

The putative lissamphibian stem-group: phylogeny and evolution of the dissorophoid temnospondyls

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Abstract.—Dissorophoid temnospondyls are widely considered to have given rise to some or all modern amphibians (Lissamphibia), but their ingroup relationships still bear major unresolved questions. An inclusive phylogenetic analysis of dissorophoids gives new insights into the large-scale topology of relationships. Based on a TNT 1.5 analysis (33 taxa, 108 characters), the enigmatic taxon *Perryella* is found to nest just outside Dissorophoidea (phylogenetic definition), but shares a range of synapomorphies with this clade. The dissorophoids proper are found to encompass a first dichotomy between the largely paedomorphic Micromelerpetidae and all other taxa (Xerodromes). Within the latter, there is a basal dichotomy between the large, heavily ossified Olsoniformes (Dissorophidae + Trematopidae) and the small salamander-like Amphibamiformes (new taxon), which include four clades: (1) Micropholidae (*Tersomius*, *Pasawioops*, *Micropholis*); (2) Amphibamidae sensu stricto (*Doleserpeton*, *Amphibamus*); (3) Branchiosauridae (*Branchiosaurus*, *Apateon*, *Leptorophus*, *Schoenfelderpeton*); and (4) Lissamphibia. The genera *Platyrhinops* and *Eoscopus* are here found to nest at the base of Amphibamiformes. Represented by their basal-most stem-taxa (*Triadobatrachus*, *Karaurus*, *Eocaecilia*), lissamphibians nest with *Gerobatrachus* rather than Amphibamidae, as repeatedly found by former analyses.

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Introduction

Modern amphibians and their ancient relatives could hardly be more distinct in size, morphology, and ecological features. Whereas the three extant clades (frogs, salamanders, and caecilians) are generally small creatures with feeble skeletons, most of which feed on small invertebrates, their likely stem-group, Paleozoic temnospondyls, encompasses 1–2 m long, heavily ossified predators. The evolutionary transition between the Paleozoic giants and the dwarfed modern forms has long been sought among the Dissorophoidea, a speciose clade of mainly terrestrial and presumably insect-eating Carboniferous–Triassic temnospondyls that were usually smaller and had less-massive skeletons than their fish-eating fellows (Watson, 1940; Romer, 1947; Reig, 1964), but alternative scenarios are still debated (Laurin and Reisz, 1997; Marjanovic and Laurin, 2008, 2009, 2013; Pardo et al., 2017). In recent time, new important stem-taxa were discovered (Bolt, 1969; Anderson et al., 2008a, b), which have made the origin at least of batrachians from within temnospondyls more likely and put the dissorophoids into focus of rigorous phylogenetic research (Milner, 1988, 1993; Trueb and Cloutier, 1991; Schoch and Milner, 2004; Ruta and Coates, 2007; Anderson, 2008; Fröbisch and Schoch, 2009a; Sigurdson and Green, 2011; Maddin et al., 2012; Pérez Ben et al., 2018).

Dissorophoids themselves were only slowly recognized as a clade, because they fall into several highly distinct groups: (1) the ~50 cm long, heavily built olsoniforms, comprising the

monophyletic armoured Dissorophidae and the Trematopidae (Carroll, 1964; Berman et al., 1985; Dilkes, 1990; Reisz et al., 2009; Schoch, 2012); (2) the 10–20 cm long terrestrial Amphibamidae (Bolt, 1969; Daly, 1994; Schoch and Rubidge, 2005; Anderson et al., 2008b; Fröbisch and Reisz, 2008); (3) the 10–30 cm long aquatic gill-bearing Micromelerpetidae (Boy, 1972, 1995, 2002a, b; Werneburg, 1991, 1994; Witzmann and Pfretzschner, 2003); and finally (4) the larval-neotenic, gill-bearing, up to 15 cm long Branchiosauridae (Boy, 1972, 1978, 1987; Werneburg, 1986, 1987, 1991, 2001; Schoch, 1992; Schoch and Milner, 2008). Although each of these groups has been hypothesized as a clade, an inclusive test of all of these hypotheses on a broader range of dissorophoid taxa has yet to be conducted.

The first dissorophoid skeleton discovered was that of a poorly preserved branchiosaurid from the lower Permian of Münsterappel in Germany, reported by Meyer (1848) as *Apateon pedestris*. This superficially salamander-like branchiosaurid was soon complemented by more frog-like skeletons from Pennsylvanian rocks from from Mazon Creek, Illinois, which were referred to as *Pelion* (*Amphibamus*) *grandiceps* (Wyman, 1858; Cope, 1865). After Moodie (1909, 1916), Romer (1930), and Steen (1931, 1938) had reported a range of similar taxa from additional Pennsylvanian deposits in the United States and Czech Republic, Watson (1940) wrote a review paper on the origin of anurans in which he sought the ancestry of frogs among amphibamids, notably the tiny, lightly built *Amphibamus grandiceps* Cope, 1865. The enlarged squamosal embayment (otic notch) of

these taxa was given particular attention, as were the delicate stapes and the reduced nature of some dermal bones. These of course were features long known by students of temnospondyls to be shared with anurans, as exemplified by Quenstedt's (1850, p. 14) exclamation "true frog ears!" (German "wahre Froschohren") upon his discovery of the otic notch in the large capitosauroid temnospondyl *Cyclotosaurus*.

Almost three decades after Watson, Bolt (1969) reported the tiny amphibamid *Doleserpeton annectens*, phylogenetically the sister taxon to *Amphibamus*, but with two additional characters then unknown from any other temnospondyl: pedicellate teeth and cylindrical pleurocentra in the trunk vertebrae (Bolt, 1977). This was an important step forward because the cylindrical pleurocentra had long been considered a major argument against a temnospondyl relationship of lissamphibians, later upheld by Gardiner (1983) who argued for a close relationship among aistopods, nectrideans, and lissamphibians on this basis. This was one reason why lepospondyls had been seen as the most plausible stem-group of extant amphibians, especially salamanders and caecilians (Romer, 1945; Schmalhausen, 1968; Carroll and Currie, 1975; Carroll and Holmes, 1980). However, this diphyletic hypothesis (temnospondyl anurans versus lepospondyl salamanders and caecilians) was never consistent in that anurans share the cylindrical vertebrae with other lissamphibians, but were nevertheless regarded temnospondyl descendants by most authors (Danto et al., 2016).

Later, Bolt (1977) discovered pedicely also in *Amphibamus*, but not *Platyrhinops*, which shares with the two others and lissamphibians tooth crowns with two cusps (bicuspidity). The clade formed by *Doleserpeton* and *Amphibamus* was since conceived the most plausible stem-group of Lissamphibia by many authors (Milner, 1988, 1993; Trueb and Cloutier, 1991; Ruta et al., 2003; Schoch and Milner, 2004; Ruta and Coates, 2007), except for those upholding a non-temnospondyl origin for lissamphibians (Laurin and Reisz, 1997; Laurin, 1998; Vallin and Laurin, 2004; Marjanovic and Laurin, 2008, 2009, 2013).

Jenkins and Walsh (1993) discovered skeletons of a Jurassic stem-caecilian with rudimentary limbs, which they named *Eocaecilia micropodia*. In the course of studying this interesting stem taxon, Jenkins et al. (2007) and Maddin et al. (2012) discovered numerous plesiomorphic characters in the skeleton of *Eocaecilia*, most notably the retention of dermal bones in the skull that are lost in extant lissamphibians. This demonstrated that batrachians (salamanders and frogs) and caecilians must have lost these elements convergently.

A further milestone was taken by Anderson et al. (2008a) who named and described the lower Permian amphibamid *Gerobatrachus hottoni* from Texas. This taxon retains a full complement of dermal bones in the skull contrasting the reduced set of dermal bones in batrachians. However, these elements cover much smaller areas than in other temnospondyls, giving expanded orbits and highly reduced strut-like palatal elements, much like in salamanders and frogs. At the same time, the cylindrical vertebrae are reduced to 17 presacrals, the lowest count among temnospondyls and just slightly higher than that of batrachians. Because of its possession of a basale commune in the tarsus, which is a feature of salamanders, *Gerobatrachus* has been considered a stem-batrachian (Anderson et al., 2008a)

rather than ancestral to all lissamphibians, which was since found as an alternative placement (Fröbisch and Schoch, 2009a).

Recently, Pardo et al. (2017) reported a tiny stereospondyl temnospondyl, *Chinlestegophis jenkinsi* Pardo, Small, and Huttenlocker, 2017, which they argued shares derived features with caecilians and *Eocaecilia*. On this basis, they proposed a new diphyletic origin of lissamphibians, with batrachians having evolved from amphibamid dissorophoids, but caecilians from brachyopoid stereospondyls. This remains an open question that needs an inclusive analysis to be carried out in the future.

At any rate, the discovery of *Eocaecilia* and *Gerobatrachus* brought an end to the long-practiced separate treatment of temnospondyls and lissamphibians, because both *Eocaecilia* and *Gerobatrachus* retained dermal bones that are not present in any extant lissamphibian, but are well known from temnospondyls. The alternative placements of *Gerobatrachus* indicate that—given the temnospondyl hypothesis of lissamphibian origin is true—some amphibamids are most probably more closely related to lissamphibians (or batrachians) than others. The Amphibamidae has therefore come increasingly into the focus of studies on the origin of lissamphibians. This is irrespective of whether all extant taxa or only batrachians were derived from amphibamids, a discussion that is still in a state of flux (Anderson, 2008; Pardo et al., 2017). Amphibamid relationships within dissorophoids therefore form a central problem that also involves the nature of the Branchiosauridae and the primitive condition of Dissorophoidea as a whole.

Objectives.—This study aims to resolve or constrain the following major questions of dissorophoid phylogeny:

What does the large-scale phylogeny of Dissorophoidea look like?

What is the position of the enigmatic taxon *Perryella*?

Do Micromelerpetidae form a clade and where do they nest?

Do Branchiosauridae form a clade and where do they nest?

What is the relationship of *Doleserpeton* and *Gerobatrachus* to Lissamphibia/Batrachia?

What is the most likely evolutionary scenario for the origin of the Dissorophoidea and the origin and early diversification of Lissamphibia/Batrachia?

The olsoniforms, which were studied in detail elsewhere (Schoch, 2012; Maddin et al., 2013; Schoch and Sues, 2013), are not represented by a complete set of known taxa, but a smaller selection representative of their diversity.

Phylogenetic analysis

Taxon sampling.—The goal of the phylogenetic analysis was to cover as wide a range of well-studied dissorophoid taxa as possible, with the focus on the origin and diversification of this clade and the in-group relationships of amphibamids. Other questions have been addressed elsewhere, such as the placement of dissorophoids among temnospondyls (Schoch, 2013) or the

evolutionary history of olsoniforms (Schoch, 2012; Holmes et al., 2013; Maddin et al., 2013; Schoch and Sues, 2013). Although kept relatively large, the number of taxa has been restricted to address the specific questions formulated above. To this end, datasets for amphibamids (Schoch and Rubidge, 2005; Anderson et al., 2008a, b; Fröbisch and Reisz, 2008; Sigurdson and Bolt, 2010), dissorophids (Schoch, 2012; Holmes et al., 2013; Maddin et al., 2013), and branchiosaurids (Schoch and Milner, 2008; Fröbisch and Schoch, 2009a) have been combined, recoded in some places (indicated in Supplemental Dataset), and additional new characters were added. Throughout, the data set from which the coding was adopted is cited. In order to elucidate the origin of lissamphibians, three representative stem taxa were chosen, interpreted to reflect the primitive condition, the salientian (stem-anuran) *Triadobatrachus*, the urodele (stem-caudate) *Karaurus*, and the apodan (stem-gymnophionan) *Eocaecilia*, which are also fairly consistent with the primitive condition, as assessed by analysis of character distribution in the crown-groups (Schoch, 2014a). The analysis of the recently reported *Chinlestegophis* is spared for a separate study in the future.

The following taxa have been included in the phylogenetic analysis.

Outgroup:

Dendrysekos helogenes Steen, 1931 (Holmes et al., 1998; Schoch and Milner, 2014).

Ingroups:

1. *Trimerorhachis insignis* Cope, 1878 (Case, 1935; Milner and Schoch, 2013).
2. *Sclerocephalus haeuseri* Goldfuss, 1847 (Boy, 1988; Schoch and Witzmann, 2009).
3. *Acanthostomatops vorax* (Credner, 1883) (Boy, 1989; Witzmann and Schoch, 2006).
4. *Perryella olsoni* Carlson, 1987 (Case, 1935; Ruta and Bolt, 2006).
5. *Micromelerpeton credneri* (Bulman and Whittard, 1926) (Boy, 1972, 1995; Schoch and Milner, 2014).
6. *Limnogyrinus elegans* Fritsch, 1881 (Werneburg, 1994).
7. *Branchierpeton amblyostomum* (Credner, 1881) (Werneburg, 1991).
8. *Eimerisaurus graumanni* (Boy, 1980) (Boy, 2002b).
9. *Branchiosaurus salamandroides* Fritsch, 1876 (Boy, 1987; Werneburg, 2012; Schoch and Milner, 2014).
10. *Apateon gracilis* (Boy, 1987; Werneburg, 1991; Schoch and Fröbisch, 2006).
11. *Leptorophus tener* Schönfeld, 1911 (Boy, 1986, 1987).
12. *Schoenfelderpeton prescheri* Boy, 1986 (Boy, 1987).
13. *Amphibamus grandiceps* (Milner, 1982; Bolt, 1969, 1979).
14. *Doleserpeton annectens* Bolt, 1969 (Bold, 1977; Sigurdson and Bolt, 2010).
15. *Gerobatrachus hottoni* Anderson et al., 2008a.
16. *Georgenthalia clavinasica* Anderson et al., 2008b.
17. *Eoscopus lockardi* Daly, 1994.
18. *Platyrrhinops lyelli* (Wyman, 1858) (Carroll, 1964; Clack and Milner, 2010).
19. *Micropholis stowi* Huxley, 1859 (Broili and Schröder, 1937; Boy, 1985; Schoch and Rubidge, 2005).

20. *Tersomius texensis* Case, 1910 (Carroll, 1964; Maddin et al., 2013).
21. *Pasawioops mayi* Fröbisch and Reisz, 2008.
22. *Fedexia striegeli* Berman et al., 2010.
23. *Acheloma durni* Polley and Reisz, 2011.
24. *Phonerpeton pricei* Dilkes, 1990.
25. *Ecolsonia cutlerensis* Berman et al., 1985.
26. *Broiliellus texensis* Williston, 1914 (Carroll, 1964; Schoch, 2012).
27. *Dissorophus multicinctus* Cope, 1895 (Carroll, 1964; DeMar, 1968; Schoch, 2012).
28. *Cacops morrisoni* Reisz et al., 2009.
29. *Conjunctio multidentis* Carroll, 1964 (Schoch and Sues, 2013).
30. *Triadobatrachus massinoti* Piveteau, 1936 (Rage and Roček, 1989; Ascarrunz et al., 2016).
31. *Karaurus sharovi* Ivakhnenko, 1978.
32. *Eocaecilia micropodia* Jenkins and Walsh, 1993 (Jenkins et al., 2007).

Character sampling.—Many previously used characters were revised and others excluded because they were uninformative for the present range of taxa. For instance, characters 30, 46, and 47 of Schoch and Rubidge (2005), and character 51 of Anderson et al. (2008b) have been omitted because they were found not to be defined clearly enough; otherwise the characters added by Anderson et al. (2008b) are included in the present in data set, albeit with some modified coding. In addition, I have added 38 characters from other sources, including some newly formulated ones. Altogether, the dataset includes 108 discrete morphological characters sampled from all over the temnospondyl skeleton. The complete list of characters is given in the Supplemental Dataset.

Analysis.—The parsimony analysis was performed in TNT 1.5 (available at <http://www.cladistics.org/tnt.html>) using the New Technology Search option, which generated 10 equally most parsimonious trees (tree length [TL] = 282 steps; consistency index [CI] = 0.429; retention index [RI] = 0.711). Multistate characters (1, 45, 53, 67, 75, 94) were ordered. When these six characters were left unordered, the topology of the strict consensus tree remained the same. *Dendrysekos* (= *Dendrerpeton* of many former authors who referred to Holmes et al., 1998) has served as only outgroup taxon. The non-dissorophoid ingroup taxa are *Trimerorhachis* (for Dvinosauria), *Sclerocephalus* (for Eryopiformes), and *Acanthostomatops* (for Zatracheidae).

Results.—The resulting 10 trees produce seven polytomies in the strict consensus. Homoplasies are common. The Dissorophoidea (sensu Schoch and Milner, 2014), Micromelerpetidae (for spelling, see Schoch and Milner, 2014), Xerodromes, Olsoniformes, Dissorophidae, and the Amphibamiformes + Lissamphibia clade each form stable nodes (see Bremer support values given in the following tree description). The main instabilities are located: (1) within the higher Micromelerpetidae (*Eimerisaurus*, *Branchierpeton*, *Limnogyrinus*); (2) the Trematopidae (*Ecolsonia*, *Fedexia*, and

advanced trematopids); (3) the base of the Amphibamidae-Lissamphibia clade (*Eoscopus*, *Platyrhinops*, all higher taxa); and (4) the higher Amphibamiformes (phylogenetic definition see next section), which encompass an unresolved

polytomy between *Georgenthalia*, Branchiosauridae, *Gerobatrachus* and Lissamphibia. The following nodes were found in the main analysis of 33 taxa and 108 characters (see also Fig. 1.1):

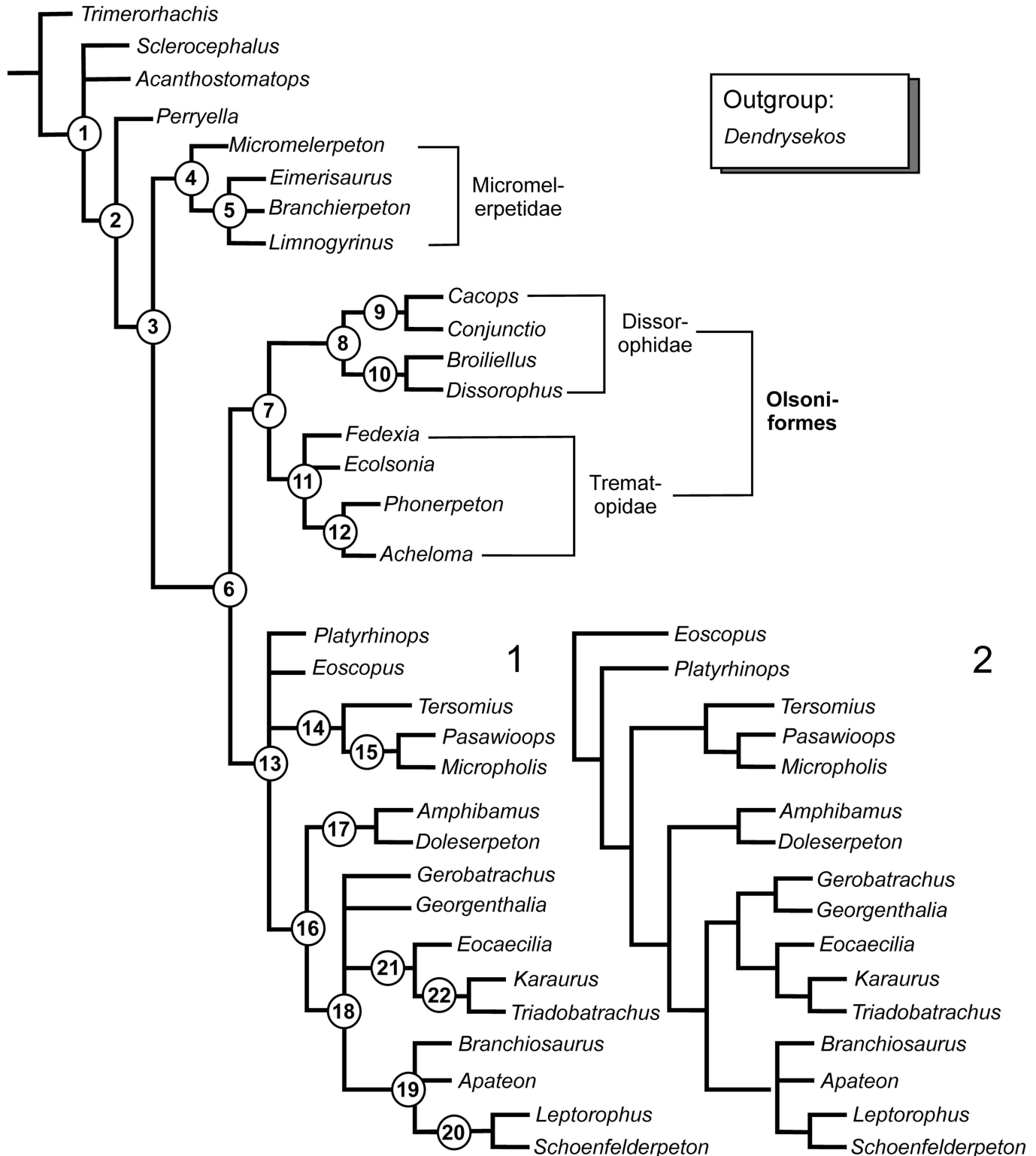


Figure 1. Phylogeny of dissorophoids as found by present analysis. Strict consensus of ten MPTs, 282 steps, CI=0.429, RI=0.711. (1) Main data set (108 characters); (2) variant analysis, giving slightly better resolved Amphibamiformes (omitting characters 17 and 30).

- (1) Rhachitomi. Bremer Support: >6. Within Rhachitomi, and above *Trimerorhachis*, *Sclerocephalus*, *Acanthostomatops*, and all higher taxa form an unresolved polytomy.
- (2) *Perryella* + Dissorophoidea. Bremer Support: 2. Synapomorphies: 9, 21, 38.
- (3) Dissorophoidea. Bremer Support: 3. Synapomorphies: 24, 30, 32.
- (4) Micromelerpetidae. Bremer Support: 3. Synapomorphies: 40, 74, 77, 84.
- (5) *Eimerisaurus* + *Branchierpeton* + *Limnogyrinus*. Bremer Support: 3. Synapomorphies: 5, 7, 38, 51. The relationships among the three genera remain unresolved.
- (6) Xerodromes. Bremer Support: 5. Synapomorphies: 18, 53, 88.
- (7) Olsoniformes. Bremer Support: > 6. Synapomorphies: 16, 19, 20, 28, 29, 35, 60, 90, 92.
- (8) Dissorophidae. Bremer Support: 5. Synapomorphies: 4, 10, 54, 57, 59.
- (9) Dissorophinae. Bremer Support: 5. Synapomorphies: 12, 22, 30, 61, 65.
- (10) Cacopinae. Bremer Support: 4. Synapomorphies: 17, 40, 49, 50, 55.
- (11) Trematopidae. Bremer Support: 1. Synapomorphies: 39. This node contains an unresolved polytomy consisting of *Ecolsonia* + *Fedexia* + higher Trematopidae.
- (12) *Phonerpeton* + *Acheloma*. Bremer Support: 3. Synapomorphies: 27, 40, 51.
- (13) Amphibamiformes. Bremer Support: 5. Synapomorphies: 11, 12, 75, 93.
- (14) Micropholidae. Bremer Support: 1. Synapomorphies: 34, 51.
- (15) *Pasawioops* + *Micropholis*. Bremer Support: 1. Synapomorphies: 33.
- (16) Post-micropholid Amphibamiformes. Bremer Support: > 6. Synapomorphies: 6, 40, 86, 91, 101.
- (17) Amphibamidae. Bremer Support: 1. Synapomorphies: 71, 89, 102.
- (18) Higher Amphibamiformes. Bremer Support: 1. Synapomorphies: 7, 30, 84, 100. This node contains an unresolved polytomy consisting of *Georgenthalia* + Branchiosauridae + *Gerobatrachus* + Lissamphibia.
- (19) Branchiosauridae. Bremer Support: 3. Synapomorphies: 12, 74, 80. This node contains an unresolved polytomy consisting of *Branchiosaurus* + Apatoninae + Leptorophinae.
- (20) Leptorophinae. Bremer Support: 3. Synapomorphies: 17, 83, 95, 96.
- (21) Lissamphibia. Bremer Support: > 6. Synapomorphies: 71, 94, 105, 107, 108.
- (22) Batrachia. Bremer Support: >6. Synapomorphies: 4, 72, 89, 98, 104, 106.

Variant analyses.—Because two of the characters (17, 30) of Schoch and Rubidge's (2005) data set were criticized as mutually overlapping or non-independent (Holmes et al., 2013), they were omitted in variant analyses, giving a significantly higher resolution (Fig. 1.2). Exclusion of character 30 alone resulted in four MPTs, whereas omission of character 17 or both

characters 17 and 30 likewise resulted in only two MPTs. In both variants, the higher Amphibamiformes were almost completely resolved, with *Gerobatrachus* + *Georgenthalia* forming the sister taxon to *Eocaecilia* + (*Karaurus* + *Triadobatrachus*), and in turn with the branchiosaurids forming the sister taxon of the latter clade; at the base of the Amphibamiformes, *Eoscopus* and *Platyrrhinops* form successive sister taxa of Amphibamidae and the aforementioned clades.

Robustness of nodes.—The robustness of nodes and sources of instability were studied by the variant analysis of selected subsets of taxa. When *Acanthostomatops* is omitted, Dissorophoidea nests with *Sclerocephalus* rather than *Trimerorhachis* (seven trees). When *Perryella* is omitted in addition to *Acanthostomatops*, the result remains the same (seven trees).

Systematic paleontology

Temnospondyli Zittel, 1888
Rhachitomi Watson, 1919, sensu Schoch, 2013
Dissorophoidea Bolt, 1969

Diagnosis.—Laterally exposed palatine (LEP) (1); dorsal quadrate process (DQP) (2); stapes directed mostly laterally (24), postparietal four times wider than long (30), posterior projection of vomer absent (32), short preorbital region (51), jugal anteriorly blunt (97); medial quadratojugal process (99).

Occurrence.—Moscovian–Induan (Moscovian–Recent, if some or all lissamphibians are dissorophoids).

Definition (node).—The least inclusive clade containing *Micromelerpeton credneri* and *Dissorophus multicinctus* (Schoch, 2013).

Diversity.—Dissorophoidea in its present form contains 52 genera and 85 species (without Lissamphibia).

Micromelerpetidae Boy, 1972

Diagnosis.—Basipterygoid joint unossified (74), elevated presacral count (77), and unossified pubis (84); additional characters: lateral line sulci present even in large adults, bony ventral scales with thickened, knob-bearing posterior margin (Boy, 1972).

Occurrence.—Moscovian–Sakmarian.

Definition (node).—The least inclusive clade containing *Micromelerpeton credneri* and *Branchierpeton amblyostomum*. Range: Moscovian–Artinskian.

Diversity.—Within the frame of the present analysis, Micromelerpetidae encompasses *Micromelerpeton* (three species), *Eimerisaurus* (two species), *Limnogyrinus* (two species), *Nyranerpeton milneri* Werneburg, 2012 (not included in analysis), and *Branchierpeton* (three species).

Xerodromes Schoch and Milner, 2014

Diagnosis.—Squamosal-supratemporal suture shortened by extension of squamosal embayment (18), distance squamosal embayment-orbit foreshortened (53), trunk ribs throughout shorter (88).

Occurrence.—Moscovian–Induan (Moscovian–Recent, if some or all lissamphibians are dissorophoids).

Definition (stem).—The most inclusive clade containing *Apateon pedestris*, *Amphibamus grandiceps*, and *Dissorophus multicinctus*, but not *Micromelerpeton credneri* (Schoch and Milner, 2014).

Diversity.—Xerodromes in its present form contains 48 genera and 76 species (without Lissamphibia).

Amphibamiformes new taxon

Diagnosis.—Palatine and ectopterygoid reduced to narrow struts (11), interpterygoid vacuity greatly expanded laterally (12), humerus length:waist ratio 6:10 (75), basioccipital and supraoccipital absent (93).

Occurrence.—Moscovian–Induan (Moscovian–Recent, if some or all lissamphibians are dissorophoids).

Etymology.—Derived from the genus *Amphibamus*, which it contains.

Definition (stem).—The most inclusive clade containing *Amphibamus grandiceps*, but not *Dissorophus multicinctus* (Schoch and Milner, 2014).

Diversity.—Amphibamiformes in its present form contains 12 genera and 15 species.

Remarks.—The new name Amphibamiformes is here proposed for three reasons: (1) it encompasses a much larger group than the family-level name Amphibamidae suggests; in its extreme version, which has become more likely with the discovery of *Doleserpeton* and *Gerobatrachus*, it includes the more than 7,846 extant lissamphibian species (www.amphibiaweb.org as of April 18, 2018); (2) it contains the Branchiosauridae, which according to all cladistic analyses so far are retained as a clade; and (3) Amphibamiformes is a more apt sister taxon to Olsoniformes than Amphibamidae, considering that even in its smallest form (without Lissamphibia) it is comparably more speciose than its sister group.

Micropholidae Watson, 1919

Diagnosis.—Vomer with additional fang pair posterior to mid-vomerine depression (8), postparietal more than four times as wide as long (30), palatine bordering interpterygoid vacuity along its entire lateral margin (34), preorbital length foreshortened (51).

Occurrence.—Sakmarian–Induan.

Definition (node).—The least inclusive clade containing *Tersomius texensis* and *Micropholis stowi*. In the present analysis, Micropholidae includes *Tersomius*, *Micropholis*, and *Pasawioops*.

Remarks.—This clade, which has been found by some former authors (Schoch and Rubidge, 2005; Sigurdson and Bolt, 2010), probably also contains *Rubeostratilia* and possibly *Plemmyradytes*. It is in many respects more plesiomorphic than the Amphibamidae and Branchiosauridae, as discussed below. *Eoscopus* may be a stem taxon of Micropholidae, sharing the additional vomerine fang pair with the clade.

Amphibamidae Moodie, 1909

Diagnosis.—Anterior trunk ribs not widened distally (71); choana medially expanded (89), patch of tiny denticles on vomer replacing tusk pair (102); additional character: squamosal posteriorly shortened, compensated by the quadratojugal.

Occurrence.—Moscovian–Artinskian.

Definition (node).—The least inclusive clade containing *Amphibamus grandiceps* and *Doleserpeton annectens*.

Remarks.—This definition is admittedly very restrictive, confining the family to a minimum of two taxa, but potentially including a still undescribed taxon from the Pennsylvanian Kinney Brick Quarry site in New Mexico (R. Werneburg, personal communication, 2017).

Branchiosauridae Fritsch, 1879

Diagnosis.—Palatine Y-shaped, posterior part separate from maxilla; ectopterygoid completely separate from maxilla; branchial denticles single, without bony base, often with brush-like tips.

Occurrence.—Moscovian–Artinskian.

Definition (node).—The least inclusive clade containing *Branchiosaurus salamandroides* and *Schoenfelderpeton prescheri*.

Diversity.—Branchiosauridae in its largest form contain five genera and 19 species (without *Eumicrerpeton*, *Milnererpeton*, and *Tungussogyrinus*).

Remarks.—The incompletely known taxa *Milnererpeton huberi* (Hunt, Lucas, and Berman, 1996) *Tungussogyrinus bergi*, and *Eumicrerpeton parvum* Moodie, 1909 have been omitted, but their possession of branchiosaurid characters makes them likely members of the clade rather than larvae of some other dissorophoid taxa, as has been proposed by former authors (Hunt et al., 1996; Werneburg, 2009; Milner in Schoch and Milner, 2014).

Evolution of major dissorophoid clades

The evolution of dissorophoids and their relationship to extant amphibians have been analyzed by numerous authors in the 20th century, with the first detailed account given by Watson (1940), who elucidated the origin of anurans based on the examination of *Amphibamus grandiceps*. Subsequent seminal studies on dissorophoids were conducted by Carroll (1964) and Bolt (1969, 1977, 1979), but these were mostly confined to aspects of character evolution, such as Bolt's discovery of pedicely in *Dolesempetron* and *Amphibamus*. Resting on Parsons and Williams' (1962, 1963) findings, Bolt (1969) proposed a common origin of Lissamphibia from within the dissorophoids. This was the first time that substantial data were provided to link extant amphibians with a single group of Paleozoic tetrapods, after lissamphibians had been considered polyphyletic by authors from different schools of thought (Holmgren, 1933; Romer, 1945; Schmalhausen, 1968). In his thorough reexamination of *Amphibamus grandiceps*, Bolt (1979) also considered developmental aspects, and developed a scenario in which *r*-selection might have driven the origin of lissamphibians.

The first inclusive phylogenetic analyses of dissorophoids, based on cladistic principles, were carried out by Jürgen Boy, initially focused on branchiosaurids and micromelerpetids (Boy, 1972), then expanded on dissorophoids as a whole (Boy, 1981). This author had studied the ontogeny of dissorophoids and other temnospondyls in unprecedented detail (Boy, 1974, 1978, 1988), and incorporated many of these data into his evolutionary scenarios. Having recognized the monophyletic status of branchiosaurids, he suggested that they either formed a neotenic clade or a group of larval forms whose terrestrial adults simply remained unknown. Boy also clearly distinguished micromelerpetids from branchiosaurids and amphibamids, although he considered the paedomorphic, aquatic micromelerpetids and branchiosaurids as close relatives. He was also the first author who recognized micromelerpetids and branchiosaurids as dissorophoids, after these groups had been neglected in other surveys (Romer, 1947; Carroll, 1964; Bolt, 1969).

More inclusive than any previous accounts were Milner's (1988, 1990, 1993) analyses and discussions, which also employed scenarios of heterochrony and paleoecology. This author was the first to widen the perspective by including all lissamphibians, finding that their closest relatives among Paleozoic tetrapods were indeed *Amphibamus* and *Dolesempetron* as forming successive sister taxa towards the crown, thus synthesizing the variant hypotheses of Watson (1940) and Bolt (1969). Milner was the first author who knew both the Central European branchiosaurids and the North American terrestrial dissorophoids by first-hand examination and in sufficient detail to find robust support for their relationship, as well as the close relationship between branchiosaurids and lissamphibians (Milner, 1993). He therefore went further in developing the first inclusive phylogeny of temnospondyls plus lissamphibians (Milner, 1990, 1993), and most of his hypotheses have been confirmed by computer-assisted cladistic analyses (Yates and Warren, 2000; Ruta and Coates, 2007; Schoch, 2013).

The first large-scale cladistic analysis of early tetrapods and lissamphibians was performed by Laurin and Reisz (1997), who

presented the most radical alternative to Bolt's (1969) and Milner's (1988) Temnospondyl Hypothesis in proposing that all lissamphibians were derived from lysorophians, a small clade within lepospondyl microsaurids. Naturally, this analysis included various dissorophoids and notably *Dolesempetron*, but omitted branchiosaurids with reference to their then-believed completely immature status. In their Lepospondyl Hypothesis, Laurin and Reisz (1997) employed numerous lepospondyl features shared by sirenid, proteid, and amphiumid salamanders as well as caecilians. The implications of this study were analyzed by Schoch and Milner (2004) who argued that the lysorophian-lissamphibian similarities constitute convergences produced by repeated events of neoteny (salamanders) and burrowing (caecilians). Although this new perspective was highly stimulating for further research, it primarily failed to explain the close resemblance particularly of anurans with temnospondyls, such as pedicely, the auditory apparatus, and various characters in the palate (Ruta and Coates, 2007; Sigurdson and Green, 2011)—despite efforts to surmount these problems (Laurin, 1998; Vallin and Laurin, 2004; Marjanovic and Laurin, 2008). At any rate, this intensified phylogenetic controversy had an enormous stimulating influence on the description and revision of dissorophoid taxa.

The last decade has brought various new and increasingly refined phylogenetic studies of dissorophoids, triggered by the examination of long-known taxa (*Micropholis*: Schoch and Rubidge, 2005; *Cacops*: Reisz et al., 2009; *Perryella*: Ruta and Bolt, 2006; *Dolesempetron*: Sigurdson and Bolt, 2010; *Mordax*: Werneburg, 2012; *Tersomius*: Maddin et al., 2013) and the discovery of new species and genera (*Eimerisaurus*: Boy, 2002b; *Plemmyradytes*: Huttenlocker et al., 2007; *Pasawioops*: Fröbisch and Reisz, 2008; *Rubeostratilia*: Bourget and Anderson, 2011; *Nyranerpeton*: Werneburg, 2012; *Reiszertpeton*: Maddin et al., 2013; *Scapanops*: Schoch and Sues, 2013). These studies provide a much broader platform of taxa, characters, and hypotheses, which will form the basis of the following discussion.

The origin of the Dissorophoidea.—Dissorophoids must have evolved well before the Westphalian D (Moscovian: 307–315 Myr), because at that time the major clades were already established (Schoch and Milner, 2014). Morphologically, there is a considerable gap between dissorophoids and other temnospondyls. Departing from the primitive condition (edopoids, *Dendrysekos*, *Balanerpeton*), dissorophoids or certain taxa among Xerodromes with plesiomorphic traits (basal amphibamiforms, basal trematopids) have a short preorbital region, a longer postorbital skull table with elongate rectangular supratemporals, and enlarged orbits in adults, which give most taxa a juvenile appearance.

Most importantly, the enlargement of the orbits, which separated the jugal and lacrimal, correlates a dorsal expansion of the palatine (LEP), a structural innovation that evolved convergently but in a rather different morphological framework within the higher dvinosaurians (Shishkin, 1973). In dvinosaurians, the orbits were not enlarged, but shifted to the side, and only in later Permian–Triassic taxa an LEP was established, whereas the lower Permian forms retained a regular jugal-lacrimal contact.

A second consistent character, the dorsal quadrate process, has been argued to have evolved in the enlarged tympanic notch of dissorophoids (Bolt and Lombard, 1985). In the palate, the interpterygoid vacuities expanded even more substantially than the orbits, pushing the pterygoid, palatine, and ectopterygoid to the margin (Watson, 1940; Carroll, 1964). This probably correlated with enlarged eye musculature (Witzmann and Werneburg, 2017). The pectoral girdle of dissorophoids is much lighter built and the limb bones are more slender and well ossified compared to in most other temnospondyls (Carroll, 1964; DeMar, 1968; Boy, 1972; Daly, 1994), although the distal end of the humerus has a modified condyle, which is still not widely known but may be another synapomorphy of dissorophoids and lissamphibians (Sigurdson and Bolt, 2009).

The placement of the Dissorophoidea within the large-scale phylogeny of temnospondyls remains controversial. According to the existing hypotheses, the immediate sister taxon of dissorophoids may either be: (1) the zatracheids (Schoch, 1997, 2013; Holmes et al., 1998), (2) some or all eryopiforms (Milner, 1990; Yates and Warren, 2000), or (3) the dvinosaurians (Ruta and Bolt, 2006, albeit with *Dendrysekos* interpreted as basal dissorophoid). In an inclusive cladistic analysis, Schoch (2013) found: (1) Dvinosauria; and (2) Zatracheidae to form successive sister taxa of Dissorophoidea, although only the support for Zatracheidae was robust. The implications of a dvinosaurian-zatracheid-dissorophoid clade would leave a somewhat paradoxical situation: features shared by dvinosaurians and basal dissorophoids, such as the long posterior skull table and the laterally placed orbits, would have to be reversed in zatracheids, but these appear to have retained the primitive condition exemplified by eryopiforms. Additional shared derived characters, such as the LEP, are as mentioned before not strictly similar in dvinosaurians and dissorophoids, do not occur in zatracheids, and evolved only late in the dvinosaurian lineage, because basal taxa (*Trimerorhachis*, *Neldasaurus*, and *Erpetosaurus*) do not have these features (Milner and Schoch, 2013; Schoch, 2018), whereas they are present even in the basal-most dissorophoids. In turn, the character-states shared by zatracheids and dissorophoids do not require reversals within dissorophoids, and otherwise do not appear anywhere in temnospondyls (see next section). Therefore, I regard the relationship between dvinosaurians and dissorophoids as unsettled, whereas the likelihood for a zatracheid-dissorophoid clade as relatively high, as measured by the number of synapomorphies shared by the two.

In line with the aforementioned evidence, in the present analysis finds that zatracheids form an unresolved trichotomy with dissorophoids and eryopiforms. Dvinosaurians are found outside this clade in all variant analyses. Interestingly, the omission of zatracheids (represented by *Acanthostomatops*) results in a closer placement of eryopiforms with dissorophoids rather than dvinosaurians. By phylogenetic definition, taxa below Micromelerpetidae fall outside Dissorophoidea (Schoch and Milner, 2014), which puts *Perryella* on a grade outside this taxon.

Zatracheidae as plausible sister group.—The lower Permian Zatracheidae encompasses only three genera: *Acanthostomatops* (Saxony, Germany), *Zatrachys* (Texas, New Mexico,

Oklahoma, and Virginia), and *Dasyceps* (Texas, England, and Czech Republic). In their examination of the basal taxon *Acanthostomatops*, Boy (1989) and Witzmann and Schoch (2006) reported most developmental changes to be consistent with eryopiforms, whereas certain morphological features are shared with dissorophoids. These include the small squarish interclavicle, slender clavicles, the wide squamosal embayment, and the intervomerine and internarial fontanelles. The shared character-states are consistent with the observation that the ontogeny of both *Acanthostomatops* and many dissorophoids involved some metamorphosis, as outlined by Schoch (2009). As in most dissorophoids, zatracheids probably had fully terrestrial adults, whereas the juveniles were gill-bearing larvae similar to those of eryopiforms and the basal dissorophoid *Micromelerpeton* (Boy, 1988, 1989, 1990; Schoch, 2003; Witzmann and Pfretzschner, 2003; Witzmann and Schoch, 2006). In *Acanthostomatops*, metamorphosis was more gradual and stretched-out than in the branchiosaurid *Apateon gracilis*, which underwent a drastic transformation that is more similar to, but less pronounced than that of extant lissamphibians (Schoch and Fröbisch, 2006). The light-built pectoral girdle, stout humerus, and the absence of lateral line sulci indicate that adults of zatracheids were terrestrial, and the remodeling of the hyobranchium as well as shortening of the trunk and extreme broadening of the skull during metamorphosis (Witzmann and Schoch, 2006), which are unique features of the clade, are quite consistent with that. Zatracheids evidently underwent some substantial separate evolution since their ancestors diverged from those of dissorophoids in the early Pennsylvanian.

Perryella as basal taxon.—The small temnospondyl *Perryella olsoni*, from the lower Permian of Oklahoma, was first described by Carlson (1987) who considered it incertae sedis, and later referred it to the Dissorophoidea by Milner (1990). Ruta and Bolt (2006) revised the taxon and provided a masterly detailed description. Schoch and Milner (2014) considered *Perryella* an “amphibamid” (amphibamiform in the present framework), based especially on the narrow ectopterygoid and short palatine ramus of the pterygoid; these features are not well exposed, however. *Perryella* instead forms a plausible candidate for a basal dissorophoid, sharing several plesiomorphies with micromelerpetids. These include the elongate posterior skull table, with supratemporal and squamosal having a long suture, despite a substantial squamosal embayment. The orbits are markedly smaller than in dissorophoids, although the general appearance of the skull roof is similar to large specimens of *Micromelerpeton* and *Eimerisaurus*. As in large *Micromelerpeton* and *Limnogyrinus*, the postorbital is elongate and posteriorly pointed. The palatal bones bear numerous small teeth, and the parasphenoid is covered by a long denticle field on the parasphenoid. The basal plate is large, and the foramina for the internal carotid artery are placed lateral to the denticle field, which is close to micromelerpetids and amphibamiforms. Moreover, the wide jugal, the supinator, and the relatively large quadrangular interclavicle are plesiomorphic features consistent with the primitive condition of dissorophoids.

Characters shared by *Perryella* and Dissorophoidea are the anteriorly reduced palatine ramus and the lost prefrontal-

postfrontal contact, admittedly two features known to have evolved in parallel repeatedly within xerodromes, as well. Due to its more elaborate squamosal embayment and the aforementioned characters, *Perryella* appears to nest outside Dissorophoidea sensu stricto, which makes it an interesting taxon to study the stepwise acquisition of dissorophoid characters. The three most important features of this clade, LEP, DQP, and medial quadratojugal process (MQP), were well established.

Micromelerpetidae.—This clade of small paedomorphic dissorophoids, known since Credner's (1881, 1886) reports on the Saxon lower Permian deposits, had long been lumped together with true branchiosaurids and larvae of seymourimorphs under the name “branchiosaurs” (Fig. 3). Micromelerpetids were first recognized as distinct from true branchiosaurids by Bulman and Whittard (1926), and their morphology was described in detail by Boy (1972, 1995, 2002a, b) and Werneburg (1988, 1991, 1994). They are mostly represented by larval or immature specimens, which co-occur with the smaller branchiosaurids and larvae of eryopiforms (*Sclerocephalus*, *Onchiodon*, *Actinodon*) in Central European lake deposits of lower Permian age (Boy and Sues, 2000). Their possession of an LEP and other dissorophoid characters was discovered rather late after larger adult specimens had been found, (*Branchierpeton*: Werneburg, 1991; *Micromelerpeton*: Boy, 1995), which clearly demonstrated their dissorophoid affinities. Unlike branchiosaurids, micromelerpetids had elongate bodies with 26–29 presacral vertebrae, elongate ribs, well-established lateral line sulci, and a larger quadrangular interclavicle. These features indicate aquatic locomotion and sensation, suggesting the clade formed an early aquatic dissorophoid offshoot. However, it remains an open question whether all taxa of this clade led an aquatic adult life, as large specimens of *Micromelerpeton credneri* lack lateral line sulci and have enlarged eyes, indicated by the sclerotic ring (Boy, 1995; F. Witzmann, personal communication, 2018).

Bulman and Whittard (1926) only knew small adults of *Micromelerpeton* from the Odernheim lake deposit in Germany, whose adult skull length was in the 20–25 mm range. After new excavations had been carried out in stratigraphically older formations, Boy (1995) reported much larger specimens, which had proportionately longer skulls (45 mm), with expanded posterior skull tables and polygonal ornamentation on the dermal bones (Schoch and Milner, 2014, fig. 27A). This morphology is consistent in many features with *Perryella*, which is nonetheless found to nest at the base of Dissorophoidea rather than with micromelerpetids.

Branchierpeton, which was confined to the eastern part of Germany, is known from a stratigraphic sequence of species that probably spanned the Pennsylvanian-Permian boundary (Werneburg, 1989, 1991). In that series, the most ancient species, *B. saalense* Werneburg, 1990, has a slightly narrower skull with a longer postorbital skull table, which resembles that of *Eimerisaurus* and *Micromelerpeton*. This pattern is consistent with the present phylogeny, which contradicts the traditional interpretation of Werneburg (1989), Milner (1993), and Boy (2002a) in which *Limnogyrinus* was the basal taxon and *Micromelerpeton* the most advanced. These early phylogenies were based on a few characters only, and in this case the prefrontal-postfrontal contact in *Limnogyrinus* was given much

weight. Yet the frequent reversals of this character, as exemplified by *Perryella*, trematopids, branchiosaurids, and amphibamiforms, cast doubt on referring exclusively to this feature. Instead, *Limnogyrinus* and its close relative *Nyranerpeton* (Werneburg, 2012), are similar in most features to *Branchierpeton*, and the phylogenetic sequence suggested by the present analysis may be read as a trend towards stronger paedomorphosis. This trend also can be traced within single lineages, such as *Branchierpeton* (*B. saalense*–*B. reinholdi* Werneburg, 1988–*B. amblyostomum* of Werneburg, 1989) and stratigraphically arranged morphotypes of *Micromelerpeton credneri* (Niederkirchen–Odernheim–Humberg of Boy, 1995). The ancient micromelerpetid is thus likely to have looked more like the large, long-skulled *Micromelerpeton credneri* from late Niederkirchen than more paedomorphic *Limnogyrinus*. However, character conflicts cannot be resolved because *Limnogyrinus* has a lower presacral vertebral count than the other taxa, which appears more plesiomorphic.

Micromelerpetids form a clade of early dissorophoids, which successfully explored larger, nutrition-rich lakes by extending the larval period or abandoning metamorphosis (neoteny). Their larvae were less evolved than those of true branchiosaurids (Witzmann and Pfretzschner, 2003), which had specialized marginal teeth, branchial dentition, and often retained an open cheek well within late larval stages (Boy, 1987; Schoch and Milner, 2008). In micromelerpetids, metamorphosis was not as drastic as in branchiosaurids and neotenic specimens are less clearly distinguished from metamorphosed ones (Schoch, 2009). As indicated by the well-known larvae of eryopiforms, micromelerpetid larvae are likely to form the primitive condition for dissorophoids, a point to which I shall return to below when discussing branchiosaurids.

Xerodromes.—Xerodromes form the higher, post-micromelerpetid dissorophoids, which share elongate limb bones, short trunk ribs, and light-built pectoral girdles with a small interclavicle (Fig. 4). Many xerodromes have a disproportionately wide skull, but not necessarily an extended interorbital distance. The postfrontal, postorbital, jugal, and squamosal are the main elements affected by this widening, contrasting markedly with the condition in micromelerpetids, which does not share these features. What is more, the orbits are proportionately larger than in micromelerpetids and the enlarged squamosal embayment wedges deeply between the squamosal and supratemporal, extending onto a large, rounded, unornamented area—this is found in taxa as different as *Dissorophus*, *Platyrhinops*, *Apateon*, and *Doleserpeton*; the trematopid squamosal embayment is to be regarded secondarily modified (Schoch, 2012).

The ancestral xerodrome morphology is not readily assessed because basal taxa are unknown and both olsoniforms and amphibamiforms fall into quite disparate groups with substantial numbers of autapomorphies (Maddin et al., 2013; Schoch and Milner, 2014). Among those representatives with the largest number of plesiomorphies are the amphibamiforms *Tersomius*, *Eoscopus*, and *Platyrhinops* and the olsoniforms *Ecolsonia*, *Mordex*, and *Phonerpeton*. These share with micromelerpetids the elongate postorbital skull table (supratemporals long rectangular, also in *Broiliellus*) and a posteriorly extended basal plate of the parasphenoid. The palatal elements

were covered almost throughout with denticles, as retained in *Platyrrhinops*, *Phonerpeton*, and *Cacops*. The pterygoid reached far anterior to contact the vomer, as exemplified by *Phonerpeton*, *Acheloma*, and *Ecolsonia*. Reduction of the anterior portion of the palatine ramus appears to have occurred in parallel in amphibamiforms and dissorophids, also because the resulting morphology of the palatine ramus differs: in amphibamiforms pushed laterally, often contacting only the ectopterygoid, and in dissorophids aligned anteriorly, with the palatine firmly sutured to the blunt anterior end of the pterygoid. Ancient xerodromes appear to have had well-sutured prefrontal and postfrontal, a feature retained in *Platyrrhinops* and *Amphibamus*, as well as *Mordex*.

Adult xerodromes were apparently advanced over micromelerpetids in having a more definitive terrestrial existence; the two only known clades represent alternative evolutionary strategies to enable enhanced terrestriality: amphibamiforms became small salamander-like insectivores that eventually developed bicuspid pedicellate teeth, whereas olsoniforms evolved into larger predators with enlarged fangs and heavy dermal armor in some forms, as the only dissorophoids to reach sizes around a meter. In both clades, ossification of the braincase and axial skeleton was reinforced as compared with outgroups, with amphibamiforms evolving the cylindrical pleurocentrum, whereas olsoniforms strengthened the intercentrum instead and evolved uncinat processes on their ribs for further reinforcement (Bolt, 1969; Berman et al., 1985; Daly, 1994). The larval life of xerodromes is only known from branchiosaurids, which evolved neoteny and specialized larval characters; the question how widespread branchiosaurid larval features were in post-micromelerpetid taxa remains controversial. Milner (2007) and Werneburg (2012) reported branchiosaurid-like larvae in *Mordex*, but these specimens are more likely to fall within the population of *Branchiosaurus salamandroides* after reexamination. At any rate, the lissamphibian characters evolved mostly within amphibamiforms, with xerodromes only sharing the shortened ribs, light-built dermal pectoral girdle, and the tendency to form a wide skull with modern amphibians—features that were much accentuated in later amphibamiform evolution.

Olsoniformes.—The monophyly of Trematopidae and Dissorophidae had long been questioned (Milner, 1993), but was eventually confirmed after reexamination of postcranial characters and the description of much better preserved skull material in both groups (Reisz et al., 2009; Polley and Reisz, 2011; Schoch, 2012; Maddin et al., 2013; Schoch and Sues, 2013). The present analysis confirms the placement and monophyletic status of the Olsoniformes, a clade that was named by Anderson et al. (2008b). A detailed discussion of olsoniform relationships and their monophyly was given by Schoch (2012).

Amphibamiformes.—This clade is rather uncontroversial and has so far mostly been referred to as Amphibamidae (Schoch and Rubidge, 2005; Huttenlocker et al., 2007; Fröbisch and Reisz, 2008; Bouget and Anderson, 2011; Maddin et al., 2013; Schoch and Milner, 2014). Basal amphibamiforms as found in the present study (*Platyrrhinops*, *Eoscopus*) have rather long presacral vertebral columns in which the pleurocentra reach

somewhat farther ventral than in *Micromelerpeton*, as Daly (1994) has shown. The interclavicle of these forms is quadrangular like in micromelerpetids, but much smaller compared with the skull or vertebral column. In the trunk, the ribs are shorter than in any other group (including olsoniforms), but the first five thoracic ribs are still substantially longer and more robust than the following ones. Basal amphibamiform skulls are heavily ornamented with many small polygonal ridges, and the circumorbital elements remain relatively large with well-established sutures. The posterior skull table houses elongate supratemporals as in outgroups, but the squamosal embayment is huge, and the distance between orbits and squamosal embayment is minimized but usually strengthened by ornament.

There are two alternative phylogenetic hypotheses found by the present analysis that produce equally plausible evolutionary scenarios: (1) *Platyrrhinops* and *Eoscopus* form basal taxa that branched off before micropholids and higher amphibamiforms diverged, or (2) *Platyrrhinops* nests at the base of Amphibamidae and *Eoscopus* at the base of Micropholidae. Because the palate of *Platyrrhinops* retains many plesiomorphies (wide pterygoid, palatine and ectopterygoid, vomer completely covered by denticles, basal plate), this taxon has a rather ancestral appearance. On the other hand, the postorbital skull table is not as elongate as in Micropholidae and *Eoscopus*, which appear more plesiomorphic in this respect. In addition, the squamosal embayment is medially expanded much like in Amphibamidae sensu stricto, resulting in a laterally restricted supratemporal and tabular. *Eoscopus*, on the other hand, has a very wide skull due to enormously enlarged orbits, which resembles that of *Georgenthalia*, branchiosaurids, and especially *Gerobatrachus*; but these are probably convergences, as indicated by multiple lines of conflicting evidence. On the other hand, *Eoscopus* shares only plesiomorphies with micropholids, such as the morphology of the basal plate and posterior skull table, and with *Platyrrhinops* it is consistent in the completely dentigerous vomer, a feature also retained in *Doleserpeton* and *Amphibamus*.

Branchiosauridae.—Over decades, the main “branchiosaur” question was: Why are branchiosaurids not simply the larvae of other amphibamiforms? The short answer to this question is: because adult branchiosaurids are known, and they differ from all other dissorophoids. Adult metamorphosed specimens were described by Werneburg (1991) and Schoch and Fröbisch (2006), who found that during metamorphosis the main branchiosaurid synapomorphies were conserved, and the adult morphology was unlike that of *Amphibamus* or *Doleserpeton*. Instead, branchiosaurids share derived features with *Gerobatrachus*, such as the morphology of the pterygoid and vomer, and the lack of tooth patches on these bones. These features were formerly referred to immaturity, but their presence in adult branchiosaurids, as well as their absence even in immature specimens of *Platyrrhinops* (Clack and Milner, 2010) and *Amphibamus* (Milner, 1982) indicate that these features were independent of developmental changes.

First recognized by Fritsch (1879) as a distinct group, then described in detail by Credner (1881, 1886), the Branchiosauridae were eventually identified as a separate clade by Boy (1972). This was an important step beyond the confusion that

existed since Romer (1939, 1947) had proposed them as larvae of eryopiform temnospondyls. Boy (1971) first suggested that branchiosaurids were not simply larvae of other clades, but that in most branchiosaurid species, adults retained a larval morphology, and concluded by analogy with extant salamanders that they were neotenic forms. Boy (1987) and Werneburg (1991) reported adult specimens in one species, *Apateon gracilis*, which were apparently metamorphosed adults. Whereas Boy (1987, p. 76) referred to them as “matured,” Werneburg (1991) highlighted their metamorphosed (transformed) appearance. The different developmental pathways of branchiosaurid ontogenies were then studied in more detail by Schoch and Fröbisch (2006: metamorphosis) and Fröbisch and Schoch (2009b: neoteny). Finally, Sanchez et al. (2010) were able to confirm the neoteny hypothesis on the basis of skeletochronological and paleohistological data, revealing that sexual maturity was reached in animals retaining a larval morphology. The Branchiosauridae, as defined by Boy (1987) on the basis of synapomorphies, were first found by a cladistic analysis of Schoch and Milner (2008) and later, in the somewhat extended analyses of Fröbisch and Schoch (2009a), they were found as a clade, nesting within the amphibamiforms. The present analysis confirms this hypothesis, based on a much larger set of taxa and characters.

Branchiosaurids have a wide, abbreviated basal plate much like in Amphibamidae, but in contrast to these it lacks a denticle field. Consequently, the internal carotid foramina are located more medially and are more closely set, whereas they are located much farther laterally in larval *Platyrhinops* and Amphibamidae, where the denticle field pushes them far away from the central part of the basal plate. Furthermore, branchiosaurids all share a wide, Y-shaped palatine, which is present throughout ontogeny and not found in other dissorophoids (Boy, 1972, 1987; Werneburg, 1987, 1989, 1991, 2001; Schoch, 1992). The broad lateral wing of the palatine forms the only contact of the palatine with the maxilla, and the ectopterygoid is completely separate from the maxilla. Consequently, the interpterygoid vacuities are narrower than in other amphibamiforms, resembling those of trematopids and micromelerpetids. The palatine ramus of the pterygoid reaches much farther anteriorly than in most other amphibamiforms, which is quite possibly a derived feature that lends more strength to the otherwise strongly reduced palate. Again, this palatal architecture does not only characterize branchiosaurid larvae and neotenes, but also metamorphosed adults, as exemplified by *Apateon gracilis* (Schoch and Fröbisch, 2006).

Milner (1982) suggested that adult *Amphibamus* developed from a branchiosaurid larva, after he had described larval specimens from Mazon Creek originally reported by Moodie (1909) and Gregory (1950). Some of the early larvae he referred to Branchiosauridae proper (*Branchiosaurus*), others to *Amphibamus*. These specimens, although potentially of high significance, are challenging because of poor preservation. My own examination of this material, especially *Eumicrerpeton parvum*, has made me doubtful of this conclusion. The large specimen of *Eumicrerpeton* (USNM 4400; Schoch and Milner, 2014) clearly has a branchiosaurid palate architecture, and development into adult *Amphibamus* is conceivable only when some kind of resorption and reorientation of the palatine, ectopterygoid, and

pterygoid had taken place. Although resorption and reorientation of palatal elements is well known in salamanders and probably forms the primitive condition of the clade (Lebedkina, 1979; Schoch, 1998), the ontogenetic changes of the palate in branchiosaurid metamorphosis were minimal, consisting of a broadening of the palatine, but no reorientation; the palatine ramus of the pterygoid was as long as in larvae, contacting most of the palatine medially.

In the present analysis, the genera *Tungussogyrinus* (?late Permian of Tunguska region, central Siberia) and *Milnererpeton* (Pennsylvanian of New Mexico) were omitted because they are not sufficiently known to be placed with reasonable confidence (Hunt et al., 1996; Shishkin, 1998; Schoch and Milner, 2008). This said, both *Tungussogyrinus* and *Milnererpeton* are now known to share the Y-shaped palatine and free ectopterygoid (Werneburg, 2009; Werneburg et al., 2013) with Branchiosauridae. Both taxa also share the single branchial denticles with Branchiosauridae, first reported by Credner (1881), restored by Stadtmüller (1936), and described in detail by Boy (1972). These form a modification of the plesiomorphic branchial dentition, which consists of small bony plates located on the branchial arches, bearing rows or patches of teeth, occurring in temnospondyl outgroups (eryopiforms, dvinosaurians), colosteid stem-tetrapods (Hook, 1983; Witzmann, 2013), and many osteichthyans. In branchiosaurids, the bony plates were lost and the single teeth were much thinner and often had brush-like tips, which has been interpreted as a filter-feeding device (Boy, 1972). This character forms a larval synapomorphy of branchiosaurids (Boy, 1987), but, unlike the palatal features, their distribution cannot be traced well within Dissorophoidea because unequivocal larval specimens are still only known from micromelerpetids and branchiosaurids (Boy, 1974; Schoch, 2009). Single branchial denticles were also mentioned by Milner (1982) to occur in larval *Amphibamus*. Branchiosaurids may thus have shared the branchial denticles with some or all amphibamiforms, and they are also known to occur in salamandroid caudate *Beiyanerpeton* from the Late Jurassic of Liaoning, China (Gao and Shubin, 2012) and in the Paleogene caudate *Seminobatrachus* (Skutschas and Gubin, 2012). The distribution of this character indicates that the single denticles were probably more widespread among dissorophoids, potentially occurring throughout amphibamiforms or even xerodromes.

Amphibamidae proper.—The two genera *Doleserpeton* and *Amphibamus* share a range of derived characters (Fig. 5), among which are the posteriorly foreshortened squamosal (compensated for by the quadratojugal), the loss of palatine and ectopterygoid tusks, extreme reduction in the width of the ectopterygoid, and reduction of the tabular to a tiny element without distinct horn. In all these features, amphibamids are more derived than branchiosaurids, but not necessarily more similar to lissamphibians. Conversely, the palatal denticles, which cover the complete vomer, pterygoid, palatine, and large areas of the parasphenoid, are a consistent feature of *Amphibamus* and *Doleserpeton*, and even found in juveniles much smaller in size than branchiosaurids. In these features, amphibamids are consistent with *Platyrhinops*, which developed a fully dentigerous palate early in ontogeny (Clack and

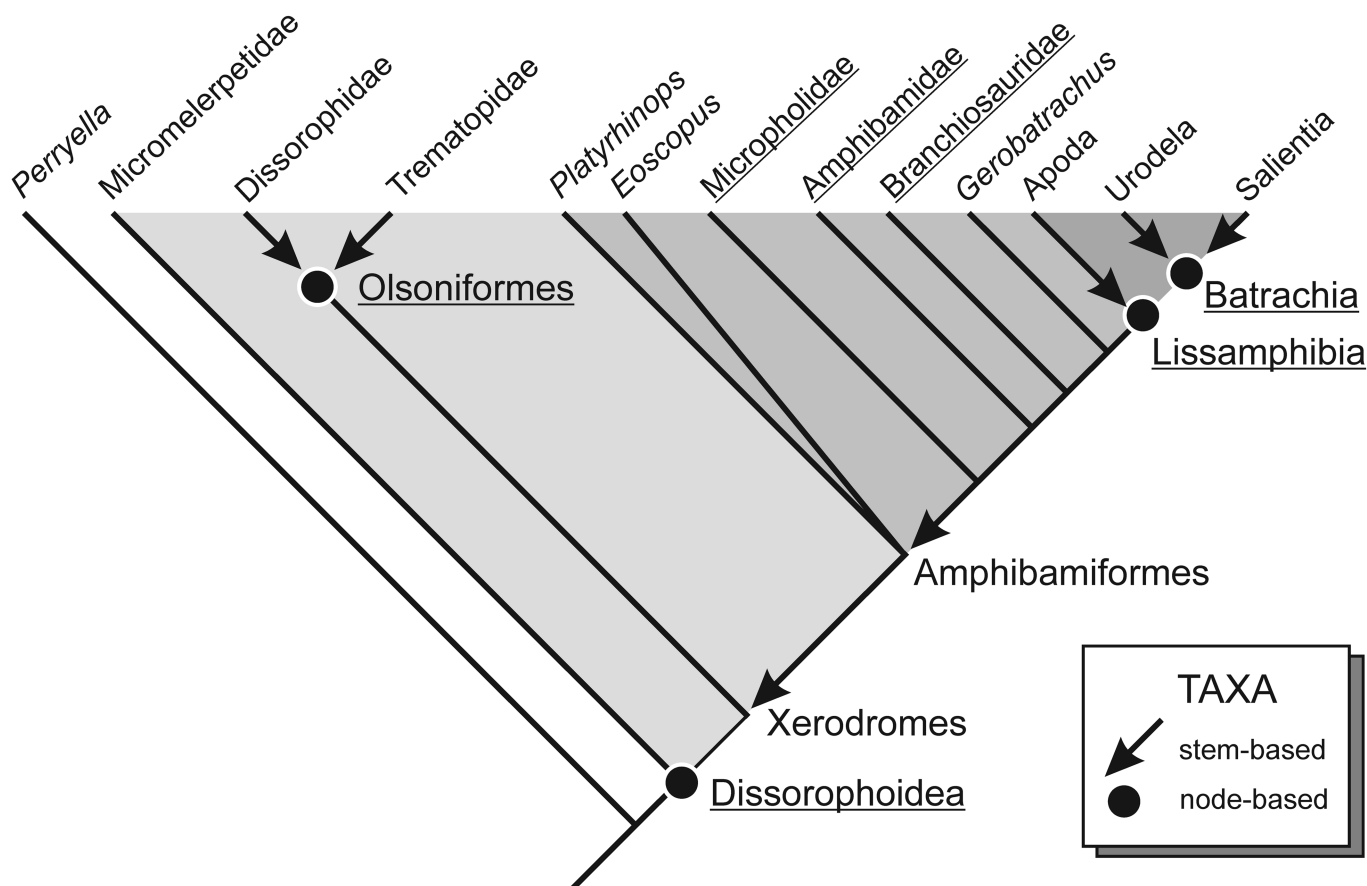


Figure 2. Phylogenetic definition of Dissorophoidea and its major constituent taxa, based on the results of the present analysis.

Milner, 2010). The small larval specimens referred to *Amphibamus* by Milner (1982) indeed have a triangular denticle field on the parasphenoid, which does not occur in branchiosaurids, and the tabular is tiny as in adults; the palatal elements are too poorly preserved to show whether the palatine was Y-shaped or not. Within Amphibamidae, the number of presacral vertebrae was reduced from 24 (*Doleserpeton*) to 18 (*Amphibamus*), a trend that was paralleled within Micropholidae (*Micropholis* having 19) and Branchiosauridae (22–19), where the presacral count is particularly variable within species (Boy, 1972). A strict identification of small larvae on the basis of presacral numbers alone, as attempted by Werneburg (2012) to distinguish taxa, may therefore be misleading. Amphibamids were more lightly built than branchiosaurids, which is also apparent in the more delicate anterior trunk ribs in the former, as well as the less-intense ornament on the dermal skull bones. At the same time, branchiosaurids contrast amphibamids in the more-reduced areas occupied by circumorbital elements, a derived character shared with *Georgenthalia* and *Gerobatrachus*. Furthermore, amphibamids are also plesiomorphic in the retention of dentigerous palatal ossicles, which is consistent with the presence of extensive tooth patches on the vomer and palatine, whereas such ossicles are absent even in large adult branchiosaurids.

Georgenthalia and *Branchiosauridae*.—The incompletely known small amphibamiform *Georgenthalia clavinasica* shares

a few characters with branchiosaurids. Although the exposed palate is more consistent with micropholids and amphibamids proper, the skull roof closely resembles that of large specimens of *Branchiosaurus* and *Apateon*, having large orbits with correlated broad postorbitals, and a much shortened posterior skull table. Also shared with branchiosaurids is the retention of substantial tabular horns. Also alternatives are equally parsimonious, *Georgenthalia* may form a plausible sister taxon to Branchiosauridae, in which the palatal architecture of branchiosaurids was not yet established.

Origin of lissamphibians.—The discovery of *Doleserpeton*, *Eocaecilia*, and *Gerobatrachus* has added substantial data to the controversy over the origin of lissamphibians, yielding some unexpected information on character evolution before and during the early diversification of lissamphibians, or at least batrachians.

Doleserpeton was essential in demonstrating that two lissamphibian characters evolved within amphibamiforms: pedicely and cylindrical pleurocentra, which gave rise to the single cylindrical centrum in Lissamphibia (Bolt, 1969). A third feature, the absence of the ectopterygoid, is now controversial: it was proposed by Bolt (1969) and retained in the new reconstruction of Sigurdson and Bolt (2010), whereas a tiny strut-like ectopterygoid was figured in the same paper (Sigurdson and Bolt, 2010, fig. 2). This is also a problematic character, because *Gerobatrachus*, which is closer to

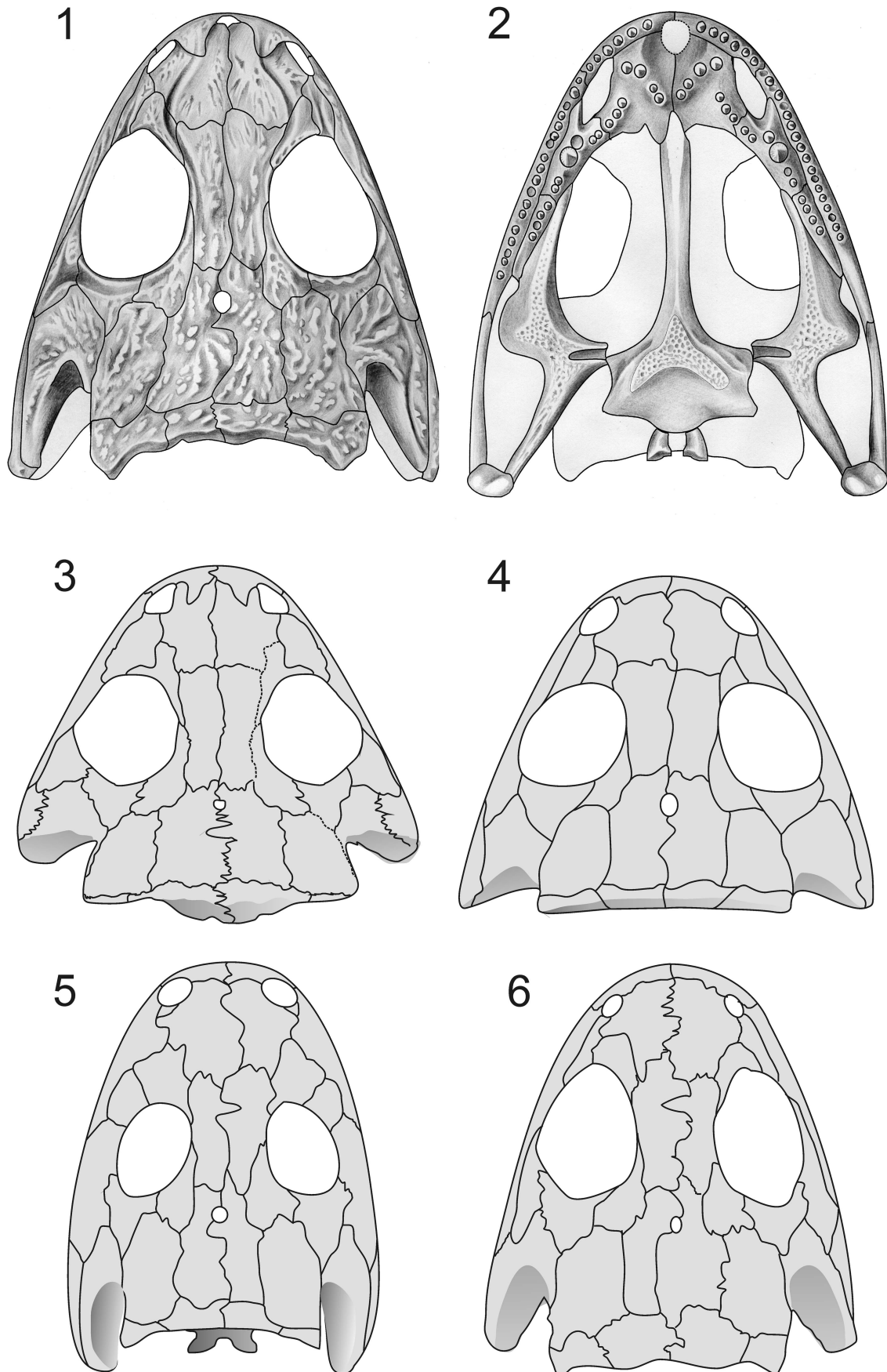


Figure 3. Morphology of basal dissorophoids (micromelerpetids) and stem taxon (*Perryella*). (1) *Micromelerpeton credneri*, skull roof (after Boy, 1995; skull length 25 mm); (2) *M. credneri*, palate (after Boy, 1995; skull length 22 mm); (3) *Limnogyrinus elegans* (personal observation; skull length 18 mm); (4) *Branchierpeton amblyostomum* (after Werneburg, 1991; skull length 20 mm); (5) *Perryella olsoni* (after Ruta and Bolt, 2006; skull length 41 mm); (6) *Eimerisaurus graumanni* (after Boy, 1981; skull length 32 mm).

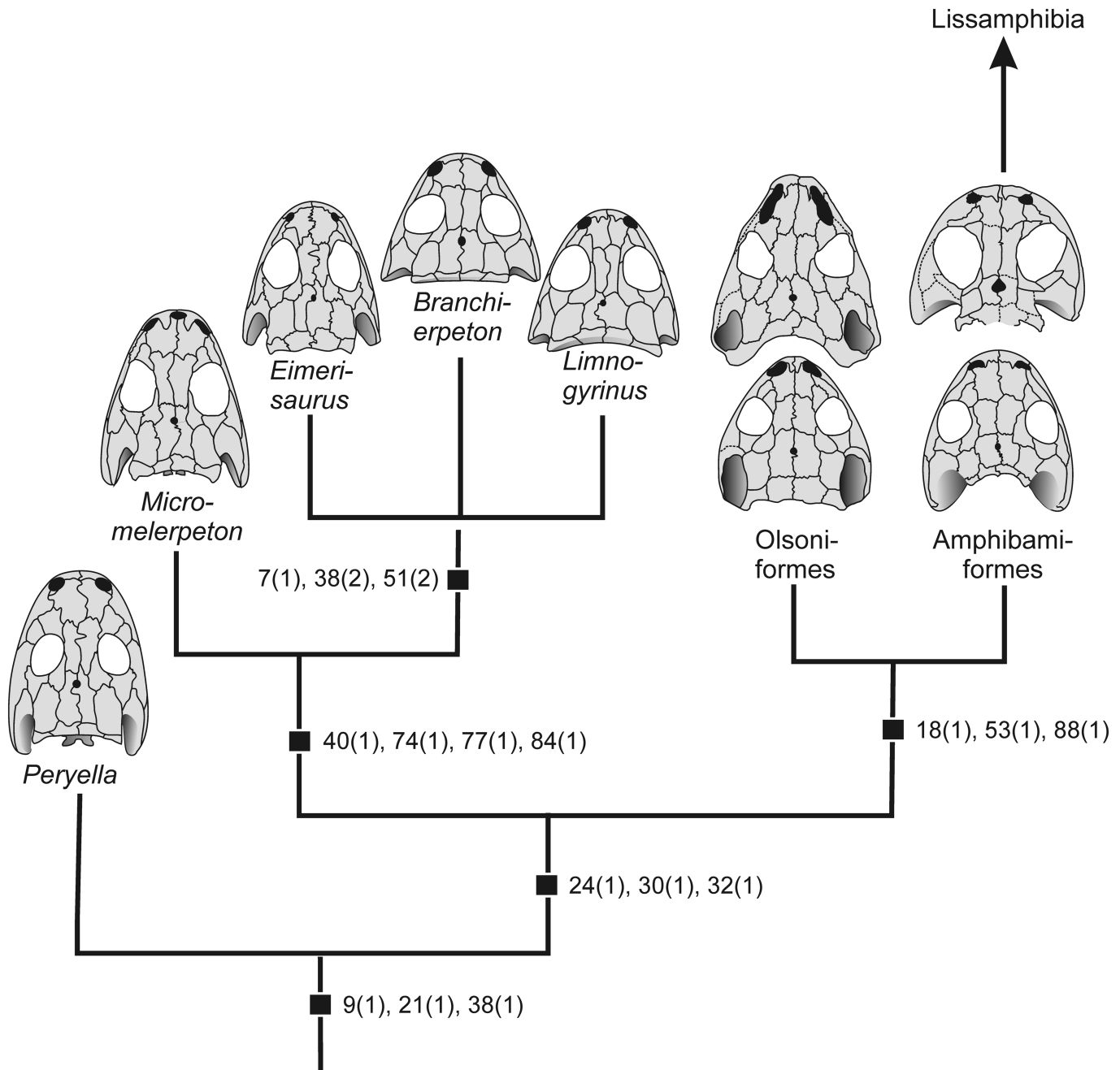


Figure 4. Phylogeny of basal dissorhoids (synapomorphies mapped, see Supplemental Dataset for character definitions).

lissamphibians in a full range of characters, retains quite a substantial ectopterygoid. The only shared character-state of amphibamids and *Gerobatrachus* is the absence of teeth on the ectopterygoid, a feature that also evolved in parallel within branchiosaurids, namely the leptorophine genera *Melanerpeton*, *Leptorophus*, and *Schoenfelderpeton* (Boy, 1987).

Pedicely and bicuspidity were later also found in *Amphibamus* (Bolt, 1979) and some ontogenetic stages of *Tersomius* (Bolt, 1977; Fig. 6). The existence of fused pedicels in the branchiosaurid *Apateon pedestris* remains unclear (Boy, 1978), whereas the report of pedicely in adults of the same taxon by Schoch and Carroll (2003) may be caused by breaks produced by an underlying mandible. Even in *Gerobatrachus*, pedicely

remains controversial; the maxilla teeth of this taxon were shown to have a narrow zone that may have formed by the fusion of pedicellate crowns with their bases (Anderson et al., 2008a).

According to the evidence from *Doleserpeton* and *Tersomius*, pedicely originated in juveniles, after the peg-like larval dentition had been replaced and before fully mineralized, labyrinthodont teeth were produced (Bolt, 1977, 1979). It is conceivable that this stage was not always established, or preserved, because it may have been confined to brief developmental phases. Unlike in many other morphological features, pedicely is a condition that may be derived by incomplete mineralization of a tooth, resulting from the

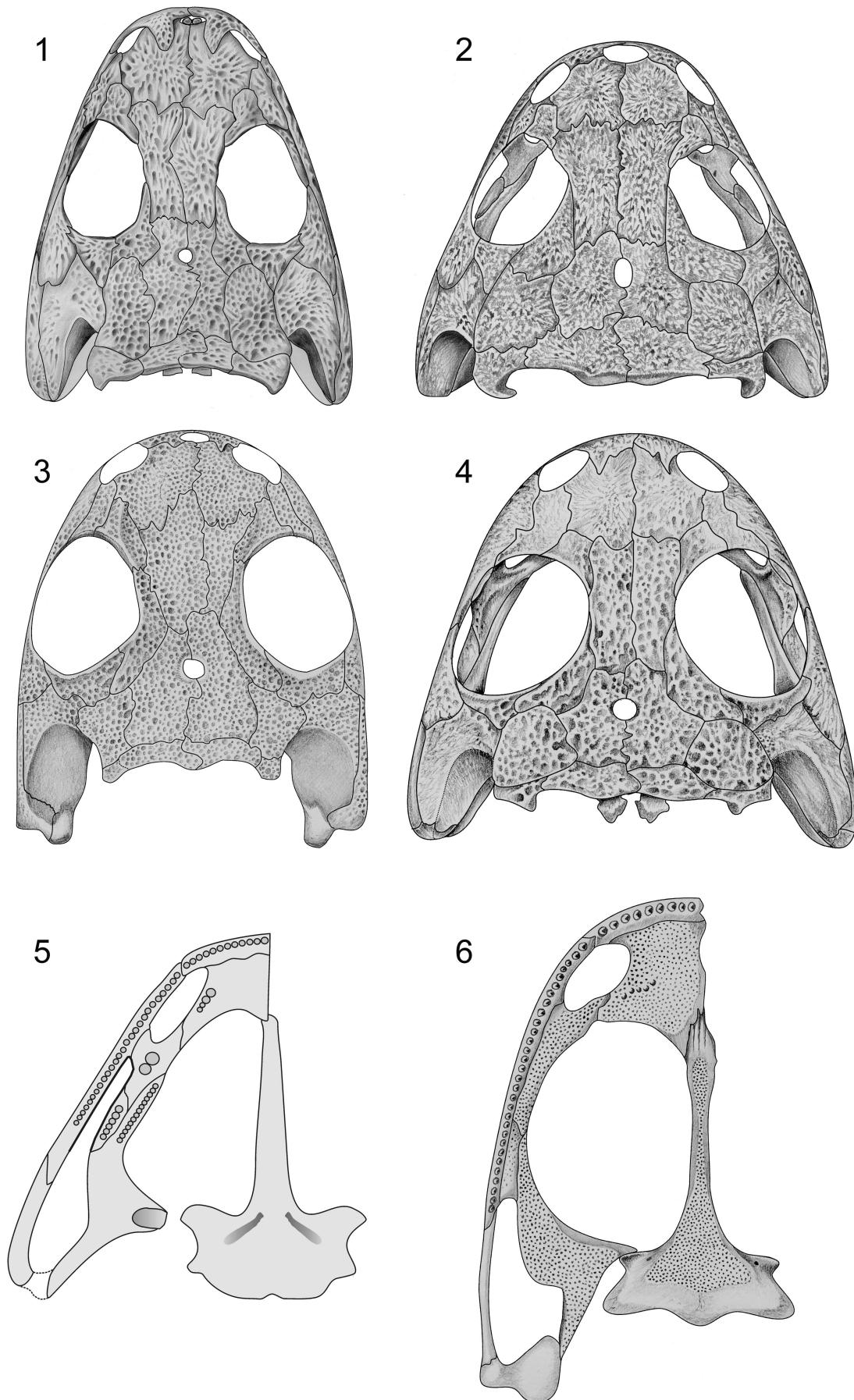


Figure 5. Morphology of amphibamiforms. (1) *Micromelerpeton credneri*, skull roof (Schoch and Milner, 2014; skull length 40 mm); (2) *Apateon caducus*, skull roof of neotenic adult (Fröbisch and Schoch, 2009; skull length 29 mm); (3) *Amphibamus grandiceps*, skull roof (personal observation; skull length 18 mm); (4) *Apateon gracilis*, palate of metamorphosed adult (personal observation; skull length 18 mm); (5) *Apateon caducus*, palate of neotenic adult (after Schoch, 1992; skull length 16 mm); (6) *Amphibamus grandiceps*, palate (personal observation; skull length 18 mm).

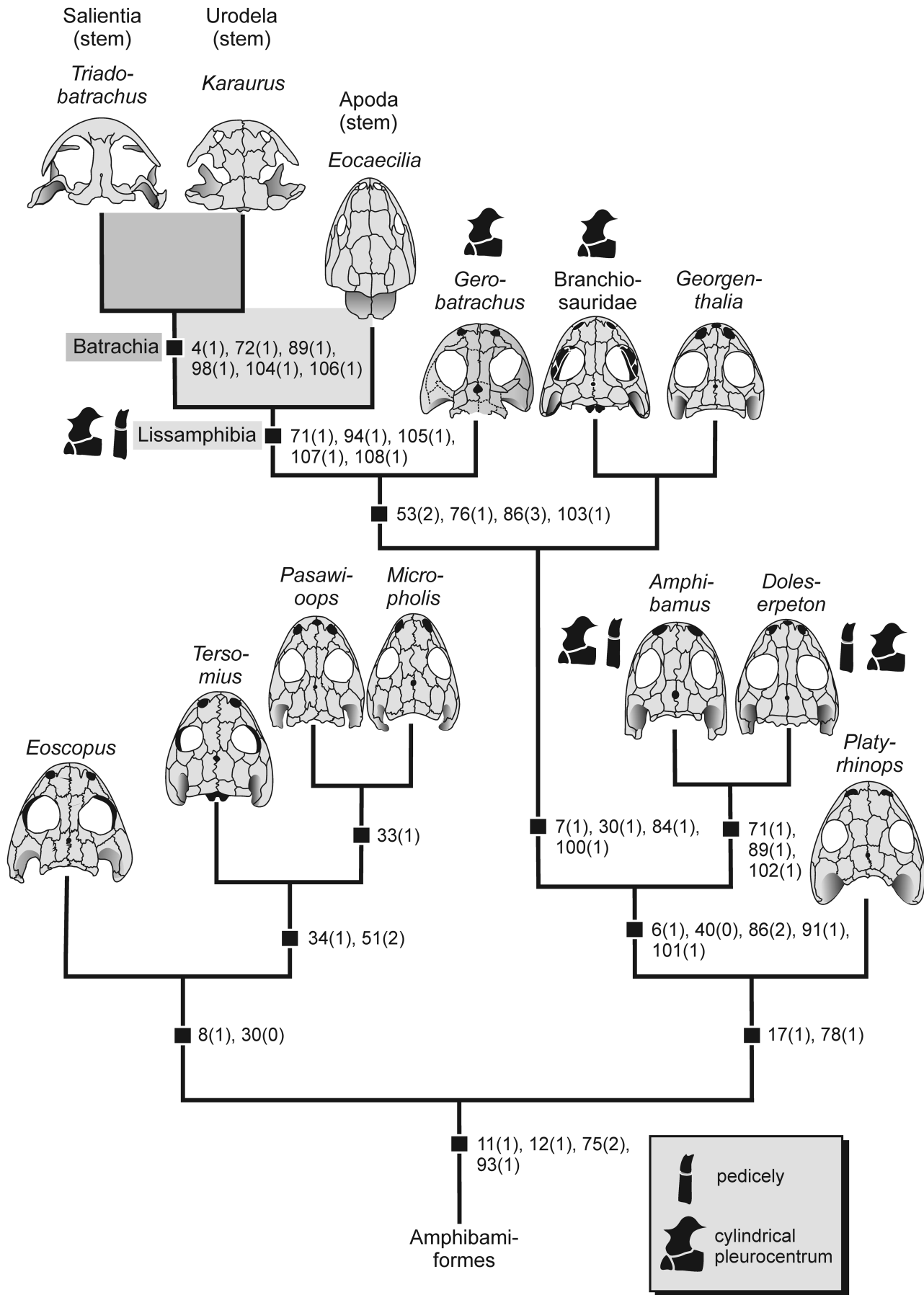


Figure 6. Phylogeny of derived dissorophoids and origin of lissamphibians according to the monophyletic temnospondyl hypothesis (synapomorphies mapped, see Supplemental Dataset for character definitions). Two lissamphibian characters (pedicely and vertebral centrum) mapped.

existence of two separate centers of mineralization (Vasilieva and Smirnov, 2001). Thus, it may well have been absent as a discrete morphology in some stem taxa, whereas the mechanism of tooth production inherently carried pedicely as an option. Eventually it became the standard condition in lissamphibians, but even in these it was repeatedly lost—either because larval teeth were retained by neoteny, or alternatively in taxa whose feeding strategy required stronger conical teeth.

In contrast to pedicely, the distribution of cylindrical pleurocentra in the current phylogenetic topology carries less conflicting evidence. Having been long known in *Doleserpeton* and *Amphibamus*, they have been found in adults of the branchiosaurid *Apateon gracilis* (R. Werneburg, personal communication, 2017), closely resembling those of *Doleserpeton*. In *Gerobatrachus*, as in lissamphibians, the pleurocentrum has become the dominating element.

Gerobatrachus has a parabolically wide skull, huge posteriorly extended orbits, and a disproportionately small skull table. These are batrachian features, and along with the basale commune in the tarsus (an exclusive salamander character) have been cited to support *Gerobatrachus* as a stem-batrachian (Anderson et al., 2008a). However, all other derived features of this taxon are consistent with the hypothesis that it forms a stem-lissamphibian, which was found alternatively by Fröbisch and Schoch (2009a) and in the present analysis.

Unlike amphibamids proper, *Gerobatrachus* retains a remarkably branchiosaurid-like tabular horn, more massive anterior trunk ribs, and a large but autapomorphic ectopterygoid. The latter is unique among dissorophoids in expanding posteromedially to support the palatine ramus of the pterygoid, which is aligned just like in anurans, salamanders, and *Eocaecilia*. Like in branchiosaurids and lissamphibians, the pterygoid is entirely edentulous and its rami are reduced to thin struts, contrasting the shelved and denticle-bearing pterygoid of *Doleserpeton* and *Amphibamus*. Conversely, the palatine is much reduced and has almost lost contact to the ectopterygoid, with its vomerine process broadened much like in anurans (in adult salamanders, the palatine is absent). The rounded maxilla with its wide shelf is consistent with that of branchiosaurids as well as lissamphibians.

Considering character-states shared only by Branchiosauridae, *Gerobatrachus*, and Lissamphibia (see below), the Amphibamidae have lost their exclusive status as immediate lissamphibian sister taxon. This was indicated by Fröbisch and Schoch (2009a) and (implicitly) by Sigurdson and Bolt (2010) who included *Gerobatrachus* in their analysis, where it nested with branchiosaurids rather than amphibamids proper. *Gerobatrachus* is clearly advanced over amphibamids proper and branchiosaurids in the reduced presacral count (17 instead of 18 or 19) and it has a wide parabolic skull shape consistent with that of *Triadobatrachus*, *Karaurus*, and many crown batrachians, although the primitive condition of caecilians is not yet fully known, but the skull of *Eocaecilia* is certainly more consistent with that of amphibamiforms in the structure of the palate than crown caecilians would suggest (e.g., the arrangement of the pterygoid, the shape of the parasphenoid, and the structure of the palatine). The teeth are much smaller, more closely set, and more numerous in *Gerobatrachus* and lissamphibians, as

contrasted with all other amphibamiforms. Moreover, the premaxilla has a thin dorsal (alar) process, again more consistent with that of lissamphibians than with branchiosaurids or amphibamids.

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Accessibility of supplemental data

Data available from the Dryad Digital Repository:
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