

REVIEW

Loss of teeth and enamel in tetrapods: fossil record, genetic data and morphological adaptations

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Abstract

Since their recruitment in the oral cavity, approximately 450 million years ago, teeth have been subjected to strong selective constraints due to the crucial role that they play in species survival. It is therefore quite surprising that the ability to develop functional teeth has subsequently been lost several times, independently, in various lineages. In this review, we concentrate our attention on tetrapods, the only vertebrate lineage in which several clades lack functional teeth from birth to adulthood. Indeed, in other lineages, teeth can be absent in adults but be functionally present in larvae and juveniles, can be absent in the oral cavity but exist in the pharyngeal region, or can develop on the upper jaw but be absent on the lower jaw. Here, we analyse the current data on toothless (edentate) tetrapod taxa, including information available on enamel-less species. Firstly, we provide an analysis of the dispersed and fragmentary morphological data published on the various living taxa concerned (and their extinct relatives) with the aim of tracing the origin of tooth or enamel loss, i.e. toads in Lissamphibia, turtles and birds in Sauropsida, and baleen whales, pangolins, anteaters, sloths, armadillos and aardvark in Mammalia. Secondly, we present current hypotheses on the genetic basis of tooth loss in the chicken and thirdly, we try to answer the question of how these taxa have survived tooth loss given the crucial importance of this tool. The loss of teeth (or only enamel) in all of these taxa was not lethal because it was always preceded in evolution by the pre-adaptation of a secondary tool (beak, baleens, elongated adhesive tongues or hypselodonty) useful for improving efficiency in food uptake. The positive selection of such secondary tools would have led to relaxed functional constraints on teeth and would have later compensated for the loss of teeth. These hypotheses raise numerous questions that will hopefully be answered in the near future.

Key words birds; enamel; evolution; mammals; toads; tooth loss; turtles.

Introduction

Teeth originated in stem gnathostomes, approximately 450 million years ago (Ma) (Reif, 1982; Donoghue, 2002; Donoghue & Sansom, 2002; Donoghue et al. 2006). Two theories have been proposed for the evolution of teeth. Firstly, that teeth derived from skin denticles/odontodes that moved from the outside of the mouth in and, alternatively, that teeth derived from pharyngeal denticle whorls and moved up into the mouth (Smith, 2003; see review by Huyseune et al. 2009). Either way the newly acquired oral structures played an important role in concert with jaws, allowing for a predatory lifestyle, and their development

would have been selectively constrained. During the further evolution of gnathostomes, these original 'teeth' diversified, changing their location, shape and ornamentation, size, mode of attachment and number of generations, such diversification permitting adaptation to a variety of diets (Huyseune & Sire, 1998). Although exhibiting morphological differences in various vertebrate taxa, teeth have mostly conserved their original structure, i.e. a pulp cavity surrounded by a dentine crown generally covered by enamel or enameloid, two hard protective tissues. Such a long-lasting conservation of tooth organization and structure is related to strong selective constraints that result from the important role played by these organs. This also explains why teeth are still present in all living gnathostome lineages [chondrichthyans (cartilaginous fish), actinopterygians (ray-finned fish) and sarcopterygians (coelacanth, lungfish and tetrapods: lissamphibians, reptiles and mammals)]. However, although teeth and their protective tissues (enamel/enameloid) seem to be crucial

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tools for animal survival, they have been lost in several vertebrate taxa. The three lineages of tetrapods mentioned above include several toothless and enamel-less taxa. This review focuses on these taxa and provides data on their evolution and adaptation. We have considered only the tetrapods because, to our knowledge, there are no examples of edentate species in chondrichthyans, coelacanths, lungfish and actinopterygian fish. In the latter lineage, however, we know of numerous species that either lack teeth in the oral cavity but retain them in the pharyngeal region only (e.g. all cypriniforms, such as carp, roach and zebrafish) or possess functional teeth in larvae and juveniles that are shed and not replaced in further stages (e.g. sturgeons, armoured catfish) (Sire et al. 2002; see also Huysseune et al. 2009). We will not describe tetrapod species that have lost teeth on one jaw but retain them in the other, such as some odontocetes (e.g. sperm whales) and most anuran amphibians (e.g. clawed toads). In mammals, however, we will comment on the platypus (monotremes) as this species possesses teeth in the embryos and juveniles but loses them in the adult.

The location of all living edentate taxa in the tetrapod phylogeny clearly indicates that, during evolution, (i) the ability to form teeth was lost independently in seven

lineages and (ii) enamel disappeared in two unrelated lineages (Fig. 1). This suggests that teeth and/or enamel could be 'easily' lost; however, we will see that species survival after tooth or enamel loss strongly depends on the recruitment of various pre-existing morphological adaptations that were secondarily retained as more efficient mechanisms for food processing.

In humans, most cases of tooth loss (agenesis) are related to syndromes that result from mutations in various genes (e.g. ectodysplasia, Mikkola & Thesleff, 2003; see Caton & Tucker, 2009; De Coster et al. 2009). Non-syndromic tooth loss in humans has so far been linked to three genes (*Pax9*, *Msx1* and *Axin2*) that lead to hypodontia (loss of a few teeth) and oligodontia (loss of at least six teeth) (Matalova et al. 2008). To our knowledge, there are no reports in the literature on completely edentate patients in which no other phenotype was identified and which are not related to a syndrome.

Here, we review the current knowledge of edentate and enamel-less tetrapods, and we present scenarios that could explain how the ability to form teeth was lost and why these taxa survived tooth loss. These topics are covered in four sections. The first section is devoted to tooth loss in chicken, as this is the only edentate species in

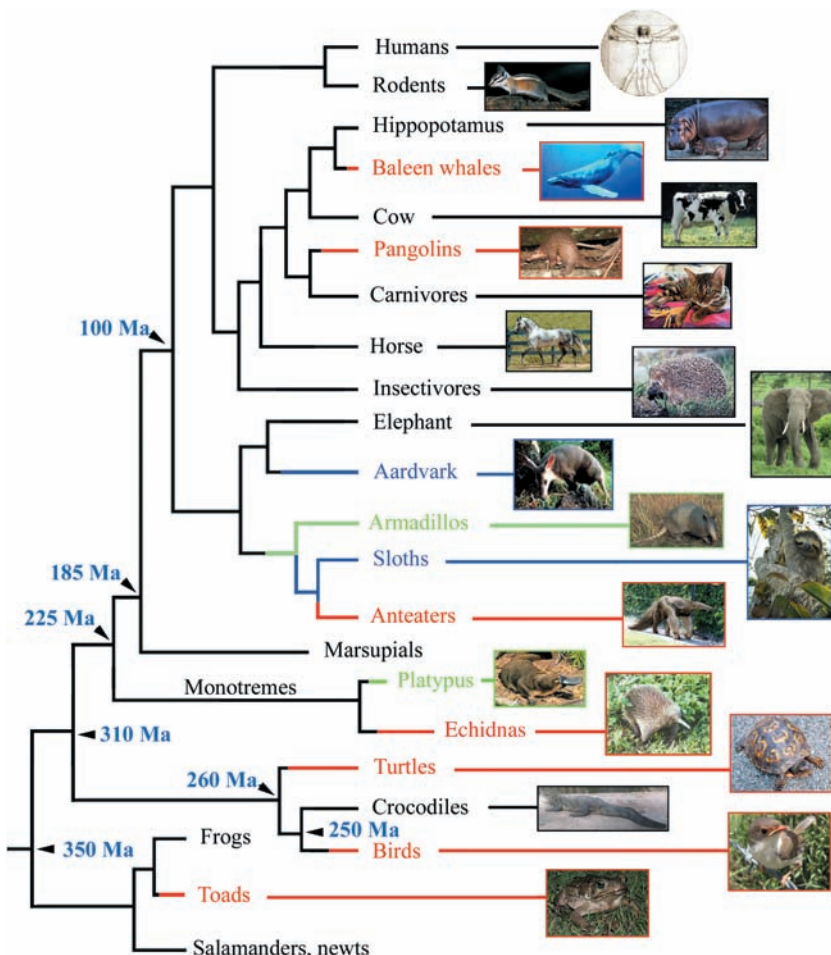


Fig. 1 Simplified tetrapodan phylogeny with indication of toothless lineages (red lines), enamel-less lineages (blue lines) and lineages with enamel reduction and tooth reduction (green lines). Tetrapodan relationships after Murphy et al. (2001) and Hedges (2002).

which developmental mechanisms underlying the loss of teeth have been studied. The second section is a review of the current knowledge of tooth loss in the other edentate lineages. For each lineage, data are compiled on tooth loss during evolution and ontogeny. The situation in the adult is presented and the specialized morphological adaptations that were selected during evolution as improving food uptake when teeth were lost are pointed out. The third section concerns enamel loss in two mammalian lineages [Xenarthra (sloths and armadillos) and Tubulidentata (Afrotheria: aardvark)] with particular attention to the nine-banded armadillo. The fourth section provides information on the platypus, a monotreme species in which teeth are lacking in adults but are present in embryos and juveniles. When available, we have included palaeontological data that can support a tentative datation of tooth loss or enamel loss in these taxa and the relationships of these taxa. These findings are then briefly discussed in the conclusion.

Tooth loss in birds and the famous hen's teeth

Fossil and living birds constitute the Avialae (*sensu* Gauthier, 1986) (Fig. 2). The living species (around 10 000 species in 26 orders) are divided into the Paleognathae and the Neognathae (Sibley & Ahlquist, 1990). Both form the clade Neornithes. All of them are toothless.

What the fossil record tells us

It is generally believed that modern birds (Neornithes) derive from theropod dinosaurs (e.g. Ostrom, 1976),

although a close relationship between birds and crocodylians has also been proposed on the basis, amongst other characters, of similarities of tooth ornamentation (Whetstone & Martin, 1979; Martin et al. 1980). The last common ancestor of Avialae was equipped with clearly functional teeth. The oldest widely accepted Avialae is the well-known *Archaeopteryx lithographica*, which lived some 150 Ma and possessed teeth. More than 70 avialan genera are known from the Mesozoic (140–120 Ma; Chiappe & Dyke, 2002), which allows a phylogenetic tree to be drawn based on anatomical characters (Fig. 2). The most recent toothed Avialae in fossil records, the ornithurine birds *Hesperornis regalis* and *Ichthyornis dispar*, are known from the late Cretaceous (93–65 Ma; Marsh, 1872; Gregory, 1952). *H. regalis* was a swimming bird. It had a long beak with a rhamphotheca covering the pre-maxilla region only and was provided with small but effective conical teeth set firmly in the jaw (Gingerich, 1975). The teeth of the upper jaws were few in number and set in the back part, whereas those of the mandibles formed a complete series (Fig. 2). *H. regalis* therefore had half a beak and teeth. To date, *I. dispar* is the closest Avialae to the common ancestor of Neornithes (Clarke, 2004). The teeth of *I. dispar* are set in a groove as in *H. regalis*, and they are broad and flattened with highly expanded roots (Martin & Stewart, 1977). *I. dispar* was a powerful flying bird and did not differ notably from the common flying birds of the present time (e.g. terns). Most molecular dates for the divergence of Neornithes imply that they existed 40 Ma (Van Tuinen & Dyke, 2004) prior to the oldest identified ornithurine fossils in the late Cretaceous (Campanian, 80 Ma; Fountaine et al. 2005). Therefore, this would place the origins of

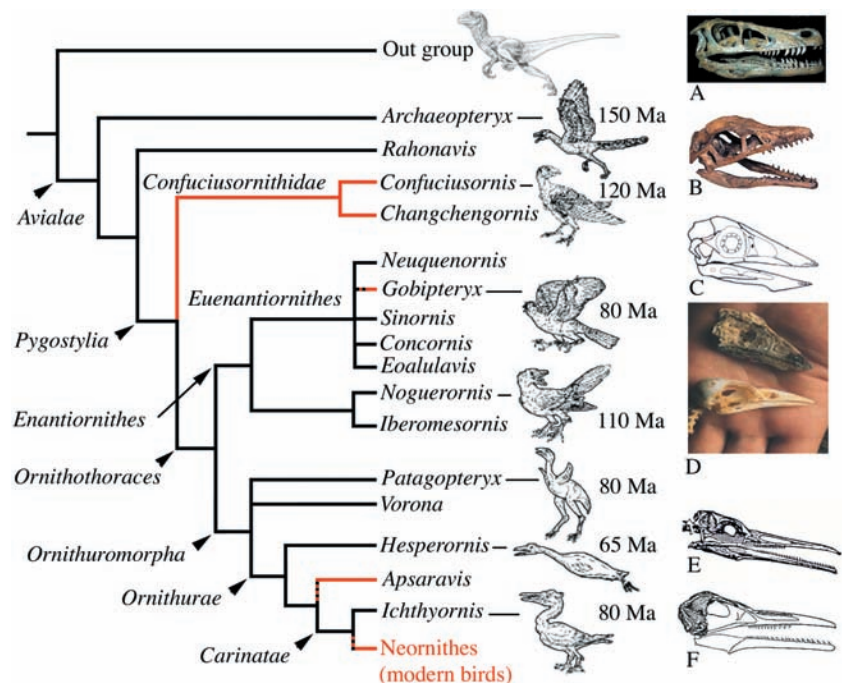


Fig. 2 Simplified phylogeny of Avialae (after Chiappe, 2002). Lineages with toothless taxa are in red. (A) Skull of *Velociraptor*. (B) Skull of *Archaeopteryx lithographica*. (C) Skull of *Confuciusornis sanctus*. (D) Comparison of the skulls of *C. sanctus* and a pigeon (bottom). (E) Skull of *Hesperornis regalis* (after Marsh, 1880). (F) *Ichthyornis dispar* (after Marsh, 1883).

modern birds in the early Cretaceous (120 Ma; Smith & Peterson, 2002). This later date is still earlier than that estimated from the fossil records but establishing accurate calibration times for molecular phylogenies on the basis of fossil data is a difficult task. However, the recent discovery of a close relative of ducks (Anseriformes) in the Maastrichtian of Antarctica (70 Ma; Clarke et al. 2005) indicates that Neornithes originated long before the Cretaceous/Tertiary boundary, probably earlier than 80 Ma, even if they diversified later, during the early Cenozoic (65 Ma; Zhou, 2004).

We do not know whether the fossil taxa closer to the most recent common ancestor of Neornithes than *I. dispar* have teeth. Therefore, we can reasonably estimate that tooth loss in crown Aves arose maximally on the stem lineage between *I. dispar* and Neornithes, and minimally in the most recent common ancestor of Neornithes, the origin of modern birds, i.e. approximately 100 Ma.

What the developmental genetics tells us

It has been known for more than a century and a half that transient epithelium thickenings, homologous to the dental lamina stage in the mouse, appear in the chick oral cavity at embryonic day (E)5 (Geoffroy Saint-Hilaire, 1820; Blanchard, 1860; Gardiner, 1884; Röse, 1892; Carlsson, 1896). This could be interpreted as the 'remnants' of an ancestral toothed condition. In birds, early tooth development arrest could be the consequence of mutations that inactivated the genetic pathways leading to tooth formation. In the last few decades, taking advantage of recent technical advances, several scientists have attempted to stimulate the 'dormant' odontogenic pathway in chicken with the ultimate goal of resuscitating teeth. This dream seemed attainable through elegant experiments involving either mouse/chick tissue recombinations aiming to reinitialize epithelial/mesenchymal interactions or using beads impregnated with various signalling molecules to mimic as closely as possible such interactions. The results of these experiments are summarized below and schematically analysed in Fig. 3.

Kollar & Fisher (1980) performed a simple experiment in which they recombined dental epithelium of E5 chick embryos with molar mesenchyme of E16–E18 mouse embryos. These recombinant tissues were cultivated within the anterior chamber of a mouse eye. Several days later they obtained teeth with a dentine cone and an enamel cover, the famous 'hen's teeth'. This meant that the cells of the E5 chick dental epithelium not only had retained the genetic potential to respond to the induction from the mouse mesenchymal cells for more than 100 Ma (tooth loss in a Neornithe ancestor) but they were also able to develop until the last developmental step (enamel matrix deposition). However, a possible contamination of mouse mesenchyme by mouse epithelium makes such

interpretation uncertain. Indeed, it is difficult to completely eliminate contamination of mouse mesenchyme with residual epithelium and such a contamination would allow tooth formation (Arechaga et al. 1983). Eighteen years later, in another series of recombination experiments, Kollar's group showed that chick epithelium was able to induce cell proliferation and the expression of key developmental genes (*Msx1*, *Msx2* and *Bmp4*) in the mouse mesenchyme, leading to odontoblast differentiation and formation of tooth germs (Wang et al. 1998) (Fig. 3D).

Another important finding was obtained by Chen et al. (2000) who showed that the early odontogenic pathway remained inducible in chick mandibles. During mandible development, they analysed the expression of crucial genes known to regulate mouse tooth morphogenesis. They discovered that the main markers for dental lamina formation in the mouse are expressed in the developing chick mandible (*Fgf8*, *Pitx2*, *Pax9*, *Barx1*, *Msx1* and *Msx2*). However, the expression of three key genes (*Bmp4*, *Msx1* and *Msx2*) was missing in the distal mandibular mesenchyme facing the presumptive chick dental lamina (Fig. 3B). When beads impregnated with *Bmp4* and *Fgf4* were implanted into the epithelium, facing cells in the distal mandibular mesenchyme expressed *Msx1* and *Msx2* (Fig. 3C). These experiments suggest not only that the odontogenic signalling pathway is conserved in the chick and can be reactivated but also that a defect in *Bmp4* signalling could be responsible for the lack of *Msx1* and *Msx2* expression. A defect in *Bmp4* signalling may therefore have occurred in a Neornithe ancestor, leading to a premature arrest of tooth development. However, exogenous addition of *Bmp4*, even in the presence of *Fgfs*, did not allow tooth development in the chick to proceed beyond the cap stage, i.e. to the stages of tooth germ differentiation. Structures that could be defined as teeth were not formed (Chen et al. 2000).

These findings obtained *in vitro* were confirmed *in vivo* by Mitsiadis et al. (2003) who performed transplantations of mouse (E8) neural crest cells in 1-day-old chick embryos. They showed that the avian dental epithelium was still able to induce a non-avian developmental programme in mouse neural crest-derived mesenchyme, resulting in tooth germ formation. Here again they did not obtain functional teeth but tooth germs developed until an advanced stage of differentiation in which some dentine-like matrix was deposited by the ectomesenchymal cells (odontoblast-like cells) and started to mineralize (Fig. 3D). No enamel-like structure was observed, however. This *in-vivo* study also indicates that endogenous factors coming from the mouse ectomesenchyme are more efficient at supporting tooth formation beyond the cap stage than exogenous factors used in the previous study (Chen et al. 2000). Unfortunately, the duration of these experiments was too short to determine whether or not tooth differentiation would have eventually reached a more advanced

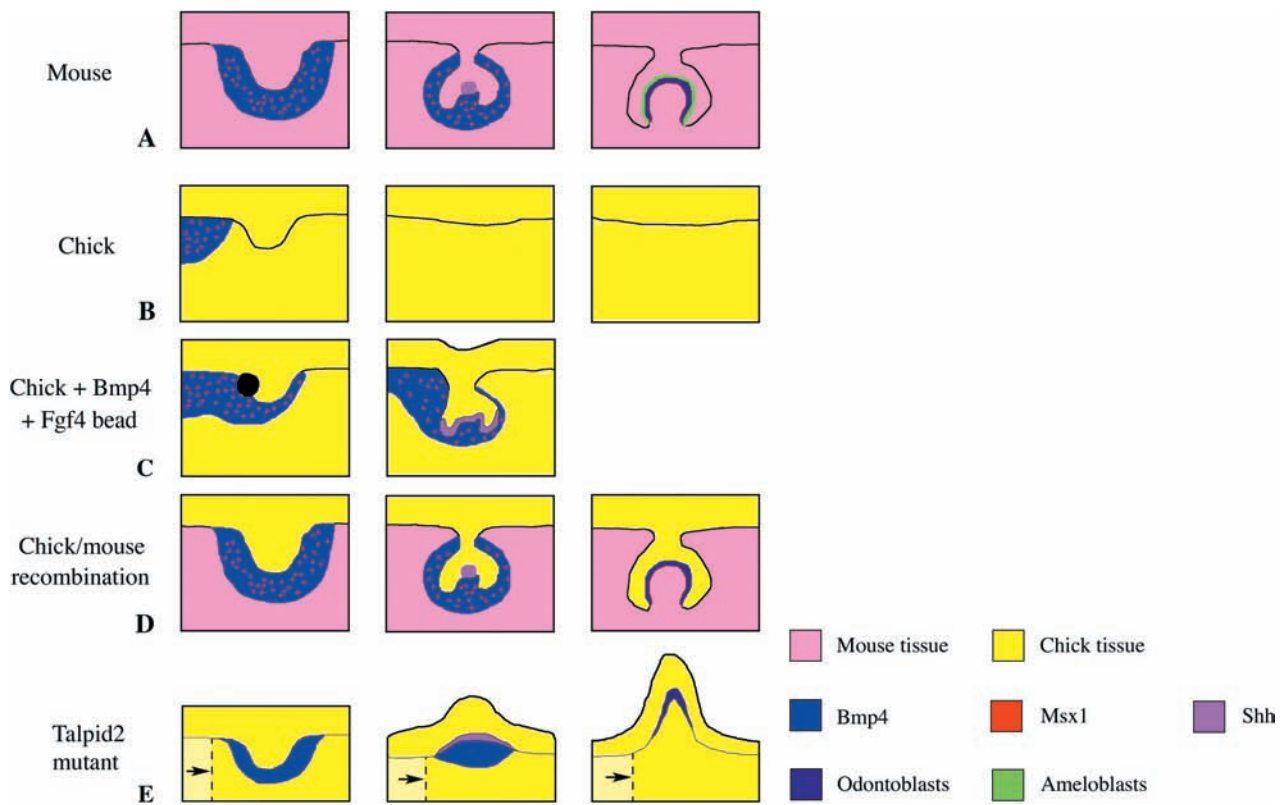


Fig. 3 A shift in the positioning of the odontogenic epithelium relative to the dental competent mesenchyme could explain the loss of the ability to form teeth in the modern bird ancestor. Schematic drawings summarizing the chick tooth experiments. (A) Mouse molar developmental stages, from bud [embryonic day (E)12.5] to cap (E14.5) to bell (E16.5). The condensing mesenchyme around the bud stage tooth germ expresses *Bmp4* and *Msx1* and induces development of the enamel knot at the cap stage, which expresses signalling molecules such as *Shh*. The inner enamel epithelium forms the ameloblasts that form enamel, whereas the adjacent mesenchyme forms the odontoblasts that form dentine (see Caton & Tucker, 2009). (B) Chick development. At Hamburger & Hamilton (HH) stage 28 a bud-like thickening of the oral epithelium is observed. Expression of *Bmp4* and *Msx1* is not, however, associated with this region. No further tooth development is observed at later stages and the thickening regresses. Note that, at an earlier stage (stage 24), *Bmp4* expression is epithelial and shifts into the mesenchyme at stage 28 (Francis-West et al. 1994). (C) When a bead impregnated with *Bmp4* and *Fgf4* is implanted into the chick epithelium, the expression of *Bmp4* and *Msx1* in the mesenchyme extends around the developing tooth bud. This leads to the extension and folding of the bud epithelium, and induction of *Shh*. No further progression of the tooth germs is observed, however (Chen et al. 2000). (D) When mouse mesenchyme is combined with chick epithelium (either by recombination of mandible tissue or by earlier neural crest grafts of mouse neural crest into a chick embryo), the chick epithelium induces *Msx1* and *Bmp4* in the mouse mesenchyme. The tooth germ progresses to the cap stage and forms an enamel knot-like structure expressing *Shh*. The mouse tissue differentiates into odontoblasts and forms a bell stage tooth germ. Tooth differentiation does not proceed beyond this stage and enamel is not deposited (Wang et al. 1998; Mitsiadis et al. 2003). (E) In the chick mutant *talpid2* a shift in the positioning of the epithelium and mesenchyme has been described (indicated by dashed lines and arrows). The chick epithelium is able to induce expression of *Bmp4* in the underlying mesenchyme and expresses *Shh*. The tooth germ develops by evagination, similar to that observed in alligator embryos. At later stages differentiated odontoblasts are identified by histology but no further differentiation occurs (Harris et al. 2006).

stage and the problem remains as to what extent tooth programmes are maintained in the chicken.

Recent observations made in a mutant chick [*talpid2* (*ta²*) affected gene unknown], in which the development of several organ systems is affected, have brought additional light to the investigation on the mechanisms underlying tooth loss in birds. Indeed, the *ta²* mutant was shown to develop rudimentary teeth under the rhamphotheca (Harris et al. 2006). These outgrowths from the distal mandible were conical and caniniform-shaped, morphological features that were similar to the so-called rudimentary teeth that develop in crocodile embryos. In the latter,

these teeth do not develop further than the stage of dentine deposition and degenerate without any enamel covering being deposited. In contrast to the *ta²* mutant, in crocodylians and lepidosaurians (lizards and snakes) when the rudimentary teeth are degenerating, first-generation teeth start to form, which are functional at hatching (and covered with enamel). Unfortunately, the oldest *ta²* mutant embryos died at E16, several days prior to hatching, and so further developmental stages beyond the early dentine-like deposition were not available. It is therefore not possible to determine whether or not these rudimentary teeth would have degenerated and been replaced, as in

crocodile embryos, by functional teeth. In ta^2 mutants, genes necessary for tooth formation in the mouse are expressed in the mandible, including *Bmp4* (Fig. 3E). The formation of advanced tooth germ in ta^2 mutants is associated with defects in the specification of the oral/aboral boundary. This leads to a developmental repositioning of the presumptive dental epithelium to overlie mesenchyme competent to form teeth. Therefore, the authors proposed that changes in the relative position of a lateral signalling centre over competent odontogenic mesenchyme led to the loss of the ability to form teeth in an ancestral Neornithe.

In summary, these elegant experiments provide strong support for the mechanism of tooth loss in birds being a consequence of a developmental shift in the oral epithelium. This resulted in signalling molecules from the epithelium no longer reaching their targets in the mesenchyme and leading to a failure in the induction of key molecules, such as *Msx1* and *Bmp4*, and an early arrest of tooth development. These experiments also indicate that, under appropriate conditions, the odontogenic capacity of the chicken dental epithelium can be reactivated. However, if the reactivation of such odontogenic pathways is a prerequisite to initiate tooth development and to proceed further until tooth differentiation, it appears insufficient to form true functional teeth, i.e. a dentine cone covered with enamel. At the end of the developmental pathway, genes encoding for structural proteins might be activated but it appears that they were not activated in any of the experiments. In fact, they could not. Indeed, all of the genes encoding dental-specific proteins (dentine sialophosphoprotein for dentine; amelogenin, ameloblastin and enamelin for enamel) have disappeared from the chicken genome after chromosomal rearrangement or are pseudogenes (Sire et al. 2008). The revival of hen's teeth will therefore remain elusive.

Adaptations to tooth loss in birds

The fossil record tells us that the ability to form teeth was lost several times in non-avian theropodan and avian lineages (Sander, 1997). Three unrelated, extinct avians lack teeth: *Confuciusornis sanctus* (early Cretaceous of China; approximately 120 Ma), *Gobipteryx minuta* and *Apsaravis ukhaana* (Campanian; approximately 80 Ma) (Chiappe et al. 1999, 2001; Clarke & Norell, 2002) (Fig. 2). However, the closest Avialae to modern birds is *I. dispar*, a toothed bird. Therefore, the loss of teeth occurred independently at least four times during Avialae evolution, including the lineage leading to modern birds. It is difficult to believe that there were four different mechanisms that led to tooth loss in Avialae and a shift in the mutual position of the competent tissues involved in epithelial/mesenchymal interactions may have been responsible in each case. However, how is it possible to survive the loss of such

important tools? Indeed, we can suppose that tooth loss would have been lethal if these taxa were not already equipped with an alternative tool, the beak.

It is noteworthy that tooth loss in Avialae coincides with the presence of a beak. Indeed, although horny beaks are not preserved in fossils there are indications, for instance in the subjacent bone, of the probable presence of a rhamphotheca. Theropod and early toothed Avialae had no keratinized beak and had teeth on both the maxilla and pre-maxilla. *H. regalis* had a keratinized beak covering the pre-maxilla only, and no teeth in this region (Gingerich, 1975). This is a good example of a morphological intermediate structure between toothed birds, which lack a horny beak, and beaked, toothless birds. *C. sanctus* is the earliest bird known in the fossil record to have a toothless, horny beak, like modern birds (Fig. 2). It was hypothesized that *C. sanctus* fed on plant materials due to its toothless beak (Zhou & Zhang, 2003) but recent findings indicate that its diet comprised fish, like modern sea birds (Dalsätt et al. 2006).

In Avialae the horny beak represented a dramatic innovation resulting from the transformation of the reptilian snout into a beak. The rhamphotheca is assumed to have evolved from reptilian keratinized scales (e.g. Zweers et al. 1997). Interestingly, the BMP pathway was shown to be involved in the fine-tuning of beak morphoregulation in birds (Wu et al. 2004, 2006). It is speculated that this innovation was retained in Avialae because the beak compensated for the dedication of forelimbs to flight, which meant that the limbs were no longer efficient tools for finding (and manipulating) food. With beaks selected as a new tool for food uptake, the strong selective constraints on teeth were relaxed and their loss could have occurred with no drastic consequence for bird survival.

A review of non-avian edentate tetrapodan taxa with a tentative datation of tooth loss in these lineages

A rapid glance at tetrapodan relationships (Fig. 1) indicates that tooth loss occurred independently at least seven times during evolution. Hereafter, we review the current knowledge, unfortunately often limited, of the six non-avian edentate tetrapod taxa, i.e. toads (Lissamphibia), turtles (Sauropsida) and echidnas, anteaters, pangolins and baleen whales (Mammalia). In tetrapods, the edentate condition is not insignificant as it concerns at least 350 toads, 257 turtles, 10 000 birds and 27 mammals, i.e. one third of the currently living tetrapod species (c. 30 000 species).

Toads

Within Anura, the most successful group of living amphibians with 4500 species (Hofrichter, 2000), 'toad' commonly

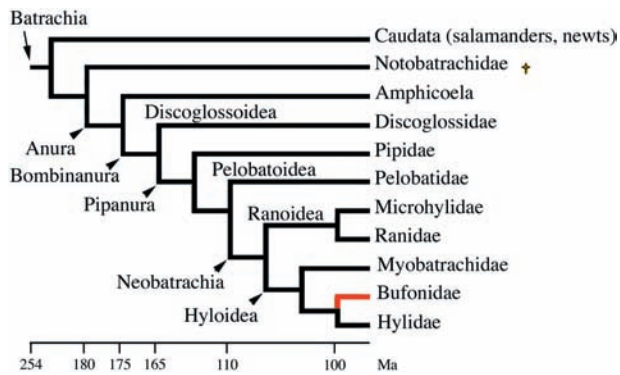


Fig. 4 Simplified phylogeny of Batrachia (after Marjanovic & Laurin, 2007). Red line: edentate toad lineage. It is worth noting that several species in various anuran lineages have also lost the ability to form teeth independently (not shown). †Extinct lineage.

refers to a number of species of unrelated families (Bombinatoridae, Discoglossidae, Pipidae, Pelobatidae, some Microhylidae and Bufonidae) (Fig. 4). In fact, 'true toads' are found only in the family Bufonidae (26 genera, more than 350 species). Not a single bufonid species has teeth, whereas most members of the other 'toad' families possess teeth. Most frogs have an edentulous lower jaw but tiny teeth are attached to the upper jaw (pre-maxillary and maxillary teeth) and on the palate (vomerine teeth) (Cannatella & Hillis, 1993; Cannatella & Graybeal, 1996). Only *Gastrotheca guentheri* is known to possess teeth on the lower jaw. The teeth are homodont (i.e. all teeth similar), bicuspid or monocuspid, and divided into a distal crown and a proximal pedicel, separated by a non-mineralized fibrous ring similar to that described in the teeth of many species of the two other lissamphibian lineages, Caudata and Gymnophiona (reviewed in Davit-Béal et al. 2007).

What the fossil record tells us

The origins of extant amphibians (Lissamphibia) are still largely debated. The fossil record indicates that they probably arose in the Permian (about 260 Ma) (Marjanovic & Laurin, 2007; Anderson et al. 2008) but molecular data infer an older date (Carboniferous, 337 Ma; Zhang et al. 2005). The earliest known Anura and Caudata date back to the Mesozoic (early Jurassic period, 190 Ma; Shubin & Jenkins, 1995) but fossil Bufonidae are only known from Paleogene deposits (70–30 Ma) (Estes & Reig, 1973; Marjanovic & Laurin, 2007). Records of fossil anurans from the early Cretaceous indicate that the lower jaw was edentulous (Gao & Chen, 2004). Within the Hyloidea, Bufonidae are the sister group of a clade comprising Telmatobinae and Hylidae (Fig. 4), which have teeth. This means that the ability to form teeth in the oral cavity was lost in the common ancestor of modern bufonids (100–70 Ma).

Adaptations to tooth loss in toads

Toads are carnivorous, feeding principally on insects, worms, spiders and other invertebrates. The lack of teeth would seem, therefore, an important handicap for these animals but this large range of diet demonstrates that it is not. Indeed, anurans have developed an efficient tool: a protractile tongue covered with a sticky substance, making it a trap for prey. It also produces mucus to help in swallowing. This organ was already present and probably useful (as we can judge by its presence in the current sister lineage, the toothed tree frogs, Fig. 4) in the common ancestor of modern bufonids. In the few toothed anuran species, in which the oral cavity has been studied at the histological level, the oral epithelium is stratified, highly folded, thick and mucous, and shows a well-elaborated tridimensional architecture (e.g. Albright & Skobe, 1965; Loo et al. 1980). Only the very tip of the tooth pierces this epithelium, suggesting that teeth in frogs are not that efficient in aiding food uptake and processing, making them less governed by selective constraints. Their loss could have no lethal consequence (as in toads), as it would be largely compensated by the powerful tool that is their sticky tongue for food uptake. The complete lack of teeth is tentatively considered a synapomorphy of Bufonidae by most authors (e.g. Ford & Cannatella, 1993) but there are also toothless species in other anuran families (e.g. some leptodactylids; Lynch, 1970). It is worth noting that many frogs have developed odontoid elements (i.e. simple tooth-like projections of bone) on various bones and even on the lower jaw (Trueb, 1973; Shaw & Ellis, 1989). These odontoids (sometimes called fangs) could be adaptations for prey handling, which occurred as convergent evolutionary events in various lineages (Fabrezi & Emerson, 2003). The genetic processes underlying odontoid development need further investigation because their bony composition suggests that they result from different tissue interactions than those involved in tooth formation.

The edentate condition in true toads and in some other anuran taxa indicates that (i) the ability to form teeth can be easily lost and (ii) an alternative tool (the tongue), which probably arose prior to tooth loss, has been selected for during evolution. The developmental changes underlying tooth loss in toads, however, remain to be found. When considering that the ability to form teeth on the lower jaw was lost long ago in anurans, we can postulate that this condition was already that of bufonid ancestors, i.e. prior to tooth loss on the upper jaw. Therefore, the loss of ability to form teeth may have occurred in several steps, i.e. first tooth formation arrested in the lower jaw and then in the upper jaw and vomerine region. The same mechanism was probably involved but it remains to be understood. Further studies could be undertaken as for instance gene expression comparison in the upper (toothed) vs. lower (toothless) jaw of clawed toads, *Xenopus laevis* or

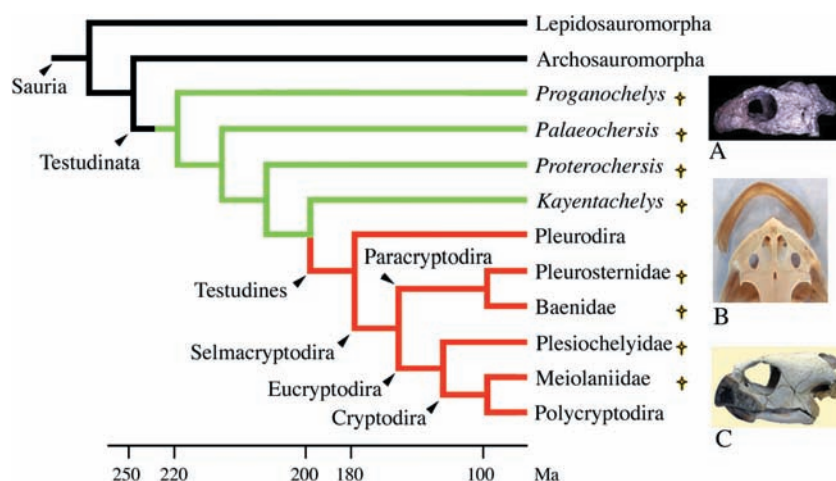


Fig. 5 Simplified turtle relationships (after Gaffney & Meylan, 1988; Rieppel, 1999; Joyce, 2007). Green lines: lineages with palatine teeth only; red lines: toothless lineages. (A) Skull of *Proganochelys quenstedti* (from Gaffney, 1990). (B) Dorsal view of the skull and beak of the snapping turtle (aquatic). (C) Skull and beak of a terrestrial turtle. †Extinct lineages.

X. tropicalis, or in the upper (toothed) jaw of these vs. the upper (toothless) jaw of a true toad.

Turtles

Fossil and living turtles constitute Testudinata (*sensu* Joyce, 2007). The living turtles are distributed within Pleurodira (side-neck turtles) and Cryptodira (hidden neck turtles) with a total of 257 living species and 12 families (Ernst & Barbour, 1992). The monophyly of Testudinata has never been questioned and all living species are toothless (Fig. 5).

What the fossil record tells us

The origin of turtles from ancestral sauropsids is still unclear and largely debated. Molecular data are partially congruent with morphological characters supporting diapsid rather than anapsid turtle relationships [Rieppel & deBraga, 1996; deBraga & Rieppel, 1997; see Laurin & Reisz, 1995 and Lee, 1997 for Parareptilia (anapsid) turtle relationships]. However, the molecular data conflict with palaeontological data as to where exactly turtles fit within diapsids (Rieppel, 1999). Phylogenetic studies either place turtles close to the lepidosauromorphs (tuatara, snakes and lizards) (e.g. Hill, 2005) or close to the archosauromorphs (crocodiles and birds) (e.g. Hedges & Poling, 1999; Iwabe et al. 2005). The turtle ancestor diverged from the other diapsids between 285 and 270 Ma (McGeoch & Gatherer, 2005) but its origin remains a mystery.

The most ancient and well-known turtle is *Proganochelys quenstedti* (late Triassic, 220 Ma). It shows primitive features absent from modern turtles that make it useful for understanding turtle evolution (Gaffney, 1990). *P. quenstedti* was roughly similar to the species that live today, except for, among other characters, the presence of several rows of conical teeth on the vomers, palatines and pterygoids (Fig. 6D), which make it unique among Testudinata as the other ancient turtles lack these teeth

(Joyce, 2007). The maxilla, pre-maxilla and dentary are edentulous but the pre-maxillary has tooth vestiges (Kordikova, 2002). Among amniotes the presence of vomerine and palatine teeth is widespread. This is interpreted as the primitive condition in Testudinata. Another fossil turtle, *Proterochersis robusta* (late Triassic) was probably contemporary to *P. quenstedti*. *P. robusta* possessed several rows of small pterygoid teeth and shared several features of pleurodires such as the pelvis fused into the shell. Its affinities to pleurodires, however, are not certain. *Palaeochersis talampayensis* (late Triassic of Argentina) appears closer to all other turtles than *P. quenstedti* and *P. robusta* but still outside the common ancestor of cryptodires and pleurodires (Rougier et al. 1995; Joyce et al. 2004; Joyce, 2007).

Unquestionable pleurodire skulls are not observed until the early Cretaceous (145 Ma). From this period onwards, Pleurodira lack palatine teeth. The oldest turtle, recognized by some authors as a Cryptodira, is *Kayentachelys aprix* (late Jurassic, 150 Ma), which possessed pterygoid teeth (Fig. 6E). The position of *K. aprix*, however, is still questioned and it is put outside the common ancestor of cryptodires and pleurodires by Joyce (2007). All other Cryptodira lack teeth. The pleurodire/cryptodire dichotomy probably took place during the early Jurassic (200–180 Ma). Depending on the position of *K. aprix* in the tree, palate teeth could have been lost either independently in Pleurodira and Cryptodira (Gaffney, 1990) or in the common ancestor of pleurodires and cryptodires. In ancient turtles, palatine teeth are characterized by a primitive, thecodont implantation, in which teeth are set in sockets that are distinct from the surrounding bone (Kordikova, 2002).

Adaptations to tooth loss in turtles

All living turtles, aquatic and terrestrial, have a keratinized beak (Fig. 6B,C). Although the common ancestor of all living turtles was aquatic, the earliest turtles clearly lived in a terrestrial environment (Joyce & Gauthier, 2004;

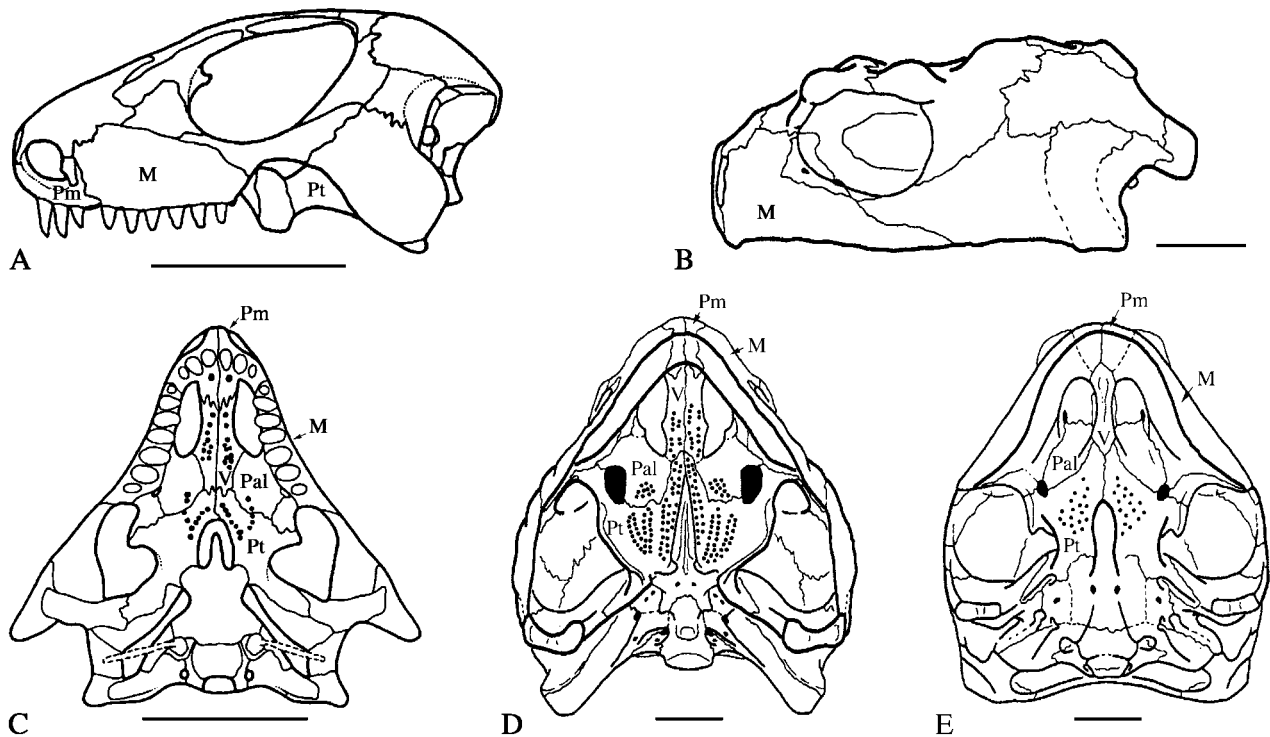


Fig. 6 (A and C) Lateral and ventral views of the skull of a primitive tetrapod, the parareptilian *Procolophon*. (B and D) Lateral and ventral views of the skull of *Proganochelys quenstedti*. (E) Ventral view of the skull of *Kayentachelys aprix*. Small dots represent teeth. M, maxillary; Pal, palatine; Pm, pre-maxillary; Pt, pterygoid; V, vomer. (A and C) From Carroll & Lindsay (1985). (B and D) From Gaffney (1990). (E) From Gaffney et al. (1987). Scale bars, 1 cm.

Scheyer & Sander, 2007). The origin of the beak in turtles was traced back as early as *P. quenstedti*, which had toothless jaws but already possessed a rigid beak (the presence of a keratinized beak is suggested by the condition of the mandibles). Turtle beaks have sharp edges for cutting food and most have strong jaws, which they use to tear food and capture prey.

Hatchlings emerge from their eggs using what is commonly known as the egg tooth or caruncle. It is located at the front of the upper jaw and disappears a few months after hatching. This is a modified scale (i.e. a keratinized structure derived from epithelial cells) and not a real tooth.

The beak covers the upper and lower jaws and is ornamented by horny ridges, similar to the beak of birds. Various adaptations of the beak and oral cavity have occurred by means of natural selection (knife-sharp ridges for carnivorous species, ridges with serrated edges for plant-eating species, bony ridges helping to crush mollusc shells) that permitted turtles to utilize a large food spectrum (leaves, fruit, mushrooms, insects, snails, crayfish, fish and jelly fish). Turtles use their tongues in swallowing food.

It seems clear from the fossil record that the ability to form teeth on the upper and lower jaw was lost before the origin of Testudinata, i.e. approximately 250–220 Ma. At this epoch, the teeth were widespread in the oral cavity of

early reptiles. We postulate that early Testudinata (yet to be found in the fossil records) possessed teeth fixed on the upper and lower jaw bones and on various bones of the palate. Teeth located on the pre-maxillary, maxillary and dentary were subsequently lost during Testudinata evolution but before the first Testudinata found in the fossil record, approximately 220 Ma. As suggested for birds, the presence of a keratinized beak that was efficient for food uptake probably relaxed the functional pressure on teeth, which were probably lost through a similar process to that described in birds (see above). In the turtle's ancestor, as in the bird's ancestor, the beak minimized the negative consequences of tooth loss.

In turtles, teeth were retained on the palate longer than in jaws. Teeth were lost in the vomers and palatines first, then later on the pterygoids. It appears that this loss was not a dramatic event as this has occurred independently in several vertebrate lineages during evolution. For instance, the Archosauria (living crocodiles and birds but also extinct non-avian dinosaurs, pterosaurs and relatives of crocodiles) have no teeth on the pterygoid, palatine or vomer.

Echidnas

Monotremes form a single order, Monotremata. These egg-laying mammals are the survivors of an early branching

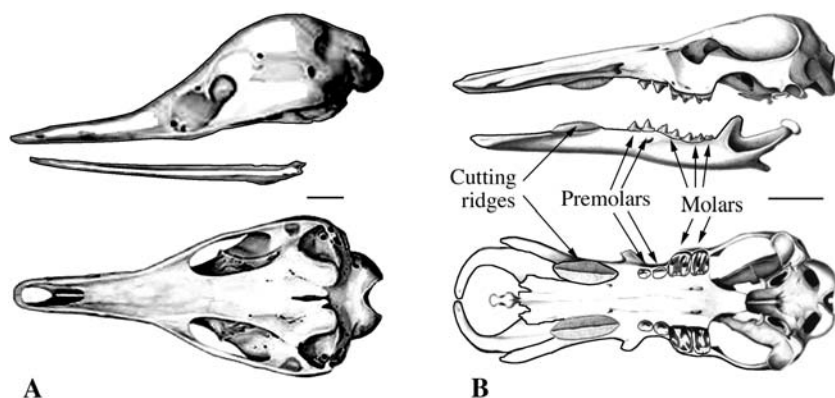


Fig. 7 Lateral and ventral views of the skull of Monotremata. (A) The echidna, *Tachyglossus aculeatus*. (B) A fossil (adult) ornithorhynchid, *Obdurodon dicksoni* (after Musser & Archer, 1998). In this species the skull morphology is very similar to that in platypus (see Fig. 13), except for the presence of teeth in adults (two pre-molars and two or three molars). Scale bars: A, 1 cm; B, 2 cm.

of the mammal tree that diverged from the other mammalian lineage, Theria, about 225 Ma (molecular data; Westerman & Edwards, 1992; van Rheede et al. 2006) (Fig. 1). Monotremata have retained reptilian characters, such as oviparity and a therapsid-like shoulder girdle, that have been lost in therian lineages. Extant and extinct monotremes are composed of four families, two extinct (Kollikodontidae and Steropodontidae, both from the early Cretaceous, 110 Ma) and two still represented in nature, the semi-aquatic Ornithorhynchidae (platypus) and the terrestrial Tachyglossidae (echidnas) (Musser & Archer, 1998). To date, monotremes are all indigenous to Australia and New Guinea but they were more spread in the past as revealed by the presence of a fossil platypus (*Obdurodon*) in the middle Miocene (10 Ma) of Argentina (Pascual et al. 1992; Musser & Archer, 1998).

Extant tachyglossids include the short-beaked echidna, *Tachyglossus aculeatus*, living in Australia and New Guinea, and three species of the genus *Zaglossus*, the long-beaked echidnas *Z. bruijnii*, *Z. bartoni* and *Z. attenboroughi*, in New Guinea. The most prominent feature of the head is the elongate, hairless and rounded snout (Fig. 7A). This tubular, elongated snout has a small and narrow opening. Juvenile and adult echidnas are edentulous and the lower jaw is extremely reduced. However, the hatchling possesses an egg tooth on the snout (a true tooth as in lizards and snakes) that is a useful tool to pierce the keratinized egg shell. This is the only tooth the echidna will ever possess and it is lost after hatching. However, it is worth noting that this egg tooth is crucial for echidna survival. This explains why, although echidnas have been toothless for many millions of years, there are still strong functional constraints acting on dental proteins (Sire, unpublished data).

What the fossil record tells us

Ornithorhynchidae are known from the Cretaceous (120 Ma), whereas Tachyglossidae seem younger, as they are only found in the mid-Tertiary (25 Ma). Their relationships are not clear. There are several arguments that suggest

that echidnas are derived from a platypus-like ancestor but they may have evolved from an unknown, more generalized Monotremata (Musser, 2003). The divergence between the two lineages is estimated to have occurred by the end of the Cretaceous (80–70 Ma; Rowe et al. 2008).

Fossil tachyglossids, such as *Megalibgwilia robusta* from the middle Miocene (15 Ma; Griffiths et al. 1991), are similar in appearance to the living *Z. bruijnii*, although larger, and were probably able to consume larger prey. The skull had a long, sturdy snout and the palate possessed a groove to accommodate tongue extension. There were no teeth. The largest extinct echidna *Z. hacketti* is known only from a few bones found in Western Australia (Pleistocene, 0.5 Ma).

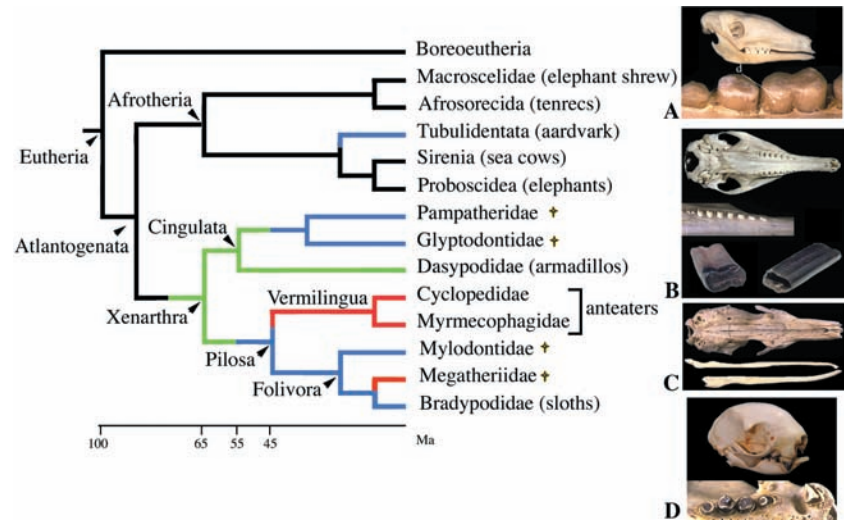
Given that ornithorhynchids have teeth (see below), it appears that the ability to form jaw teeth in tachyglossids was lost after the echidna ancestor diverged from the platypus ancestor, i.e. by the end of the Cretaceous/beginning of the Tertiary (70 Ma).

Adaptations to tooth loss in echidnas

Echidnas are insectivores. The diet of *T. aculeatus* consists of ants and termites, whereas *Z. bruijnii* prefers insect larvae. They use their long sticky tongue to catch prey from its hole. Capturing and grinding prey with the tongue occur due to a unique jaw mechanism; the echidnas open and close their mouth by rotating their mandibles around their long axes (Murray, 1981). Histological observations and scanning electron microscopy of the palatal epithelium of *T. aculeatus* revealed the presence of keratinized palatal spines. It is suggested that these horny teeth-like projections are highly differentiated filiform papillae, which have developed as a compensatory mechanism of mastication, as this animal is edentulous (Doran, 1975).

The large temporal gap between the earliest echidna in the fossil record (around 25 Ma) and the molecular datation of the differentiation of the lineage (estimated at 70 Ma) does not allow for clear relationships to be defined. However, we can speculate about the events that might

Fig. 8 Simplified phylogeny of Xenarthra and Afrotheria (after Hallström et al. 2007; Seiffert, 2007). Red lines: toothless lineages; blue lines: enamel-less lineages; green lines: enamel reduction. (A) Lateral view of the skull and detail of the upper cheek teeth of an armadillo, *Oryzomys afer*. (B) Ventral view of the skull and of the upper right jaw, and detail of extracted teeth of a nine-banded armadillo, *Dasyus novemlineatus*. (C) Ventral view of the skull of a giant anteater, *Myrmecophaga tridactyla*. (D) Ventral view of the skull and detail of the upper jaw of a three-toed sloth.



have led to tooth loss. It appears that the earliest toothed monotremes, such as kallikodontids from the early Cretaceous (110 Ma; Flannery et al. 1995), were beaked species. Again, as described for birds and turtles (see above), it seems that the presence of a keratinized beak prior to tooth loss could account for relaxing the functional constraints on teeth. Simultaneously, the elongated morphogenesis of the lower jaw could have reduced the ability of the jaw to accommodate teeth. Snout elongation (like in anteaters, see below) was probably concomitant with the presence of an elongate tongue, a tool that compensated, along with the beak, for tooth loss. Further adaptations improving food processing, such as the spines on the tongue, developed later through natural selection.

Anteaters

Anteaters belong to Xenarthra, which are principally characterized from the other placentals in having additional vertebral articulations (one of the five skeletal synapomorphies of Xenarthra). The clade Xenarthra is composed of Cingulata (shelled Xenarthra, armadillos and extinct pampatheres and glyptodonts) and Pilosa [composed of Vermilingua, anteaters and Folivora (arboreal sloths and their extinct relatives)] (Fig. 8). The 31 species of living xenarthrans are distributed into five families, all of which are restricted to Central and South America (with the exception of one species, the nine-banded armadillo, which ranges north to the USA). The five families include the Dasypodidae (armadillos, 21 species), Myrmecophagidae and Cyclopedidae (anteaters, four species), Bradypodidae (three-toed sloths, four species) and Megalonychidae (two-toed sloths, two species) (Vizcaino, 1995; Anderson & Handley, 2001). Anteaters are toothless, whereas sloths lack enamel on their teeth and, in armadillos, enamel is present only in the first generation of teeth, at least in the nine-banded armadillo (see below). In order to better

understand the origin of the dental features observed in Xenarthra, prior to dealing with anteaters specifically, we briefly review our current knowledge on Xenarthra origin and on the evolution of their dentition.

What the fossil record tells us

Since the redistribution of the species, which comprised the polyphyletic Edentata into the three, now undisputed, monophyletic clades, Xenarthra, Tubulidentata and Pholidota, xenarthrans are considered as a major clade of placental mammals (e.g. Murphy et al. 2001; Madsen et al. 2001; Delsuc et al. 2002, 2004; Möller-Krull et al. 2007). However, their position among placentals is still in debate, as illustrated by two recent studies based on large data sets of nuclear genes (Hallström et al. 2007; Nikolaev et al. 2007). The current trends supported by these molecular phylogenies are that Xenarthra are the sister group of Afrotheria (Figs 1, 8). The probable reason for the previous uncertainty is that the divergence of the placental lineages took place a long time ago within a narrow temporal window. Molecular analyses provide estimates of the origin of Xenarthra within the late Cretaceous (100 Ma; Delsuc et al. 2004). These molecular data are, as usual, at odds with the palaeontological data, which indicate that the earliest records of Xenarthra come from the Paleocene (58–55 Ma; Scillato-Yané, 1976). However, because these fossil species show features that are already proper to this order, this strongly suggests that the origin of the xenarthrans is more ancient and probably located in the late Cretaceous/early Paleocene (80–61 Ma), a date that substantially reduces the gap between molecular and palaeontological data. It is probable that the evolutionary radiation of Xenarthra occurred around 65 Ma, after South America became isolated from the other continents (Patterson & Pascual, 1972).

In the early Paleocene (65–61 Ma) there are no fossil species that could be considered with certainty as a stem

Xenarthra (e.g. proto-Xenarthra). Given their similar anatomy to extant armadillos, the Palaeonodonta are often proposed as related to the Xenarthra (Hoffstetter, 1982; Rose et al. 2004). However, palaeonodonts are also considered to be closer to Pholidota (pangolins) than to the other lineages (see below). It is believed that the ancestral lineages that lead to armadillos, sloths and anteaters diversified during the Eocene (55–33 Ma). However, phylogenetic relationships of modern species to fossil records are difficult and the absence of representatives of the main xenarthran lineages in the South American fossil record of the late Cretaceous/early Paleocene period is still an enigma (Carlini & Scillato-Yané, 2004).

The first group consisted of land-dwelling armoured plant-eaters, whose descendants developed into modern-day armadillos. Armadillos are the oldest true Xenarthra known in the fossil record (late Paleocene, 59–57.5 Ma; Scillato-Yané, 1976; Bergqvist et al. 2004). A second group consisted of insectivores, which specialized in ants and termites (anteaters). There are only a few fossils available for early anteaters. The oldest fossil of this group is from the early Miocene, and probably closely related to *Tamandua* (Carlini et al. 1992). Specimens, such as *Protamandua*, begin to be more abundant during the early/middle Miocene (17.5–16.3 Ma; Hirschfeld, 1976; Gaudin & Branham, 1998). Given the close relationships between *Vermilingua* and *Folivora* (sloths), anteater ancestors were certainly older and could have differentiated during the early Eocene (Pujos & De Iuliis, 2007; Pujos et al. 2007). The third group is represented by modern-day sloths, a tree-dwelling group of animals lacking armour. The oldest sloths are reported from the Eocene of Antarctica (Vizcaino & Scillato-Yané, 1995) but most records are from the Miocene period. Recently, the description of two *Pseudoglyptodon* species from the early and late Oligocene (36–24.5 Ma) revealed that this typical ground sloth shared several dental characters with armoured *Glyptodontoidea* and may represent not only a pre-folivoran stage but also the earliest known *Pilosa* (McKenna et al. 2006; Pujos et al. 2007; Pujos & De Iuliis, 2007).

Evolution of the dentition in Xenarthra

The condition of xenarthran ancestors was a continuous dentition, including incisors and canines (Hoffstetter, 1982). However, in all true Xenarthra fossils known to date, the upper teeth are only located on the maxilla, the pre-maxilla being always edentulous. The early fossils (early Miocene; Carlini et al. 1992) relating to *Vermilingua* (anteaters) lack teeth (a synapomorphy of this group) and a study of the skull structure in *Neotamandua* and *Palaeomyrmidon* clearly shows that the fundamental specializations of the skull, related to ant and termite eating, had been achieved prior to the Miocene.

The general dental morphology of extant and extinct *Cingulata* (armadillos) and *Folivora* (sloths) is different

from that of the other placental mammals, including their closest relatives, the Afrotheria, i.e. absence of the most anterior teeth, incisors and canines; presence of a few molariform cheek teeth ('cheek' means that pre-molars and molars are hardly morphologically distinguishable one from another), which are cylindrical, generally homodont in *Cingulata* but heterodont (i.e. teeth of various shapes such as incisors, canines and molars) in *Folivora*, and hypsodont (i.e. teeth with a high crown and tall with respect to their occlusal area) and hypselodont (ever growing with open roots). In addition to lacking enamel (see the armadillo description), xenarthran teeth are built of different dentine tissues (Vizcaino & Scillato-Yané, 1995). In a comparative study of xenarthran dental structure, Ferigolo (1985) described the teeth as being generally composed of a small layer of cementum covering the dentine crown that is constituted of a hard layer of modified orthodentine (osteodentine in some *Glyptodontidae*). The upper region of the pulp cavity is occupied by vasodentine. There is a large difference in resistance to wear between the hard outer and soft inner layers of dentine. Vasodentine is absent from non-abraded teeth, which means that either this typical dentine is deposited late, after tooth eruption, or is a reaction to abrasion and contributes to the protection of the pulp cavity from external pathogens.

There is no information on teeth in the oldest *Cingulata* from the late Paleocene but subsequent representatives already show the dental characters of living species and a similar statement is true for the early fossil sloths recorded. This means that ever-growing, high-crowned teeth differentiated early in the evolution of these lineages, i.e. probably during late Cretaceous/early Paleocene, and that the adaptation to plant feeding was secondary to this condition.

The Xenarthra appear marked by strong disadvantages, with a reduced dentition and thin enamel that would have provided poor tooth protection against abrasion. Therefore, one would have expected rapid extinction, rather than the observed rapid radiation. Apparently, early Xenarthra found, in the recently geographically isolated South America, free niches, which allowed their diversification, with hypsodonty and hypselodonty permitting teeth to compensate for tooth erosion.

Anteaters have lost the ability to form functional teeth

The four living species of anteater (giant and pygmy anteaters, and two tamanduas) and all of their fossil relatives are toothless. The ability to form functional teeth was lost long before the Miocene period (i.e. > 25 Ma) but in tropical forests fossilization is not favourable and skeletal remains of *Vermilingua* from earlier periods have probably not been conserved.

Although anteaters have no functional teeth, tooth germs have been described in anteater embryos, in many

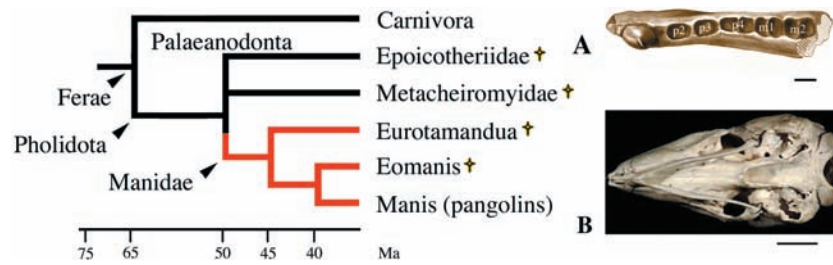


Fig. 9 Simplified phylogeny of Pholidota (after McKenna & Bell, 1997; Nowak, 1999). Red lines: toothless lineages. (A) Dorsal view of the right dentary of a primitive palaeanodont (Eocene) showing the large canine (no incisors) and the alveoli for three pre-molars p2–p4 (p1 absent) and two molars. Modified after Rose et al. (2004). (B) Ventral view of the skull and lower jaw of a living pangolin, *Manis javanica*. Note the extremely narrow and weak blade-like mandible. †Extinct lineages. Scale bars: A, 1 mm; B, 1 cm.

if not all situations. These remnants of teeth, however, are resorbed prior to birth (Peyer 1968). This is evidence that Vermilingua ancestors had functional teeth and that early odontogenetic processes are conserved long after the ability to form functional teeth is lost, as observed in modern birds (see above).

Adaptations to tooth loss in anteaters

Anteaters ingest food without chewing using a slender, elongated tongue, which can project to a distance greater than the cranial length. The tongue is ornamented with little keratinized spines that point backwards and is covered with sticky saliva. A large and elongated hyoid apparatus, with articular surfaces permitting great freedom of movement, supports the tongue. The hyoid muscle enables the tongue to project out with great speed and precise positional control. This feature combines with an elongated secondary palate, accommodating the retracted tongue within the oropharynx. The giant anteater, *Myrmecophaga tridactyla*, has a long and thin head, and its tongue is 60 cm in length and can extend and retract at the incredible rate of 150 times per minute. With their special insect-catching tongue, anteaters can eat thousands of ants or termites in a day. Instead of teeth, edentulous anteaters have horny protrusions called papillae on the roofs of their mouths and strong, muscular stomachs (Vizcaíno & Loughry, 2008).

When considering the features of the dentition in Xenarthra (but see later and Fig. 8), it appears that (i) these highly adapted features in anteaters (especially the skull and tongue) were acquired after the Vermilingua differentiated from the Tardigrada, and (ii) the ancestral condition in the Vermilingua was a reduced dentition with hypsodont and hypselodont teeth either covered or not with enamel and a diet based on insects. One can also imagine that a sticky tongue was present before the ability to form teeth was lost (as also supposed for the Pholidota ancestor, see below). We can deduce that the absence of an efficient protective layer of enamel (no enamel or only a thin layer that is rapidly worn away) led to strong abrasion of the teeth. This excessive abrasion could have

resulted in a specialization towards a diet that does not necessitate teeth (ants and termites) and the presence of a sticky tongue was therefore necessary to fulfil this function. Given such a specific adaptation, the functional constraint on teeth was relaxed, allowing for the loss of teeth, perhaps through changes in the odontogenic pathways resulting, for instance, from the shift of competent odontogenic territories linked to a change in the shape of the jaw.

Pangolins

Pangolins, also called scaly anteaters, have their back covered with large overlapping keratinized scales made up of agglutinated hairs. They constitute the clade Pholidota, with a single family (Manidae) and a single genus (*Manis*) including several extinct species. Eight species are still living, four in Africa and four in Asia (Gaubert & Antunes, 2005).

Living pangolins and extinct manids lack teeth as adults. They have incomplete zygomatic arches and possess an extremely reduced, bladelike mandible (Fig. 9B). Each dentary has a single bony protrusion (Nowak, 1999). The ability to form functional teeth appears to have been lost in an ancestral manid. It is worth noting that vestigial teeth start to form in embryos but are resorbed prior to birth. In *M. javanica*, Röse (1892) described a dental lamina in the anterior part of the lower jaw and tooth germs forming rounded buds. The observation of vestigial teeth was confirmed by Tims (1908).

What the fossil record tells us

The relationships of Pholidota are still debated. Phylogenetic analyses based on morphological data placed pangolins as the sister group to xenarthrans (Novacek & Wyss, 1986; Novacek, 1992a,b) and, even within them, close to anteaters. However, it seems that these close relationships are due to high anatomical convergences in relation to a similar diet. Molecular analyses based on large data sets clearly indicate that Pholidota is the sister group to Carnivora (Murphy et al. 2001; Scally et al. 2001) (Fig. 9).

Modern-looking fossil pangolins (e.g. *Eomanis* and *Eurotamandua*) were found in the middle Eocene of Germany (40 Ma; Horovitz et al. 2005). A single fossil was found in the lower Oligocene of North America (30 Ma) and more recent remains of scaly anteaters come from the Pliocene of Europe, the Plio-Pleistocene of South Africa and the Pleistocene of Asia (Emry, 1970).

Palaeontological studies support that manids are possibly related to Palaeanodonta, a group of extinct toothed anteater-like mammals that lived from the late Paleocene (55 Ma) to the early Oligocene (35 Ma; Gunnell & Gingerich, 1993) (Fig. 9). Both manids and palaeanodonts constitute the Pholidota. The best-known extinct Pholidota is *Metacheiromys*, from the mid-Eocene of North America, that already looked like a modern species (Simpson et al. 1931). Palaeanodonts had a reduced dentition with no incisors, one large canine and five to eight post-canine teeth not strongly rooted in the jaw (Fig. 9A). They probably fed mainly on ants and termites as suggested by the morphology of their post-canines. In some palaeanodonts the post-canines are covered with enamel (e.g. epoicotheriids), whereas others have no enamel or only a thin layer that is rapidly worn away (e.g. metacheiromyids; Gunnell & Gingerich, 1993).

Adaptations to tooth loss in pangolins

Pangolins are covered by tough scales that protect them from their aggressive prey, ants and termites. The pangolins open the anthills and termite mounds using their long claws on their forelimbs and catch the prey with their long tongue; the myrmecophagous diet and the elongate, sticky tongue are efficient adaptations to toothlessness. Their tongue is vermiform and can reach 25–40 cm. It is covered with viscous saliva, secreted by a large salivary gland. The tongue extends to the abdominal cavity and is attached to the pelvis. In addition, the stomach possesses a muscular, gizzard-like pyloric region with keratinized spines (the so-called 'pyloric teeth'; Krause & Lesson, 1974) and contains small stones and sand. All of the processes of grinding are done in this gizzard, compensating for the lack of teeth. The stomach lumen is lined with a cornified, stratified squamous epithelium and there are numerous glands (a mucous gland spread in three regions, an oxyntic gland and a pyloric gland) responsible for the digestion of their highly chitinous diet (Nisa et al. 2005).

It remains uncertain which developmental failure was responsible for tooth loss in the toothed ancestral Pholidota but, given the palaeontological data, this event probably occurred during the late Paleocene/early Eocene period (55–50 Ma), when the first manid ancestors differentiated. Indeed, edentate fossil manids are found as early as 40 Ma in the fossil record (Horovitz et al. 2005). We postulate that tooth loss could have been preceded by a drastic reduction of the enamel cover, as observed in some palaeanodont taxa. Such a loss of protective hard

cover resulted in a strong abrasion of the teeth, as described in some species (Secord et al. 2002), and hence in a specialization towards a different diet based on insects. Whether these early manids already had a protactile sticky tongue as observed in recent species remains uncertain but such a tool could have contributed greatly to improve food uptake and compensated when the ability to develop teeth was lost.

Baleen whales

The Cetacea is composed of 'Archaeoceti', which is a stem polyphyletic and exclusively extinct assemblage, Odontoceti (toothed whales) and Mysticeti (baleen whales). Living odontocetes contain six families (69 species): Delphinidae, Monodontidae, Phocoenidae, Physeteridae, Platanistidae and Ziphiidae. All fossil and modern toothed whales are thought to be monophyodont, i.e. they have a single tooth generation. Most species are homodont but some extinct Odontoceti, e.g. Agorophiidae and Squalodontidae, were heterodont (Fordyce, 1982). In contrast to delphinids and phocoenids, in which teeth are numerous on both jaws, modern physeterids, ziphiids and monodontids have reduced dentitions: no teeth or unerupted teeth on the upper jaw and a few teeth on the lower jaw. Some females can even be edentate. A toothless ziphiid-like fossil has been described (Fordyce et al. 2002). Male (rarely female) narwhals (monodontids) have a single long spiralled tusk derived from the incisor on the left maxilla and a small tooth on the right maxilla; they have no teeth on the lower jaw.

Living mysticetes comprise at least 14 species distributed within four families: Balaenidae, Balaenopteridae, Neobalaenidae and Eschrichtiidae (Fig. 10). All living species of baleen whales are toothless but they derive from toothed mysticete ancestors.

What the fossil record tells us

The origin and phylogeny of the Cetacea is still debated but most authors now agree (with a large input from molecular phylogenies) that cetaceans are nested with artiodactyls (hence the name of the new order, Cetartiodactyla) and closely related to anthracotheres, the ancestors of hippopotamuses (Gatesy et al. 1996; Nikaido et al. 1999). Modern cetaceans are supposed to have arisen from 'Archaeoceti', a group of early cetaceans, which were large toothed predators that appeared in the Eocene (50 Ma). The archaeocetes themselves are believed to derive from aquatic even-toed ungulates (artiodactyls) that diverged at the Cretaceous/Tertiary boundary and during the early Paleocene (60 Ma; Bajpai & Gingerich, 1998; Thewissen et al. 2007; Uhen, 2007). Archaeoceti, such as *Basilosaurus* from the middle Eocene, were diphyodont, i.e. they had milk teeth, shed and replaced by permanent teeth. They were also heterodont separated from the

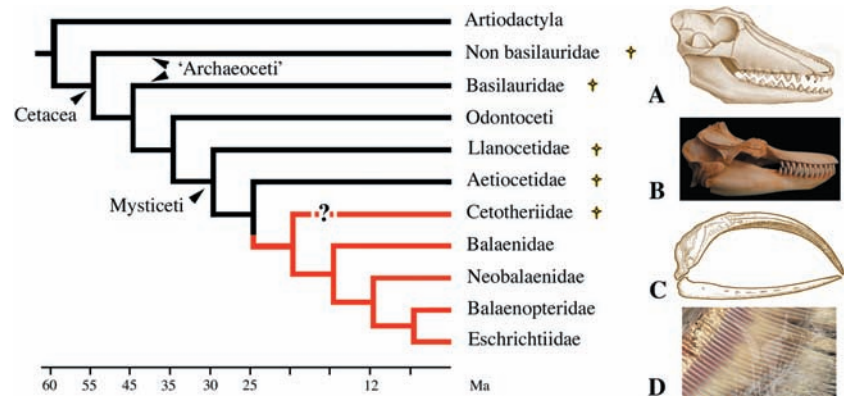


Fig. 10 Simplified phylogeny of Cetacea (after Uhen, 2002; Deméré et al. 2008). (A) Skull of an early archaeocete, *Pakicetus* from the early Eocene. (B) The toothed jaws of an odontocete, the orca *Orcinus orca*. (C) The edentulous jaw of a mysticete, the bowhead whale *Balaena mysticetus*. (D) Detail of a baleen plate of a gray whale *Eschrichtius robustus*. †Extinct lineages.

cheek teeth (pre-molars and molars) by a diastema (Uhen, 2002).

Early fossil mysticetes were also found as early as the Eocene period. They possessed teeth, although their cranial architecture was similar to that of modern baleen whales (Barnes & Sanders, 1996). *Aetiocetus cotylalveus* (Aetiocetidae) appears to be the most primitive mysticete and possessed a heterodont dentition on both jaws (incisors, canines, pre-molars and molars). It is considered a transitional, toothed mysticete that gave rise to toothless primitive types (Barnes, 1984). The small multicusped teeth of aetiocetids were used to filter food from the seawater rather than for selecting individual prey (Fordyce & Barnes, 1994). Aetiocetids are thought to have had rudimentary baleen plates on the upper jaw (Deméré, 2005; Deméré et al. 2008). Being made of keratin, baleens do not fossilize. However, the bones that support the baleens (i.e. the palate bones) show additional features that bear witness to the presence of baleens in toothed mysticeti from the early Miocene (20 Ma; Kellogg, 1965; Fordyce & Muizon, 2001) and even earlier as recently reported in Oligocene aetiocetids by Deméré et al. (2008). These authors described the simultaneous presence of teeth and baleens on the jaw, which supports a progressive transition from a condition with teeth to a condition with baleens during further evolution in mysticetes. Such a transition is still observed in the embryonic series of modern species. Indeed, although mysticetes are toothless at birth, tooth buds form and grow in the embryos, then baleens develop and the teeth are resorbed (Karlsen, 1962).

Most post-Oligocene mysticetes are baleen-wearing and toothless (Fordyce & Barnes, 1994). The most ancient family of edentulous mysticetes are the Cetotheriidae that lived from late Oligocene (25 Ma) to Pliocene (5 Ma). They have been positioned close to Balaenopteridae but their relationships with modern Mysticeti are unclear (Uhen, 2002) (Fig. 10).

Vestigial teeth in baleen whale embryos

The presence of tooth buds in baleen whale embryos was first reported by Geoffroy Saint-Hilaire (1807) and Cuvier

(1836) but no information was available about the structure of the hard tissues. Van Dissel-Scherft & Vervoort (1954) and then Karlsen (1962) described the successive steps of vestigial tooth development in fin whale *Balaenoptera physalus* embryos. A total of 110 tooth germs (53 at the upper jaw and 57 at the lower jaw) develop up to the advanced bell stage but the teeth do not erupt. It is noteworthy that 23 and 15 of these tooth germs, respectively, are double teeth developing by fusion of tooth germs. On the lower jaw, the first tooth germs develop close to Meckel's cartilage before the bone starts to ossify. In *B. physalus*, tooth germs start to develop in 2-cm embryos from a dental lamina and then tooth morphogenesis and differentiation take place until an advanced bell stage. The enamel organ is well differentiated and faces well-differentiated odontoblasts, which deposit pre-dentine matrix. The pre-dentine, however, does not mineralize (Fig. 11A–D). Two types of dentine are deposited by the odontoblasts: one is acellular orthodentine, whereas the other contains enclosed mesenchymal cells. Some tooth germs produce cellular dentine only, whereas both dentine types occur in others. Dentinal tubules are not present and no enamel matrix is produced. This is the most advanced stage that tooth germs can reach (in 63-cm embryos) and then they start to be resorbed. The odontoblasts disappear and the enamel organ reduces. In an 82-cm embryo, dentine resorption has started (Fig. 11E,F). In a 212-cm embryo, all tooth germs on the upper jaw have been resorbed. A marked proliferation of epithelial cells, indicating the formation of the first baleen plates (also called whalebones), is identified in the region of the tooth germs as the latter start to be resorbed. In a 330-cm embryo, tooth germs are no longer seen in the lower jaw. It is worth noting that a newborn fin whale is 6 m long.

Working on *Balaenoptera acutorostrata* (a baleen whale that is 3 m long at birth), Ishikawa & Amasaki (1995) similarly found that vestigial teeth develop until the embryos reach 90 cm, then degenerate and were no longer detected in 182-cm embryos. As described above, degenerate tooth buds and rudiments of baleen plates were simultaneously present on the upper jaw, a situation

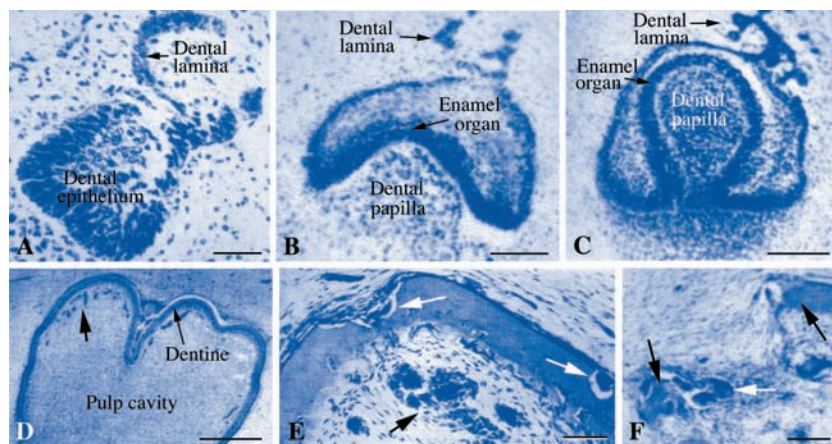


Fig. 11 Histological evidence of embryonic teeth developing in the embryos of the baleen whale *Balaenoptera physalus*. Most tooth germs attain an advanced bell stage, until dentine deposition, and are then progressively resorbed. (A–D) Tooth morphogenesis and differentiation. (E and F) Resorption. (A) Cap stage. (B) Bell stage; dental epithelium starts to fold around dental papilla cells. (C) Advanced bell stage, in which the dental epithelium entirely surrounds the dental papilla. (D) A thin layer of dentine has been deposited; enamel is not identified. The arrows point to numerous capillary blood vessels located close to the dentine layer. (E) Initiation of resorption process. White arrows indicate osteoclasts. Black arrow points to blood vessels. (F) Advanced stage of resorption. Black arrows indicate the dentine; white arrow points to osteoclasts. Modified after Karlsen (1962). Bars: A, 50 μm ; B, C, E and F, 100 μm ; D, 500 μm .

that is reminiscent of the ancestral condition in Oligocene mysticetes.

Adaptations to tooth loss in baleen whales

Baleen plates are a unique filtering structure used to consume prey. The baleen plates are keratinized modified elements of the oral epithelium, rooted in and suspended from the palate and extending into the oral cavity (Utrecht, 1965). They consist of many fibrous tubules packed together to form plates (Fig. 10D), arranged in a row on either side of the mouth. The ends of the plates become frayed and interwoven, resembling a comb, creating an efficient sieve for trapping small fish and zooplankton.

Trends in tooth evolution in baleen whales can be summarized as follows. The primitive heterodont and diphyodont dentition of the ancestral aquatic artiodactyls was replaced by a homodont and monophyodont dentition (as currently observed in toothed whales). In the ancestors of mysticetes the teeth became smaller and the animals adapted to filter-feeding, which allows access to small prey. This change in feeding was facilitated by the simultaneous presence of rudimentary baleen plates. As these tools provided more efficient filtering they were positively selected during further mysticete evolution and the functional constraints were relaxed on teeth. The small teeth were no longer useful and their loss was largely compensated by the presence of baleens. This event could have occurred during the Oligocene period, around 30 Ma. It is still difficult to establish solid relationships between the Oligocene toothed baleen whales and the modern mysticete. However, it seems more parsimonious to postulate that the common ancestor (probably toothed)

of all the modern mysticete lineages possessed baleens rather than baleens appearing independently in these lineages.

In modern mysticetes there is a close relationship between the vestigial tooth bud being resorbed and baleen plate development, and we could speculate that the development of the baleen plates could have played a role in tooth loss. However, baleen plates and functional teeth were simultaneously present, in the same locus, in the upper jaw of archaic mysticetes. The process that could have led to the loss of the ability to form teeth in an ancestral mysticete therefore remains unknown but, as discussed for birds (see above), the loss of the ability to form functional teeth could have been the consequence of a developmental shift of the oral epithelium preventing signalling molecules from reaching their mesenchymal targets.

Some comments on species lacking enamel only

Three clades of living mammals, i.e. Cingulata (armadillos) and Folivora (sloths) within Xenarthra and Tubulidentata (aardvark) within Afrotheria, are said to possess enamel-less teeth (Figs 1, 8). This claim is not correct as enamel has been described in the nine-banded armadillo.

The case of armadillos

The nine-banded armadillo (*Dasypus novemcinctus*) has 32 teeth but giant armadillos (South America) have up to 100 teeth. Armadillos are a key group in the story of xenarthran

dentition because, in addition to generalized features shared with sloths, in juveniles the teeth are covered with a thin layer of enamel, at least in *D. novemcinctus*, the best known species in the Cingulata lineage. The presence of enamel needs further attention as this condition is generally neglected in most text books.

Tomes (1874) was the first to describe an enamel organ in embryonic nine-banded armadillos. He believed that this enamel organ belonged to vestigial (rudimentary) teeth, as a general condition observed in several unrelated species, and that its presence was independent of enamel deposition. Nevertheless, 30 years later Spurgin (1904) reported that a thin layer of true enamel was present on the dentine surface of the first-generation teeth. This observation was confirmed by Martin (1916), who described, using a large growth series of embryos and juveniles, that not only is a thin enamel layer present in milk teeth but also that it covers permanent cheek teeth. The enamel organ is well differentiated and the four typical layers are present (inner and outer enamel epithelium, stratum intermedium and stellate reticulum). The differentiated ameloblasts are short but with well-defined Tomes' processes. However, in permanent teeth the possibly prismatic nature of the 100- μ m-thick enamel layer was not identified, if present. In both dentitions the thin enamel layer progressively disappears by abrasion.

Unfortunately there are no similar studies in the other 20 living cingulatan species. It is now agreed that the sub-family Dasypodinae (with the only genus *Dasypus*) is the most basal lineage of Cingulata (Delsuc et al. 2002). One can therefore speculate that the presence of a thin enamel cover was the ancestral condition in Cingulata and that enamel is either still present (but not yet described) in the three other sub-families (Euphractinae, Chlamyphorinae and Tolypeudinae) or was secondarily lost in these lineages after they separated from the Dasypodinae. This should be clarified in the future through detailed studies focusing on embryos.

In some species, the lack of enamel is compensated by a well-developed cementum, which allows a better resistance to abrasion than dentine. The cementum is thin but very hard in species of the genera *Euphractus* and *Cabassous*. In *Dasypus*, the cementum is not hard but it is thick, cellular and lamellar (Keil & Venema, 1963).

The presence of enamel in the nine-banded armadillo is evidence that its ancestor had typical enamel-covered crowns. Simpson (1932) found true enamel on the permanent teeth of an extinct armadillo, *Utaetus buccatus*, from the Eocene. The enamel layer was, however, thin and not present in all teeth due to wear. The presence of enamel in the Cingulata lineage means that this tissue was present in the common ancestor of the two Xenarthra lineages, Cingulata and Pilosa (see above). After Pilosa differentiated, enamel could have been lost either in their common ancestor [i.e. before tooth loss occurred in the

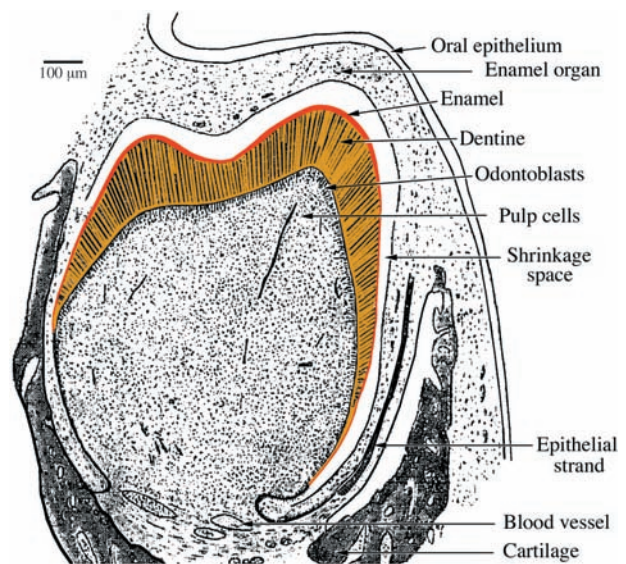


Fig. 12 Tooth developing in the nine-banded armadillo. Note the presence of a thin layer of enamel covering the orthodontine crown. Modified after Martin (1916).

Vermilingua (anteaters) lineage] or in the ancestor of Folivora (sloths) as, to our knowledge, enamel has never been reported in sloths, living or extinct (Fig. 7). Loss of enamel occurred after the acquisition of a typical xenarthran tooth form.

In the nine-banded armadillo, in addition to cheek teeth, it is worthy of note that four to five tooth buds form in the anterior region of the lower jaw (probably homologous to three to four incisors and one canine). The two to three most anterior tooth germs (incisors) do not develop beyond the bud stage and degenerate before birth (Martin, 1916). The canine and sometimes the last incisor develop normally and erupt. At birth they are covered by a thin enamel layer (10–20 μ m thick) (Fig. 12). In contrast to the cheek teeth they are not replaced. These descriptions support the cingulatan (and probably xenarthran) ancestor possessing a continuous dentition and the teeth being covered by enamel.

Given that previous descriptions of tooth development are from the beginning of the last century (and illustrated only with drawings) and only devoted to *D. novemcinctus*, new studies should be undertaken in representatives of the various armadillo sub-families, with particular attention being paid to enamel development and mineralization.

Adaptations to enamel reduction or loss in armadillos

Armadillos are generally myrmecophagous but some have a broadly omnivorous behaviour. Like anteaters, armadillos possess an elongate tongue ornamented with spiny projections and well supplied with sticky saliva that helps to catch insects. In armadillos, as in sloths (see below), teeth are hypsodont and hypselodont, and this feature

compensates for their upper surface being worn after the thin enamel cover is abraded. It is also clear that the presence of a well-developed and well-mineralized cement around the crown improves the resistance to wear. These adaptations compensate for enamel being reduced or lacking (see above). The trend toward enamel reduction is characteristic of xenarthran ancestors and there is evidence that this process started (for unknown reasons) after teeth had acquired the hypsodont/hypselodont condition. It is well known that this condition was acquired independently in numerous mammalian lineages and is a convergent adaptation for highly abrasive diets, notably in herbivorous species (Jernvall & Fortelius, 2002).

Sloths

Living sloths are represented by two genera, *Bradypus* (four species) and *Choloepus* (two species), but their extinct relatives were more diverse (mylodontids, megatheriids, megalonychids and nothrotheriids) (Gaudin, 2004). They have up to 10 teeth; incisors and canines are absent but the anterior cheek tooth is triangular in cross-section and canine-like (called caniniform). It is reduced in *Bradypus* and separated from the rest of the cheek teeth by a short diastema. The anterior surface of the lower caniniform meets the posterior surface of the upper, wearing against each other and continuously sharpening their edges.

To our knowledge there is no information on the presence of either vestigial teeth or tooth germs in the anterior region of the jaws of sloth embryos. If we consider what occurs in other taxa lacking teeth as adults (see below), tooth germs would be expected to form in this region, to be either resorbed or lost before birth. Even the possible presence of advanced tooth germs with formation of an enamel organ could be envisaged when considering the condition existing in armadillos, their closest toothed relatives. These hypotheses should be tested in the near future and, in particular, the presence of an enamel organ and ameloblast differentiation.

Adaptations to enamel loss in sloths

The dental condition observed in living sloths can be traced back to ground sloth species that lived in the Pleistocene (2 Ma) (Bargo et al. 2006). These ancient species probably dug for food and ingested abrasive soil particles, a condition that could have played a major role in selection of dental characteristics. Living sloths feed mainly on leaves but also include some fruits, buds and even small vertebrates in their diet. As described for armadillos above, the teeth of extant and extinct sloths are hypsodont and hypselodont (Fig. 8D). This well-developed hypsodonty is considered as an evolutionary adaptation to an abrasive herbivorous diet, either as a direct (abrasive grasses) or secondary (accumulation of grit or dust on plants) consequence, and to the absence of enamel, which would make the teeth

less durable and wear down faster (Vizcaino & De Iuliis, 2003; Bargo et al. 2006). To date, our poor knowledge of tooth structure and evolution in extinct and extant sloths cannot help us to date tooth loss in this lineage.

The absence of enamel, or its extreme reduction in the Folivora ancestors, has certainly been a strong developmental constraint that has influenced the morphology of sloth dentition in such a way as to reduce the number of possible morphological responses compared with those occurring in the other placentals.

Aardvark

The aardvark, *Orycteropus afer*, is the only living representative of Tubulidentata and lives in sub-Saharan Africa (Shoshani et al. 1988). Due to convergent evolution with Pholidota and Xenarthra (as a result of a similar diet), Tubulidentata were long considered to be related to these clades (Edentata; e.g. Anthony, 1934). However, biochemical and molecular analyses (Dene et al. 1983; Hallström et al. 2007) have demonstrated that the aardvark is closer to elephant shrews, hyraxes, elephants, manatees and dugongs, golden moles and tenrecs. These taxa, including the aardvark, now constitute the clade Afrotheria (Fig. 8). The extant representatives of these seven afrotherian lineages differ dramatically in their morphology but this can be easily understood as the roots of each lineage extend back into the late Cretaceous or early Paleocene (75–65 Ma; Springer et al. 2003). Indeed, all afrotherian lineages had probably diverged within the first 5 million years of crown afrotherian evolution (Murphy et al. 2007). This could explain why the sister group of Tubulidentata is not well defined, although some analyses indicate a preferred relationship with Paenungulata (which include hyraxes, elephants and sea cows) (Seiffert, 2007).

The aardvark has a heterodont and diphyodont dentition. In the adult there are five permanent teeth (two pre-molars and three molars, also called cheek teeth due to a tendency for molarization of pre-molars) located toward the back of the long slender, pig-like snout (Fig. 8A). There are no incisors or canines. The tooth structure is characteristic of Tubulidentata (Anthony, 1934). As the name implies, instead of the classical pulp cavity, the teeth have numerous (around 1000) tubular pulp cavities running vertically, forming hexagonal columns (prisms) perpendicular to the occlusal plane and held together by cementum. Each prism is pierced by a tubule that does not enclose Tomes' fibers but is homologous to the pulp cavity (Duvernoy, 1853). The teeth have no hard enamel covering and are hypselodont. Teeth are worn but the continuous growth compensates for abrasion.

In contrast to the condition observed in adults, embryos and juveniles have a complete dentition. Of the 40 tooth germs that form in the embryos, 20 are vestigial and 20 erupt. The front teeth, incisors and canines, are shed after

birth and are not replaced (Lönnerberg, 1906). The initial cusps of the cheek teeth are rapidly worn leaving two flat planes, with no indication of the presence of an enamel cover (Heuvelmans, 1939).

What the fossil record tells us

Afrotheria and Xenarthra are thought to have differentiated from a common ancestor during the Cretaceous and diverged approximately 100 Ma (see above). Afrotheria probably derive from primitive ungulates such as Condylarthra (Protoungulata), which lived from the late Cretaceous to the end of the Miocene. Aardvarks originated in Africa (Arambourg, 1959) and they first appear in the fossil record in the early Miocene of Kenya (20–16 Ma), a period during which the representatives of this group were more diversified and more widespread than today (fossils have been found in Europe and Asia) (Patterson, 1975). Morphologically, the genus *Orycteropus* has been remarkably conservative since at least the early Miocene (Pickford, 2005). All of these fossils already possess a reduced, tubulidentate dentition, which means that this structure appeared prior to the Miocene. The fossil record of Tubulidentata in older periods is scanty, however, and it is therefore difficult to date precisely when enamel was lost in this lineage (probably during the Paleocene, 60–20 Ma).

Most of the living members of the closest afrotherian lineage to Tubulidentata, Paenungulata (hyraxes, elephants and manatees and dugongs), possess two kinds of teeth, incisors and molars, separated by a diastema. The incisor in the upper jaw may be developed into a tusk that grows continuously or may be quite rudimentary. The molars erupt late in life, which is a uniting feature of the Afrotheria (Asher & Lehmann, 2008). None of these species form tubulidentate teeth or hypselodont molar teeth and enamel is present.

Adaptations to enamel loss in aardvark

Aardvark mainly feed on ants and termites, and can eat 50 000 insects in a night. They possess an elongated, flattened, protactile tongue (as long as 30 cm) that is covered by a strongly adhesive saliva that helps in catching prey (Kingdom, 1971). The continuous growth of the teeth compensates for the lack of enamel and subsequent abrasion, which is slowed down due to the presence of cementum around the tubular pulp cavities. Their strong gizzard-like stomach contains a high percentage of sand and small stones that help food digestion, as described in pangolins (see above).

The tubulidentate teeth and the absence of enamel are specific features of the lineage that led to aardvarks and it is difficult to understand how they evolved given the long period that separates the common afrotherian ancestor from the first Tubulidentata. It is agreed that the ancestor possessed a complete dentition (incisors, canines, pre-molars

and molars), a condition that is strongly suggested by the milk dentition in young aardvarks. We postulate that hypselodont/hypselodont teeth appeared in an ancestral afrotherian and were positively selected for as being more adapted to resist an abrasive diet. Somewhere in the Tubulidentata lineage the ability to form enamel was lost, resulting in an increased abrasion process. This was compensated by the positive selection of tubular pulp cavities held by cementum, which are more resistant to wear. The change of diet towards ants and termites was also less abrasive and the sticky tongue allowed such a specialization.

Platypus: a species lacking teeth secondarily during ontogeny

Platypuses belong, with the echidnas, to Monotremata (see above). *Ornithorhynchus anatinus* is the only living representative of the Ornithorhynchidae and is only present in eastern Australia. The platypus deposits eggs similar to those of lizards and snakes, possesses a keratinized beak similar to a duck's bill, has a beaver-like tail and otter-like feet, is venomous (males), is covered by hairs and the female feeds the babies with her milk. Juvenile platypuses have three-cusped molars that are shed and substituted in adults by keratinized pads (Fig. 13A,B).

Tooth development

In platypus embryos, a dental lamina develops in the two jaws but it disappears rapidly in the most anterior region (incisor field) of the upper jaw, whereas it is retained in the lower jaw (Green, 1937). Tooth germs form as follows: five incisors, one canine, two pre-molars and three molars in the lower jaw, whereas incisors are lacking in the upper jaw. The anterior teeth do not erupt and are small and fragile but an enamel covering is present (Lester & Boyde, 1986). A single pre-molar (which is the only tooth to have a replacement) and two tribosphenic (three cusps arranged in a triangle) molars erupt (Fig. 13C,D). All erupted teeth

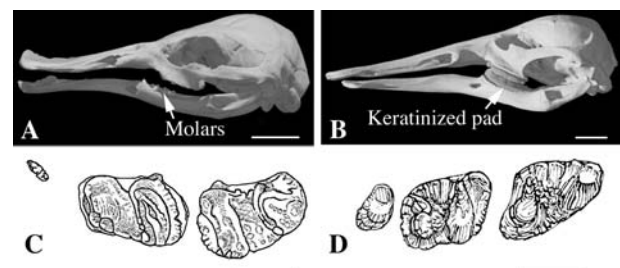


Fig. 13 *Ornithorhynchus anatinus*, the platypus. (A) Lateral view of the skull in a juvenile. (B) Lateral view of the skull in adult. (C) The three teeth (a small pre-molar and two molars) on the upper left maxilla of a juvenile. (D) The three opposite teeth on the lower left jaw. Bars: A and B, 1 cm; C and D, 1 mm.

have cusps and the dentine and enamel have the typical mammalian characters (Lester et al. 1987).

At hatching the platypus possesses both a bluntly conical os caruncle, i.e. a nodule of bone covered by a cornified epithelium (Hughes & Hall, 1998), and an egg tooth, consisting of a pulp cavity surrounded by dentine covered by an outer layer of presumptive enamel (Hill & De Beer, 1949). These tools help hatchlings to extricate themselves from the egg. The egg tooth is lost at 2 days post-hatching but the oscaruncle lasts for 11–14 weeks (Manger et al. 1998).

The three erupted teeth are lost before the young platypuses leave the breeding burrow and are not replaced. We do not know by which process these are lost in the absence of replacement teeth. Is there important remodelling of the supporting bones or is this lost related to the formation of thick heavily keratinized pads in their place (Fig. 13B)? The beak starts to form after hatching and reaches the adult size at 6 months (Manger et al. 1998). It consists of a large upper plate that arises from a cartilaginous extension of the maxillary and pre-maxillary bones, and a small lower plate originating from a cartilaginous extension of the mandibular bone.

What the fossil record tells us

Late Triassic/early Jurassic fossil remains (205 Ma) have probable monotreme affinities (Musser, 2003) but the oldest established extinct monotremes are from the early Cretaceous of Australia. They are *Teinolophos trusleri* (123 Ma) and *Steropodon galmani* (110 Ma), two steropodontids (closely related to modern ornithorhynchids), and *Kallikodon richieri*, a kallikodontid (100–104 Ma). Only a single molar was recovered in *T. trusleri* but the teeth were functional as indicated by wear facets in this two-rooted molar (Rich et al. 2001). *S. galmani* possessed at least three functional molars and the mandibular canal suggests the presence of a beak (Archer et al. 1985; Woodburne, 2003). One pre-molar and two molars were found on the dentary of *K. richieri* (Flannery et al. 1995). The oldest reported ornithorhynchid is *Obduron insignis* from the late Oligocene of Australia (25–15 Ma). Its close Miocene relative, *O. dicksoni*, had a wide flattened beak, two upper and two lower pre-molars, and two upper and three lower molars (Musser & Archer, 1998) (Fig. 7B). It is worth noting that fossil ornithorhynchids had a functional dentition in adults; the retained teeth were well developed despite the presence of a beak. During further evolution of this lineage, the tooth roots were progressively reduced and the oral epithelium took more and more importance in food processing through the formation of horny pads (Musser & Archer, 1998).

Adaptation to tooth loss in adult platypus

Platypuses are semi-aquatic animals and food is collected in the water by means of the beak. The diet consists mainly

of invertebrates, the cuticle of which is broken up between the tongue and the horny grinding plates and shearing ridges on the upper and lower jaws. The edges of the beak are ornamented with sharp transverse ridges that help in catching prey. Epithelial ridges on the palate aim to secure and cut the prey in the absence of teeth. The tongue is ornamented by keratinized spines, which work against the palate to help mastication (Griffiths, 1978). All of these features are secondary adaptations that were selected during evolution as they improve food uptake and processing. They compensate for tooth loss that was, again, probably related to a relaxed functional constraint when the beak was positively selected.

Conclusion

The main question that led to us preparing this review was how to reconcile the view of a strong functional pressure acting on vertebrate teeth, on the one hand, with the existence of numerous tetrapod species lacking teeth or enamel, on the other hand. Is it so easy to lose teeth or enamel? After our comparative analysis we can answer yes, provided a pre-adapted 'tool' is present prior to tooth loss, otherwise tooth or enamel loss would be lethal. In fact in all tetrapod lineages examined herein, the existence of a pre-adapted tool appears to explain tooth loss (or enamel loss). Four tools were identified: beaks (birds, turtles, echidnas and platypus), elongated sticky tongues (toads, pangolins and anteaters), baleens (baleen whales) and hypsodonty/hypselodonty (armadillos, sloths and aardvark). At the beginning of the story (i.e. in a common ancestor) the tool was of secondary importance for food uptake and/or processing. However, it was certainly of some help, which explains its natural selection. Subsequently, the tool increased in importance by allowing more efficient food uptake, enhancing the ability of its owner to occupy a specific ecological niche. The positive selection of the secondary tool then relaxed the functional constraints on the teeth and they disappeared either suddenly or progressively.

To date, we have only a single explanation of what could be part of the story of tooth loss in birds: a displacement of the appropriate location of the competent (dental) epithelial cell population resulting in signalling molecules missing their target. However, tooth induction can be resurrected as has been described in the chick *ta²* mutant. The mechanism responsible for this displacement is, however, unknown. Is there a mutation leading to an incorrect patterning of the oral epithelium? Such questions still remain unanswered and need to be revisited. Concerning the other beaked species (turtles and monotremes), we can only speculate that a similar mechanism was probably acting.

For the other species we could think that the relaxed pressure on teeth favoured the occurrence of various mutations. Mutations leading to tooth loss could concern

a local disruption in the epithelial/mesenchymal interactions. Such a local disruption could explain loss of teeth in a particular region of the jaw only [e.g. the loss of palatine teeth in turtles, or teeth of the lower jaw in anurans and, more generally, reduction of tooth location during evolution, first in the number of bone supports concerned and then in the same bone support (Huyseune & Sire, 1998)].

Loss of enamel while conserving dentine is another interesting character that concerns Xenarthra and Tubulidentata. It appears that hypsodont/hypselodont teeth was the primary condition that occurred early in evolution in these unrelated lineages. These features are considered pre-adaptations, which allowed further adaptation to plant feeding and to an abrasive diet. They confirm the limited alternatives for tooth adaptation to an abrasive diet and an apparent facility for teeth to acquire hypsodonty/hypselodonty from a normal structure. Molars with numerous tubular pulp cavities held with cementum, open roots (hypselodonty) and lacking enamel have also been described in a basal mammal from the late Jurassic, which shows no relation to Xenarthrans (Luo & Wible, 2005). The combination of continuously growing teeth and a lack of enamel thus again goes hand in hand.

But why was enamel lost? It seems clear that such a loss was not a consequence of hypsodonty/hypselodonty as numerous, unrelated herbivorous species (e.g. some rodents and horse) have a similar adaptation but have conserved enamel. Enamel reduction was probably a first step toward complete loss of enamel but we do not know the process leading up to this reduction then loss. It is noteworthy that tooth loss (or enamel loss) is consecutive to the positive selection of a similar but convergently acquired tool (e.g. an elongated, sticky tongue) and to similar food adaptation in four unrelated lineages, i.e. feeding on small prey, ants and termites: anteaters in Xenarthra, pangolins in Pholidota, aardvark in Afrotheria and echidna in Monotremata. In addition, engulfing such prey does not require teeth but numerous secondary adaptations to help digestion, e.g. a gizzard-like stomach (pangolins and aardvark) containing small stones like in a bird's gizzard.

Eventually, although it seems common (under appropriate conditions) to lose or modify teeth (a reminiscence of an ancestral plasticity), it appears very difficult to remove the evolutionary imprints as illustrated by the presence of vestigial teeth in all toothless taxa; these rudimentary tooth germs bear witness to the ancestral condition of each species and are of considerable help for the evolutionary biologist.

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