Cannibalism in a small growth stage of the Early Permian branchiosaurid *Apateon gracilis* (Credner, 1881) from Saxony

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Abstract

An almost complete specimen of the branchiosaurid temnospondyl *Apateon gracilis* (Credner, 1881) with a skull length of approximately 7 mm from the Early Permian Döhlen Basin in Saxony is described that possesses a nearly complete conspecific specimen of approximately 4 mm skull length in its digestive tract. This is the first evidence of cannibalistic behaviour in small growth stages of branchiosaurids, whose dentition and hyobranchial morphology suggest adaptations to the capture of small invertebrates and feeding on small phyto- and zooplankton by filtering from the water. Therefore, cannibalism in small branchiosaurids certainly represents an exceptional case, possibly triggered by unfavourable environmental conditions.

Introduction

Cannibalism has been observed in a variety of extant amphibians. In some species, cannibalism is a common phenomenon, whereas in others, it is an optional behaviour (Warburg et al. 1979; Crump et al. 1992; Degani 1993; Bressi et al. 1996; Thiesmeier 2004). Most frequently, amphibian eggs and larvae are subjected to cannibalistic predation, and most cannibals are found among larvae of urodelans and anurans. Superficially similar to extant urodelan larvae are the branchiosaurids, a group of temnospondyl amphibians that are known from Late Carboniferous to Permo-Triassic strata and reached an adult body size of 150 mm. Branchiosaurids exhibit larval characters such as branchial dentition and external gills, and most of them are considered to be paedomorphic, water-dwelling animals (Bystrow 1939; Boy 1972; Boy & Sues 2000). Their branchial denticles, attached to the mostly cartilaginous ceratobranchials, are often split distally into several tips and formed a fine trap in the gill region, suggesting that branchiosaurids relied on filter feeding of phyto- and zooplankton at least to some extent (Boy 1993; Boy & Sues 2000). However, large growth stages of branchiosaurids also fed on small vertebrates such as other branchiosaurid species or small fishes. Werneburg (1988a) reported large specimens of *Melanerpeton eisfeldi* (Werneburg, 1988) from Early Permian strata of the Thuringian Forest that preyed on small growth stages of *Apateon kontheri* (Werneburg, 1988), as indicated by gut contents. Rare cases of cannibalism have been reported in which small conspecifics have been discovered in the digestive tract of large branchiosaurids (Werneburg 1989). In the present study, I describe a small growth stage of the branchiosaurid *Apateon gracilis* from the Early Permian Döhlen Basin in Saxony with a smaller, nearly complete conspecific preserved in its digestive tract.

Werneburg (1991) revised *Apateon gracilis* and referred it to *Melanerpeton gracile*. A recent comprehensive phylogenetic analysis of branchiosaurids by Schoch & Milner (2008), however, suggests the assignment of this species to the genus *Apateon* (see also a discussion of characters in Schoch & Fröbisch 2006). Therefore, the binomen *Apateon gracilis* is preferred in the present study. *Apateon gracilis* is a medium-sized branchiosaurid that is represented by different growth stages ranging from 6 to 23 mm skull length, and is the only branchiosaurid species known so far of which metamorphosed, land-dwelling stages have been found (Werneburg 1991; Schoch & Fröbisch 2006). Apart from Niederhäslich in the Döhlen Basin, *Apateon gracilis* has been demonstrated in the southwestern Saale Basin in Thuringia (Werneburg 1988b, 1991). The Niederhäslich de-
positional setting is interpreted as a small, shallow lake with restricted inflow of freshwater (Boy 1977). Fishes have not been found in the lake sediments, whereas a rich tetrapod fauna has been reported. Apart from *A. gracilis*, the micromelerpetontid *Branchierpeton amblystomus* (Credner, 1881), the zatracheid *Acanthostomatops vorax* (Credner, 1883), the eryopid *Onchiodon labyrinthicus* Geinitz, 1861, as well as discosauriscids and microsaurids have been discovered (Boy 1977; Schneider 1994). The specimen described here is the first evidence of cannibalism of a small growth stage of a branchiosaurid and the first documentation of this behaviour in the Döhlen Basin.

**Material**

The specimen described here consists of an individual of *Apateon gracilis* (skull length approximately 7 mm) from the lower limestone horizon of Niederhändisch near Dresden (Döhlen Basin; Asselian) with a smaller conspecific individual (skull length approximately 4 mm) preserved as its gut content. It consists of slab and counterslab that are deposited in the Staatliches Museum für Mineralogie und Geologie Dresden, Saxony, under inventory number SaP 717. 1 and 2, labelled as ‘*Branchiosaurus gracilis*’.

**Description of Apateon gracilis** (Credner, 1881)

The larger of the two specimens of *Apateon gracilis* (Credner, 1881) is represented by an almost complete skeleton including the anterior portion of the tail (Figs 1, 2). The skull is badly crushed so that individual bones cannot be discerned, and the marginal teeth are not preserved. The right stapes is visible at the posterior end of the skull table. Between skull and pectoral girdle, the branchial denticles are well preserved. Five rows of denticles are visible. As described by Werneburg (1991), individual denticles have a proximal gumbolar base and several distal tips (up to four tips can be distinguished). The vertebral column, in which only the neural arches are ossified, is curved, and the short, straight ribs are visible only in the anterior three quarters of the trunk and in the anteriormost portion of the tail. In the posterior portion of the preserved part of the tail, ossified haemapophyses are visible. The pectoral girdle is poorly preserved, only the ill-defined outlines of one cleithrum and clavicle are visible, and the pelvic girdle is not preserved. Fore- and hindlimbs are present and almost complete.

A second, smaller specimen of *A. gracilis* is visible ventral to the vertebral column of the larger specimen, with its longitudinal axis aligned parallel to that of the larger specimen. Its fragmentary skull is located immediately posterior to the forelimbs of the large specimen. Its fragmentary skull is located immediately posterior to the forelimbs of the large specimen. Individual bones of the skull can hardly be identified except for a portion of the dislocated premaxilla with three preserved teeth. The cleithrum is preserved posterior to the skull table. In the anterior trunk region, two rows of branchial denticles are present. The branchial denticles correspond in their morphology to – but are smaller than – those of the large specimen. From the presacral vertebral column, approximately 20 vertebrae are preserved. They continue to the presumed

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**Figure 1.** Photograph of *Apateon gracilis* (Credner, 1881) on slab MMG SaP 717.2.
cloacal region of the large specimen. Thereby, the column extends between the proximal parts of the femora of the large specimen (Fig. 3). Based on the proportions of the vertebral column, the neural arches, and the cleithra, one can estimate the size of the small specimen as being 50–60% of that of the larger one, thus having a reconstructed skull length of approximately 4 mm.

Discussion

Morphological features indicate that the two specimens are conspecific. The position of the smaller specimen relative to the larger one – specifically the fact that its backbone is ventral and parallel to the backbone of the larger specimen and situated between the femora of the latter – suggest that it was preyed upon by the larger specimen, and this testifies to the first record of cannibalistic behaviour in this branchiosaurid. In the stomachs of a sample of *Salamandra salamandra* (Linnaeus, 1758) larvae, the proportions of which are approximately comparable to those of branchiosaurids, Bressi et al. (1996) found conspecifics of about 40% of the size of the predators. However, larvae that measure more than half the size of their cannibalistic predators are also preyed on (Fig. 4; Thiesmeier 2004). As in this extant example of *S. salamandra* larvae, the cannibalistic *A. gracilis* attacked its smaller conspecific from behind and swallowed it tail first.

In branchiosaurids, direct evidence of cannibalism has been described by Werneburg (1989) in two large specimens of *Melanerpeton sembachense* Werneburg, 1989 with a skull length of 26 mm from the Early Permian of the Thuringian Forest, with small specimens of
9 mm skull length located just anterior to and within the mouth and in the trunk, respectively. Boy (1993) and Boy & Sues (2000) reported Early Permian specimens of A. caducus (Ammon, 1889) from the Saar-Nahe Basin with skull lengths exceeding 12 mm that have branchiosaurid prey items preserved in their digestive tracts. It is, however, unclear if the gut remains can be assigned to the same species or to the smaller A. pedestalris Meyer, 1844 (J. A. Boy, pers. comm. 2008). Each of these examples of cannibalistic branchiosaurids apply to specimens of large growth stages. In contrast, the cannibalistic A. gracilis reported here represents a small growth stage with a skull length of only 7 mm. In small growth stages of branchiosaurids, no identifiable gut contents are generally preserved (Boy & Sues 2000). Solely, Werneburg (1986) described a specimen of Apateon flagrifer (Whittard, 1930) from Friedrichroda (Thuringia) with a skull length of 10 mm that had preyed upon four small conchostracan crustaceans. As Boy (1993) and Boy & Sues (2000) pointed out, branchiosaurids of small growth stages probably ingested very small prey items by filter-feeding, especially phyto- and zooplankton, which are, nevertheless not preserved. Additionally, small invertebrates from the lake bottom might have been captured.

The morphology of small growth stages of A. gracilis and other branchiosaurids of less than approximately 12 mm skull length indicate that they were not carnivorous: the small, slender and closely packed marginal teeth (Werneburg 1991) are not suitable for holding large, struggling prey; in addition, a tongue that could be used to facilitate transport of larger prey towards the oesophagus was almost certainly absent, as is the case in extant salamander larvae (Boy & Sues 2000). Therefore, the capture of larger prey items like conspecifics and other vertebrates must have been rare exceptions for A. gracilis of the size class described here, since it must have been obviously very difficult and laborious to capture, fix and swallow this comparatively large prey. In extant anuran and urodelan larvae, cannibalism is obviously under partial genetic control, but triggered by environmental factors. This behaviour can be observed mostly in harsh environments like ephemeral water bodies that are overcrowded by conspecific amphibian larvae, where other food resources are limited or absent (Crump 1992; Degani 1993; Bressi et al. 1996). Boy (1977) presumed that the Niederhäslich lake in the Döhlen Basin may have been temporarily reduced to residual water bodies, in which the aquatic tetrapods were crowded together. Similar to the situation in extant amphibian larvae, the high density of conspecifics and the presumable scarcity of food might have induced the cannibalistic behaviour in our example. However, because of the poor preservation of the skull, I cannot determine if the larger specimen had developed into a morph with enlarged teeth, a larger head, and a wider gape, as has been observed in some cannibalistic larvae of extant amphibians (Crump 1992).

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References


