Palatal morphology predicts the paleobiology of early salamanders

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Abstract

Ecological preferences and life history strategies have enormous impacts on the evolution and phenotypic diversity of salamanders, but the yet established reliable ecological indicators hinder investigations on the paleobiology of early salamanders. Here we statistically demonstrate using geometric morphometric analysis that both the shape of the palate and many non-shape variables particularly associated with vomerine teeth are ecologically informative in early stem- and basal crown-group salamanders. The morphology of the palate is heavily impacted by convergence constrained by feeding mechanisms and also exhibits clear stepwise evolutionary patterns with alternative phenotypic designs to cope with similar functional demand. Paleoecological disparities in early salamanders had took place before the Middle Jurassic and have achieved all ecological preferences in the Early Cretaceous. Metamorphosis is significant in the expansion of ecomorphospace of the palate in early salamanders. The common ancestor of salamanders is metamorphosed and terrestrial, and share unified lifestyles with other modern amphibians.
Introduction

Salamanders, anurans and caecilians are highly distinctive from one another in their morphology in both living and geologically the oldest known representatives from the Triassic (Ivachnenko, 1978; Ascarrunz et al., 2016; Pardo et al., 2017a; Schoch et al., 2020). As a result, the evolutionary origin(s) of modern amphibians have remained controversial since the late 19th century (Haeckel, 1866), with a number of extinct tetrapod groups in different ecological types at adult stages having been hypothesized as their purported ancestors, including: amphibamid (terrestrial) and branchiosaurid (terrestrial and aquatic) dissorophoid temnospondyls (Fröbisch and Schoch, 2009; Maddin and Anderson, 2012), stereospondylian (semiaquatic) temnospondyls (Pardo et al., 2017b), and several groups of lepospondyls (aquatic, semiaquatic or terrestrial; Marjanović et al., 2013). The specialized morphologies in modern amphibians are greatly impacted by ecology and their complex life history strategies (e.g., Wake, 2009), for example, even the earliest anuran Triadobatrachus and caecilian Chinlestegophis from the Triassic were already as morphologically well-adapted as their living relatives for aboveground and subterranean terrestrial living settings, respectively. Salamanders (or Caudata, the total group), on the other hand, have a more conservative body plan and more diversified ecological preferences when compared to anurans and caecilians (Deban and Wake, 2000; Bonett and Blair, 2017; Fabre et al., 2020), and have been frequently used as comparative analogues for inferring the paleoecology of extinct tetrapods (Schoch and Fröbisch, 2006; Fröbisch and Schoch, 2009). However, the evolutionary history of paleoecology in modern amphibians and particularly in early salamanders has received insufficient attention.
Cryptobranchoidea is the most primitive clade of the crown group salamanders (Urodela) and contains two subclades: Pancryptobrancha (crown + stem cryptobranchids; Vasilyan et al., 2013) and Panhynobia (crown + stem hynobiids; Jia et al., 2021a). The two subclades are united by a set of synapomorphies as sister-group taxa (Dunn, 1922; Estes, 1981; Jia et al., 2021a), but are different from each other in life history strategies and ecological preferences at their respective adult stage: most pancryptobranchans are neotenic or partially metamorphosed and live in water permanently by retaining larval features (e.g., gill slits), albeit the pancryptobranchan Aviturus from the Paleocene was interpreted as semiaquatic with an unknown life history strategy (Vasilyan and Böhme, 2012). Alternatively, panhynobians are predominantly metamorphosed, except that the stem hynobiid Regalerpeton from Early Cretaceous (Rong, 2018) and some populations of the living hynobiid Batrachuperus londongensis are neotenic (Jiang et al., 2018). Post-metamorphosed hynobiids have lost larval features and are characterized by an anterolaterally directed palatal ramus of the pterygoid, and are able to live in water (e.g., Paradactylodon), on land (e.g., Hynobius) or are semiaquatic (Ranodon) outside of breeding season (Kuzmin and Thiesmeier, 2001; Fei et al., 2006; Materials and Methods).

Cryptobranchoidea are critical in understanding the paleoecology of early salamanders because the earliest fossil cryptobranchoids represent the oldest known crown urodeles; and disparities in both life history strategies and ecological preferences have been underway among these earliest crown urodeles from the Middle Jurassic Bathonian, including ‘Kirtlington salamander B’ from the UK, Kiyatriton krasnolutskii from Russia, and Chunerpeton, Neimengtriton, and Jeholotriton from China (Evans and Milner, 1994; Gao et
al., 2013; Skutschas, 2016; Jia et al., 2021a). Both *Chunerpeton* and *Jeholotriton* are neotenic as confirmed by the presence of external gills and high caudal dorsal fin in adult specimens (Gao and Shubin, 2003; Wang and Rose, 2005), whereas *Neimengtriton* is the oldest metamorphosed and semiaquatic cryptobranchoid (Jia et al., 2021a; see below). The ‘Kirtlington salamander B’ and *K. krasnolutskii* are both represented by fragmentary materials and their paleoecology unfortunately remains unknown. In contrast, other contemporaries (e.g., *Kokartus, Marmorerpeton*) from the Middle Jurassic Bathonian of UK, Russia and Kyrgyzstan are all neotenic and aquatic at their adult stage, and have been classified as stem urodeles by the absence of spinal nerve foramina in the atlas that characterizes crown Urodela (Ivachnenko, 1978; Evans et al., 1988; Skutschas and Krasnolutskii, 2011; Skutschas and Martin, 2011; Skutschas, 2013; Skutschas et al., 2020).

The only known pre-Jurassic salamander, *Triassurus* from the Middle/Upper Triassic of Kyrgyzstan, is merely represented by two larval specimens with no clue to its paleoecology at adult stage (Schoch et al., 2020).

To date, seven other basal cryptobranchoids have been reported from the Upper Jurassic to Lower Cretaceous of northern China: *Laccotriton, Liaoxitriton, Linglongtriton, Nuominerpeton, Pangerpeton, Regalerpeton* and *Sinerpeton*, most of which are represented by articulated specimens and have been recently recovered as stem hynobiids or hynobiid-like taxa (see Gao et al., 2013; Jia and Gao, 2016; 2019; Jia et al., 2021a). Besides the neotenic *Regalerpeton* as aforementioned, living habitat preferences of these metamorphosed taxa and paleoecological disparity patterns of Cryptobranchoidea remain largely unexplored mainly due to yet established osteological indicators for ecology (see
Discussion). The configuration of vomerine teeth has long been identified as useful for the classification of living cryptobranchoids (Zhao and Hu, 1984) and was recently claimed to be ecologically informative (Jia et al., 2021b), but such statements have not received rigorous tests with inclusion of fossil taxa.

Our series of studies on living and fossil cryptobranchoids noticed that besides vomerine teeth, the palate varies in shape and proportion, and could potentially serve as an indicator for paleoecological reconstruction (Jiang et al., 2018; Jia et al., 2019; 2021a; 2021b; Figure 1 and Figure 1—figure supplements 1–23). To test these hypotheses and to address the constraints underlying the morphological disparity of the palate, here we conducted a 2D landmark-based geometric morphometric analysis on the palate of all living and most aforementioned basal genera of cryptobranchoids, stem and other basal crown urodeles based primarily on micro-CT scanned specimens. We statistically investigated disparity patterns within the morphospace of the palate with respect to ecological preferences, life history strategies and taxonomic affiliations. Based on a time-calibrated cladogram we established for fossil and living cryptobranchoids (Jetz and Pyron, 2018; Jia et al., 2021a), we further quantified the evolutionary rate of the palate and reconstructed the ancestral states for ecological preferences, life history strategies, palate shape and vomerine tooth configurations of the respective common ancestor of Panhynobia, Pancryptobrancha, Cryptobranchoidea, Urodela, and Caudata. We demonstrate that the palate is a reliable proxy in ecological reconstructions for early salamanders, and the morphospace of the palate is predominantly shaped by ecological constraints and also displays stepwise evolutionary pattern. Within Urodela, morphological disparities of the palate start ahead of the
Panhyobia-Pancryptobrancha split before the Middle Jurassic Bathonian and came to
occupy most of the current morphospace by the Early Cretaceous. Our results show that the
common ancestor of Caudata is metamorphosed and terrestrial and support modern
amphibians share similar lifestyles in their early evolution.

Results
In ventral view of the palate, the anteromedial fenestra is present between the vomer and the
upper jaw in most early salamanders and is only absent in living cryptobranchids (Figure
1—figure supplements 1–23). The paired vomers medially articulate with each other in most
taxa and posteriorly to different extents overlap the anterior part of the cultriform process of
the parasphenoid and/or the orbitosphenoid. The teeth are closely packed as a continuous
tooth row positioned along the anterolateral periphery of the vomer in cryptobranchids but
have diversified configurations in other taxa. The parasphenoid is a sword-like, azygous bony
plate with its anterior part articulating dorsally with the orbitosphenoid and its posterior part
flooring the otic capsule.

Morphospace and shape disparity patterns of the palate in early salamanders. The
palate is symmetric about the midsagittal plane of the skull, with symmetric shape
components accounting for 96.15% of the total shape variation and the left-right asymmetry
accounting for the remaining 3.85%. The shape and the size of the palate, with the latter
represented by the centroid size (CS), are significantly correlated as revealed by the standard
multivariate regression between log (CS) (independent variable) and symmetric shape
components (dependent variable) across 70 specimens ($R^2 = 9.3331\%; p < 0.001; F = 6.9977; Z = 3.918$) and 34 species ($R^2 = 13.276\%; p < 0.001; F = 4.8985; Z = 3.1691$). However, when phylogenetic relationships of the 34 species were factored in, the association between size and shape of the palate is no longer significant as shown in the evolutionary allometry analysis ($p = 0.1583$). Such inconsistency between standard and evolutionary allometry analyses is related to the fact that the CS of the palate has a strong phylogenetic signal (Blomberg’s $K = 0.997, p = 0.001, Z = 4.1921$) and, hence, the CS accounts for an even smaller amount of shape variations of the palate ($R^2 = 4.494\%; F = 1.5059; Z = 1.0141$) when evolutionary history among species were counterbalanced in the evolutionary allometry analysis. To eliminate impacts from both asymmetry and allometry on the spatial patterns of the palate, residuals from the multivariate regression of symmetric shape components on log (CS) were retained for downstream statistical analyses.

To visualize the spatial patterns of the palate in the morphospace, on the basis of the size-corrected (allometry-free) 24-landmark dataset, we conducted a standard PCA across 70 specimens; and we also conducted three other types of PCA across 34 species with the time-calibrated cladogram and ancestral internal nodes projected into the morphospace (Figure 2—figure supplements 1–3), including: a phylomorphospace analysis (PA), phylogenetic principle component analysis (Phylo-PCA) and a phylogenetically aligned components analysis (PaCA). The first three PC axes in each of the four PCAs collectively measure up to about 70% of total shape variances (Supplementary file 1A). Within the phylomorphospace defined by PCs 1–2 (Figure 2B and Figure 2—figure supplement 1b), aquatic and terrestrial living species of Cryptobranchoidea are generally located along the
positive and negative interval of the PC 2 axis (21.35%), respectively. Most taxa are ecologically exclusive at the genus level, except that *Liua* and *Paradactylodon* are the only two living hynobiid genera with species occupying both the aquatic (*Liua shihi, Paradactylodon mustersi*) and terrestrial (*Liua tsinpaensis, Paradactylodon persicus*) zones. The living semiaquatic hynobiid *Ranodon sibiricus* occupies at a location intermediate between the aquatic and terrestrial zones. Shape changes of the palate relative to the mean shape of all terminal and internal taxa along the positive values of PC 2 (aquatic zone) involve an anteromedial extension of the vomer and therefore a reduction of the anteromedial fenestra, a shrinkage of the anterolateral and posterolateral borders of the vomer and the width of the cultriform process of the parasphenoid, and an anterior extension of the parasphenoid. Oppositely, the negative values of PC 2 (terrestrial zone) characterize a shrinkage of the anteromedial border of vomer and therefore a posterolaterally expanding anteromedial fenestra, a laterally widening retrochoanal process of the vomer and the cultriform process of the parasphenoid, and an anteroposteriorly shortening parasphenoid. On the other hand, basalmost crown urodeles (e.g., *Beiyanerpeton, Chunerpeton, Pangerpeton*) and karaurids are largely separated from living taxa along the PC 1 axis (37.49%), with stem hynobiids widely scattered in the phylomorphospace and lying within either the aquatic (*Liaoxitriton*) or terrestrial (*Linglongtriton and Nuominerpeton*) zone. The only known aquatic stem hynobiid *Regalerpeton* is situated at a region of the aquatic zone occupied by other aquatic fossil taxa (e.g., *Beiyanerpeton, Chunerpeton*), whereas the semiaquatic stem hynobiid *Neimengtriton* occupies a location closer to terrestrial than either aquatic zone or the semiaquatic zone of extant hynobiids. From the largest to the smallest value of PC 1, both the
vomer and the parasphenoid have an anteroposterior extension and the cultriform process of
the parasphenoid changes from an anteriorly widened plate with an indented anterior edge
into a bilaterally narrowed plate with a pointed anterior edge.

The evolutionary history among species has a moderate but significant contribution in
the formation of the spatial patterns in the phylomorphospace (phylogenetic signal: observed
$K_{mult} = 0.4154$, $p = 0.001$, $Z = 4.9856$). When the multivariate shape data of the palate were
maximumly aligned with phylogenetic signal (Figure 2C and Figure 2—figure supplement
1c), fossil taxa can be roughly divided from living taxa along PaCA-C 1. Both crown and
stem taxa of Panhynobia are more compactly clustered than that seen in the
phylomorphospace created by PA, and are distinct from the Pancryptobrancha and the region
occupied by karaurids, basal cryptobranchooids and salamandroids; but taxa in different types
of ecological preference are mixed together. By contrast, when the phylogenetic signals of the
 palate were eliminated by the Phylo-PCA (Figure 2D and Figure 2—figure supplement 1d),
the overall spatial patterns among species are essentially preserved, albert slightly rotated
clockwise, as observed in the phylomorphospace. In this phylogeny-free morphospace, the
common ancestors of Pancryptobrancha and of Hynobiidae lie within the aquatic zone and
are tightly associated with living cryptobranchids and the stem urodele *Kokartus*, respectively.
Alternatively, the common ancestors of Panhynobia, Cryptobranchoidea, Urodela and
Caudata lie within the terrestrial zone and are adjacent to three highly derived living
hynobiids, respectively, *Pseudohynobius shuichengensis*, *Pseudohynobius jinfo* and *Liua
tsinpaensis*.

Our pairwise comparison (Supplementary file 1B–E) and phylogenetic procrustes
ANOVA reinforce that the shape of the palate is significantly different among groups of species that are classified by ecological preference ($R^2 = 28.079\%; p = 0.001; F = 4.3740; Z = 3.4838$) and life history strategy ($R^2 = 6.797\%; p = 0.006; F = 3.1766; Z = 2.4673$), but not by taxonomic affiliations neither at the genus ($R^2 = 68.711\%; p = 0.248; F = 1.2549; Z = 0.78492$) nor family ($R^2 = 12.858\%; p = 0.581; F = 0.8263; Z = -0.21234$) level. With regard to the life history strategy, living cryptobranchids are separated from all Mesozoic neotenic taxa including Beiyanerpeton, Chunerpeton, karaurids and Regalerpeton by the vast majority of the metamorphosed taxa, which occupy most of the morphospace of the palate (Figure 2—figure supplement 4). The single living neotenic hynobiid Batrachuperus londongensis is situated between the two neotenic groups as aforementioned and lies alongside its metamorphosed conspecifics and those all collectively overlap with other aquatic hynobiids. When the seven-landmark-dataset for the right vomer was analyzed following the same procedure, the purported semiaquatic basal pancryptobranchian Aviturus is nested within the terrestrial zone along with living metamorphosed taxa (Figure 2—figure supplement 5).

**Evolutionary rates of the palate.** The palate exhibits considerable differences in morphological disparity and evolutionary rates across regions represented by the 24 landmark points as calculated under a Brownian motion evolutionary model (Figure 3; Supplementary file 1B–H). The vomer evolves about two times faster than the parasphenoid, and such a pattern is also supported by the fact that the vomer has a higher phylogenetic signal ($K_{mult} = 0.6407$) than that of the palate ($K_{mult} = 0.4154$). The posterior border of the vomer and the medial most part of the choanal notch are the fastest and the slowest evolving parts in the
palate, respectively. In the parasphenoid, however, the highest evolutionary rates are
concentrated on the anterior end, posterior end and the lateral alae, and the lowest
evolutionary rates are located at the posterior contact between the cultriform process and the
orbitosphenoid and the junction area where the cultriform process merges with the lateral
alae.

Species in different ecological ($p = 0.002, Z = 2.6284$) and taxonomic ($p = 0.035, Z = 1.7594$) groups vary greatly in evolutionary rate of the palate (Supplementary file 1H).

Namely, aquatic and terrestrial taxa are comparable with each other in the evolutionary rate
of the palate, the vomer and the parasphenoid, and both of which evolve at a rate that is less
than half of that in semiaquatic taxa championed by *Neimengtriton*. Stem hynobiids have the
highest evolutionary rates, followed successively by basal cryptobranchoids,
pancryptobranchans, crown hynobiids and non-cryptobranchoid basal salamanders. Neotenic
and metamorphosed taxa are not significantly different from each other in evolutionary rates
($p = 0.843, Z = -1.1341$).

**Non-shape covariates from the vomer.** Similar to shape variables, allometry has a
significant impact on most of the five continuous non-shape covariates of the palate (length
ratios between parasphenoid and palate, and that between vomer and palate; width ratios
between vomerine tooth row [VTR] and vomer, and between outer and inner branch of the
VTR; and vomerine tooth number) when each of them is statistically regressed on the log
(CS). We thus use residuals from the regression for boxplots to visualize the distribution
patterns of the covariates in three ecological groups across 70 specimens, and use the Mann
Whitney U test to analyze if the covariates are reliable ecological indicators (Figure 3). In line with shape changes revealed from PCA, aquatic and terrestrial species are significantly different in all of the five covariates, with aquatic taxa having proportionally a longer parasphenoid (median: ~0.84), a shorter vomer (median: ~0.28), a higher ratio in outer/inner branch of the VTR (median: ~1.6), a narrower VTR (except neotenic taxa; median: ~0.37) and fewer vomerine teeth (median: 7) than that of terrestrial taxa (i.e., ~0.78, ~0.33, ~0.58, ~0.64, 16). Semiaquatic species cannot be confidently differentiated from aquatic or terrestrial taxa in any of the five covariates, but interestingly their parasphenoid length and VTR width resemble those in terrestrial taxa and their vomer length, outer/inner branch VTR width ratio and teeth numbers are more similar to aquatic taxa. The contingency table and Cochran-Mantel-Haenszel statistical test (Figure 3—figure supplement 1 and Supplementary file 1I) show that the arrangement and position of VTR are both significantly correlated to ecological preference and life history strategy. The VTR in anterior and middle position of the vomer or being arranged obliquely or parallel to the marginal tooth row are only present in aquatic taxa, with the VTR located anteriorly exclusively present in neotenic species. The VTR in the mid-posterior position of the vomer or are transversely arranged is only present in metamorphosed taxa with all kinds of ecological preferences. Species with VTR in the posterior position or being transversely arranged are more likely to be terrestrial than aquatic or semiaquatic.

Ancestral states of the palate. The ancestral states in both the shape and non-shape covariates of the palate were reconstructed for the common ancestors of Hynobiidae,
Panhynobia, Pancryptobrancha, Cryptobranchoidea, Urodela, and Caudata, respectively, using maximum likelihood (Supplementary file 1J). When compared to the mean shape of the palate, the configurations of the palate in the respective common ancestors of Caudata, Urodela and Cryptobranchoidea are characterized by a shortened parasphenoid with an anteriorly widened cultriform process, an anteriorly shortened but posteriorly widened vomer. This configuration coincides with our reconstructions that the respective common ancestors of Caudata, Urodela and Cryptobranchoidea are metamorphosed and terrestrial (Figure 3—figure supplement 2). Upwards from the common ancestor of Cryptobranchoidea, the vomer is first elongated anteroposteriorly towards the common ancestor of Panhynobia and is followed by an anteroposterior shrinkage and a lateral shift from its counterlateral towards the common ancestor of Hynobiidae; whereas the vomer is elongated anteroposteriorly with a reducing trend of the anteromedial fenestra towards the common ancestor of Pancryptobrancha. The parasphenoid, however, is anteriorly elongated and the cultriform process is anteriorly narrowed bilaterally stepwise from the common ancestor of Cryptobranchoidea towards that of Hynobiidae and Pancryptobrancha. The parasphenoid in the common ancestor of Pancryptobrancha also has a unique lateral expansion at the base of the cultriform process and an extension posteriorly. The vomerine tooth row for the respective common ancestors of Caudata, Urodela and Cryptobranchoidea is in parallel with the marginal tooth row and is located in the posterior part of the vomer with the outer longer than the inner branch. The VTR remains the similar configuration except is transversely arranged in the common ancestors of Panhynobia and Pancryptobrancha, then the VTR shifted to the middle part of the vomer in the common ancestor of Hynobiidae.
Discussion

Among modern amphibians, anurans and caecilians are predominantly metamorphosed and their postmetamorphosed adults are terrestrial. In contrast, salamanders have their ecological preference decoupled from life history strategy, especially in metamorphosed taxa such as living and extinct hynobiids where postmetamorphosed adults live in water, on land or are semiaquatic (Fei et al., 2006; Jia and Gao, 2016; Jia et al., 2021a). The discrepancy in ecological preference between salamanders and anurans and caecilians is unhelpful in understanding the purported monophyletic origin of modern amphibians from Temnospondyli.

Previous studies on fossil salamanders generally focus on the taxonomy, whereas the paleoecology and the evolutionary history of the ecological decoupling particularly in metamorphosed taxa had received insufficient attention, which to some extent are contributed by the lack of taphonomic analyses (Wang et al., 2019). The main obstacles are (1) sufficient and reliable ecological indicators in the bony skeleton have not yet been established for extant cryptobranchoids and early salamanders (Xiong et al., 2016; Jia et al., 2021b) and (2) soft anatomical structures (e.g., labial fold, caudal fin) and stomach contents that are ecologically informative, as commonly seen in lacustrine deposits of certain neotenic species (Dong et al., 2012), are rarely preserved in metamorphosed taxa. Here we quantitively investigate the turnover of paleoecology in the earliest known salamanders based on the shape and non-shape variables of the palate. Our results shed light on the interactions between morphology and paleoecology, and the underlying mechanisms governing ecomorphological diversity among early salamanders along the rise of modern amphibians.
As shown in the phylomorphospace, the shape of the palate in early salamanders is heavily impacted by convergence resulted from ecological disparity, as extant cryptobranchoids and fossil taxa in different ecological preferences occupy distinctive zones in the morphospace. Such spatial patterns remain essentially unaltered after phylogenetic signals were eliminated by the Phylo-PCA, along the first axis of which aquatic and terrestrial taxa regardless of their phylogenetic closeness are separated from each other, confirming our hypothesis that the shape of the palate is a reliable ecological indicator in early salamanders as reported in other living salamanders (Fabre et al., 2020). As shown above, non-shape variables of the palate particularly with regard to the vomerine tooth row are also ecologically informative. The palate in aquatic species is characterized by a reduction of the anteromedial fenestra, a shrinkage of the vomer laterally and anteroposteriorly with few vomerine teeth, a high ratio in the outer/inner branch of the VTR, a narrow VTR, and an elongate parasphenoid. On the other hand, the palate in terrestrial species is characterized by a posterolateral expansion of the anteromedial fenestra, an anteroposteriorly elongated vomer and a widened retrochoanal process with more vomerine teeth, a low ratio in outer/inner branch of the VTR, a widened VTR, and an anteroposteriorly shortened parasphenoid. The palate in extant semiaquatic taxa is transitional in both the morphospace and many non-shape variables between aquatic and terrestrial taxa as mentioned above, whereas the palate of the only known fossil semiaquatic Neimengtriton (Middle Jurassic) is configured more like that of terrestrial taxa as argued in the original study (Jia et al., 2021a), indicating that Neimengtriton is probably more adapted to terrestrial environment despite being semiaquatic. The respective common ancestors for Caudata, Urodela, Cryptobranchoidea and Panhynobia,
the basal cryptobranchoid *Pangerpeton* (Late Jurassic) and the stem hynobiids *Linglongtriton* (Late Jurassic) and *Nuominerpeton* (Early Cretaceous) are terrestrial, whereas other stem hynobiids *Liaoxitriton* and *Regalerpeton* (Early Cretaceous) and the respective common ancestors for Hynobiidae and Pancryptobrancha are aquatic, demonstrating that ecological shifts occurred frequently in the evolution of early salamanders. The earliest known fossil pancryptobranchian *Aviturus* is likely metamorphosed and terrestrial based on the shape and non-shape variables of the vomer, but this hypothesis awaits to be tested by the discovery of a more completely preserved palate.

Morphological adaptations in the palate and perhaps in other cranial features (e.g., hyobranchial apparatus; see Jia et al., 2021b) to aquatic and terrestrial living settings are likely constrained by feeding mechanisms (Regal, 1966; Reilly and Lauder, 1990). In extant cryptobranchoids, terrestrial species grasp prey items with their jaws and/or by their tongue protruding from the mouth; the captured prey is then transported from the snout into the esophagus with the assistance of several cyclical movements of tongue and hyobranchial apparatus that press and reposition the prey against the palate (Deban and Wake, 2000; Wake and Deban, 2000). Such feeding mechanisms in terrestrial cryptobranchoids demand a sticky tongue moisturized by the intermaxillary or internasal gland that is housed above an expanded anteromedial fenestra, a wide snout contributed by lateral expansions of the vomer, many vomerine teeth to efficiently subdue the prey, and a shortened parasphenoid to reduce the distance of intraoral transportation. The posteriorly elongated inner branch of the VTR as represented by derived terrestrial hynobiids *Hynobius* and *Salamandrella* would serve as toothed surfaces that can further facilitate tongue manipulations and transportations of small
prey items posteriorly as in other more sophisticated terrestrial feeders (e.g., salamandrids and plethodontids; Regal, 1966). By contrast, aquatic salamander species capture their prey primarily by suction feeding in which the prey is carried into the biting range of the jaws by a current of water created by retraction and expansion of buccal walls, and giant salamanders can even perform asymmetrical strikes in water via unilateral jaw and hyobranchial movements (Gillis and Lauder, 1994). Intraoral transportations of the prey in aquatic species are more efficient than that of terrestrial species, and typically involve repeating the same procedure as in the initial suction strike and are occasionally assisted by tongue manipulations against the palate (Regal, 1966). The limited usage of the tongue in aquatic prey capture diminishes the need for moisturization from the intermaxillary or internasal gland which, in turn, may lead to reduction and closure of the anteromedial fenestra as seen in aquatic species. An anteroposterior expansion of the parasphenoid increases buccal volume, and the shrinkage of the vomer along both the mediolateral and anteroposterior dimensions reduce the snout size; both features act in concert to boost the success of feeding attempts by creating a high negative buccal pressure. A narrow VTR with few teeth, most of which are located on the outer branch, likely serves to reduce hindering the influx of water and prey.

Whichever mode of prey capture and intraoral transportation cryptobranchoids perform, the palate undergoes a similar mechanical stress distribution pattern as found in species with different types of ecological preference and life history strategy (Fortuny et al., 2015: Andrias davidianus; Zhou et al., 2017: Salamandrella keyserlingii). Such findings reinforce that alternative morphological designs of certain features in the palate cope well with similar functional demands, but do not necessarily indicate that other feeding-related features of the
palate and other regions of the cranium would also exhibit such extensive convergence. For example, the ceratohyal has been argued to be ecologically informative in extant cryptobranchoids, as the ceratohyal is consistently ossified at its posterior end in aquatic species but remains cartilaginous in terrestrial taxa (Xiong et al., 2016; Jia et al., 2021b); however, the ceratohyal remains cartilaginous in all known early fossil stem and crown urodeles, and ossifications of the ceratohyal must represent a derived feature independently evolved in crown cryptobranchoids (Jia et al., 2021b). Interestingly, the palate of extant cryptobranchoids as shown in biomechanical analyses (Fortuny et al., 2015; Zhou et al., 2017) has the highest stress level concentrated in the most posterior part of the vomer and its sutural area with the parasphenoid and orbitosphenoid, which surprisingly correspond to the place with the highest evolutionary rate in the palate as revealed by our study. Here we recognize a clear stepwise evolutionary pattern at the vomer-parasphenoid-orbitosphenoid sutural area in early salamanders (Figure 4). In stem \textit{(Karaurus, Kokartus)} and many basal crown urodeles from the Jurassic and the reconstructed respective common ancestors for Panhynobia, Cryptobranchioidea, Urodela, and Caudata, the vomer has a limited contact with the parasphenoid, the cultriform process of the parasphenoid is anteriorly expanded bilaterally, and the orbitosphenoid has no anteroventral processes (see below) or in the case of \textit{Qinglongtriton} remains completely cartilaginous. In three stem hynobiids \textit{(Liaoxitriton, Nuominerpeton, Regalerpeton)} and the reconstructed ancestors for Hynobiidae and Pancryptobrancha from the Cretaceous and another stem hynobiid \textit{Linglongtriton} from the Late Jurassic, the vomer has an increased contact with the parasphenoid, the cultriform
process of the parasphenoid is anteriorly constricted, and the orbitosphenoid lacks the anteroventral process. In living cryptobranchoids, the vomer is in more extensive contact with the parasphenoid and orbitosphenoid (Trueb, 1993: Cryptobranchus; Kuzmin and Thiesmeier, 2001: Ranodon; Jiang et al., 2018: Batrachuperus; Jia et al., 2021b: Pseudohynobius), the cultriform process of the parasphenoid is anteriorly constricted bilaterally, and the orbitosphenoid has a thick anteroventral process ossified from neighboring cartilages in the nasal capsule that projects medially and firmly overlaps the cultriform process of the parasphenoid. If similar biomechanical stress patterns existed in early salamanders, the stress and strain applying to the vomer/parasphenoid sutural area would be equally offset by either a wide and thin cultriform process of parasphenoid with limited support from the vomer and the absence of anteroventral process of the orbitosphenoid as seen in Jurassic taxa, or a narrow cultriform process thickened by more extensive overlapping with the vomer as seen in Cretaceous and living taxa, and by ossification of the anteroventral process of the orbitosphenoid as seen solely in living taxa.

It is worth emphasizing that the phenotypic diversity of the palate is variously affected by neoteny and metamorphosis. Neoteny is notable in producing convergent characters through truncation of normal developmental trajectories and its indirect role associated with constraints from the aquatic environment (Wiens et al., 2005). Most early stem and crown urodeles are neotenic (e.g., Chunerpeton, Kokartus) and their phenotypic diversity of the palate is truly confined to a restricted area in the spatial-temporal morphospace. However, extant neotenic cryptobranchids and the hynobiid Batrachuperus londongensis bear an increased phenotypic diversity of the palate as compared to fossil neotenic taxa, and are
separated from one another within the morphospace with the neotenic *B. londongensis* even overlapping with its metamorphosed conspecifics, indicating that constraints imposed by ecology have more influence than neoteny in the morphogenesis of the palate. The eco-morphospace of the palate is indeed greatly and rapidly expanded by metamorphosed taxa represented by basal cryptobranchoids and most stem hynobiids as these taxa have the highest evolutionary rate in the palate. The common ancestors of Caudata, Urodela and Cryptobranchoidea are reconstructed to be metamorphosed and terrestrial as evidenced by the shape and non-shape variables of the palate. Disparities of ecological preference among metamorphosed taxa must have taken place before the Middle Jurassic Bathonian, because one of the oldest-known crown urodeles, *Neimengtriton*, is metamorphosed and semiaquatic as evidenced by the presence of a low but pliable dorsal caudal fin at adult stage. Most of the current scope in the phenotypic diversity of the palate possessed by modern cryptobranchoids have been achieved by Early Cretaceous. Our results show that metamorphosis is the ancestral life history strategy in salamanders, and we demonstrate that metamorphosis patterned with the biphasic ecological preference (aquatic larvae + terrestrial adults) as shared with anurans and caecilians is significant in the rise and diversification of modern amphibians.

**Materials and methods**

**Experimental design, specimens, palate.** Our study includes 60 wet specimens (preserved in formalin) that represent 25 living species (29% of Hynobiidae, 75% of Cryptobranchidae) in all 12 living genera of Cryptobranchoidea (*Andrias, Batrachuperus, Cryptobranchus*, *...*)
Hynobius, Liua, Onychodactylus, Pachyhynobius, Paradactylodon, Protohynobius, Pseudohynobius, Ranodon, and Salamandra), and one fossil skeleton for each of five stem hynobiids (Liaoxitriton zhongjiani, Linglongtriton daxishanensis, Neimengtriton daohugouensis, Nuominerpeton aquilonaris, and Regalerpeton weichangensis), one basal panchryptobranchan (Aviturus exsecratus), two basal cryptobranchoids (Chunerpeton tianyiensis and Pangerpeton sinensis), one basal salamandroid (Beiyanerpeton jianpingensis) and two stem urodeles (Karaurus sharovi and Kokartus honorarius; Dataset in online Dryad).

In this study, each species is represented by one to three specimens, except for the only facultatively neotenic cryptobranchoid B. londongensis where both neotenic and metamorphosed populations are each represented by three specimens. Most of the specimens are accessioned in the following seven institutional collections: Capital Normal University (CNU), Beijing, China; Chengdu Institute of Biology (CIB), Chengdu, Sichuan Province, China; Field Museum of Natural History (FMNH), Chicago, USA; Liupanshui Normal University (HNUL), Liupanshui, Guizhou Province, China; Peking University of Paleontological Collections (PKUP), Beijing, China; Zhejiang Museum of Natural History (ZMNH), Hangzhou, Zhejiang Province, China; and Zunyi Medical University (ZMU), Zunyi, Guizhou Province, China. For specimens that we were not able to examine firsthand, we used publicly available micro-CT scan data of 12 living cryptobranchoid specimens (Dataset in online Dryad) from the MorphoSource platform (MorphoSource.org) and published images for five fossil taxa, including Aviturus (Vasilyan and Böhme, 2012: fig. 3), Chunerpeton (Gao and Shubin, 2003: fig. 1), Karaurus (Ivachnenko, 1978: fig. 1), Kokartus (Skutschas and Martin, 2011: fig. 9) and Regalerpeton (Rong, 2018: fig. 5).
Besides the CT scan data obtained from MorphoSource, four fossil and all living specimens were micro-CT scanned using the following three high-resolution CT scanners: a Nikon XT H 320 LC scanner in the Industrial Micro-CT Laboratory at China University of Geosciences (Beijing); a GE Phoenix v/tome/x 240kv/180kv scanner in the PaleoCT Lab at The University of Chicago; and a Quantum GX micro-CT Imaging System (PerkinElmer\textsuperscript{®}, Waltham, USA) at CIB (Dataset in online Dryad). No filter was used because beam hardening artifacts were not encountered during CT scanning. The voxel size of the volume files generated from CT scans ranges between 14.52–87.32 μm (see detailed parameters in Dataset in online Dryad). File processing including segmentation and rendering were accomplished by VG Studio Max (version 2.2; Volume Graphics, Heidelberg, Germany), and images in jpeg format showing the ventral view of the skull were exported for landmark acquisition after digital removal of the mandible and the hyobranchium from the virtual models.

The palate in adult specimens of cryptobranchoids consists of the partes palatina of the premaxilla and maxilla, paired vomers and a single median parasphenoid. An independently ossified palatine is present in few fossil taxa and is absent in the remaining, and is therefore not included in this study as a compromise between taxa sampling and landmarks collection. The pas palatina of both the premaxilla and maxilla is a narrow bony ledge and invariably contributes to a small portion of the palate by posteriorly articulating with the vomer, and hence is not considered in this study (Figure 1A). Both the vomer and parasphenoid are dorsoventrally flattened bony plates and are homologous across species studied here, and thus are ideal for landmark collections for both living and fossil specimens, considering that the
palate in all available fossil cryptobranchoids is dorsoventrally preserved. All anatomical terms used here follow Trueb (1993) unless otherwise stated.

Cladograms. For multivariate phylogenetic comparative analyses (evolutionary allometry analysis, phylogenetic signal analysis, phylogenetic principle component analysis, phylomorphospace analysis, phylogenetic alignment component analysis and phylogenetic ANOVAs; see below), a maximum clade credibility tree (MCC; a target tree with maximum sum of posterior probabilities on its internal nodes) with 24 living taxa in our dataset was created by the software TreeAnnotator (version 1.10.4; Drummond et al., 2012) based on 10,000 random, time calibrated trees from a recent Bayesian posterior distribution analyses of living amphibians (Jetz and Pyron, 2018). To maximally match the terminal taxa in the MCC tree with our dataset, the living hynobiid *Batrachuperus taibaiensis* was eliminated from multivariate phylogenetic comparative analyses, considering that *B. taibaiensis* was recently synonymized to *B. tibetanus* (Fei and Ye, 2016). The MCC tree was configured in Newick format and was read and visualized in the package ‘ape’ (version 5.5; Paradis and Schliep, 2019) for the software R (version 4.0.5; R Core Team). The remaining fossil taxa in our dataset with age ranges determined from published literature (Dataset in online Dryad) and the Paleobiology Database (www.paleobiodb.org) were incorporated into this MCC tree based on a recent cladogram (Jia et al., 2021a: fig. 6) we constructed for living and fossil cryptobranchoids (Figure 1D). Adding fossil taxa increases the accuracy for multivariate phylogenetic comparative analyses and ancestral state reconstruction (see below; Soul and Wright, 2021), but will inevitably bring zero-length branches when the age of internal nodes
was considered equal to that of its immediate oldest descendant, resulting in difficulties in multivariate phylogenetic comparative analyses. To circumvent this problem, any zero-length branch in the cladogram was treated to have a same branch length with its first none zero-length ancestral branch by using the “equal” dating method in the “DatePhylo” function from the R package ‘strap’ (version 1.4; Bell and Lloyd, 2015). We followed the suggestion from the ‘strap’ tutorial by setting the root length as 60 Ma by using the age difference between the oldest taxon in the cladogram (Kokartus; ~170 Ma) and the first older outgroup taxon known to date (Triassurus; ~230 Ma). Considering that the palate of Aviturus is only represented by the right vomer, the cladogram (Figure 1—figure supplement 24) including Aviturus was only used for the phylogenetic signal analysis and principle component analysis (PCA; see below) of the right vomer; whereas the cladogram without Aviturus was mapped onto the morphospace for a number of PCA and evolutionary rate analysis.

Data acquisition, geometric morphometrics and symmetry and asymmetry. The geometry of the paired vomers and parasphenoid is represented on the ventral view of the palate by 24 type I (intersections of biological structures) and type II (maximum of curvature) 2D landmark points (Figure 1B; Bookstein, 1991; Slice et al., 1996) digitized by using tpsUtil (64 bit; version 1.76; Rohlf, 2015) and tpsDig2 (version 2.31; Rohlf, 2015) based on images of 70 specimens (the 24-raw-landmark dataset thereafter; see above; Dataset in online Dryad). Most specimens have completely-preserved palate except that the Paleocene Aviturus exsecratus is known only by a right vomer (Vasilyan and Böhme, 2012: fig. 3). In order to investigate the paleobiology of Aviturus exsecratus, we created a separate dataset with seven
corresponding landmarks (#1–#7) for the right vomer across all 71 specimens (termed as the seven-raw-landmark dataset thereafter; Dataset in online Dryad). To reduce measurement errors, specimens were arranged alphabetically by the name of the image files, then landmarks in each specimen were digitized by the same author (J.J.) following the same order (landmark 1 to 24; Supplementary file 1K). Procrustes coordinates (24-procrustes-landmark dataset and seven-procrustes-landmark dataset) for the geometry of the palate were obtained in the software MorphoJ (version 1.07a; windows platform; Klingenberg, 2011) after superimposing the raw landmark coordinates of all specimens through the Generalized Procrustes Analysis (GPA; Bookstein, 1991; Figure 1—figure supplement 25), which serves to minimize non-shape variations (size, location and orientation) among specimens by rescaling, repositioning and rotating raw landmark configurations. After superimposition, centroid size of the palate for each specimen is calculated as square root of the sum of squared distances of all procrustes landmarks of the palate from their corresponding centroid. The mean shape of specimen/species are calculated as the consensus from the superimposed landmark coordinates. Procrustes variances/distance between specimens are calculated as the square root of the sum of the squared distances between corresponding landmarks (Klingenberg, 2016). The mean value for each species (the species mean thereafter) in the composite cladogram was obtained by averaging procrustes coordinates and centroid sizes when there are more than one representative specimens.

The geometry of the palate shows object symmetry (Mardia et al., 2000) with landmarks #15 and #20 lying in the midsagittal axis of the skull and the remaining 11 pair landmarks being generally symmetric about the midsagittal axis (Figure 1A). To calculate respective
contributions to the total amount of shape variation from symmetric and asymmetric shape
components, we used the function ‘C1v’ in the software R to create a new double
24-raw-landmark dataset that contains all original raw landmark coordinates and their
mirrored copy reflected about the midsagittal axis and relabeled to match the original
landmark numbers (Savriama, 2018). This doubled 24-raw-landmark dataset was then
imported into MorphoJ for superimposition, covariation matrix buildup and PCA.

Considering that the symmetric and asymmetric components of shape variations occupy
complementary subspaces in the shape tangent space defined by the doubled datasets for
landmark configurations with object symmetry (Mardia et al., 2000; Klingenberg et al., 2002),
the relative amount of contributions to shape changes from symmetric and asymmetric
components equals to the sum of the eigenvalues of their corresponding PCs derived from the
PCA. This procedure does not apply to the seven-landmark dataset because the right vomer
has no object symmetry.

**Allometric analysis.** To verify if allometry, covariation of shape with size, plays a role in
shape variations, we first performed a regular multivariate regression (Monteiro, 1999;
Klingenberg, 2016) of the symmetric shape components (dependent variables) from the
24-procrustes-landmark dataset on the log-transformed centroid size (log [CS]; independent
variable) across all 70 specimens and the 34 species by using the “procD.lm” function in the
‘geomorph’ R package (version 4.0.0; Adams et al., 2021; Collyer and Adams, 2021). Then
we conducted a permutation test of 10,000 iterations using the same function to test the
significance of these two regular regressions, respectively, considering that our sample size
(N = 70 for specimens; N = 34 for species) is not considerably larger than the number of variables (N = 49) for the 24-procrustes-landmark dataset. Statistical results from these regular regression analyses may be inaccurate given that variations (or residuals) in the shape components among species are correlated with their evolutionary relationships (“phylogenetic nonindependence” in Felsenstein, 1985). To account for impacts from the phylogeny, we conducted a phylogenetic regression (or evolutionary allometry analysis) of the shape components on log (CS) across species by using the function “procD.pglsl” of ‘geomorph’, which is a distance-based phylogenetic generalized least squares method (Adams, 2014a) that uses our composite cladogram (without Aviturus) to remove phylogenetic covariances among species under a Brownian motion model of evolution. To remove impacts from allometric shape components, shape variables (procrustes landmark coordinates in shape tangent space) were transformed by using residuals from the regular multivariate regression of shape on log (CS) across 70 specimens and 34 species, respectively, by using the function “procD.lm” in ‘geomorph’ (Klingenberg, 2016). Non-allometric portion of the symmetric shape components are used for downstream statistical analyses.

**Phylogenetic signal.** The phylogenetic signal is termed based on a pattern that closely related species tend to have similar character values stemmed from their shared evolutionary history (Felsenstein, 1985; Blomberg et al., 2003; Adams, 2014b). Whether the shape of the palate bears any phylogenetic signal was tested by using the $K_{\text{null}}$ method in the function “physignal” of ‘geomorph’, which uses the composite cladogram to account for phylogenetic
nonindependence under a Brownian motion model of evolutionary divergence, and
permutes shape components of terminal taxa against the composite cladogram (without

Aviturus). A phylogenetic signal analysis was also conducted for the right vomer
(nonallometric seven-procrustes-landmark dataset) using the composite cladogram including

Aviturus. The resulting $K_{\text{mult}}$ value for each of the two analyses measures the phylogenetic
signal as a ratio of observed to expected shape variation obtained with and without
considering phylogenetic nonindependence (Adams and Collyer, 2019). When the $K_{\text{mult}}$ value
is larger than and equals to 1, closely related taxa resemble each other phenotypically more
than expected under Brownian model, whereas when the value is smaller than 1, closely
related taxa are less similar to one another phenotypically than expected under Brownian
model (Adams, 2014b).

Morphological disparity and ecological indication of the palate. Taxonomically, the five
groups considered here are Pancryptobrancha, Hynobiidae, stem hynobiids, basal
cryptobranchoids and non-cryptobranchoid early salamanders. The Paleocene Aviturus was
classified into the clade Pancryptobrancha (Gubin, 1991; Vasilyan and Böhme, 2012). The
Late Jurassic Beiyanerpeton represents a basal member of Salamandroidea (Gao et al., 2013)
and the two stem urodeles Karaurus and Kokartus were classified into the family Karauridae
(Skutschas and Martin, 2011). The family Hynobiidae is designated here as crown group
Panhynobia, and we consider crown and stem taxa of Panhynobia separately. As mentioned
above, living hynobiids are mainly metamorphosed because individuals typically go through
a short period of development during which larval features (e.g., external gills, gill slits) are
resorbed (e.g., Kuzmin and Thiesmeier, 2001; Kami, 2004; Fei et al., 2006; Poyarkov et al., 2012), and *Batrachuperus londongensis* represents the single facultatively neotenic hynobiid species; living cryptobranchids are generally referred to as neotenic or partially metamorphosed as their individuals live in water at adult stage and have few paedomorphic features (Deban and Wake, 2000), except that the life history strategy of *Aviturus* remains unknown (Vasilyan and Böhme, 2012). It is noteworthy that many fossil taxa investigated here (e.g., *Chunerpeton*) bear more apparent neotenic features (e.g., external gills, gill rakers) than living cryptobranchids, however we did not differentiate subgroups within neotenic taxa due to our sampling scope. Living habitats of cryptobranchoids at adult stage out of breeding season have been broadly classified into three types, including aquatic, semiaquatic, and terrestrial (Fei et al., 2006; Rong, 2018; Fei and Ye, 2016; AmphibiaWeb, 2021). In order to understand if factors like taxonomic affiliations at the genus and family level, life history strategy and living habitat would contribute to morphological disparities of the palate, we conducted phylogenetic Procrustes ANOVAs for the nonallometric symmetric shape components of the 24-procrustes-landmark dataset using the function “procD.pglsls” of ‘geomorph’. Pair-wise comparisons for morphological disparity (procrustes variance) for the palate, vomer, parasphenoid (Supplementary file 1B) and each landmark point (Supplementary file 1C–E) across each of the aforementioned category were performed using “morphol.disparity” in ‘geomorph’.

**Principle component analyses.** To visualize the patterns of shape changes in the morphospace of palate and to investigate the mechanisms framing up the disparity patterns,
we conducted four types of PCA by using the function “gm.prcomp” in ‘geomorph’: a standard PCA, a phylogenetically aligned component analysis (PaCA; Collyer and Adams, 2020), a phylomorphospace analysis (PA; Rohlf, 2002), and a phylogenetic principal component analysis (Phylo-PCA; Revell, 2009). The standard PCA is based on a covariance matrix of the non-allometric symmetric shape components across all 70 specimens, and the resulting first few PCs (eigenvalues) reveal predominant trends in shape disparity patterns.

The other three PCA are based on the composite cladogram (without Aviturus) and a covariance matrix of the non-allometric symmetric shape components across all 34 species, with the ancestral PC values (eigenvalue) for internal nodes (Figure 1—figure supplement 26) estimated by using the Brownian motion model of evolution. The resulting first few PCs derived from the PA reveal predominant trends in shape disparity patterns with the phylogeny superimposed in the morphospace. On the other hand, the first PC derived from PaCA reveal shape disparity patterns mostly associated with phylogenetic signal, and the first few PCs from Phylo-PCA are phylogenetic independent and therefore reveal factors (e.g., ecological preferences) other than phylogenetic signal that account for disparity patterns of the palate (see Collyer and Adams, 2020). A same analysis strategy was conducted for the nonallometric seven-procrustes-landmark dataset to address the paleobiology of Aviturus (Figure 2—figure supplement 5). Visualizations of the results were achieved by using the functions “plot.gm.prcomp”, “make_ggplot” in ‘geomorph’, and “ggplot” in the package ‘ggplot2’ (version 3.3.5; Wickham, 2016).

Covariates of the palate. Configurations of vomerine tooth row have long been identified as
informative in the classification of Cryptobranchoidea (Estes, 1981; Zhao and Hu, 1984; Fei et al., 2006) and were recently argued to be ecological informative (Jia et al., 2021b). To test if non-shape variations of the palate are reliable indicators in reconstructing paleoecology of early salamanders, we chose the following five continuous and two categorical covariates: length ratio of the vomer in the palate; length ratio of the parasphenoid in the palate; vomerine tooth number on a single vomer; width ratio of vomerine tooth row (VTR) in the vomer; width ratio of the outer and inner branch of VTR; and both the position (relative to the vomerine plate: anterior, middle, mid-posterior or posterior) and arrangement (oblique, transverse, curved) of VTR. Measurements of these continuous and categorical features were collected by using VG Studio Max and the Fiji platform (version 1.8.0_172; Schneider et al., 2012), with missing values for *Aviturus*, *Kokartus*, and *Pangerpeton* reconstructed by using the function “phylopars” in ‘Rphylopars’ R package (version 0.3.2; Goolsby et al., 2021).

Values and states for the five continuous covariates were visualized by boxplot in ‘ggplot2’ and were compared among different ecological groups by using Mann Whitney U test (Figure 3). Association of the two categorical covariates with ecology were investigated by contingency table and Cochran-Mantel-Haenszel test (Figure 3—figure supplement 1 and Supplementary file 1II).

**Ancestral state reconstruction and evolutionary rate.** The ancestral shapes of the palate for several internal nodes including the respective common ancestors of Hynobiidae, Panhynobia, Pancryptobrancha, Cryptobranchoidea, Urodela, and Caudata were estimated by using maximum likelihood method in the “anc.recon” function in the ‘Rphylopars’ R package.
The ancestral value of the five continuous covariates for the internal nodes were estimated by using “phylopars” in the same package (Supplementary file 1J). On the other hand, the ancestral states of the two categorical covariates and life history strategy were estimated using the function “ace” in ‘ape’, with their likelihood for each of the internal node depicted as a pie chart (Figure 3). Wire plots for the ancestral shape of the internal nodes with reference to the mean shape were created by using the functions “mshape” and “plotRefToTarget” in ‘geomorph’.

Evolutionary rates for each of the 24 landmarks collected on the palate, the vomer, the parasphenoid, and the five continuous covariates from the palate were calculated by using the function “compare.multi.evol.rates” in ‘geomorph’. Then evolutionary rates of the palate, the vomer and the parasphenoid among species across groups in ecology, life history strategy and taxonomic affiliation were calculated and compared by using the function “compare.evol.rates” in ‘geomorph’ (Supplementary file 1H).

Results derived from this study were exported from R and were illustrated and assembled in Adobe Photoshop® CC. Source codes for R used in this study is available at GitHub (https://github.com/PaleoSalaman).

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Additional information

Competing interests

The authors declare no competing interests.

Author contributions

Jia Jia, Conceptualization, Data curation, Formal analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Software, Supervision, Validation, Visualization, Writing – original draft, Writing – review & editing; Guangzhao Li, Formal analysis, Methodology, Software, Validation; Ke-Qin Gao, Data curation, Funding acquisition, Resources, Writing – review & editing.

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Additional files

Supplementary files 1. A. Scores of the principle components generated from the standard principle component analysis (PCA) on the 24-landmark-dataset across 70 specimens, and the phylomorphospace analysis (PA), the phylogenetically aligned component analysis (PaCA), and the phylogenetic principal component analysis (Phylo-PCA) across 34 species. Abbreviations: Cumu. RV, Cumulative RV; Cumu. P., Cumulative Proportion; Eigenva., Eigenvalue; Prop. Co., Proportion of Covariance; Prop. V., Proportion of Variance; RV by Co., RV by Component; Sing. V., Singular Value. B. Pairwise comparison and corresponding p-values of morphological disparity of the palate calculated as procrustes variances for 34 species grouped by ecological preference, life history strategy and taxonomic affiliation. *, p value <= 0.05. Abbreviations: cryptobran., cryptobranchoids. C, Pairwise comparison and corresponding p-values of single landmark point of the palate calculated as procrustes variances (× E-4) for 34 species grouped by ecological preference. *, p <= 0.05; **, p <= 0.01; ***, p <= 0.001. D, Pairwise comparison and corresponding p-values of single landmark point of the palate calculated as procrustes variances (× E-5) for 34 species grouped by life history strategy. *, p <= 0.05. E, Pairwise comparison and corresponding p-values of single landmark point of the palate calculated as procrustes variances (× E-5) for 34 species grouped by
taxonomic affiliation. *, $p \leq 0.05$; **, $p \leq 0.01$. F, Absolute morphological disparity
of the palate calculated as procrustes variances for 34 species grouped by ecology, life
history strategy and taxonomic affiliations. G, Absolute morphological disparity
calculated as procrustes variances and evolutionary rates for each landmark point of the
palate across 34 species grouped by ecology, life history and taxonomic affiliations. H,
Comparison of evolutionary rates of the palate and centroid size for 34 species grouped
by ecology, life history and taxonomic affiliations. *, $p \leq 0.05$; **, $p \leq 0.01$; ***, $p \\
\leq 0.001$. I, Contingency table showing the association between ecological preference
and life history strategy and two discrete characters of vomerine tooth row (position and
arrangement pattern) across 34 species represented by 70 specimens. J, Non-shape
covariates of the palate of 35 species and the ancestral states for internal nodes
reconstructed by the “Rphylopars” R package. K, Definition of the 24 landmarks for the
palate (vomer and parasphenoid) used in this study.

Data availability

All data needed to evaluate the conclusions are present in the manuscript and the
Supplementary file 1. Details of specimens, CT parameters and raw landmark coordinates
and centroid sizes are available in three CSV files in the online Dryad repository
(https://doi.org/10.5061/dryad.c59zw3r8x). Source codes for R and SAS used in this study is
available at GitHub (https://github.com/PaleoSalaman).

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**Figure and figure supplement caption and legend:**

**Figure 1.** The palate and phylogenic relationships of early salamanders. (A) The vomer (gold) and parasphenoid (purple) of the palate in ventral view of the skull in living hynobiid *Pseudohynobius flavomaculatus*. (B) Dorsal view of the palate showing the articulation patterns with the paired orbitosphenoid (whitish). (C) Enlarged view of the palate in ventral view with red circles corresponding to the 24 landmarks used for the geometric morphometric analysis. (D) Palatal configurations of early salamanders in ventral view, with color-coded life history strategies (square block) and ecological preferences (line) plotted on the time-calibrated tree modified from Jetz and Pyron, 2018 and Jia et al.,...
Figure 2. Spatial patterns of the palatal shape in the morphospace defined by the first two components generated from four principle component analyses (PCA). (A) standard PCA across 70 specimens. (B) Phylomorphospace analysis, (C) phylogenetically aligned components analysis and (D) phylogenetic PCA across 34 species with ancestral states for internal nodes (open circles) and phylogenetic relationships (black lines) plotted in the morphospace. The color and shape of each point represent the ecological type and taxonomic affiliation, respectively. Extreme values of the palatal shape along both PCs 1 and 2 are represented by wireframes color-coded to ecological types against the mean shape (gray) of both terminal and internal taxa.

Figure 3. Evolutionary patterns of both the shape and non-shape covariates of the palate and their association with ecological disparity in early salamanders. Ancestral shape (wireframes color-coded to ecological types superimposed with mean shape [gray]) and vomerine tooth row (zigzag black lines) configurations are reconstructed for respective common ancestors of Hynobiidae, Panhynobida, Pancryptobrancha, Cryptobranchoidea and Urodela. A complete list of evolutionary rates for each of the 24 landmarks, the vomer, parasphenoid, the palate and the continuous covariates of the vomer across 34 species is available in Supplementary file 1G,H,J. The two pie charts at each internal node of the time-calibrated cladogram are likelihoods of the position (left) and arrangement (right) of the vomerine tooth row reconstructed in this study. Continuous covariates of the vomer were subjected to Mann Whitney U test for their association with the three ecological groups with corresponding \( p \) values labeled above the boxplots.
Figure 4. Spatial-temporal patterns of phenotypic diversities of the palate and their associations with ecological preference and life history strategy in early salamanders and the stepwise evolutionary patterns at the sutural area of vomer, parasphenoid and orbitosphenoid. The morphospace of the palate is defined within a space formed by geological time scale (Z axis) and PCs 1 (X axis) and 2 (Y axis) derived from phylogenetic principal component analysis across 34 species. All silhouettes and images of salamander species are original.

Figure 1—figure supplement 1. CT rendering of the palate and orbitosphenoid of extant cryptobranchids in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. CNU, Capital Normal University, Beijing, China; FMNH, Field Museum of Natural History, Chicago, Illinois, USA.

Figure 1—figure supplement 2. Images showing the ventral view of the right vomer (PIN 4357/13) of the Paleocene pancryptobranchan Aviturus exsecratus, with landmarks represented by red circles. Modified from Vasilyan and Böhme, 2012. PIN, Paleontological Institute, Russian Academy of Sciences, Moscow, Russia.

Figure 1—figure supplement 3. CT rendering of the palate and orbitosphenoid of two species of living hynobiid Batrachuperus in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. Note that representative specimens of both neotenic (middle row) and metamorphosed (lower row) Batrachuperus londongensis are included in this study. CIB, Chengdu Institute of Biology, Chinese Academy of Sciences, Chengdu, Sichuan.
Province, China; FMNH, Field Museum of Natural History, Chicago, Illinois, USA.

Figure 1—figure supplement 4. CT rendering of the palate and orbitosphenoid of three species of living hynobiid *Batrachuperus* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. CIB, Chengdu Institute of Biology, Chinese Academy of Sciences, Chengdu, Sichuan Province, China; FMNH, Field Museum of Natural History, Chicago, Illinois, USA; MVZ, Museum of Vertebrate Zoology, Berkeley, California, USA.

Figure 1—figure supplement 5. CT rendering of the palate and orbitosphenoid of two species of living hynobiid *Hynobius* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. ZMNH, Zhejiang Museum of Natural History, Hangzhou, Zhejiang Province, China; KU, University of Kansas Center for Research Inc., Lawrence, Kansas, USA.

Figure 1—figure supplement 6. CT rendering of the palate and orbitosphenoid of two species of living hynobiid *Liua* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. CIB, Chengdu Institute of Biology, Chinese Academy of Sciences, Chengdu, Sichuan Province, China.

Figure 1—figure supplement 7. CT rendering of the palate and orbitosphenoid of the living hynobiid *Onychodactylus japonicus* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. FMNH, Field Museum of Natural History, Chicago, Illinois, USA.
Figure 1—figure supplement 8. CT rendering of the palate and orbitosphenoid of the living hynobiid *Pachyhynobius shangchengensis* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. CIB, Chengdu Institute of Biology, Chinese Academy of Sciences, Chengdu, Sichuan Province, China.

Figure 1—figure supplement 9. CT rendering of the palate and orbitosphenoid of two species of the living hynobiid *Paradactylodon* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. FMNH, Field Museum of Natural History, Chicago, Illinois, USA; MVZ, Museum of Vertebrate Zoology, Berkeley, California, USA.

Figure 1—figure supplement 10. CT rendering of the palate and orbitosphenoid of the living hynobiid *Protohynobius puxiongensis* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. CIB, Chengdu Institute of Biology, Chinese Academy of Sciences, Chengdu, Sichuan Province, China.

Figure 1—figure supplement 11. CT rendering of the palate and orbitosphenoid of three species of the living hynobiid *Pseudohynobius* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. CIB, Chengdu Institute of Biology, Chinese Academy of Sciences, Chengdu, Sichuan Province, China; LTHC, Liupanshui Normal University, Liupanshui, Guizhou Province, China.

Figure 1—figure supplement 12. CT rendering of the palate and orbitosphenoid of two
species of the living hynobiid *Pseudohynobius* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. LTHC, Liupanshui Normal University, Liupanshui, Guizhou Province, China; ZMU, Zunyi Medical University, Zunyi, Guizhou Province, China.

Figure 1—figure supplement 13. CT rendering of the palate and orbitosphenoid of the living hynobiid *Ranodon sibiricus* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. FMNH, Field Museum of Natural History, Chicago, Illinois, USA.

Figure 1—figure supplement 14. CT rendering of the palate and orbitosphenoid of the living hynobiid *Salamandrella keyserlingii* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. FMNH, Field Museum of Natural History, Chicago, Illinois, USA.

Figure 1—figure supplement 15. CT rendering of the palate of the Late Jurassic basal salamandroid *Beiyanerpeton jianpingensis* in ventral view. Landmarks are represented by red circles in the second column. PKUP, Peking University Paleontological Collections, Beijing, China.

Figure 1—figure supplement 16. CT rendering of the palate and orbitosphenoid of the Middle Jurassic stem hynobiid *Neimengtriton daohugouensis* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. PKUP, Peking University Paleontological Collections, Beijing, China.

Figure 1—figure supplement 17. CT rendering of the palate and orbitosphenoid of the
Late Jurassic stem hynobiid *Linglongtriton daxishanensis* in dorsal (first and third columns) and ventral (second and fourth columns) views. Landmarks are represented by red circles in the last column. PKUP, Peking University Paleontological Collections, Beijing, China.

**Figure 1**—figure supplement 18. Line drawings of the palate and orbitosphenoid of the Early Cretaceous stem hynobiid *Liaoxitriton zhongjiani* in ventral view. Landmarks are represented by red circles in the second column. PKUP, Peking University Paleontological Collections, Beijing, China. Modified from Gao et al., 2013.

**Figure 1**—figure supplement 19. Line drawings of the palate and orbitosphenoid of the Early Cretaceous stem hynobiid *Nuominerpeton aquilonaris* in ventral view. Landmarks are represented by red circles in the second column. PKUP, Peking University Paleontological Collections, Beijing, China. Modified from Jia and Gao, 2016.

**Figure 1**—figure supplement 20. Line drawings of the palate and orbitosphenoid of the Early Cretaceous stem hynobiid *Regalerpeton weichangensis* in ventral view. Landmarks are represented by red circles in the second column. IVPP, Institute of Vertebrate Paleontology and Paleoanthropology, Chinese Academy of Sciences, Beijing, China. Modified from Rong, 2018.

**Figure 1**—figure supplement 21. Line drawings of the palate and orbitosphenoid of the Middle Jurassic basal cryptobranchoid *Chunerpeton tianyiensis* in ventral view. Landmarks are represented by red circles in the second column. CAGS, Chinese Academy of Geological Sciences, Beijing, China. Modified from Gao and Shubin, 2003.
Figure 1—figure supplement 22. Line drawings of the palate of the Late Jurassic basal cryptobranchoid *Pangerpeton sinensis* in ventral view. Landmarks are represented by red circles in the second column. PKUP, Peking University Paleontological Collections, Beijing, China. Modified from Gao et al., 2013.

Figure 1—figure supplement 23. Ventral view of the palate of two stem urodeles, the Late Jurassic *Karaurus sharovi* (upper row) and the Middle Jurassic *Kokartus honorarius* (lower row). Landmarks are represented by red circles in the second column. Modified from Ivachnenko, 1978; Skutschas and Martin, 2011.

Figure 1—figure supplement 24. The palate and phylogenetic relationships of early salamanders with the inclusion of *Aviturus*. (a), The vomer (gold) and parasphenoid (purple) of the palate in ventral view of the skull in living hynobiid *Pseudohynobius flavomaculatus*; (b), Dorsal view of the palate showing the articulation patterns with the paired orbitosphenoid (whitish); (c), Enlarged view of the palate in ventral view with red circles corresponding to the 24 landmarks used for our geometric morphometric analysis; (d), Palatal configurations of early salamanders in ventral view, with color-coded life history strategies (square block) and ecological preferences (line) plotted on the time-calibrated tree with the inclusion of *Aviturus* modified from Jetz and Pyron, 2018 and Jia et al., 2021a.

Figure 1—figure supplement 25. Configuration and superimposition of the 24 landmarks of the palate resulted from generalized procrustes analyses across 70 specimens (a) and 34 species (b). The wireframes connecting the landmarks represent the mean shape of the terminal taxa.
Figure 1—figure supplement 26. Time calibrated cladograms used in this study with both terminal and internal taxa numerically labeled. (a), the cladogram of 34 species; (b), the cladogram of 35 species with the inclusion of Aviturus.

Figure 2—figure supplement 1. Spatial patterns of the palatal shape in the morphospace defined by the first two components generated from four principle component analyses (PCA). (a), standard PCA across 70 specimens labeled by specimen catalogue numbers; (b), Phylomorphospace analysis, (c), phylogenetically aligned components analysis, and (d), phylogenetic PCA across 34 species (labeled by species names) with ancestral states (labeled by internal node numbers as in fig. S33a) for internal nodes (open circles) and phylogenetic relationships (black lines) plotted in the morphospace. The color and shape of each point represent the ecological type and taxonomic affiliation, respectively. Extreme values of the palatal shape along both PCs 1 and 2 are represented by wireframes color-coded to ecological types against the mean shape (gray) of both terminal and internal taxa.

Figure 2—figure supplement 2. Spatial patterns of the palatal shape in the morphospace defined by PC 1 and PC 3 generated from four principle component analyses (PCA). (a), standard PCA across 70 specimens; (b), Phylomorphospace analysis, (c), phylogenetically aligned components analysis, and (d), phylogenetic PCA across 34 species with ancestral states for internal nodes (open circles) and phylogenetic relationships (black lines) plotted in the morphospace. The color and shape of each point represent the types (refer to Figure 2) of ecology and taxonomy, respectively.

Figure 2—figure supplement 3. Spatial patterns of the palatal shape in the morphospace
defined by PC 2 and PC 3 generated from four principle component analyses

(PCA). (a), standard PCA across 70 specimens; (b), Phylomorphospace analysis, (c), phylogenetically aligned components analysis, and (d), phylogenetic PCA across 34 species with ancestral states for internal nodes (open circles) and phylogenetic relationships (black lines) plotted in the morphospace. The color and shape of each point represent the types (refer to Figure 2) of ecology and taxonomy, respectively.

Figure 2—figure supplement 4. Spatial patterns of the palatal shape in the morphospace defined by the first two components generated from four principle component analyses (PCA). (a), standard PCA across 70 specimens; (b), Phylomorphospace analysis, (c), phylogenetically aligned components analysis, and (d), phylogenetic PCA across 34 species with ancestral states for internal nodes (open circles) and phylogenetic relationships (black lines) plotted in the morphospace. The color and shape of each point represent the ecological type and life history strategy, respectively. Extreme values of the palatal shape along both PCs 1 and 2 are represented by wireframes color-coded to ecological types against the mean shape (gray) of both terminal and internal taxa.

Figure 2—figure supplement 5. Spatial patterns of the shape of the right vomer in the morphospace defined by first two PCs generated from four principle component analyses (PCA). (a), standard PCA across 71 specimens labeled by catalogue numbers; (b), Phylomorphospace analysis, (c), phylogenetically aligned components analysis, and (d), phylogenetic PCA across 35 species (labeled in species names) with ancestral states for internal nodes (open circles; numbered according to fig. S33b) and phylogenetic relationships (black lines) plotted in the morphospace. The color and shape of each point...
represent the types (refer to Figure 2) of ecology and taxonomy, respectively.

**Figure 3—figure supplement 1.** Probability of the indicativeness of ecological preference (a, b) and life history strategy (c, d) from two discrete characters of the vomerine tooth row, the position (VTRP) and arrangement (VTRA) based on contingency table (Supplementary file 1I).

**Figure 3—figure supplement 2.** Time calibrated cladogram of 34 species showing the ancestral states of the life history strategy for internal nodes reconstructed by maximum likelihood.
Figure 3.

Legend

| ecological type in boxplot/wireframe | VTR position | VTR arrangement | disparity/evolutionary rate |
|--------------------------------------|--------------|----------------|----------------------------|
| terrestrial                          | anterior     | parallel       | evolution rate             |
| aquatic                              | middle       | transverse     | morphological disparity    |
| semiaquatic                          | mid-posterior| oblique        |                            |
|                       | posterior    |                |                            |

Regalerpeton weichangensis

Onychodactylus japonicus
Hynobius amjiensis
Hynobius nebulosus
Batrachuperus karlschmidtii
Batrachuperus tibetanus
Batrachuperus londongensis
Batrachuperus pinchonii
Batrachuperus yenyuanensis
Liua shihi
Liua tsinpaensis
Pseudohynobius flavomaculatus
Pseudohynobius jinfo
Pseudohynobius guizhouensis
Pseudohynobius kuankoushuiensis
Pseudohynobius shuichengensis
Protohynobius puxiongensis
Salamandraella keyserlingii
Pachyhynobius shangchengensis
Paradactylophon persicus
Paradactylophon mustersi
Ranodon sibiricus

Liaoxitrion zhongjiani
Nuominerpeton aquilonaris
Neimengtrion daohugouensis
Linglongtrion daxishanensis

Pangerpeton sinensis
Chunerrpeton tianyiensis
Beiyanerpeton jianpingensis
Kokartus honorarius
Karaurus sharovi

Cryptobranchus alleganiensis
Andrias japonicus
Andrias davidianus

Aviturus exsecratus

Hynobiidae
Panhynobia
Pancryptobrancha
Cryptobranchoidea
Urodela
Caudata

Triassic: 201.3
Jurassic: 145.0
Cretaceous: 66.0
Cenozoic: 0

Evolutionary rate

Morphological disparity

Tooth number

Parasphenoid/parotid length ratio
Vomer/parotid length ratio
Outer/Inner VTR length ratio
VTR/vomer width ratio

Ecology

*= p < 0.05
**= p < 0.01
***= p < 0.001
Aviturus exsecratus PIN 4357/13 based on Vasilyan and Böhme, 2012: fig. 3B
Batrachuperus karlschmidtii FMNH 49380

Batrachuperus londongensis neoteny CIB 14380

Batrachuperus londongensis metamorphosis CIB 14504
Nuominerpeton aquilonaris PKUP V0414 based on Jia and Gao, 2016: figs. 3 and 4
*Regalerpeton weichangensis* based on Rong, 2018: figs. 5D and 7
Chunerpeton tianyiensis CAGS-IG-02051 based on Gao and Shubin, 2003: fig. 1b
Pangerpeton sinensis PKUP V0228 based on Gao et al., 2013: fig. 4
