Seychelles reptile species have been found to display high levels of variation within and among islands, with potential implications for taxonomy and conservation management (Rocha et al., 2011, 2013, 2016; Valente et al., 2014; Harris et al.,

2015). Thus, an inspection of possibly substantial intraspecific genetic variation in *L. seychellensis* is warranted.

Currently there are 252 recognised species in the colubrid subfamily Natricinae, with the majority of this diversity found in Asia (183 species ca. 72%), only 13 species (ca. 5%) in mainland sub-Saharan Africa, and none in Madagascar (Uetz et al, 2020). Previous molecular phylogenetic analyses (Vidal et al., 2008; Dubey et al., 2012; Guo et al., 2012; Pyron et al., 2013; Zaher et al., 2019) supported the natricine colubrid (natricid according to some classifications: e.g., Zaher et al., 2019) identity of *L. seychellensis* proposed by Dowling (1990) based on vertebral and hemipenial anatomy, and indicated a close relationship with (and possible membership of) a mainland sub-Saharan African natricine radiation. However, those studies sampled only three of the six extant sub-Saharan African natricine genera (see Fig. 1 for distribution map) and only four of the 13 currently recognised species. Additionally, those previous studies disagreed as to whether *L. seychellensis* is more closely related to *Afronatrix* (Dubey et al., 2012; Pyron et al., 2013; Zaher et al., 2019) or to *Natriciteres* (Guo et al., 2012). As far as we are aware, there are no phylogenetic hypotheses that include all mainland sub-Saharan African natricine genera such that, beyond the close inferred relationship of the only two previously sampled genera in molecular phylogenetic studies (*Afronatrix* and *Natriciteres*), there appears to have been no previous consideration as to whether the mainland sub-Saharan African natricines constitute a single radiation and what their biogeographic origins might be. Finally, there are no published estimates of the age of the split between *L. seychellensis* and its closest extant relative. Having this temporal framework would allow a test of the hypothesis that the *L. seychellensis*

lineage arrived in the Seychelles via overseas dispersal rather than being a Gondwanan relic.

Here we present analyses of mitochondrial and nuclear DNA sequence data in order to test the following hypotheses: (1) mainland sub-Saharan African natricines are monophyletic, (2) *L. seychellensis* lies within the sub-Saharan African radiation, (3) intraspecific genetic variation within *L. seychellensis* is low across multiple Seychelles islands, and (4) the *L. seychellensis* lineage arrived in the Seychelles via oceanic dispersal rather than being an ancient, Gondwanan relic.

2. Materials and Methods

2.1. DNA sequence data generation and alignment

To infer phylogenetic relationships and to assess intraspecific genetic variation within the Seychelles endemic *Lycognathophis seychellensis*, we generated mitochondrial (mt) and nuclear (nu) gene sequence data for a total of 33 specimens of this species from across five of the six islands of its known distribution (Table 1). This included 14 individuals from Mahé, six from Praslin, six from Silhouette, two from La Digue, and five from Frégate. The eight partial genes sequenced were mt *16s* (n = 30), *cytb* (n = 29) and *nd4* (n = 2), and nu *cmos* (n = 8), *nt3* (n = 8), *bdnf* (n = 6), *rag1* (n = 2), and *prlr* (n = 21). In addition to the new *L. seychellensis* data, we generated 119 sequences for the three mt genes (*16s*, *cytb*, *nd4*) and four of the nuDNA loci (*cmos*, *nt3*, *bdnf*, *rag1*) for 27 other natricine snakes representing ten mainland sub-Saharan African species and five Asian species (Table 1).

Genomic DNA was extracted from muscle, liver, tail tip or ventral scale clip tissue samples stored in absolute ethanol at -20°C, using the DNeasy (Qiagen™)

blood and tissue kit. Primers and annealing temperatures for PCR amplification and for sequencing are reported in Table S1. Bidirectional sequence chromatograms were edited and assembled using Chromas Lite v. 2.1.1 (Technelysium, 2012). Possibly heterozygous positions in nuDNA sequences were given ambiguity codes. MEGA v. 7.0 (Kumar et al., 2016) was used to assemble sequences. Protein-coding sequences were checked for unexpected stop codons by translating nucleotide alignments to amino acids. Newly generated data were combined with data from GenBank for other natricines and outgroups (Table 1), and aligned using the ClustalW algorithm (Thompson et al., 1994) with default settings as implemented in MEGA.

2.2. Molecular phylogenetics

To infer the phylogenetic relationships of *L. seychellensis* and mainland sub-Saharan African natricines, we assembled a dataset of sequences from GenBank (Benson et al., 2017) and our newly generated sequences for three mitochondrial (*16s*, *cytb*, *nd4*) and four nuclear (*cmos*, *nt3*, *bdnf*, *rag1*) genes. Initially we ran analyses with a 157-leaf dataset, comprising 155 natricines and two colubroid outgroups; the grayiine (grayiid) *Grayia ornata* and sibynophiine (sibynophiid) *Sibynophis subpunctatus* (Table S2). Outgroup selection was based on the results of more broadly sampled molecular phylogenetic studies (Figueroa et al. 2016; Zaher et al. 2019; Lalronunga et al. 2020). The 155 natricines comprised 26 sub-Saharan African (including Seychelles) leaves representing all six genera and eleven of the 14 currently recognised species. The other 129 natricine leaves were selected from across the global geographic and phylogenetic diversity of the group based on each leaf being represented by at least three of the seven genes included in the dataset.

This sampling covered 30 of the 36 genera and 115 of the 245 species of extant natricines currently recognised (Uetz et al., 2020). Based on results of maximum likelihood (ML) (Felsenstein, 1981) analyses of concatenated data for this 157-leaf dataset, in which sub-Saharan and Seychelles natricines were together maximally supported as monophyletic, we subsequently ran phylogenetic analyses for a more restricted 21-leaf dataset (Table 1, Table S2) in an attempt to run phylogenetic analyses using (hopefully, more realistic) models estimated for a more focused taxon set with more complete data. This comprised 17 Seychelles and mainland sub-Saharan African natricines plus four outgroup natricines selected from among the likely closest relatives of the Seychelles + mainland sub-Saharan African clade. This dataset is also a combination of GenBank and newly generated sequences (Table S2).

Phylogenetic analyses were carried out using the CIPRES Science Gateway v3.3 (Miller et al., 2010). We applied ML using RAxML-HPC2 on XSEDE v. 8.2.12 (Stamatakis, 2014), and Bayesian inference (BI) analysis using MrBayes v. 3.2.6 on XSEDE (Ronquist et al., 2012). PartitionFinder2 (Lanfear et al., 2016) was used to determine best-fit gene and codon partitioning strategies using the greedy algorithm and BIC model selection, departing from gene and codon position blocks. RAxML analyses applied the GTRGAMMA model (following Stamatakis, 2006) to each partition. For the BI analysis we executed two runs with four Markov chains initiated from random trees and ran these for 10,000,000 generations each, sampling every 1000 generations. When the BI analysis was terminated, the standard deviation of split frequencies was less than 0.005, and the first 25% trees were discarded as “burn-in”. Convergence was assessed for all parameters using Tracer v1.6 (Rambaut et al., 2014) and the RWTY package (Warren et al., 2017). BI analysis used best-fit

models as determined by PartitionFinder. Support for internal branches was quantified using bootstrap proportions (calculated from 500 replicates) and Bayesian posterior probabilities, respectively.

We compared the fit of the data to optimal and best suboptimal trees for both ML and BI analyses for the 21-leaf dataset. For ML, we assessed differences in likelihood between optimal and suboptimal trees using the Approximately Unbiased (AU; Shimodaira, 2002) and Shimodaira-Hasegawa (SH; Shimodaira and Hasegawa, 1999) tests as implemented in the package CONSEL (Shimodaira and Hasegawa, 2001). For BI, we used Bayes factors (Kass and Raftery, 1995), with Marginal Likelihood estimate (MLE) scores estimated using path sampling and steppingstone sampling methods as described by Baele et al. (2012, 2013). Using BEAST v 1.10.4, analyses applying each of the alternative tree topologies as constraints were each run for 150,000,000 generations and sampled every 5,000. MLE scores were generated from 100 path steps for each run for 500,000 generations.

A species tree analysis of the 21-leaf dataset was carried out using a Bayesian multispecies coalescent approach as implemented in StarBEAST2 (BEAST v2.5, Bouckaert et al., 2019). Input files for StarBEAST2 were prepared using BEAUti 2 (Drummond et al., 2012). Each terminal was assigned to a described species. Priors were set as follows: linked tree model, strict clock, constant root population size model on species tree, GTR model for each locus, analyses initiated with a random starting tree for each locus, population mean set to a uniform prior and a lognormal prior for population size. The analysis was run for 100,000,000 generations sampling every 10,000 generations following a 10% burn-in. Convergence was checked in Tracer V1.6 with ESS \geq 200. Resulting trees were

summarised in LogCombiner 2.4.4 and TreeAnnotator 2.5. (Bouckaert et al., 2019). Results were visualised in FigTree 1.4.3 (Rambaut, 2016) and DensiTree (Bouckaert and Heled, 2014).

We compared support for alternative resolutions of the relationships of *L. seychellensis* under ML and BI analyses of the 21-leaf, concatenated mt and nu dataset. Support values were read from majority rule partition tables in PAUP* (Swofford, 2000) for 500 RAxML bootstrap trees, 2,000 sampled (post burn-in) MrBayes BI trees, and 1,500 sampled (post burn-in) BEAST2 BI trees. For RAxML, analyses were carried out with data partitioned by gene and by gene and codon position.

2.3. Divergence time estimation & historical biogeography

Our main aim was to test the hypothesis that the Seychelles natricine lineage reached the Seychelles via overseas dispersal, rather than being a Gondwanan relic. The geological timeframe for this test is as follows. Following the initial break up of eastern and western Gondwana from ca. 165 million years before present (Ma), Indo-Madagascar split from Antarctica+Australia ca. 130–112 Ma, and Madagascar split from India+Seychelles from ca. 88 Ma (see Samonds et al., 2012). Finally, India and Seychelles separated ca. 71–63 Ma (Collier et al., 2008; Minshull et al., 2008; Armitage et al., 2011; Ganerød et al., 2011: see Gower et al., 2016).

To estimate the age of the divergence between *L. seychellensis* and its closest extant relative, we analysed a taxonomically reduced dataset of 73 leaves to decrease computational time required for analyses (Table 2). This includes 12 natricines, including one *L. seychellensis*, and nine mainland sub-Saharan African natricines (eight currently recognised species). In addition, we included 61 non-

natricines in the dataset in order to make use of some of the best fossil calibrations available for snakes (see Head 2015, 2016; Zaher et al., 2019). Beyond the seven genes for which we generated new data, we also included an additional region in *rag1* (Wiens et al., 2008) because sequences of this marker were available for almost 40% of the samples. This dataset included in total 5,021 base pairs (Table 2). The data were newly aligned and the best-fit gene and codon partitioning scheme (Table S3) was selected using PartitonFinder2 as reported above.

We applied seven fossil calibrations, largely those recommended by Head (2015) and Head et al. (2016): (1) oldest divergence within crown Alethinophidia based on *Hassiophis terrasanctus* (Tchernov et al., 2000); minimum age 93.9; (2) divergence between non-xenodermid colubrids and their closest living relative (Xenodermidae in our tree) based on *Procerophis sahnii* (Rage et al., 2008); minimum age 50.5 Ma; (3) divergence between Boinae and its sister taxon (Erycinae + Candoiinae in our tree) based on *Titanoboa cerrejonensis* (Head et al., 2009); minimum age 58 Ma; (4) divergence between *Corallus* and (*Chilabothrus*, (*Epicrates*, *Eunectes*)); *Corallus priscus* (Rage, 2001); minimum 50.2 Ma; (5) divergence between Viperinae and Crotalinae based on *Vipera aspis* complex (Szyndlar and Rage, 1999); minimum age 20.0 Ma; (6) divergence between *Acrochordus javanicus* and (*A. arafurae*, *A. granulatus*) based on *Acrochordus dehmi* (Hoffstetter, 1964); minimum age 18.1 Ma and (7) oldest divergence between *Naja* (*Afronaja*) and *Naja* (*Boulengerina*) based on *Naja romani* (Hoffstetter, 1939) (A.B. Quadros pers. comm. 2019, based on work presented by Quadros et al., 2019); minimum age 17 Ma. Input values applied to each calibration prior are reported in Table S4.

Dating analyses were carried out using the CIPRES Science Gateway v3.3 (Miller et al., 2010). We estimated divergence times using BEAST2 on XSEDE v2.6 (Bouckaert et al., 2019) under a Yule tree prior. A relaxed lognormal clock was assigned for each partition of the concatenated BEAST2 analysis. We set up two independent runs, the Markov Chain Monte Carlo (MCMC) was run for 200,000,000 generations sampling every 10,000 trees, and effective sample size (ESS) values were evaluated using Tracer v. 1.7 (Rambaut et al., 2014). Initial runs applied the best-fit partitioning scheme and models, but this resulted in poor mixing of the GTR model parameters for partitions 1, 2, 3 & 4. We therefore repeated BEAST2 analyses implementing the less complex HKY model for these partitions. Additionally, we estimated divergence dates using Penalized Likelihood (Sanderson, 2002) as implemented in TreePL (Smith and O'Meara, 2012), applying the fossil dates above for calibrations 1–7 as hard minimum and maximum ages. Penalized likelihood uses a tree with branch lengths and calibrations without prior parametric distributions. We ran TreePL with 10,000 iterations and a smoothing parameter of 100.

Historical biogeography was investigated by estimating ancestral areas for the dated phylogeny using dispersal-extinction-cladogenesis (DEC; Ree et al., 2005, 2008) and dispersal-vicariance analysis (DIVALIKE; Ronquist, 1997; Ronquist and Sanmartín, 2011) models implemented in BioGeoBEARS v.1.1 (Matzke, 2013) in R v 4.0 (R Core Development Team, 2020). We also implemented these models using the peripatric speciation “*J*” (‘jump’) parameter, which although criticised by Ree and Sanmartín (2018), is often the most biologically realistic scenario for colonization and subsequent divergence on remote islands (Tsang et al., 2020). We defined three areas: Asia, Seychelles, and mainland sub-Saharan Africa, and set the maximum number of ancestral areas to three, because this was the maximum number of areas

occupied by any clade in our analysis. We compared the fit of these four models (DEC, DEC + j , DIVALIKE and DIVALIKE + j) using Akaike Information Criterion (AIC) scores and weights.

2.4. Genetic diversity within *Lycognathophis seychellensis*

PopArt (Leigh and Bryant, 2015) was used to generate haplotype networks to display intraspecific variation for *L. seychellensis* for *16s* and *cytb* under the median-joining algorithm (Bandelt et al., 1999).

3. Results

3.1. Molecular phylogenetics

The best-fit partitioning scheme for the 21-leaf dataset comprised six partitions, by gene and by codon position (Table S5). Mainland sub-Saharan African natricines plus *L. seychellensis* are strongly supported as monophyletic (Fig. 2A), and almost all intergeneric relationships within this lineage are well supported. In particular, and denoting the mainland sub-Saharan African genera with the first two letters of their name, the two sets of relationships (*Li,Na*) and (*Af,(He,Hy)*) are maximally supported (Fig. 2B). The relationship of *L. seychellensis* to these two lineages is less clearly resolved in analyses of the concatenated nu and mt data. In the 157-leaf dataset, *L. seychellensis* is sister to (*Af,(He,Hy)*), with ML bootstrap and Bayesian posterior probability support of 63 and 0.97, respectively (Fig. S1). The same relationship was recovered in ML(RAxML) analysis of the 21-leaf dataset, with lower support (37%), but in the BI (MrBayes) analysis of this dataset, *L. seychellensis* is instead sister to

all mainland sub-Saharan African natricines (support for monophyly of the latter clade 0.59) (Fig. 2; Table 3).

Examination of majority rule partition tables in PAUP* (Swofford, 2000) for the 500 RAxML ML bootstrap trees and 2,000 post-burnin MrBayes BI trees for the 21-leaf dataset shows that support is lowest for the third alternative resolution of the position of *L. seychellensis* with respect to (*Li,Na*) and to (*Af,(He,Hy)*), that is, as sister to the former clade (*Li,Na*), which receives support of only 27% (RAxML) and 0.12 (MrBayes) (Table 3). For the ML analysis, these support values are very similar when the same dataset is partitioned by gene instead of by gene and codon (Table 3). Both the AU and SH tests indicate that differences in likelihood between the fit of the three alternative topologies to the data are not significant (Table 4). Bayes factors indicate that the differences in fit to the data among the three alternative topologies are also not substantial, but differences in fit to data between the trees with *L. seychellensis* as sister to (*Af,(He,Hy)*) and as sister to either (*Li,Na*) or to (*(Li,Na)(Af,(He,Hy))*) are greater than differences in fit to data between the trees with the latter two sets of relationships (Table 4).

Analysis of the 21-leaf dataset for the nu and the mt data separately reveals that incongruence between these partitions explains a substantial part of the lack of compelling resolution of the relationships of *L. seychellensis* in analyses of the concatenated data. For the mt data, *L. seychellensis* is most strongly supported as sister to (*Af,(He,Hy)*), whereas for the nu data this is the least well supported of the three alternative resolutions, with instead *L. seychellensis* as sister to (*Af,(He,Hy)*) receiving the most support (Table S6). The Bayesian multispecies coalescent analysis using StarBEAST yielded 17,998 trees with 43 different topologies. Among these trees, 98% included *L. seychellensis* as sister to (*Af,(He,Hy)*).

3.2. Divergence dating & historical biogeography

When applying the HKY model in BEAST2 analyses, to those partitions for which good mixing was not achieved when applying GTR, good mixing was achieved for all parameters, for posteriors and priors (ESS > 200). Highest posterior density (HPD) intervals are wide for divergence times estimated in BEAST2 analyses. In unconstrained BEAST2 analysis, *L. seychellensis* is recovered as sister to the mainland sub-Saharan African (*Af,(He,Hy)*), with the split between these two lineages estimated to have occurred with a 95% HPD interval of 43.2–24.6 Ma (mean 35.3 Ma) (Fig. 3; Fig.S2), which is similar to the TreePL estimate of this divergence time for this node, at 35.7 Ma.

Enforcing the two alternative competing resolutions (from separate mt and nu data analyses) of the relationships among *L. seychellensis* and (*Af,(He,Hy)*) and (*Li,Na*) yields similar time estimates for the split between *L. seychellensis* and its closest extant (mainland sub-Saharan African) relative (Table 5). For BEAST2 runs applying GTR, as suggested by PartitionFinder2 results, ESS values were low for posterior (46) and prior (37) distributions, though the divergence date in question was similarly estimated (Fig. S3).

All of the BioGeoBEARS analyses agree with each other and with parsimony (carried out manually; results not shown) in their estimation of all ancestral areas. Asia is the estimated ancestral area for the sampled taxa, with a dispersal/vicariance event to sub-Saharan Africa. Dispersal rather than vicariance is the most likely explanation for origin of the sub-Saharan African clade because the phylogenetic split with Asian natricines is estimated to be much younger (48.7 Ma; HPD: 34.1–56.6) than the Gondwanan breakup. However, the route of dispersal to sub-Saharan

Africa and whether this was via land or sea is open to further investigation. A single dispersal event from mainland Africa to the Seychelles occurred relatively early on in the history of the group (Fig. 3). The divergence between the Seychelles natricine and its closest mainland African relative (35.7 Ma; HPD: 43.2–24.6 Ma) is much younger than the break-up between Seychelles and other Gondwanan landmasses (63 Ma or older, see section 2.3), strongly supporting overseas dispersal. Among the four models implemented, DEC + *j* was the best fitting, closely followed by DIVALIKE + *j* and DEC (Table 6).

3.3. Intraspecific genetic diversity

Intraspecific genetic diversity across the range of *Lycognathophis seychellensis* is low. Except for five single-site ambiguities (possible heterozygosity) in *prlr* and one in *rag1* there was no variation for any of the five nuclear gene sequences generated. For *16s* and *cytb* there are only two and four segregating sites, and zero and one parsimony informative sites, respectively. Therefore, we generated haplotype networks but not phylogenetic trees to examine spatial patterns in genetic variation (Fig. 4). There is only one instance of a single island having a haplotype not found on other islands, though this single *cytb* haplotype from Silhouette differs from those found on other islands by only a single step.

4. Discussion

4.1. Molecular phylogenetics and genetic diversity

Our molecular phylogenetic analyses are the first to comprehensively sample extant sub-Saharan African (including Seychelles) natricine genera (all six genera,

and 11 out of 14 species), and we find compelling support for their monophyly. Our results are consistent with previous work based on morphology that proposed that *Helophis* is a natricine and is closely related to *Hydraethiops* (de Witte, 1922; Broadley, 1998; Nagy et al., 2014), and that *Limnophis* and *Natriciteres* are closely related (Dowling and Duellman, 1978; Zaher, 1999). We were unable to identify with precision and reasonable support the closest living relatives of the sub-Saharan African (including Seychelles) natricine radiation. However, our results point to this radiation arising from within one of the Asian radiations (the clade excluding New World natricines plus (Eur)Asian *Natrix*, *Sinonatrix*, *Smithophis*, and *Opisthotropis*).

The phylogenetic results regarding the identity of the sister group of *L. seychellensis* are not compelling from the analyses of the concatenated nu and mt sequence data. However, partitioned analyses provide evidence of substantial nu-mt incongruence, and analysis using a multispecies coalescent approach that is designed to cope with incomplete lineage sorting strongly supports the hypothesis that *L. seychellensis* is sister to *Hydraethiops* + *Helophis* + *Afronatrix*.

The almost complete lack of molecular genetic variation within *L. seychellensis* across five Seychelles islands is consistent with there being only a single extant species of natricine occurring on the Seychelles. The very low intraspecific variation contrasts with that found for other native Seychelles squamate reptiles and other taxa (e.g., frogs: Labisko et al., 2019; caecilians: Maddock et al., 2020; crabs: Daniels, 2011), which have substantial and strongly spatially structured, intraspecific molecular genetic variation. This structured variation often includes a distinct genetic lineage on Silhouette, a hint of which is also found here for *L. seychellensis* *cytb*.

The lack of genetic variation suggests that the current genetic diversity within *L. seychellensis* is the product of a strong founder effect for a relatively recent overseas arrival (not supported by our dating results) and/or within-Seychelles dispersal from a substantially contracted refugial population representing some form of secondary founder effect. Due to the relatively deep phylogenetic divergence of *L. seychellensis* and no record of this taxon occurring beyond Seychelles, a very recent, human mediated dispersal to Seychelles is highly unlikely and can be discounted with confidence. The lack of genetic variation among populations of *L. seychellensis* from different islands is consistent with this species being a relatively good disperser among the major granitic islands of the Seychelles (perhaps the best of Seychelles squamates studied thus far), across relatively shallow, narrow seas and/or during multiple low sea level stands during the Pleistocene, when these islands would have been repeatedly dis- and reconnected (Warren et al. 2010). A closer examination of *L. seychellensis* intraspecific genetic variation within and among islands using rapidly evolving nuclear markers and a denser sampling would be worthwhile.

4.2. Historical biogeography

Our results are consistent with the *L. seychellensis* lineage reaching the Seychelles via overseas dispersal from Africa, approximately 43–25 Ma. Probably this is a similar mode and timing of origin to all of the other Seychelles native squamate reptiles for which data are available. For example, the endemic Seychelles chameleon *Archaius tigris* is most closely related to East African *Rieppeleon* spp., with the two diverging an estimated 43–27 Ma (Tolley et al., 2013: table S4). The Seychelles skinks *Trachylepis seychellensis* and *T. wrightii* are estimated to have

diverged from their closest extant relatives, mostly occurring in Africa, approximately 41–24 Ma (Lima et al., 2013; Weinell et al., 2019). Dating estimates are not yet available for other Seychelles squamates, but native and endemic Seychelles species of geckos of the genera *Phelsuma*, *Urocotyledon* and *Ailuronyx* have closest extant relatives in Africa and/or Madagascar (Rocha et al., 2010; Gamble et al., 2012). The monotypic Seychelles fossorial skinks *Janetaescincus* spp. and *Pamelaescincus gardineri* are sister to a large clade of extant African, Arabian and Indian Ocean island taxa (Brandley et al., 2005).

The origins of Seychelles squamates contrast with those for most of the native amphibians. Although the hyperoliid frog *Tachycnemis seychellensis* arrived from overseas dispersal from Madagascar approximately 35.3–9.8 Ma (Crottini et al., 2012; Maddock et al. 2014; ca. 20–12 Ma based on Portik et al. [2019: fig. S2]), the two ancient Seychelles amphibian lineages that comprise eight indotyphlid caecilian species (Maddock et al., 2017, 2018) and at least four sooglossid frog species have their closest extant relatives in India, and are Gondwanan relics (Wilkinson et al., 2002; Biju and Bossuyt, 2003; Gower et al., 2011, 2016; Kamei et al., 2012; Janani et al., 2017; Labisko et al., 2019).

Globally, natricines are ecologically diverse, utilising ground, shrub and tree, soil, semiaquatic and/or aquatic habitats. Mainland sub-Saharan African natricines, as well as being relatively species poor, are seemingly ecologically more conservative despite being the product of approximately 30 million years of evolution, with perhaps all species being predominantly semiaquatic or even aquatic (Gibbons and Dorcas, 2004; Vitt and Caldwell, 2009). Additional and more detailed ecological data would be useful, but *L. seychellensis* appears to be the only member of the sub-Saharan African (including Seychelles) radiation that is predominantly

terrestrial. It is typically encountered on the ground, though can be observed in vegetation at heights of at least 5 m as well as on the margins of pools and streams (STM, JL, RK & DJG pers. obs.). It is not clear whether this disparate ecology is ancestral to the *L. seychellensis* lineage and a factor that might have promoted its overseas dispersal to Seychelles, or whether it is a derived feature acquired since the lineage reached Seychelles. All mainland sub-Saharan African natricines are oviparous. *Lycognathophis seychellensis* is probably also oviparous (Nussbaum, 1984) but direct observations have yet to be reported (Gerlach and Ineich, 2006).

Extant mainland sub-Saharan African natricines are relatively species-poor, comprising only ca. 5% of global natricine diversity. Natricines are unknown from Madagascar or Indian Ocean islands other than Seychelles. As argued for other Indian Ocean and Madagascar taxa, the overseas dispersal of the *L. seychellensis* lineage from mainland Africa eastwards was perhaps assisted by sporadic but strong eastward oceanic currents and large, continental freshwater outflows during the Paleogene (65.5–23 Ma: Ali and Huber, 2010; Townsend et al., 2010).

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1 **Table 1.** Sequences generated in this study. Sequences used in the 21-leaf dataset are indicated with *, en-dash (–) = no data,
 2 different Islands in Seychelles are marked bold.

3

Sampl e	Species	voucher number/extracti on ID	Latitude	Longitud e	Sequenc e ID	Location	16s	cytb	nd4	cmos	nt3	bdnf	rag1	prlr
1	<i>Afronatrix anoscopus*</i>	NHMUK 2013:383	6.05370	9.47322	AE-40	Akwa, Mamfe division, Cameroon	MW699965	MW711484	–	MW711535	MW71145 2	MW71155 8	MW71158 9	–
2	<i>Afronatrix anoscopus*</i>	Lake Kuk, Cameroon	6.41072	10.2081 2	AE-41	LakeKuk, Cameroon Mount Nimba mining concession, Forestiere Region, Guinea	MW699966	MW711485	–	MW711536	MW71145 3	MW71155 9	MW71159 0	–
3	<i>Afronatrix anoscopus*</i>	BMNH2008.637	7.69960	-8.39880 22.7869	AE-42		MW699967	MW711486	MW711522	MW711537	–	MW71156 0	MW71159 1	–
4	<i>Helophis schoutedeni*</i>	CRT3853	2.03306	22.7869 4	AI-89	DR Congo, Kona	MW699968	–	MW711523	MW711538	MW71145 4	MW71156 1	MW71159 2	–
5	<i>Helophis schoutedeni*</i>	CRT3855	2.03306	22.7869 4	AJ-90	DR Congo, Kona	MW699969	–	MW711524	MW711539	MW71145 5	MW71156 2	MW71159 3	–
6	<i>Helophis schoutedeni</i>	CRT3859	2.03306	22.7869 4	AJ-91	DR Congo, Kona	MW699970	–	–	MW711540	MW71145 6	MW71156 3	–	–
7	<i>Helophis schoutedeni</i>	CRT3860	2.03306	22.7869 4	AJ-92	DR Congo, Kona	MW699971	–	–	MW711541	MW71145 7	MW71156 4	–	–
8	<i>Hydraethiops melanogaster*</i>	NMP-P6V 75491	1.05000	26.8300 0	AI-84	Bafwabianga village, DR Congo	MW699972	–	MW711525	MW711542	MW71145 8	MW71156 5	MW71159 4	–
9	<i>Hydraethiops melanogaster</i>	NMP-P6V 75491	0.54000	12.8300 0	AI-85	Makokou, Gabon	MW699973	–	MW711526	MW711543	MW71145 9	MW71156 6	MW71159 5	–
10	<i>Hydraethiops melanogaster*</i>	NMP-P6V 75772	1.81000	14.5500 0	AI-86	Dia near Sembe, Congo Rep.	MW699974	–	MW711527	MW711544	MW71146 0	MW71156 7	MW71159 6	–
11	<i>Limnophis bangweolicus*</i>	ANG-438	-17.57333	23.2266 7	AI-82	West of Sashae, Cuando Cubango, Angola	MT586815	MT587279	MT587306	MT587292	MW71146 1	MW71156 8	MW71159 7	–
12	<i>Limnophis bicolor*</i>	WC-6238	-12.33012	18.4047 9	AI-83	upstream of Lungue- bungue River bridge, Moxico, Angola	MT586817	MT587281	MT587307	MT587294	MW71146 2	MW71156 9	MW71159 8	–
13	<i>Limnophis branchi</i>	ANG-056	-7.75308	19.9568 6	AI-80	Lulele River area, Lunda Norte, Angola	MT586818	MT587283	MT587308	MT587296	MW71146 3	MW71157 0	MW71159 9	–
14	<i>Limnophis branchi</i>	ANG-058	-7.75308	19.9568 6	AI-81	Lulele River area, Lunda Norte, Angola	MT586819	MT587284	MT587309	MT587297	MW71146 4	MW71157 1	MW71160 0	–
15	<i>Natriciteres fuliginoides*</i>	CAS258133	-0.05171	11.1662 1	AD-33	Moyen-Ogooue Prov., Gabon	MW699975	MW711487	MW711528	MW711545	MW71146 5	MW71157 2	MW71160 1	–
16	<i>Natriciteres olivacea*</i>	CAS220640	–	–	AD-34	DR Congo	MT586821	MT587285	MT587311	MT587299	MW71146 6	MW71157 3	–	–
17	<i>Natriciteres olivacea*</i>	WRB-636	-5.93333	29.2000 0	AI-87	Kalemie, DR Congo Udzungwa Mts., Uzungwa Scarp F. R., Mkalazi, Tanzania	MW699976	MW711488	MW711529	MW711546	MW71146 7	MW71157 4	MW71160 2	–
18	<i>Natriciteres sylvatica*</i>	MTSN5485	-8.39750	35.9786 1	AD-36	Nguru South Mts., Nguru South (Pemba), Tanzania	MW699977	MW711489	MW711530	MW711547	MW71146 8	MW71157 5	MW71160 3	–
19	<i>Natriciteres sylvatica</i>	MTSN8183	-6.03044	37.5256 4	AD-37	MW699978		MW711490	–	–	–	–	–	–

Sample	Species	voucher number/extract ID	Latitude	Longitude	Sequence ID	Location	16s	cytb	nd4	cmos	nt3	bdnf	rag1	prlr
20	<i>Natriciteres sylvatica</i> *	MUSE13712	-	-	AD-38	Rungwe Mt. (Nkuka), Southern Highlands, Tanzania	MW699979	MW711491	MW711531	MW711548	MW711469	MW711576	MW711604	-
21	<i>Natriciteres sylvatica</i>	WRB-642	-16.28789	36.41244	AI-88	Mount Mabu, Zambezi, Mozambique	MW699980	MW711492	MW711532	MW711549	-	MW711577	-	-
22	<i>Natriciteres variegata</i>	MUSE13726	-	-	AD-39	Rungwe Mt. (Nkuka), Southern Highlands, Tanzania	MW699981	-	-	-	-	-	-	-
23	<i>Fowlea flavipunctatus</i> *	FMNH250122	-	-	AC-21	Thailand	MT586822	MT587286	MT587312	MT587300	MW711470	MW711578	MW711605	-
24	<i>Amphiesma stolatum</i> *	CAS213646	20.31286	94.74683	AC-29	Myanmar	MT586823	MT587287	MT587313	MT587301	MW711471	MW711579	MW711606	-
25	<i>Rhabdops olivaceus</i>	NCBS-AU164	11.53932	76.02188		Pookode, Wayanad District, Kerala, India	MF352831	MF352838	MF352842	MF352834	MW711472	MW711580	MW711607	-
26	<i>Smithophis atemporalis</i>	BNHS 3527	23.76338	93.09916		Mizoram University Campus, Aizawl, Mizoram, India	MK350255	MK350262	MK350258	MK350265	MW711473	MW711581	MK350257	-
27	<i>Smithophis bicolor</i>	BNHS 2369	23.71083	92.93194		Mizoram University Campus, Aizawl, Mizoram, India	MK350254	MK350261	MK350259	MK350264	MW711474	MW711582	MK350256	-
28	<i>Lycognathophis seychellensis</i>	SM251	-4.65894	55.43744		Morne Blanc Trail, Mahé, Seychelles	MW699982	MW711493	-	-	MW711475	-	-	MW711610
29	<i>Lycognathophis seychellensis</i> *	SM165	-4.63738	55.41212	AM-126	Mare Aux Cochons, Mahé, Seychelles	MW699983	MW711494	MW711533	MW711550	MW711476	MW711583	MW711608	-
30	<i>Lycognathophis seychellensis</i>	SM252	-4.65350	55.43778		Casse Dent, Mahé, Seychelles	MW699984	MW711495	-	-	-	-	-	MW711611
31	<i>Lycognathophis seychellensis</i>	SM286	-4.64493	55.43363		Congo Rouge, Mahé, Seychelles	MW699985	MW711496	-	MW711551	MW711477	MW711584	-	MW711612
32	<i>Lycognathophis seychellensis</i>	SM287	-4.65043	55.43765		Congo Rouge, Mahé, Seychelles	MW699986	MW711497	-	-	-	-	-	-
33	<i>Lycognathophis seychellensis</i>	SM290	-	-		Mahé, Seychelles	MW699987	MW711498	-	-	-	-	-	MW711613
34	<i>Lycognathophis seychellensis</i>	R9458	-4.63487	55.41158		Mare Aux Cochons, Mahé, Seychelles	MW699988	-	-	-	-	-	-	-
35	<i>Lycognathophis seychellensis</i>	R6716	-4.70760	55.50081		La Reserve, Mahé, Seychelles	MW699989	-	-	-	-	-	-	MW711614
36	<i>Lycognathophis seychellensis</i>	R6909	-4.65716	55.43421		Trail to Morne Blanc, Mahé, Seychelles	MW699990	MW711499	-	-	-	-	-	MW711615
37	<i>Lycognathophis seychellensis</i>	SM292	-4.64007	55.41602		Mare Aux Cochons, Mahé, Seychelles	MW699991	MW711500	-	MW711552	-	-	-	MW711616
38	<i>Lycognathophis seychellensis</i>	SM293	-4.62853	55.41139		Mare Aux Cochons, Mahé, Seychelles	MW699992	MW711501	-	-	MW711478	-	-	-
39	<i>Lycognathophis seychellensis</i>	SM339	-4.64883	55.42858		Between Mare Aux Cochons and Casse Dent, Mahé, Seychelles	MW699993	MW711502	-	-	-	-	-	MW711617
40	<i>Lycognathophis seychellensis</i>	SM340	-4.65114	55.41194		Between Mare Aux Cochons and Casse Dent, Mahé, Seychelles	MW699994	MW711503	-	-	-	-	-	MW711618
41	<i>Lycognathophis seychellensis</i>	SM385	-4.64675	55.43450		Congo Rouge trail, Mahé, Seychelles	MW699995	MW711504	-	-	-	-	-	MW711619

Sample	Species	voucher number/extraction ID	Latitude	Longitude	Sequence ID	Location	16s	cytb	nd4	cmos	nt3	bdnf	rag1	prlr
42	<i>Lycognathopsis seychellensis</i>	SM87	-4.33180	55.73885		Vallée de Mai, Praslin , Seychelles	MW699996	MW711505	-	-	-	-	-	MW711620
43	<i>Lycognathopsis seychellensis</i>	SM88	-4.33180	55.73885		Vallée de Mai, Praslin , Seychelles	MW699997	MW711506	-	-	-	-	-	MW711621
44	<i>Lycognathopsis seychellensis</i>	SM89	-4.33077	55.73763		Vallée de Mai, Praslin , Seychelles	MW699998	MW711507	-	MW711553	-	-	-	-
45	<i>Lycognathopsis seychellensis</i>	SM112	-4.30359	55.69084		Anse Kerlan River, Praslin , Seychelles	MW699999	MW711508	-	-	-	-	-	MW711622
46	<i>Lycognathopsis seychellensis</i>	R9459	-4.33475	55.73384		Praslin , Seychelles	MW700000	-	-	-	-	-	-	MW711623
47	<i>Lycognathopsis seychellensis</i>	R9469	-4.33143	55.73762		Vallée de Mai, Praslin , Seychelles	MW700001	MW711509	-	-	-	-	-	MW711624
48	<i>Lycognathopsis seychellensis</i>	SM206	-4.48528	55.23781		Trail to Jardin Maron, Silhouette , Seychelles	MW700002	MW711510	-	-	-	-	-	-
49	<i>Lycognathopsis seychellensis</i>	SM213	-4.47890	55.24621	AM-127	Hotel Hilton LeBreeze, Silhouette , Seychelles	MW700003	MW711511	MW711534	MW711554	MW711479	MW711585	MW711609	MW711625
50	<i>Lycognathopsis seychellensis</i>	SM225	-4.48474	55.23967		Trail to Jardin Maron, Silhouette , Seychelles	MW700004	MW711512	-	MW711555	MW711480	MW711586	-	-
51	<i>Lycognathopsis seychellensis</i>	R9467	-4.48450	55.24260		Silhouette , Seychelles	MW700005	-	-	-	-	-	-	-
52	<i>Lycognathopsis seychellensis</i>	R6680	-4.48704	55.23544		Trail to Jardin Maron, Silhouette , Seychelles	MW700006	MW711513	-	-	-	-	-	MW711626
53	<i>Lycognathopsis seychellensis</i>	R6723	-4.48497	55.23800		Trail to Jardin Maron, Silhouette , Seychelles	MW700007	MW711514	-	-	-	-	-	MW711627
54	<i>Lycognathopsis seychellensis</i>	R9460	-4.35872	55.84062		Belle Vue, La Digue , Seychelles	MW700008	MW711515	-	-	-	-	-	-
55	<i>Lycognathopsis seychellensis</i>	R9464	-4.37145	55.82741		Anse Source d'Argent, La Digue , Seychelles	MW700009	MW711516	-	MW711556	MW711481	MW711587	-	-
56	<i>Lycognathopsis seychellensis</i>	R9465	-4.58581	55.94387		Frégate , Seychelles	MW700010	MW711517	-	MW711557	MW711482	MW711588	-	-
57	<i>Lycognathopsis seychellensis</i>	R9466	-4.58581	55.94387		Frégate , Seychelles	MW700011	MW711518	-	-	MW711483	-	-	MW711628
58	<i>Lycognathopsis seychellensis</i>	SM505	-4.58475	55.93317		Frégate , Seychelles	-	MW711519	-	-	-	-	-	-
59	<i>Lycognathopsis seychellensis</i>	SM506	-4.58475	55.93317		Frégate , Seychelles	-	MW711520	-	-	-	-	-	MW711629
60	<i>Lycognathopsis seychellensis</i>	SM507	-4.58475	55.93317		Frégate , Seychelles	-	MW711521	-	-	-	-	-	MW711630

Table 2. GenBank accession and voucher numbers for gene sequences used in molecular dating analysis. – = no data

Sample	Species	Family	Sequence ID	16s	cytb	nd4	cmos	nt3	bdnf	rag1 (part 'a')	rag1 (part 'b')
1	<i>Acrochordus granulatus</i>	Acrochordidae		AF544786	AF217841	HM234054	HM234057	FJ434082	FJ433981	–	–
2	<i>Acrochordus javanicus</i>	Acrochordidae		AF512745	KX694897	HM234055	HM234058	KX694991	AY988036	–	HM234061
3	<i>Afronatrix anoscopus</i>	Colubridae (Natricinae)	AE 41	MW699966	MW711485	–	MW711536	MW711453	MW711559	MW711590	–
4	<i>Afronatrix anoscopus</i>	Colubridae (Natricinae)	AE 42	MW699967	MW711486	MW711522	MW711537	–	MW711560	MW711591	–
5	<i>Agkistrodon contortrix</i>	Viperidae (Crotalinae)		AF156566	EU483383	AF156577	–	–	EU402623	–	EU402833
6	<i>Ahaetulla pulverulenta</i>	Colubridae (Ahaetuliinae)		KC347339	KC347454	KC347512	KC347378	–	–	–	KC347416
7	<i>Anilius scytale</i>	Aniliidae		FJ755180	U69738	FJ755180	AF544722	FJ434066	EU402625	–	AY988072
8	<i>Anomochilus leonardi</i>	Cylindrophiidae+Anomochiliidae		AY953431	–	–	–	–	–	–	–
9	<i>Aparallactus capensis</i>	Atractaspididae (Aparallactinae)		AY188045	AY188006	FJ404331	AY187967	–	–	–	–
10	<i>Aplopeltura boa</i>	Pareidae (Pareinae)		AF544787	JF827673	JF827650	JF827696	FJ434085	FJ433984	–	–
11	<i>Aspidura ceylonensis</i>	Colubridae (Natricinae)		KC347361	KC347477	KC347527	KC347400	–	–	–	KC347438
12	<i>Asthenodipsas malaccanus</i>	Pareidae (Pareinae)		KX660197	KX660469	KX660597	KX660336	–	–	–	–
13	<i>Azemios feae</i>	Viperidae (Azemiopinae)		AF057234	AY352747	AY352808	AF544695	KX694977	EU402628	–	EU402836
14	<i>Bitis nasicornis</i>	Viperidae (Viperinae)		AY188048	DQ305457	DQ305475	AY187970	–	–	–	KC330012
15	<i>Boa constrictor</i>	Boidae		AB177354	AB177354	AB177354	AF544676	–	KC330044	–	KC347423
16	<i>Boaedon fuliginosus</i>	Lamprophiidae (Lamprophiinae)		AY188079	AF471060	FJ404365	FJ404270	FJ434094	EU402646	KM519732	EU402849
17	<i>Bothrolycus ater</i>	Lamprophiidae (Lamprophiinae)		AY611859	AY612041	AY611950	FJ404347	–	–	–	–
18	<i>Buroma depressiceps</i>	Lamprophiidae <i>incertae sedis</i>		AY611860	AY612042	–	AY611951	–	–	–	–
19	<i>Buroma procterae</i>	Lamprophiidae <i>incertae sedis</i>		AY611818	AY612001	DQ486328	AY611910	–	–	–	–
20	<i>Bungarus fasciatus</i>	Elapidae		EU579523	EU579523	EU579523	AY058924	KX694998	FJ433989	JF357954	–
21	<i>Calabaria reinhardtii</i>	Calabariidae		Z46494	AY099985	–	AF544682	–	EU402631	–	EU402839
22	<i>Calamaria pavimentata</i>	Colubridae (Calamariinae)		KX694624	AF471081	–	AF471103	KX694999	FJ434005	–	–
23	<i>Candoia carinata</i>	Candoiidae		EU419850	AY099984	–	AY099961	FJ434077	FJ433974	–	AY988065
24	<i>Cantoria violacea</i>	Homalopsidae		KX694627	EF395897	EF395922	–	KX695001	KX694704	–	–

Sampl e	Species	Family	Sequenc e ID	16s	cytb	nd4	cmos	nt3	bdnf	rag1 (part 'a')	rag1 (part 'b')
25	<i>Casarea dussumieri</i>	Bolyeridae		AF544827	U69755	–	AF544731	FJ434069	EU402632	–	EU402840
26	<i>Charina bottae</i>	Charinidae (Charininae)		AF544816	AY099986	AF302959	AY099971	FJ434079	FJ433978	–	AY988076
27	<i>Chilabothrus striatus</i>	Boidae		–	–	KC329966	KC329991	DQ465554	KC330056	–	KC330027
28	<i>Contia tenuis</i>	Colubridae (Dipsadinae)		AY577030	GU112384	GU112419	AF471134	–	GU112346	–	–
29	<i>Corallus annulatus</i>	Boidae		–	KC750012	KC750018	KC750007	–	JX576167	–	KC750047
30	<i>Cylindrophis ruffus</i>	Cylindrophidae+Anomochilidae		AB179619	AB179619	AB179619	AF471133	–	AY988037	AY662613	AY988071
31	<i>Daboia russellii</i>	Viperidae (Viperinae)		EU913478	EU913478	EU913478	AF471156	–	EU402636	–	EU402843
32	<i>Dityophis vivax</i>	Lamprophiidae		–	–	–	–	–	JQ073079	–	JQ073200
33	<i>Epicrates cenchria</i>	Boidae		–	HQ399501	KC329975	KC330008	JX576186	KC330073	–	–
34	<i>Eryx colubrinus</i>	Erycidae		AF544819	U69811	–	AF544716	DQ465569	EU402639	–	DQ465571
35	<i>Eryx conicus</i>	Erycidae		AF512743	GQ225658	GQ225672	–	–	–	–	AY988074
36	<i>Eunectes notaeus</i>	Boidae		AM236347	HQ399499	KC329978	HQ399536	–	KC330076	–	HQ399516
37	<i>Gerrhopilus mirus</i>	Gerrhopilidae		AM236345	AM236345	AM236345	–	GU902566	GU902394	–	–
38	<i>Grayia ornata</i>	Colubridae (Grayiinae)		AY611866	AY612048	AF544663	AF544684	KX695019	FJ434002	–	–
39	<i>Helophis schoutedeni</i>	Colubridae (Natricinae)	AI 89	MW699968	–	MW711523	MW711538	MW711454	MW711561	MW711592	–
40	<i>Hologerrhum philippinum</i>	Lamprophiidae (Cyclocorinae)		–	MG458758	–	MG458766	–	–	–	–
41	<i>Homoroselaps lacteus</i>	Lamprophiidae (Atractaspidinae)		AY611809	AY611992	FJ404338	AY611901	KX695021	JQ599029	MK621524	–
42	<i>Hydraethiops melanogaster</i>	Colubridae (Natricinae)	AI 84	MW699972	–	MW711525	MW711542	MW711458	MW711565	MW711594	–
43	<i>Liasis mackloti</i>	Pythonidae		EF545051	U69839	–	AF544726	FJ434075	FJ433959	–	–
44	<i>Limnophis bangweolicus</i>	Colubridae (Natricinae)	AI 82	MT586815	MT587279	MT587306	MT587292	MW711461	MW711568	MW711597	–
45	<i>Limnophis branchi</i>	Colubridae (Natricinae)	AI 81	MT586819	MT587284	MT587309	MT587297	MW711464	MW711571	MW711600	–
46	<i>Liopholidophis sexlineatus</i>	Lamprophiidae (Pseudoxyrhophiinae)		AY188063	DQ979985	FJ404373	AY187985	–	–	–	–
47	<i>Liotyphlops albirostris</i>	Anomalepididae		AF366762	AF544672	–	AF544727	–	EU402650	–	–
48	<i>Loxocemus bicolor</i>	Loxocemidae		AF544828	AY099993	–	AY444035	FJ434072	EU402651	–	–
49	<i>Lycognathophis seychellensis</i>	Colubridae (Natricinae)	AM 126	MW699983	MW711494	MW711533	MW711550	MW711476	MW711583	MW711608	–
50	<i>Micrelaps bicoloratus</i>	Lamprophiidae (Aparallactinae)		–	DQ486349	–	DQ486173	–	–	–	–

Sampl e	Species	Family	Sequenc e ID	16s	cytb	nd4	cmos	nt3	bdnf	rag1 (part 'a')	rag1 (part 'b')
51	<i>Mimophis mahfalensis</i>	Lamprophiidae (Psammophiinae)		AY188070	DQ486461	–	AY187992	KX695030	JQ073081	–	–
52	<i>Naja (Afronaja) mossambica</i>	Elapidae		AY611813	AY611996	DQ897723	AY611905	–	–	–	–
53	<i>Naja (Boulengerina) melanoleuca</i>	Elapidae		AY611812	AY611995	KX130765	AY611904	–	–	–	–
54	<i>Natriciteres fuliginoides</i>	Colubridae (Natricinae)	AD 33	MW699975	MW711487	MW711528	MW711545	MW711465	MW711572	MW711601	–
55	<i>Natriciteres olivacea</i>	Colubridae (Natricinae)	AI 87	MW699976	MW711488	MW711529	MW711546	MW711467	MW711574	MW711602	–
56	<i>Natriciteres sylvatica</i>	Colubridae (Natricinae)	AD 36	MW699977	MW711489	MW711530	MW711547	MW711468	MW711575	MW711603	–
57	<i>Oxyrhabdium leporinum</i>	Lamprophiidae (Cyclocorinae)		–	AF471029	–	DQ112081	–	–	–	–
58	<i>Oxyuranus scutellatus</i>	Elapidae		EU547149	EU547051	EF210827	EU546916	–	–	EU546877	–
59	<i>Pareas carinatus</i>	Pareidae (Pareinae)		AF544802	JF827677	JF827653	JF827702	FJ434086	FJ433985	–	–
60	<i>Prosymna janii</i>	Lamprophiidae (Prosymninae)		FJ404222	FJ404319	FJ404389	FJ404293	–	–	–	–
61	<i>Pseudaspis cana</i>	Lamprophiidae (Pseudaspidinae)		AY611898	AY612080	DQ486319	DQ486167	–	–	–	–
62	<i>Pseudoxenodon karlschmidti</i>	Colubridae (Pseudoxenodontinae)		JF697330	AF471080	–	AF471102	KX695042	JQ599045	–	–
63	<i>Python bivittatus</i>	Pythonidae		KF010492	JX401131	–	AF435016		XM743302 2	XM – 007441886	–
64	<i>Rhinophis drummondhayi</i>	Uropeltidae		AY701028	AF544673	–	AF544719	FJ434071	FJ433966	–	–
65	<i>Sanzinia madagascariensis</i>	Sanziniidae		AY336066	U69866	–	EU403580	–	AY988033	–	AY988067
66	<i>Sibynophis subpunctatus</i>	Colubridae (Sibynophiinae)		KC347373	KC347471	KC347516	KC347411	–	–	–	KC347449
67	<i>Tropidophis feicki</i>	Tropidophiidae		AF512733	KF811124	–	KF811110	–	KF811074	–	–
68	<i>Ungaliophis continentalis</i>	Charinidae (Ungaliophiinae)		AF544833	U69870	–	AF544724	FJ434081	EU402665	–	EU402867
69	<i>Fowlea flavipunctatus</i>	Colubridae (Natricinae)	AC 21	MT586822	MT587286	MT587312	MT587300	MW711470	MW711578	MW711605	–
70	<i>Xenodermus javanicus</i>	Xenodermidae		AF544810	–	U49320	AF544711	–	EU402667	–	EU402869
71	<i>Xenopeltis unicolor</i>	Xenopeltidae		AB179620	AB179620	AB179620	AF544689	FJ434073	EU402668	–	DQ465564
72	<i>Xenophidion schaeferi</i>	Xenophidiidae		–	AY574279	–	–	–	–	–	–
73	<i>Xylophis perroteti</i>	Pareidae (Xylophiinae)		MK340908	–	MK340910	MK344193	–	MK344197	–	MK340913

Table 3. Quantitative support (ML bootstrap and Bayesian posterior probability values) for three incompatible clades, representing the three best supported resolutions of the relationships of *Lycognathophis seychellensis* under ML and BI analyses of the 21-leaf, concatenated mt and nu dataset. Values in bold indicate best-supported resolution for each analysis. Abbreviations of genera: *Af* = *Afronatrix*; *He* = *Helophis*; *Hy* = *Hydraethiops*; *Li* = *Limnophis*; *Ly* = *Lycognathophis*; *Na* = *Natriciteres*.

Analysis	(<i>Ly</i> (<i>Hy</i> , <i>He</i> , <i>Af</i>))	((<i>Na</i> , <i>Li</i>)(<i>Hy</i> , <i>He</i> , <i>Af</i>))	(<i>Ly</i> (<i>Na</i> , <i>Li</i>))
RAxML (gene & codon position)	37	35	27
RAxML (gene)	40	36	24
MrBayes	0.28	0.59	0.12
BEAST	0.88	0.08	0.04

Table 4. Comparisons of fit of data to three trees with alternative best-supported resolutions of the sister-group relationships of *Lycognathophis seychellensis* for the 21-leaf concatenated mt and nu data. Above the diagonal are SH and AU test p values, respectively, for differences between ML trees. See Table 3 for support for each phylogenetic resolution. Below the diagonal are Bayes Factor scores for path (and stepping stone) sampling results under Bayesian analysis. * indicates best-supported resolution under MrBayes analysis; ** best-supported resolution under BEAST analysis. See Table 3 for abbreviations of genera.

	(<i>Ly</i> ((<i>Na</i> , <i>Li</i>)(<i>Hy</i> , <i>He</i> , <i>Af</i>)))	(<i>Ly</i> (<i>Na</i> , <i>Li</i>))	(<i>Ly</i> (<i>Hy</i> , <i>He</i> , <i>Af</i>))
(<i>Ly</i> ((<i>Na</i> , <i>Li</i>)(<i>Hy</i> , <i>He</i> , <i>Af</i>)))*		0.418; 0.298	0.501; 0.401
(<i>Ly</i> (<i>Na</i> , <i>Li</i>))	1.2 (1.1)		0.772; 0.682
(<i>Ly</i> (<i>Hy</i> , <i>He</i> , <i>Af</i>))**	4.0 (3.9)	2.8 (2.8)	

Table 5. Estimates of age (in Ma) of divergence between *Lycognathophis seychellensis* and its closest living (mainland sub-Saharan African) relative, for trees consistent with the three best-supported resolutions of the sister group of *L. seychellensis*. Results reported for TreePL and BEAST2 analyses of 73-leaf dataset. Values are means, with 95% highest posterior density interval in square brackets; values in and not in parentheses are from analyses implementing GTR and HKY models of sequence evolution, respectively.

Analysis	<i>L. seychellensis</i> sister group		
	(<i>Hy, He, Af</i>)	((<i>Na, Li</i>)(<i>Hy, He, Af</i>))	(<i>Na, Li</i>)
TreePL	35.68 (38.57)	39.31 (39.31)	41.27 (41.53)
BEAST2	35.31 [24.57–43.19] (43.09 [28.14–47.06])	43.34 [27.83–47.18] (42.41[27.06–47.82])	29.35 [24.11–37.84] (29.42 [25.51–39.28])

Table 6. Comparison of four models of ancestral area estimations implemented using BioGeoBEARS. Abbreviations: Ln L, log-likelihood; *d*, dispersal; *e* extinction; *j* jump parameter; *k*, number of parameters. AIC scores (sc) and weights (wt) are reported.

Model	Ln L	<i>K</i>	<i>d</i>	<i>e</i>	<i>j</i>	AIC sc	AIC_wt
DEC	-8.55	2	0.0000	0.0000	0.00	21.110	0.210
DEC+ <i>j</i>	-7.08	3	0.0000	0.0000	0.05	20.170	0.330
DIVALIKE	-11.14	2	0.0033	0.0029	0.00	26.280	0.016
DIVALIKE+ <i>j</i>	-7.3	3	0.0000	0.0000	0.06	20.600	0.270

FIGURE CAPTIONS

Fig. 1. Global distribution of sub-Saharan African (including Seychelles) natricine genera based on data from GBIF, Nagy et al. (2014), Chippaux & Jackson (2019), Conradie et al. (2020) and specimen vouchers from Port Elizabeth Museum (PEM). These genera comprise 14 currently recognised extant species as follows: *Afronatrix* (1 species), *Helophis* (1), *Hydraethiops* (2), *Limnophis* (3), *Lycognathophis* (1), and *Natriciteres* (6). Occurrence data for the one Seychelles and six of the mainland sub-Saharan African natricine genera (all except *Helophis*) were downloaded from the GBIF database (GBIF, 2019). Distribution data for *Helophis* were taken from Nagy et al. (2014). Spatial data were mapped using ArcGIS v. 10.5. (ESRI, The Redlands, CA). Sampling localities in mainland sub-Saharan Africa are labelled with numbers corresponding to those in Table 1. Precise locality information for *L. seychellensis* is presented in Table 1.

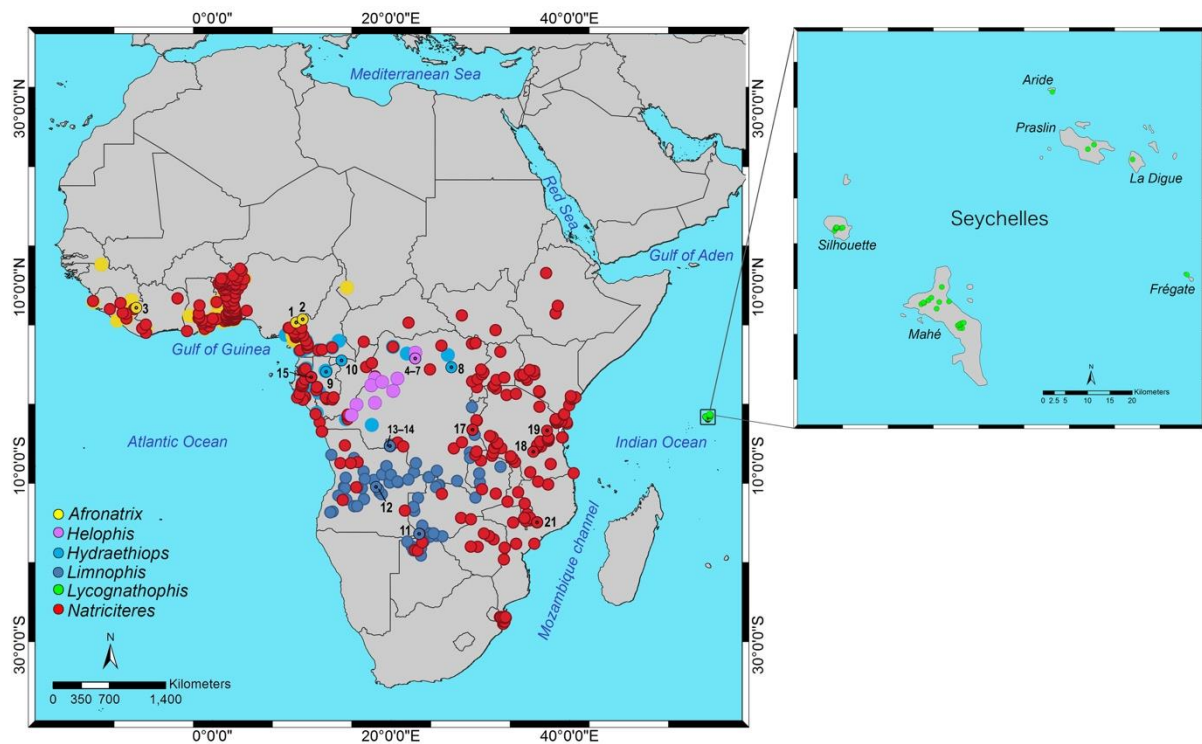


Fig. 3. BEAST chronogram showing divergence times for Seychelles and other (mainland) sub-Saharan African natricine snakes inferred from the 73-leaf, concatenated mt and nu dataset. Numbers at internal branches indicate mean divergence ages, with blue bars showing 95% highest posterior density intervals. Seychelles-Africa divergence is indicated with a *. Numbers in parentheses are sample codes (see Tables 1 and 2). See Fig. S2 for complete tree, including non-natricine alethinophidians, and values of 95% highest posterior density intervals. Ancestral area estimations (DEC + *j* model) at nodes represent areas before an inferred instantaneous speciation event, coloured boxes at tips indicate the current distribution of extant species.

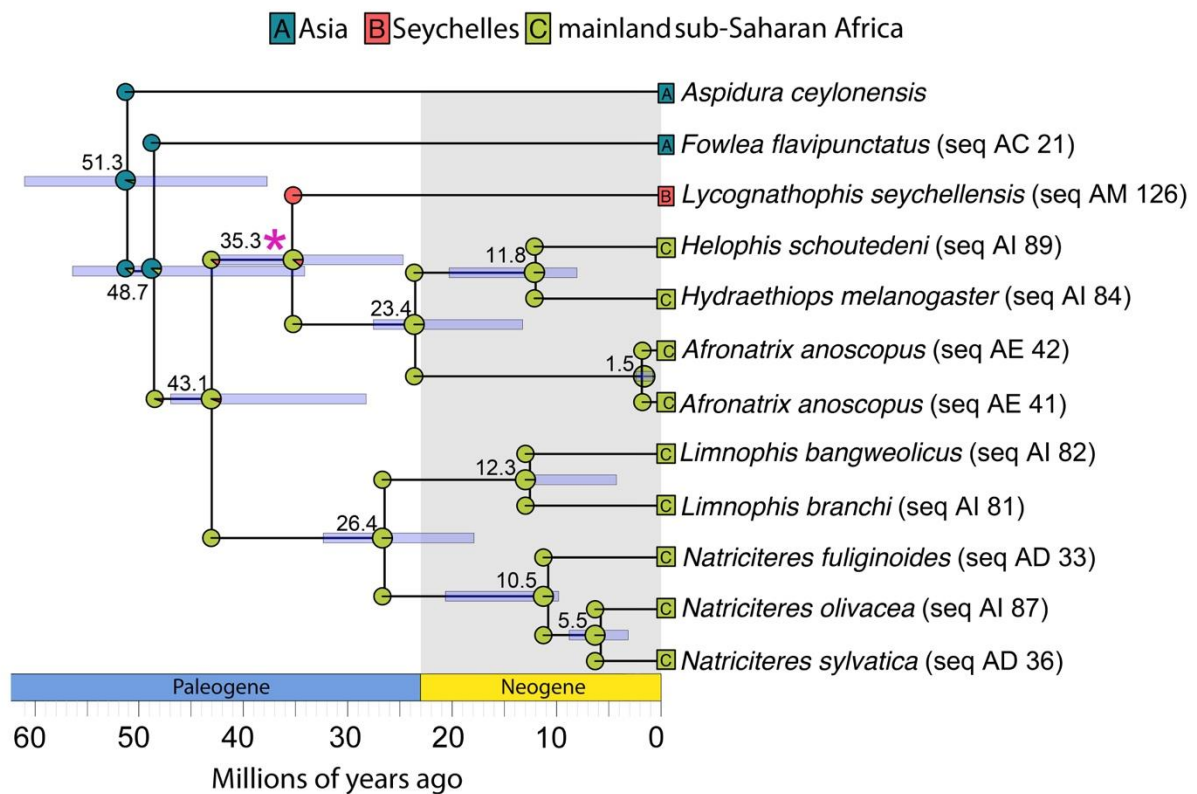
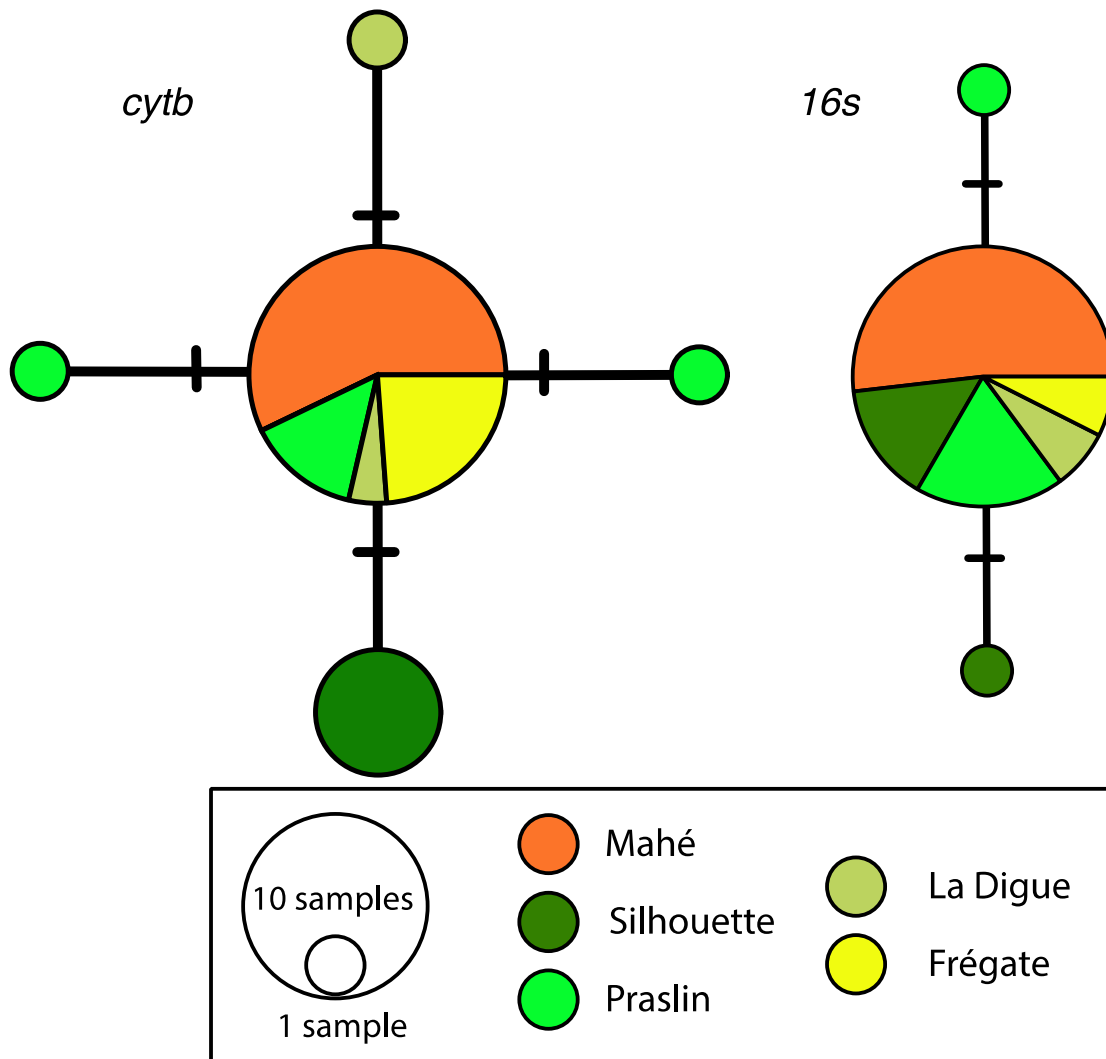


Fig. 4. Haplotype networks for mitochondrial *cytb* (n = 29) and *16s* (n = 30) for *Lycognathophis seychellensis* from five Seychelles islands using the median-joining (MJ) network algorithm. See Table 1 for sample and locality details.



Electronic supplementary material

Table S1. Primers used in this study for amplifying and sequencing target genes.

Table S2. Sequences used in the 157-leaf dataset, plus the two *Lycognathophis seychellensis* from GenBank (Vidal et al, 2008), which were used in this study only to calculate genetic distances. Sequences used in the 21-leaf dataset are indicated with *, outgroups are indicated with †, "-" = no data available

Table S3. Partitions and models of sequence evolution employed in analyses of the 73-leaf dataset. 1st, 2nd and 3rd denote codon position for protein-coding genes.

Table S4. Parameter values for fossil calibrations used in the BEAST divergence dating analysis. Ages in Ma. All maximum ages soft, except hard maximum for calibration 6.

Table S5. Partitions and models of sequence evolution employed in analyses of the 21-leaf dataset. 1st, 2nd and 3rd denote codon position for protein-coding genes.

Table S6. Support values for three incompatible clades representing the three competing resolutions of the relationships of *L. seychellensis* and other (mainland) sub-Saharan African natricines for the 21-leaf dataset. Values in each cell are given separately for analyses of mt - nu - mt+nu data (in that order). Highest support values for each analysis are highlighted in bold.

Fig. S1. A. Maximum Likelihood phylogeny and B. Bayesian phylogeny for the 157-leaf dataset.

Fig. S2. BEAST2 chronogram generated using concatenated-gene showing divergence times for natricine snakes and other alethnophidian snakes. Numbers at internal branches indicate posterior probabilities. Error bars indicate 95% highest posterior densities for node ages. Nodes C1–C7 are the seven calibrated nodes.

Fig. S3. BEAST2 chronogram generated using concatenated-gene with models of sequence evolution as suggested by partition finder. Numbers at internal branches indicate node ages. Error bars indicate 95% highest posterior densities for node ages.