General Palaeontology, Systematics and Evolution (Vertebrate Palaeontology)

Adaptation of the vertebral inner structure to an aquatic life in snakes: Pachyophiidi peculiarities in comparison to extant and extinct forms

Adaptation de la structure interne vertébrale à la vie aquatique chez les serpents : particularités des achyophiides en comparaison des formes actuelles et éteintes

Alexandra Houssaye a,*, Anthony Herrel a, Renaud Boistel a, b, Jean-Claude Rage c

a UMR 7179 CNRS, Département "Adaptations du Vivant", Muséum national d’histoire naturelle (MNHN), 57, rue Cuvier, CP 55, 75005 Paris, France
b IPHEP-UMR CNRS 6046, UFR SFA, Université de Poitiers, 40, avenue du Recteur-Pineau, 86022 Poitiers, France
c UMR 7207 CNRS, Sorbonne Université, Département "Origines et Évolution", Muséum national d’histoire naturelle (MNHN), 57, rue Cuvier, CP 38, 75005 Paris, France

A B S T R A C T

Bone microanatomy appears strongly linked with the ecology of organisms. In amniotes, bone mass increase is a microanatomical specialization often encountered in aquatic taxa performing long dives at shallow depths. Although previous work highlighted the rather generalist inner structure of the vertebrae in snakes utilising different habitats, microanatomical specializations may be expected in aquatic snakes specialised for a single environment. The present description of the vertebral microanatomy of various extinct aquatic snakes belonging to the Nigerophiidae, Palaeophiidae, and Russelophiidae enables to widen the diversity of patterns of vertebral inner structure in aquatic snakes. A large-scale comparative analysis with extant snakes, including numerous semi-aquatic and aquatic forms, and additional extinct taxa, highlights that, even for snakes specialised for a single environment, vertebral microanatomy does not correlate well with the ecology. Thus, it cannot be used as a proxy for ecological inferences in snakes. In addition, the study emphasizes the strong difficulty in characterizing "osteoosclerosis" in snake vertebrae. Finally, it points out and discusses the peculiarity of the marine hind-limbed snakes, the only snakes showing pachyosteosclerosis.

© 2019 Académie des sciences. Published by Elsevier Masson SAS. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

R É S Ù M É

La microanatomie osseuse apparaît fortement liée à l'écologie des organismes. Chez les amniotes, l'augmentation de la masse osseuse est une spécialisation microanatomique souvent rencontrée chez les taxons aquatiques qui réalisent de longues plongées à faible profondeur. Si les travaux antérieurs ont mis en évidence le caractère plutôt généraliste de

Mots clés :
Serpent
Vertèbre
Microanatomie

* Corresponding author.
E-mail addresses: houssaye@mnhn.fr (A. Houssaye), anthony.herrel@mnhn.fr (A. Herrel), rboistel@gmail.com (R. Boistel).

https://doi.org/10.1016/j.crpv.2019.05.004
1631-0683/© 2019 Académie des sciences. Published by Elsevier Masson SAS. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Please cite this article in press as: Houssaye, A., et al., Adaptation of the vertebral inner structure to an aquatic life in snakes: Pachyophidi peculiarities in comparison to extant and extinct forms. C.R. Palevol (2019), https://doi.org/10.1016/j.crpv.2019.05.004
1. Introduction

Extant aquatic snakes are represented within several taxa, e.g., Acrochordidae, Homalopsidae, Natricidae, Elapidae, which (non-exclusively) adapted to a great variety of freshwater and marine habitats (Ineich, 2004). Adaptation to an aquatic life occurred independently within these taxa and even several times within Elapidae (Ineich, 2004).

In addition, five extinct families of snakes adapted to various freshwater and marine environments: the Anomalogophiidae, Nigerophiidae, Palaeophiidae, Palaeophiidae, and Russelophiidae. These various extinct aquatic snake taxa are known from the Cenomanian to the end of the Eocene (that is, from about 100 to 34 Myr ago).

Various adaptive features associated with an aquatic life are encountered in extant aquatic snakes, such as a permeable skin to facilitate gaseous exchange, a sublingual gland for salt excretion, a streamlined head shape minimizing drag, a paddle-like tail to increase propulsion (Aubret and Shine, 2008; Babonis and Brischoux, 2012; Ineich, 2004; Segall et al., 2016, 2019). However, most of these anatomical features are not preserved in the fossil record. Conversely, one major adaptation to an aquatic lifestyle is observed in the vertebrae and ribs of the Cenomanian extinct Pachyphiiidae: bone mass increase (see Houssaye, 2009; Ricqlès and de Buffrénil, 2001). In these taxa, bone mass increase consists of pachyostosis, i.e. the combination of an increase in periostal bone deposits (pachyostosis), which confers to the bones a bloated aspect, and an increase in inner bone compactness (osteosclerosis; Houssaye, 2013). This specialization is thought to be associated with a passive control of buoyancy and body trim and it is generally encountered in poorly active swimmers performing long dives in shallow water environments (Houssaye, 2009; Ricqlès and de Buffrénil, 2001; Taylor, 2000). Pachyostosis has, however, not been mentioned for any other extant or extinct snake taxon. A high bone compactness has been documented in vertebrae of only a few extant aquatic snakes (Houssaye et al., 2013a). This raises the question of:

• whether bone mass increase, observed in many amniotes that have secondarily adapted to an aquatic life (see Houssaye, 2009; Houssaye et al., 2016; Ricqlès and de Buffrénil, 2001 for a review), is indeed absent in these other lineages of aquatic snakes;
• how snakes differently adapted to an aquatic lifestyle in terms of their vertebral inner structure.

Vertebrae and ribs are the bones the most commonly affected by bone mass increase when it occurs (Houssaye, 2009). Moreover, vertebrae correspond to most of the snake skeleton and are the fossil snake bones the most frequently discovered. The vertebral microanatomy (i.e. the distribution of the osseous tissue in the bone) of numerous extant snakes has previously been investigated (Buffrénil and Rage, 1993; Buffrénil et al., 2008; Houssaye et al., 2010, 2013a). As for extinct snakes, data are conversely rare. They essentially pertain to pachyphiiids (Buffrénil and Rage, 1993; Houssaye, 2013; Houssaye et al., 2011) and to a lesser extent to Palaeophiidae (Buffrénil and Rage, 1993; Houssaye et al., 2013). The current study analyses the inner structure of vertebrae from several extinct snakes considered aquatic or possibly aquatic. Coupled with the addition of diverse aquatic snakes in the comparative sample of extant taxa, it will enable:

• to illustrate the occurrences of bone mass increase in aquatic snakes and to obtain a broader picture of the diversity of vertebral inner structure patterns in aquatic snakes;
• to correlate these patterns with ecological features in extant forms and to possibly make inferences about the paleoecology of the extinct snakes sampled;
• to discuss the different types of adaptation of the vertebrae to an aquatic lifestyle in snakes throughout their evolutionary history.

2. Material and methods

2.1. Material

2.1.1. Fossil material

Anomalogophiidae, Nigerophiidae, Palaeophiidae, and Russelophiidae are only represented by postcranial elements (Rage, 1984). Their systematic assignment was made
through vertebral characters that are diagnostic within snakes. Pachyophiidae are defined as including Pachyophis, Pachyrhachis, and all taxa more closely related to these genera than to extant snakes (Lee et al., 1999). Their status and phylogenetic position are strongly debated (Martill et al., 2015; Palci et al., 2013a, b; Reeder et al., 2015).

Concerning the fossil material, taxa from three of the five families encompassing aquatic forms were sampled (Table 1). Some data relative to Pachyophiidae are already available (Buffrénil and Rage, 1993; Houssaye, 2013). The Anomalophiidae are represented by a single species, Anomalophis (Archaeophis) bolcensis Auffenberg, 1959, from the lower Eocene of Italy, whose remains are scarce.

Two Nigerohiphidae were sampled: the genera Nigrophis and Indophis. Nigerohiphidae are known from the Cenomanian and Palaeocene of Africa and Asia. Moreover, possible nigerophiids were discovered in the latest Cretaceous of India and in the Eocene of Europe (Rage and Werner, 1999). Three precolacal vertebræ of Nigrophis mirus Rage, 1975a, from the Palaeocene outcrop Krebb de Sessao, in Niger, were analysed (Fig. 1). Their original position along the vertebral column cannot be determined in the absence of comparative material. The depositional milieu corresponds to a shallow marine environment (Capetta; pers. com. 2007). We also analysed four vertebræ of Indophis sahnii Rage and Prasad, 1992, from the Maastrichtian of Naskal, Andhra Pradesh, India. VPL/JU Unnumb.3–4 are isolated centra, so that their original position along the vertebral column cannot be determined. As for VPL/JU Unnumb.2, it is considered a posterior trunk vertebra and VPL/JU Unnumb.1 a mid-trunk one (JC. R. pers. obs.). The environment corresponds to a floodplain/brackish water (Rage et al., 2004), which is in agreement with the vertebral characters illustrating adaptation to an aquatic life (Rage and Prasad, 1992). These bones are particularly small, so that only one section per vertebra could be made (except for VPL/JU Unnumb.2; Fig. 2). Moreover, correct sectional planes (see below) were difficult to obtain.

Within Palaeophiidae, species from the two genera Palaeophis (Maastrichtian–Eocene) and Pterosphenus (Eocene) were investigated. These snakes, which display various degrees of adaptation to an aquatic life, lived in marine or marginal marine waters (e.g., estuaries, deltas, lagoons, mangroves; Parmley and Devore, 2005). Palaeophis colossaeus Rage, 1983a and P. maghrebianus are considered among the Palaeophiidae, and these species are thought to be less adapted to an aquatic life (Rage, 1983b). The P. colossaeus sections analysed, which were previously broadly described by Buffrénil and Rage (1993), were available in the collections of the MNHN. They were made from vertebræ discovered in the Lutetian of Tamagnuité, Mali. Sections from vertebræ of P. maghrebianus A Bamourg, 1952, from the Ypresian of the Phosphates of Morocco were previously described by Houssaye et al. (2013b). Five vertebræ assigned to Palaeophis typaeus Owen, 1850 (Fig. 3), two from Palaeophis toliapiscus Owen, 1841, and one of an undetermined Palaeophis species, from the Ypresian of Prémontré, Aisne, northern France and Egem, Pittem, Belgium, were analysed. These two species are considered more highly adapted to an aquatic life than P. maghrebianus (Rage et al., 2003). We also analysed two vertebræ from Pterosphenus schucherti Lucas, 1899, from the late Eocene of Georgia, USA. This species, assumed to have lived in estuarine or low-salinity environments (Hutchison, 1985; Westgate and Ward, 1981), is thought to be one of the most adapted to an aquatic life within Palaeophiidae (Rage et al., 2003).

As for Russellophiidae, we sampled the species Russellophis tenus Rage, 1975b, from the early Eocene of France and India. The two vertebræ come from the late Ypresian of Condé-en-Brie, Aisne, northern France. This taxon was thought to live in rivers, lakes, and estuaries (Rage, 1983b).

For comparative purposes, sections of two unnumbered vertebræ of the pachyophiid snake Similophis rochbrunei from Les Renardières, in Charente-Maritime, France, described in Buffrénil and Rage (1993), and three vertebræ of Palaeophis maghrebianus from the Ypresian Phosphates of Morocco, described in Houssaye et al. (2013b), were also analysed, but not described.

2.1.2. Extant taxa

Our sample includes the various semi-aquatic and aquatic extant snakes from Houssaye et al. (2013a) and additional semi-aquatic and aquatic taxa (Table 2; Fig. 4). These taxa were added in order to better represent the diverse ecologies encountered within aquatic snakes, with deep divers (e.g., Hydrophis peronii, Hydrophic elegans) and shallow swimmers (most Aipysurus species), the unique pelagic species (Hydrophis platius), taxa occupying diverse habitats (e.g., Hydrophis stokesi and Hydrophis curtus) and others with a more specialized niche (Enydocephalus annulatus), live-bearers (all homalopsids) and oviparous taxa (Laticauda species), forms living almost exclusively in water (hydrophids, acrochordids), and some regularly coming onto land (Laticauda species; Heatwole, 1999).

All vertebræ of extant taxa are mid-precolacal ones, where bone mass increase is supposed to be the most intense when it occurs (see Houssaye, 2013 for data relative to Pachyophiidae). Fossil vertebræ are all precolacal, but their position along the vertebral column is variable (see above) as the sampling choice was limited. The strongest variation in the intensity of bone mass increase in pachyophiids, however, essentially concerns pachyostosis, whereas osteosclerosis is more homogeneous along the precolacal region (Houssaye, 2013). The impact of this potential bias for compactness comparisons is therefore considered limited.

3. Methods

Various methods were used to investigate the vertebral microanatomy of our sample [for the material already digitized for Houssaye et al. (2013a), see details therein]. If some classical thin sections could be made for a few specimens, virtual ones, through the use of microtomography, were preferred for most specimens, especially because of the rarity of this material and the wish to use a non-destructive technique (Tables 1 and 2). Classical thin sections were made in the mid-sagittal and the neutral transverse planes using standard techniques (see Houssaye et al., 2008). However, as Indophis vertebræ
Table 1
List of the material of extinct taxa analysed. Abb.: abbreviation [used in Fig. 14]; Resol.: resolution (in μm). All fossil specimens were scanned at the ESRF, Grenoble, France. Cts and Cls: compactness (in %) in longitudinal and transverse sections, respectively; CL: centrum length (in mm); USTL: Université des Sciences et Techniques du Languedoc, Montpellier, France; VPL: Vertebrate Palaeontology Laboratory, University of Jammu, Jammu, India; MNHN: Muséum national d’histoire naturelle, Paris, France; Unnumb.: unnumbered.

<table>
<thead>
<tr>
<th>Family</th>
<th>Taxon</th>
<th>Abb.</th>
<th>Age</th>
<th>Locality</th>
<th>Collection reference</th>
<th>Resol.</th>
<th>Cts</th>
<th>CL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nigerophiidae</td>
<td>Nigrophiops mirus</td>
<td>Nm</td>
<td>Palaeocene</td>
<td>Krebb de Sessao, Niger</td>
<td>USTL SES 105</td>
<td>x</td>
<td>93.9</td>
<td>75.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>USTL SES 106</td>
<td>x</td>
<td>93.9</td>
<td>75.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>USTL SES 107</td>
<td>x</td>
<td>93.9</td>
<td>75.0</td>
</tr>
<tr>
<td>Indophiidae</td>
<td>Indophiops sahnii</td>
<td>X</td>
<td>Maastrichtian</td>
<td>Naskal, India</td>
<td>VPL/JU Unnumb. 1</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>VPL/JU Unnumb. 2</td>
<td>x</td>
<td>x</td>
<td>2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>VPL/JU Unnumb. 3</td>
<td>x</td>
<td>x</td>
<td>1.9</td>
</tr>
<tr>
<td>Palaeophiidae</td>
<td>Palaeophis colosseus</td>
<td>P</td>
<td>Lutetian</td>
<td>Tamaguilelt, Mali</td>
<td>MNHN Unnumb. 1</td>
<td>x</td>
<td>x</td>
<td>69.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MNHN Unnumb. 2</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MNHN Unnumb. 3</td>
<td>x</td>
<td>x</td>
<td>1.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MNHN Unnumb. 4</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MNHN Unnumb. A</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MNHN Unnumb. B</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Palaeophis maghrebianus</td>
<td>Pm</td>
<td>Ypresian</td>
<td>Phosphates of Morocco</td>
<td>OCP DEK/GE 644</td>
<td>x</td>
<td>92.1</td>
<td>90.4</td>
<td>11.6</td>
</tr>
<tr>
<td>Palaeophis typhaeus</td>
<td>Pt</td>
<td>Ypresian</td>
<td>Prémontré, France</td>
<td>MNHN Unnumb.</td>
<td>20.2</td>
<td>54.4</td>
<td>60.9</td>
<td>13.8</td>
</tr>
<tr>
<td>Palaeophis toliacicus</td>
<td>Pto</td>
<td>Ypresian</td>
<td>Egem, Belgium</td>
<td>MNHN Unnumb. C</td>
<td>5.0</td>
<td>53.7</td>
<td>71.5</td>
<td>5.3</td>
</tr>
<tr>
<td>Palaeophis sp.</td>
<td>Ps</td>
<td>Ypresian</td>
<td>Egem, Belgium</td>
<td>MNHN Unnumb.</td>
<td>5.0</td>
<td>53.7</td>
<td>71.5</td>
<td>5.3</td>
</tr>
<tr>
<td>Pterosphenus schucherti</td>
<td>Ps</td>
<td>Late Eocene</td>
<td>Georgia, USA</td>
<td>MNHN Unnumb. A</td>
<td>5.0</td>
<td>53.7</td>
<td>71.5</td>
<td>5.3</td>
</tr>
<tr>
<td>Russelliophiidae</td>
<td>Rustelliophis tenuis</td>
<td>Rt</td>
<td>Late Ypresian</td>
<td>Condé-en-Brie, France</td>
<td>MNHN Unnumb. A</td>
<td>5.0</td>
<td>53.7</td>
<td>71.5</td>
</tr>
<tr>
<td>Pachyophiidae</td>
<td>Simoliophis rochebrunii</td>
<td>Sr</td>
<td>Cenomanian</td>
<td>Charente-Maritime, France</td>
<td>MNHN Unnumb. A</td>
<td>5.0</td>
<td>53.7</td>
<td>71.5</td>
</tr>
</tbody>
</table>

Fig. 1. Nigrophiops mirus. Krebb de Sessao, Niger. Palaeocene. A–B. Vertebra USTL SES 105. C. Vertebra USTL SES 106. D–F. Vertebra USTL SES 107; in A, D, left lateral; B, dorsal; C, F, anterior; E, ventral views. The scale bars equal 1 mm.

Fig. 2. Indophiops sahnii. Naskal, Andhra Pradesh, India. Maastrichtian. VPL/JU Unnumb. 2 in A, anterior; B, left lateral; C, ventral; and D, dorsal views. The scale bar equals 1 mm.

Please cite this article in press as: Houssaye, A., et al., Adaptation of the vertebral inner structure to an aquatic life in snakes: Pachyophiida peculiarities in comparison to extant and extinct forms. C.R. Palevol (2019), https://doi.org/10.1016/j.crpv.2019.05.004
were particularly thin, only one section per vertebra could be made for three of them. High-resolution computed tomography was used for numerous extant specimens at:

- the Steinmann Institut, University of Bonn, Germany (GE PhoenixX-ray v(tome)xs 180 and 240; resolution between 7.5 and 89.9 µm; reconstructions performed using datax/res software);
- the Montpellier Rio Imaging (MRI; Microtomograph RX SkyScan 1076; resolution: 9.4 µm; reconstructions performed using NRecon software);
- the University of Poitiers, France, using a X8050-16 Viscom model (resolution: between 15.7 and 34 µm; reconstructions performed using Feldkamp algorithm with DigiCT software, version 1.15 [Digisens SA, France]) at the laboratory Études–Recherches–Matériaux (ERM, Poitiers, France; http://www.erm-poitiers.fr).}

For the fossil ones, we resorted to synchrotron microtomography on the ID 19 beamline (resolution of either 7.5 or 20.2 µm) at the European Synchrotron Radiation Facilities (ESRF, Grenoble, France); reconstructions were performed using the filtered back-projection algorithm with the ESRF PyHST software.

Three measurements were taken on the sections using ImageJ (Abramoff et al., 2004):

- global compactness in transverse section (Cts), calculated as the total sectional area minus the area occupied by cavities and the neural canal multiplied by 100 and divided by the total area minus the area occupied by the neural canal;
- global compactness of the centrum in longitudinal section (Cls), calculated as the total area of the centrum minus the area occupied by cavities multiplied by 100 and divided by the total area of the centrum;
- centrum length (CL), considered as an indicator of size.

Welch’s t-tests (Welch, 1947) were performed on compactness data to test for differences pending on ecological categories (semi-aquatic, aquatic, terrestrial and arboreal, fossorial).

4. Description

This section describes the inner structure of the fossil vertebrae sampled. It then proposes a comparative analysis based on data from extant snakes and extinct taxa previously described.

4.1. Extinct taxa

None of the specimens analysed displays any sign of pachyostosis.

4.1.1. Nigrophis mirus

In longitudinal sections, the relative surface occupied by primary periosteal bone varies between vertebrae (Fig. 5). Indeed, primary periosteal bone resorption is restricted to the area surrounding the neutral point (point where the centrum growth starts; sensu Buffrénil et al., 2008) in USTL SES 105 and especially USTL SES 106 (Fig. 5A) so that remodelling almost exclusively occurs in the endochondral territory. However, periosteal bone resorption is more intense in USTL SES 107, where a wide cavity occupies the core of the centrum (Fig. 5C). Compactness is generally relatively high (78% < Cts < 94%); Cavities are randomly shaped so that there is no true trabecular network in the endochondral territory. In the three vertebrae, remains of calcified cartilage are important in the core of trabeculae close to the neutral point (Fig. 5D).

In transverse sections, compactness is very high (95% < Cts < 98%). Remodelling is extremely limited. Lacking in vertebrae USTL SES 105–106 (Fig. 5B), it is observed from the core of the centrum to the base of the neural canal in USTL SES 107.

4.1.2. Indophis sahnii

In longitudinal sections, various patterns are observed. Some vertebrae (VPI/JU Uunnub.2-3; Fig. 6A) are very compact and show a strong inhibition of primary periosteal bone resorption. Conversely, another vertebra (VPI/JU Uunnub.4; Fig. 6B) appears highly remodelled in both endochondral and periosteal territories.

In transverse sections, vertebrae appear very compact (Fig. 6C). The neural canal is wide and surrounded by a unique layer almost exclusively consisting of primary periosteal bone, except at the base of the neural canal and around some small cavities.

4.1.3. Palaeophis colossaeus

In most longitudinal sections, both endochondral and periosteal territories are occupied by a remodelled spongiosa (Fig. 7A), whereas the primary periosteal bone
displaying radially oriented vascular canals (like in all palaeoophids) is restricted to the ventral edge of the centrum and the upper part of the cotyle. Inter trabecular spaces are irregularly shaped and randomly oriented. They are much smaller toward the epiphyses. Global compactness indices of 57.3% and 69.4% were calculated for the two sections with a complete centrum. One longitudinal section (Fig. 7B) displays a relatively limited resorption of primary periosteal bone, so that compactness is higher.

In transverse sections, compact bone generally consists of two layers surrounding the neural canal and the periphery of the vertebras (Fig. 7C and E). They are connected by a spongiosa that occupies most of the section. Cavities are randomly shaped and distributed. In the biggest specimens, they are much more numerous and relatively smaller than in the smallest ones, conferring a honeycomb structure (Fig. 7D), which is in agreement with the description of Befrérin and Rage (1993). A global compactness index of 69.8% was calculated for MNHN Unnumb. 1. In the biggest sections, remodelling is very limited at the base of the neural arch causing a sharp interruption of the structure in “double-rings enclosing a
spongiosa” (Fig. 7D). Furthermore, one section (Fig. 7E) displays a particularly thick layer of primary periosteal bone surrounding its periphery. This section appears thus characterized by a relative inhibition of primary periosteal bone resorption.

The differences highlighted in both longitudinal and transverse sections suggest significant intraspecific and/or intracolumnar variability in this taxon.

4.1.4. *Palaeophis typhaeus*

All vertebrae display a wide cavity in the core of the centrum (Fig. 8). However, the thickness of the ventral layer of compact cortex and the tightness of the spongiosa vary between the vertebrae in longitudinal section, from a thick compact cortex (Fig. 8A) to a thinner cortex surrounding a loose spongiosa (Fig. 8C). In transverse section, whereas some vertebrae show a compact neural arch and neural spine (Fig. 8B), others show a hollow neural spine sometimes associated with large cavities in the ventral part of the neural arch (Fig. 8D). As a result, some vertebrae are much more compact (MNHN Unnumb. A-B) than others (MNHN Unnumb. C-D).

4.1.5. *Palaeophis toliapicus*

The structure observed in *P. toliapicus* (Fig. 9) is very similar to that previously described in *P. typhaeus*.

4.1.6. *Palaeophis sp.*

This is also the case for the vertebra of *Palaeophis sp.* (Fig. 10).

4.1.7. *Pterosphenus schucherti*

The two longitudinal sections display similar features. Like in the other palaeophiids, a wide cavity occupies the core of the centrum, whereas the rest of the section consists of a spongiosa (Fig. 11). The latter is loose in the periosteal territory, with wide randomly shaped intertrabecular spaces, but much tighter in the endochondral territory, with relatively small numerous cavities (Fig. 11A). In transverse section, MNHN Unnumb. A displays a relatively feebly remodelled structure. Wide resorption cavities are restricted to the neural spine and the core of the centrum, whereas the rest of the section is relatively compact, with rather small and scarce cavities (Fig. 11B). Conversely, resorption is much more intense in MNHN Unnumb. B, which displays the structure in “double-rings connected by very few trabeculae” characteristic of extant squamates (Houssaye et al., 2010; Fig. 11C). Compactness is thus much lower in the latter (Cts = 49.1% versus 82.5%).

4.1.8. *Russellospis tenuis*

In longitudinal sections, vertebrae are highly compact (mean Cts = 81.2%). Periosteal bone remodelling appears very inhibited. There is no true trabecular network in the endochondral territory, but a few wide sub-sagittal cavities occur (Fig. 12A–B). Vascularization appears lacking in primary periosteal bone. In transverse sections, cavities are almost lacking. Compactness is thus very high (mean Cts = 98.9%). The neural canal is rather wide in this species, like in *Indophis*.

---

Please cite this article in press as: Houssaye, A., et al., Adaptation of the vertebral inner structure to an aquatic life in snakes: Pachyphid peculiarities in comparison to extant and extinct forms. C.R. Palevol (2019), https://doi.org/10.1016/j.crpv.2019.05.004
Fig. 5. *Nigerophis mirus*. Krebb de Sessao, Niger. Palaeocene. A, B. Vertebra USTL SES 106; A, longitudinal section (LS) of the centrum in polarized light (PL); arrows indicate the limit between periosteal and endochondral territories; B, half transverse section showing the absence of remodelling. Scale bars: 500 \( \mu \text{m} \). C, D. vertebra USTL SES 107; C, longitudinal and half transverse sections in natural light (NL); scale bars: 1 mm; D, remains of calcified cartilage (indicated by arrows) inside the trabeculae of endochondral origin in LS; scale bar: 500 \( \mu \text{m} \).

Fig. 5. *Nigerophis mirus*. Krebb de Sessao, Niger. Paléocène. A, B. Vertèbre USTL SES 106 ; A, coupe longitudinale (LS) du centrum en lumière polarisée (PL) ; les flèches indiquent la limite entre les territoires périostiques et endochondraux ; B, demi-coupe transversale illustrant l’absence de remaniement. Barres d’échelle : 500 \( \mu \text{m} \). C, D. Vertèbre USTL SES 107 ; C, coupes longitudinale et demi-transversale en lumière naturelle (NL) ; échelle: 1 mm ; D, restes de cartilage calcifié (pointé par des flèches) à l’intérieur des travées d’origine endochondrale en LS ; échelle : 500 \( \mu \text{m} \).

Fig. 6. *Indophis sahni*. Naskal, Andhra Pradesh, India. Maastrichtian. A. VPL/JU Unnumb. 3. B. VPL/JU Unnumb. 4. C. VPL/JU Unnumb. 1; A–B, longitudinal sections in NL; C, transverse section in PL. The scale bars equal 300 \( \mu \text{m} \).

Fig. 6. *Indophis sahni*. Naskal, Andhra Pradesh, Inde. Maastrichtien. A. VPL/JU sans numéro. 3. B. VPL/JU sans numéro. 4. C. VPL/JU sans numéro. 1 ; A–B, coupes longitudinales en NL ; C, coupe transversale en PL. Barres d’échelle : 300 \( \mu \text{m} \).

Please cite this article in press as: Houssaye, A., et al., Adaptation of the vertebral inner structure to an aquatic life in snakes: Pachyphid peculiarities in comparison to extant and extinct forms. C.R. Palevol (2019), https://doi.org/10.1016/j.crpv.2019.05.004
4.2. Comparisons with extant taxa

Many extant aquatic snakes display a high inner compactness. Average compactness values for the extant semi-aquatic and aquatic forms are 72.9% (Cls) and 86.9% (Cts). There are no striking differences between occasionally aquatic and semi-aquatic species on the one hand (mean Cls = 72.1%; mean Cts = 84.1%) and predominantly aquatic ones (mean Cls = 73.9%; mean Cts = 90.7%), though the difference in transverse section, made at the neutral point, indicates a higher thickness of the compact cortical bone in the latter. Calculations based on data from Houssaye et al. (2013a) indicate lower values for terrestrial (generalist and arboreal) taxa (mean Cls = 71.3%; mean Cts = 76.5%), whereas fossorial species show high compactness values (mean Cls = 74.9%; mean Cts = 86.0%), close to those observed in aquatic snakes (Fig. 13). There is no significant difference in the compactness in longitudinal section for these four ecological categories (semi-aquatic, aquatic, terrestrial and arboreal, fossorial; \(P=0.60\) for a Welch’s \(t\)-test), but it is significant in transverse section (\(P<0.001\)).

Comparisons of the compactness values obtained for the extinct and extant taxa from our sample and Simolophis...
and *Palaeophis maghrebianus* sections (see Table 1) reveal that all fossil specimens sampled exhibit compactness indices in the range of those observed in the extant aquatic snakes (Fig. 14). Moreover, no clear distinction is observed between semi-aquatic and almost exclusively aquatic forms (Fig. 13). Palaeophiids display a wide range of compactness values with some particularly light forms and the *Palaeophis maghrebianus* specimens exhibiting rather high compactness values. *Nigerophis*, *Russelophis* and *Simoliophis* exhibit high compactness values, but they are not distinct from some extant snakes (e.g., *Enhydris*, *Cerberus rynchos*, *Homalopsis buccata*). What is of particular interest is that these snakes display various ecologies, from mangrove mud flats for *Acrochordus granulatus*, *Cerberus rynchos*, and *Bitia hydroides*, to a purely marine lifestyle, for *Hydphis peronii*, *H. gracilis*, *H. elegans*, *H. schistosus*, *H. stokesii*, and *H. curtus*, through freshwater environments for *Enhydris*, *Erpeton tenteculatum*, and *Homalopsis buccata*.

5. Discussion

5.1. Histological and microanatomical features of the extinct taxa

*Nigerophis*. Vertebrae are characterized by the inhibition of both periosteal bone and calcified cartilage resorption, which evokes osteosclerosis. Whereas vertebrae USTL SES 105-106 are considered mid-trunk vertebrae, USTL SES 107 could correspond to a more anterior trunk vertebra (JCR; pers. obs.) The variation in compactness observed between
these vertebrae is consistent with a maximal osteosclerosis intensity in the mid-trunk region. Buffrénil and Rage (1993) described the inner organization of specimens of both Nisterophis mirus and Palaeophis colossaeus as a honeycomb weave structure characterized by an intense remodelling activity. As for N. mirus, this result is not in accordance with the sections analysed here. No illustration of these sections is available in the literature; however, two longitudinal sections assigned to this taxon were loaned by V. de Buffrénil. The microanatomy is relatively comparable to what is observed in vertebra USTL SES 107. Unfortunately, no cast or illustration of the original vertebrae is available, so that no inference about the original position of these vertebrae along the vertebral column can be made.

Indophis. Osteosclerosis has been observed in most sections analysed, although VPL/JU Unnumb. 4 displays a pattern similar to that of most extant squamates. Osteosclerosis in this taxon is probably restricted to a peculiar region of the vertebral column that cannot be determined based on this sample.

Palaeophiidae. None of the Palaeophis colossaeus vertebrae displays osteosclerosis. Their inner structure is rather spongious. The various patterns observed show an important intraspecific variation in vertebral microanatomy. Like

---

*Fig. 9. Palaeophis toliapicus.* Egem (Belgium). Ypresian. MNHN CBL 3-6. A, C. Longitudinal virtual section of the centrum. B, D. Transverse virtual section. The scale bars equal 1 mm.

*Fig. 9. Palaeophis toliapicus.* Egem (Belgique). Yprésien. MNHN CBL 3-6. A, C. Coupe longitudinale virtuelle du centrum. B, D. Coupe transversale virtuelle. Les barres d’échelle représentent 1 mm.
for P. colossaeus, no vertebra of Palaeophis typhaeus displays osteosclerosis, though some vertebrae show compact neural arches and spines, whereas others do not, reflecting a rather high intraspecific variability. These vertebrae, however, all display a large central cavity in the core of the centrum. The P. toliapicus and Palaeophis sp. inner structure appears very similar to that of P. typhaeus. This is also the case for Pterosphenus. The endochondral territory in the Palaeophis vertebrae consists in a rather loose spongiosa, which becomes tighter in some (but not all) of the largest vertebrae. Within Palaeophiidae, important intraspecific variation in microanatomical organization is observed, which probably reflects important intracolumnar and size variation in the vertebral microanatomy of these taxa. Further analyses on vertebrae of various sizes illustrating diverse clearly defined positions along the vertebral column would be required to explain this variability.

Most Palaeophis vertebrae, except those of P. colossaeus, show an open cavity in the core of the centrum, even
the vertebrae of *P. maghrebianus*. This feature often occurs in snake vertebrae and even more widely in squamates (Houssaye et al., 2010, 2013a), but is not at all general to amniotes (Dumont et al., 2013; Houssaye et al., 2014).

*Russelophis tenuis*. The two vertebrae appear osteosclerotic, with only a few large cavities in the endochondral territory.

For the extinct taxa for which several specimens are available, some display rather distinct compactness values, which highlights that intraspecific variability can be high as compared to interspecific variability.

5.2. The occurrence of osteosclerosis

The coupled inhibitions of both calcified cartilage resorption and primary periosteal bone resorption are clearly observed in two specimens of *Nigerophis*, but only to a lesser extent in USTL SES 107, which shows a wide cavity in the core of the centrum and a compactness not particularly high. Osteosclerosis also occurs in most vertebrae of *Indophis*, with a clear inhibition of primary bone resorption. In *Russelophis*, the observation of growth marks enables to assume a similar inhibition of primary bone resorption. However, the resolution of the virtual sections does not
Fig. 14. Graphs illustrating the compactness values for our sample with color indications for various ecologies and the family of the extinct taxa. A. Cts values. B. Cts values. C. Their covariation. Species name abbreviations are listed in Tables 1 and 2.
allow us to conclude on the amount and extent of calcified cartilage remains. Osteosclerosis has been described in the anterior and mid-precloacal regions in *Palaeophis maghrebianus*, though compactness is not extreme, based on the combination of an inhibition of primary bone (but not calcified cartilage) resorption with excessive secondary bone deposits during remodelling (Houssaye et al., 2013b). The other *Palaeophis* vertebrae show compactness values that do not suggest osteosclerosis. Different processes relative to cartilage and bone resorption/remodelling are thus encountered within these aquatic snakes. If primary bone resorption is limited in all compact vertebrae, the inhibition of calcified cartilage resorption and excessive secondary bone deposits during remodelling occur only in some taxa.

Snake vertebrae (whatever the ecology) are rather compact, as compared to those of other amniotes. The small sample of non-aquatic and non-flying amniotes from Houssaye et al. (2014) shows average *Cls* and *Cts* values of 43.4 and 45.9%, respectively, as compared to the 71.3 and 76.5% values obtained for terrestrial and arboreal squamates, excluding fossorial taxa that would increase these numbers. It has been emphasized that snakes generally display no clear microanatomical specialization in their vertebrae (neither in their ribs; Canoville et al., 2016), except a few ecologically highly specialized taxa (Houssaye et al., 2013a). Their generalist inner morphology was interpreted as resulting from their general use of different habitats and locomotor modes and the necessity for snakes to move efficiently in different environments. Bone mass increase could nevertheless be expected in aquatic taxa with extremely limited terrestrial locomotion and utilising a single environment. Houssaye et al. (2013a) indeed suggested the occurrence of osteosclerosis in *Erpeton tentaculatum*, which usually remains suspended or anchored to vegetation while hunting in rather shallow water (Smith et al., 2002) and, to a lesser extent, in *Enhydris bocourtii*. Conversely to these taxa confined to a single milieu, *Laticauda* moves in both deep water and across land, which could explain the absence of bone mass increase in this taxon. As for surface swimmers not requiring buoyancy control, like *Hydrophis platurus*, the absence of bone mass increase was expected. The addition of numerous semi-aquatic and marine taxa to the sample from Houssaye et al. (2013a) enables a wider comparative analysis. Our results show no strong difference in snake vertebral compactness between an aquatic lifestyle and the other ecologies. It also highlights a high variation in compactness for both semi-aquatic and aquatic snakes with no distinction between the two habitat preferences.

Therefore, no good correlation is highlighted between the compactness values and the ecology in snake vertebrae. It is not possible to assume an aquatic lifestyle based on high compactness, as for other amniotes (e.g., Amson et al., 2015; Buffrénil et al., 2010; Hayashi et al., 2013; Houssaye et al., 2016). Among aquatic forms, which occupy a great diversity of habitats, compactness also does not enable to distinguish the various ecologies. This comparative analysis thus shows that it is not possible to make reliable palaeoecological inferences for snakes based on vertebral microanatomy, although this tool appears very efficient for other amniotes and/or other bones (e.g., Dumont et al., 2013; Houssaye and Botton-Divet, 2018; Laurin et al., 2011). This result is all the more surprising because snakes do not have limbs and the axial skeleton is thus more strongly involved in locomotion as compared to limbed-amniotes. However, as previously suggested, snakes seem to display a rather generalist vertebral inner structure (Buffrénil and Rage, 1993; Houssaye et al., 2013a). Osteosclerosis needs to be determined based on the relative condition in “non-affected” sister taxa. Compactness being rather high in numerous snakes, this prevents the use of the term osteosclerosis in the most compact bones. The peculiarity of snakes strongly raises the question of the characterization of osteosclerosis: should it be based on a compactness threshold or on a mechanism? Our comparative analysis highlights the extreme difficulty in finding a threshold value. As for the mechanism, the few taxa for which classical sections were available show different processes of bone mass increase, with inhibition or enhancement of different processes. The use of the term “osteosclerosis” appears thus extremely complex and ambiguous for snakes.

5.3. The peculiarity of the marine hind-limbed snakes

One group remains peculiar among snakes: the marine hind-limbed snakes. These Cenomanian snakes are the only snakes displaying pachyostosis. It occurs in all trunk vertebrae and ribs except the anteriormost and posteriormost ones, with a maximal intensity in the mid–trunk region (see Houssaye, 2013 for a review on this question). This osseous specialization is intense in these taxa, though less strong in smaller specimens within each species (as observed in *Eupodophis*, *Pachyrhachis*, and *Pachyophis*; Houssaye, 2013).

Bone mass increase is considered to enable an increase in oxygen store and to control buoyancy and body trim (Taylor, 2000). Though more intense in the mid-trunk region, bone mass increase seems to affect almost the entire vertebral column in pachyophids (Houssaye et al., 2011), which suggests a limited, if present, role in body trim control. In marine snakes, oxygen store is likely not a problem, since they can absorb oxygen through their skin. Other morphological features, e.g., cardiac shuntage, increase in blood volume, the sacular lung, are also suggested to increase immersion duration in snakes (Heatwole, 1999; Ineich, 2004). But cutaneous respiration is considered one major adaptation of sea snakes to a marine lifestyle (Heatwole, 1999) and is limited to these taxa within squamates. There is currently no consensus about the status and phylogenetic position of the pachyophids (Caldwell, 2000; Martill et al., 2015; Palci et al., 2013b; Reeder et al., 2015; Rieppel and Zaher, 2000). If pachyophids are among the first snakes, they might not show the morphological specializations observed in other sea snakes, notably cutaneous respiration, which might explain the need for strong bone mass increase in order to increase oxygen store. Compactness in most snakes is often rather high. The occurrence of pachyostosis might be a mean to further increase bone mass. The combination of osteosclerosis and pachyostosis (pachyosteosclerosis) is similarly observed in the precloacal skeleton of some “dolichosaurs” or stem-oophidians that
display a morphology rather similar to that of hind-limbed snakes (elongated skeleton with a high number of vertebrae and a reduction of the limbs) and are supposed to have lived in similar environments as the pachyphoids, i.e. coastal shallow waters (Houssaye, 2013). Pachyostosis is assumed to reduce movements between adjacent vertebrae and might have caused a reduction in lateral undulation in these taxa, whose aquatic locomotion might have thus slightly differed from that of other snakes.

6. Conclusion

The description of the inner structure of vertebrae of various extinct aquatic snakes and the interpretation of these microanatomical data in the light of a large comparative sample of extant snakes shows:

• important intraspecific variation in the vertebral microstructure in some snake taxa;
• that the vertebral microstructure cannot be used as a reliable ecological proxy in snakes, even for specialized species;
• that the use of the term “osteoclerosis” in snakes is extremely complex;
• the peculiarity of marine hind-limbed snakes that are the sole snakes showing pachyostosis.

Acknowledgments

We would like to thank M. Lemoine and P. Loubry (MNHN, Paris, France) for the making of the thin sections and vertebra pictures, respectively. We are thankful to G.V.R. Prasad (University of Delhi, India) and B. Marandet (ISEM, Montpellier, France) for the loan of the Indophis and Nigersophis vertebrae, respectively, and V. de Buffrénil (MNHN, Paris, France) for the loan of the P. colossaeus sections. We thank K. Lim (Lee Kong Chian Natural History Museum, Singapore) for loaning us extant material for scanning. We thank the Steinmann Institute (University of Bonn, Germany), R. Lebrun (ISEM, Montpellier, France) and the MRI platform member of the national infrastructure France-Biobraining (supported by the ANR-10-INBS-04, “Investments for the future”, the labex CEMEB [ANR-10-LABX-0004] and NUMEV [ANR-10-LABX-0020]), the “Centre de microtomographie” of the Université de Poitiers, P. Tafforeau and the beamline ID19 of the ESRF (Grenoble, France), for providing access, beamtime, and support. This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors. This manuscript is naturally dedicated to the memory of Jean-Claude who supervised part of this work during the Ph.D. thesis of A. Ho. To the memory of a wonderful and inspiring scientist!

References

Abramoff, M.D., Magelhaes, P.J., Ram, S.J., 2004. Image processing with ImageJ. Biophoton Int. 11, 36–42.

Please cite this article in press as: Houssaye, A., et al., Adaptation of the vertebral inner structure to an aquatic life in snakes: Pachyphoid peculiarities in comparison to extinct and extant forms. C.R. Palevol (2019), https://doi.org/10.1016/j.crpv.2019.05.004


Welch, B.L., 1947. The generalization of ‘Student’s’ problem when several different population variances are involved. Biometrika 34, 28–35.