Resolving the evolution of the mammalian middle ear using Bayesian inference

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Abstract

Background: The minute, finely-tuned ear ossicles of mammals arose through a spectacular evolutionary transformation from their origins as a load-bearing jaw joint. This involved detachment from the postdentary trough of the mandible, and final separation from the dentary through resorption of Meckel’s cartilage. Recent parsimony analyses of modern and fossil mammals imply up to seven independent postdentary trough losses or even reversals, which is unexpected given the complexity of these transformations. Here we employ the first model-based, probabilistic analysis of the evolution of the definitive mammalian middle ear, supported by virtual 3D erosion simulations to assess for potential fossil preservation artifacts.

Results: Our results support a simple, biologically plausible scenario without reversals. The middle ear bones detach from the postdentary trough only twice among mammals, once each in the ancestors of therians and monotremes. Disappearance of Meckel’s cartilage occurred independently in numerous lineages from the Late Jurassic to the Late Cretaceous. This final separation is recapitulated during early development of extant mammals, while the earlier-occurring disappearance of a postdentary trough is not.

Conclusions: Our results therefore suggest a developmentally congruent and directional two-step scenario, in which the parallel uncoupling of the auditory and feeding systems in northern and southern hemisphere mammals underpinned further specialization in both lineages. Until ~168 Ma, all known mammals retained attached middle ear bones, yet all groups that diversified from ~163 Ma onwards had lost the postdentary trough, emphasizing the adaptive significance of this transformation.

Keywords: Australosphenida, Middle ear detachment, Meckel’s groove, Postdentary trough, Theria

Abbreviations: BPP, Bayesian posterior probability; DMME, Definitive mammalian middle ear; MMEC, Mandibular middle ear of cynodonts; PMME, Partial mammalian middle ear

Background

One of the most famous and complicated transformations in vertebrate evolution is the origin of the mammalian middle ear bones (ectotympanic, incus and malleus) from load-bearing post-dentary elements (angular, articular and quadrate) during the evolution of synapsids [1]. This is also the oldest and best-documented example of developmental recapitulation of an evolutionary transformation [2–5], as the final separation of the middle ear from the dentary through disappearance of Meckel’s cartilage occurs in mammalian development as in evolution [6, 7] (Fig. 1).

The freely suspended middle ear of extant adult mammals is considered a derived pattern termed the definitive mammalian middle ear (DMME) [8]. A DMME has also been reported in other extinct mammal lineages, including multituberculates and several cladotheres (close therian relatives) [9]. Although a detached middle ear has been considered a defining feature of living mammals [10, 11], there is mounting evidence that the DMME was in fact acquired independently in monotremes and therians [7, 10, 12–15], which suggests strong selection for a sensitive auditory system adapted to high-frequency sounds [16, 17]. However, current hypotheses on the convergent evolution of middle ear
bones are complex and controversial, partly because of a lack of phylogenetic resolution and partly because the interpretation of the fossil evidence is difficult [7, 18].

Some clues as to the sequence in which the mammalian middle ear evolved come from two additional patterns of the mammalian middle ear in the fossil record [7]. Both of these include a permanent connection of the middle ear bones to the dentary. The most plesiomorphic is the mandibular middle ear of cynodonts (MMEC; Fig. 1a), in which the middle ear bones are fully attached to the posterior part of the dentary, and are housed in a postdentary trough and angular fossa. In contrast, the “partial mammalian middle ear” (PMME; Fig. 1b), also termed “transitional mammalian middle ear” [9], has the middle ear bones connected to the dentary by an ossified or possibly cartilaginous Meckel’s cartilage [6, 7, 9, 19]. In the PMME, the ectotympanic ring and the malleus have no direct contact with the mandible, which therefore lacks a postdentary trough and angular fossa [20]. The PMME has been observed in eutriconodont and spalacotheroid mammals. It is also expected more generally across fossil mammals that lack a postdentary trough, but retain a prominent Meckel’s groove [6, 19]. The PMME is found in early stages of middle ear development in extant mammals (Fig. 1c), thus reinforcing the impression that it represents an intermediate evolutionary condition [6].

Hypotheses of independent acquisitions of the DMME are based on (1) differences between monotremes and therians regarding the origins of the jaw-opening muscles [13]; (2) hypotheses of a retention of the angular, articular and prearticular bones directly attached to the lower jaw in Cretaceous monotremes (particularly Teinolophos trusleri), inferred from the suspected presence of a postdentary trough to accommodate these bones, and (3) the polyphyletic distribution of the DMME across Mesozoic mammals, as inferred from the absence of a postdentary trough and Meckel’s groove.

Unfortunately, the historical emphasis on whether middle ear bone migration occurred independently in therian and monotreme ancestors has often overshadowed a broader taxonomic perspective on this evolutionary
process. Phylogenetic and coding variation among recent studies allow for up to seven such independent migrations, based on the loss of the postdentary trough, along lineages leading to monotremes, euhamarimyidans, multituberculates, *Hadrocodium*, *Fruitafossor*, eutriconodonts, and trechnotheres (derived symmetrodonts and cladotheres). In Fig. 2 we provide a summary of parsimony inference for postdentary trough transformations based on four recent phylogenies [21–24] with varying taxonomic composition and character coding. Most remarkably, in three of these analyses the parsimony solution requires one or two reversals back to postdentary attached middle ear bones. Only the phylogeny of the most recent dataset [22] unambiguously supports losses of the postdentary trough (four in total), although that study did not include the basal symmetrodont *Kuehneotherium* which could add an extra loss of the postdentary trough connection, depending to its phylogenetic position.

Each of the proposed independent origins of the DMME is to some extent controversial or ambiguous. In some cases the phylogeny is controversial, such as for the relationships of haramiyidans (e.g. [22, 25]). Character coding is also contentious. Notably, there has been increasing debate about the apparent absence of a postdentary trough in the early mammaliaform *Hadrocodium*, which now appears to be a juvenile specimen with limited preservation of poorly ossified areas having erased evidence of a postdentary trough [20, 26–28]. The controversies surrounding character coding and preservation artifacts also affect the interpretation of key Cretaceous fossil monotremes. There has been a long debate regarding the presence of a postdentary trough and Meckel’s groove in *Teinolophos* and *Steropodon* [14, 18, 29, 30]. Comprehensive reviews now tend to support the absence of a postdentary trough from both species [18, 29, 30]. The suggested Meckel’s groove in

**Fig. 2** A summary of parsimony inference for postdentary trough transformation based on four recent phylogenies [21–24] with varying character coding. The reconstructions minimize the number of postdentary trough losses (blue) and regains (red). Note that taxon sampling differs between the studies (a–d). Placements of monotreme and therian mammals are represented by the platypus and ocelot, respectively. In Krause et al. [23] the symbols I and II indicate independent origins for australosphenidian taxa.
Steropodon has been alternatively explained as “pre-depositional breakage” of the only known specimen [31].

Historically, two main alternative scenarios [19] have been described for the apparent multiple origins of the DMME during evolution: (1) the DMME was present in the common ancestor of Mammalia, but some clades (eutriconodonts and spalacotheroids) re-evolved a Meckel’s cartilage middle ear attachment to the mandible through paedomorphosis, and hence a PMME; or (2) the DMME was absent in the mammalian common ancestor and evolved once in extant monotremes, for a second time in multituberculates, and again in the ancestors of both marsupials and placentals [7, 19]. The latter scenario does not clearly identify whether the common ancestor of modern mammals had a PMME or a MMEC, and resolution of these paths towards the DMME depends on phylogenetic relationships and the status of characters in several fossil taxa [7].

As noted, both of the above scenarios are partially based on controversial evidence and have also exclusively relied on parsimony analyses that minimize rather than estimate the number of transformations. Here, we infer the phylogenetic history of middle ear migration in mammals, and test which scenario better explains the evolution of the DMME, using Bayesian methods to reconstruct the ancestral states of the two key characters associated with middle ear detachment. We also revisit the possibility of preservational artifacts influencing the scoring of Meckel’s groove and the postdentary trough in morphological matrices, using simulations of erosions in virtual 3D models of monotreme lower jaws.

Results
Virtual reconstruction of possible diagenetic artifacts and character re-scoring for the postdentary trough and Meckel’s groove
We found that the presence of Meckel’s groove and its disappearance can be traced in development for the platypus and echidna (see also [6]). However, we could not identify any developmental evidence for the presence of a postdentary trough at any stage. Both our virtual erosion and thresholding of 3D reconstructions of a juvenile platypus dentary with recently detached middle ears showed that an anatomical feature resembling a substantial Meckel’s groove could be easily produced from a slight lingual indentation (Fig. 3d and e). This indentation could be a filled-in trace of Meckel’s groove, or a ventral “lip” below the developing mandibular canal (Fig. 3a). Notably, except for our most extreme erosion scenario (Fig. 3e), this artifact could be produced without compromising the anatomical detail in other parts of the dentary, suggesting that erosion of a dentary in the fossil record might not be noticed. This suggests that the coding for the degree of development of Meckel’s groove (0: Well developed; 1: Weakly developed; 2: Vestigial or absent) that has been used as a character in phylogenetic analyses may in practice be difficult to distinguish based on sub-optimally preserved, eroded fossils, especially for the state ‘weakly developed’ and ‘vestigial or absent’. Using this observation we re-scored Meckel’s groove in both datasets as: (0) Meckel’s groove developed (including former character state 1: “weakly developed”), (1) Meckel’s groove not developed or vestigial. A developed Meckel’s groove, in this case, is taken as evidence of a PMME. Using this observation and based on new fossil information [21, 29, 30], we recoded the presence/absence of the postdentary trough in Hadrocodium and several australosphenidans (Ausktribosphenos, Bishaps, Steropodon, and Teinalophos). These recoded datasets were then used for the phylogenetic analyses to trace the origins of the DMME.

In the first dataset [22], all euharamiyidans, multituberculates and cimolodontans were re-scored as Meckel’s groove “not developed or vestigial”. For eutriconodonts, spalacotheroids, and cladotheres that were coded as
Meckel's groove “Weakly developed” (see [22]), this character was re-coded as “Meckel's groove developed” in our analyses. Similarly, Meckel's groove was re-coded for eutriconodonts and cladotheres in the second dataset [21] to match our re-coding for the first dataset [22].

For Hadrocodium, the postdentary trough has been scored as “absent”, and Meckel's groove as “weakly developed” [21] or “vestigial or absent” [22] respectively. These characters were coded as “present” and “developed” respectively in our analyses based on new evidence that suggests Hadrocodium is a juvenile specimen with limited preservation of poorly ossified areas largely erasing evidence of its postdentary trough [20, 26–28], and Meckel's groove.

For Asfaltonylos we scored the postdentary trough as “present”, and Meckel's groove as “uncertain”. Although Meckel's groove has been coded as “vestigial or absent” [22], and as “well developed” [21], this character cannot be observed in the only known specimen in which this region of the dentary is damaged [32]. This new interpretation also changes the codification of the curvature of Meckel's sulcus as “uncertain”. Postdentary trough was coded as “present” in our analyses.

For Aukstribosphenos, the postdentary trough has been considered as “present” [33], or “absent” [21], and Meckel's groove as “well developed”. The dentary of Aukstribosphenos has been illustrated including a remnant of Meckel's groove that extends until the m3 [34]. However, similar remnants were observed when thresholding was applied to virtually reconstructed models of juvenile platypus (O. anatinus) to simulate removal of layers of bone, to mimic diagenetic bone attrition (Fig. 3). The postdentary trough seems to be absent based on one illustration (see [34]). We concurred with the interpretation of the postdentary trough as “absent” [21], but recoded Meckel's groove as “developed”, based on our virtual reconstructions and thresholding comparison of a juvenile platypus that lost Meckel's groove during development. This shows the possibility of the character in Aukstribosphenos being an artefact. In an alternative analysis the postdentary trough was coded as “uncertain”, but the topologies of the trees and posterior probabilities were not affected (Additional file 1: Table S1).

For Bishops, we concurred with the interpretation for scoring the postdentary trough as “absent” [21], but we considered Meckel's groove as “developed”. Although the postdentary trough has been coded as present based on previous datasets [22], the description of the illustrations of the taxon [35] show no trace of the presence of postdentary trough.

In Steropodon, the postdentary trough might be stated as “absent” and Meckel's groove as “weakly developed” [21], contrary to recent datasets (see [22, 33]) in which these characters were coded as “present” and “well developed”, respectively. The illustration of the holotype of this taxon [31] shows no clear presence of a postdentary trough. Although the authors mentioned that a depression in the lingual side of the dentary extends until m3, they are unsure whether this represents a Meckel's groove or pre-depositional breakage. We expect this feature results from “pre-depositional breakage", but coded the character as “uncertain”. We scored the postdentary trough alternatively, as either “absent” or “uncertain” in our analyses.

For Teinolophos the postdentary trough has been considered “present” [14], or “absent” [21, 33]; and Meckel's groove as “well developed” [22], or “weakly developed” [21]. We consider the postdentary trough to be “absent” and Meckel's groove “developed” (following [29]). Based on our virtual reconstructions and because the postdentary trough is clearly identifiable in taxa for which this structure is present, we concur with Teinolophos lacking a postdentary trough.

**Phylogenetic ancestral state reconstructions**

Our Bayesian ancestral state reconstruction based on the first dataset [22] suggests that the postdentary trough was lost independently, once in australosphenidans, and again in the common ancestor of Fruitafossor and Theriiformes (eutriconodonts, euharamiyidans, multituberculates, spalacotheriids, and cladotheres), regardless of the placement of the pseudotribosphenic shuotheriids (Fig. 4a and c). Only when all haramiyidans and multituberculates were forced together was the postdentary trough lost a third time (Fig. 4b), in the common ancestor of multituberculates and advanced haramiyidans (euharamiyidans). The ancestral condition of the postdentary trough does not re-evolve on the tree after the postdentary bones are detached from the dentary and the postdentary trough is lost (Fig. 4). The Bayesian posterior probability (BPP) for the ancestor of australosphenidans retaining a postdentary trough (Additional file 1: Table S1) is high (BPP > 0.92), except when Shuotheriidae was forced outside Mammalia (~0.63), while being very low for the ancestor of stem and crown monotremes (BPP < 0.01). Similarly, the posterior probability of the ancestor of Mammalia possessing a postdentary trough was high (BPP > 0.85), except when Shuotheriidae was forced outside Mammalia (~0.59).

The analyses of the second dataset [21], which does not include multituberculates or haramiyidans, also favours the postdentary trough being lost independently only twice, once in monotremes and once in the common ancestor of Theriiformes, regardless of the position of Kuehneotherium (Fig. 4e and f). All analyses (Additional file 1: Table S1) support the ancestor of australosphenidans retaining a postdentary trough (BPP > 0.92), and the ancestor of monotremes...
Fig. 4 Bayesian inference of the loss of the postdental trough among mammals (blue), based on the re-coded datasets (a-d [22]; e-f [21]). Each tree summarizes the following analyses: a, b) Shuotheriidans unconstrained (allowed to group with australosphenidans); b) constraining monophyly of the multicuspate haramiyidans and multituberculates; c) constraining Shuotheriidans to fall outside Mammalia; d) excluding cheek teeth characters; e) forcing Kuehneotherium outside Mammalia; f) allowing Kuehneotherium to fall freely in the phylogeny. Postdental trough is lost twice in all the analyses except in (b) when the earliest haramiyidans (e.g. Haramyavia) are forced together with Euharamiyida and multituberculates. The analyses excluding cheek teeth characters (d) place primitive haramiyidans outside Mammalia, with euharamiyidans and multituberculates both falling on the therian stem lineage—negating any requirement for a third loss of the postdental trough in mammals. In (d) Haramyavia is not included but likely forms a clade with Megaconus as observed in several of these trees (not b).
lacking a postdentary trough (BPP = 0.99). The presence of a postdentary trough at the origin of crown mammals in this dataset was also high (BPP > 0.94) in all the analyses.

Meckel’s groove is more variable in analyses of both datasets, with a DMME evolving from a PMME several times since the earlier loss of the postdentary trough. Meckel’s groove is well developed in adults at the crown ancestor of mammals (BPP > 0.95), then is independently lost in adults of recent australosphenidans (crown monotremes), multituberculates, and in several cladotheres, including along the stem lineages of both marsupials and placentals. In all analyses the ancestor of australosphenidans (and the ancestor of stem monotremes) was found to have a developed Meckel’s groove during adulthood (BPP > 0.84) (Additional file 1: Table S1).

Discussion

Our Bayesian analysis supports a simple, directional two-step scenario of mammalian middle ear evolution. The first step includes the development of postdentary bones from the dentary to form a partial mammalian middle ear (PMME); this occurred convergently in the northern hemisphere ancestors of therians and the southern hemisphere ancestors of monotremes (Fig. 4). Second, the transition from a PMME to a definite mammalian middle ear (DMME) occurred multiple times, including at least three cases of independent evolution within extant mammals (in monotremes, metatherians and eutherians), between the late Jurassic (~163 Ma) and the Late Cretaceous (~80 Ma). In addition, our results clarify the ancestral mammalian condition, suggesting that the ancestor of all mammals possessed a plesiomorphic, fully attached mandibular middle ear of cynodonts (MMEC).

The parallel evolution of the MMEC to the transitional PMME in therians and monotremes appears to represent a continuation of a mammaliform trend towards reduced postdental jaw articulation, decoupling the auditory and feeding systems of the skull [36]. We show that this transformation only occurred twice, if basal haramiyidans and multituberculates are placed as members of the therian stem lineage as supported by our analyses excluding cheek teeth. This is a far simpler scenario than the up to seven losses and gains of the post-dental trough in our parsimony-based reconstructions of recently published coding and phylogenies (Fig. 2). MMECs in taxa such as Haramiyavia, Kuehneotherium and Shuhotherium are thus plesiomorphic retentions, and not reversals from the DMME or the PMME as suggested by previous studies [7, 9].

In addition, our ancestral state reconstruction substantially pushes back the inferred date of MMEC loss in the australosphenidans (monotreme ancestors). This was considered to have occurred relatively recently, in the common ancestor of living monotremes, while our analysis suggests that this transformation occurred at least 120 million years ago in the ancestors of the Australian australosphenidans (monotremes and ausktribosphenidans, and Bishops).

Intriguingly, our results suggest a fast transition from a MMEC to a PMME: all known mammals up until ~168 Ma retained fully attached middle ear bones, while all mammalian families that diversified from ~163 Ma onwards had lost the postdentary trough. The rapidity of the MMEC/PMME transition emphasizes the key adaptive significance that has been attributed to this transformation [37].

Our analyses unambiguously identify the PMME as a phylogenetic stage preceding the evolution of a DMME [9, 19], regardless of whether Meckel’s cartilage was cartilaginous or ossified (as observed as a possibly plesiomorphic trait [1, 6, 7, 19] in some adult eutriconodonts and spalacotheroids). This conclusion is surprisingly robust to the phylogenetic position and presence or absence of postdentary troughs in controversial taxa (e.g. Hadrocodium and Teinolophos), which have previously represented debated obstacles in the interpretation of mammalian middle ear evolution (see Introduction).

Our results demonstrate that Bayesian inference represents a powerful tool for inferring ancestral character states, especially in complicated cases such as the evolution of the mammalian middle ear. Bayesian ancestral state reconstruction is standard in analyses of extant taxa, but has rarely been used with the inclusion of fossils, which in turn offers the possibility to overcome extinction biases that affect inferences from extant taxa alone [38].

Aside from the more nuanced approach of modelling evolutionary processes compared to the binary decision-making of parsimony, Bayesian posterior probabilities can provide predictive power to inform character state assignments in debated fossils. This is perhaps best illustrated with the use of ancestral reconstruction to provide us with a prior probability for a postdentary trough in Hadrocodium, which until recently had been considered to have a DMME. Consistent with recent finds of a MMEC in this species, Bayesian inference analysis retrieved a high probability (BPP > 0.82) of MMEC presence.

The coding of presence or absence of Meckel’s cartilage has caused some controversy, particularly in Hadrocodium and australosphenidans such as Teinolophos, which appear to have a Meckel’s groove imprint (see [39]). Our virtual simulation of erosion showed that we could easily create artifacts that are near-identical to the suspected Meckel’s groove in Steropodon. Another
issue is the possibility that the presence of Meckel's groove in juvenile fossils might indicate a later time of detachment during development, rather than partial attachment of the middle ear to the dentary in adulthood. This suggests that the current coding of different states of Meckel's groove (absent or vestigial/weakly developed/strongly developed) incurs the risk of overestimating the presence of Meckel's groove and affecting the estimation of when the DMME evolved. Coding Meckel's groove as present only where it is unambiguously well-developed (see Methods) therefore appears to be the best strategy.

Conclusions
This study used Bayesian analysis to provide the first probabilistic evidence that the most recent common ancestor of mammals originally possessed a cynodont-like middle ear with a loadbearing function. Detachment from the postdentary trough and functional uncoupling of the auditory and feeding apparatus occurred independently in therian and monotreme ancestors. The final detachment of the middle ear through disappearance of Meckel's cartilage was rapid in some lineages and long delayed in others, matching the overall pattern of rapid diversification of mammals during this time. Functional and ecological comparisons in lineages with different detachment timeframes might provide information as to whether detachment was driven by selective pressures [13, 40], or represents a side-effect of developmental change elsewhere [7].

Methods
Virtual reconstruction of possible diagenetic artifacts and character re-scoring
To test whether the apparent presence of a postdentary trough and a weakly developed or vestigial Meckel's groove in stem monotremes (e.g. *Teinolophos*, *Steropodon*) might be artifacts of poor fossil preservation, we used μCT scans of the dentary and middle ear bones of a juvenile platypus with a detached middle ear, but faint scarring on the lingual side of the dentary (Fig. 3). We used the Materialise Mimics software digitally removed the outer bone layers of the 3D reconstructed models obtained from the μCT scans by increasing the thresholding values of the model mesh, and also using the “erosion” tool, which removes the external layer of “3D pixels” (voxels). This was to simulate a situation where diagenetic erosion of bone could produce an artifact that simulates the presence and degree of development of Meckel's groove (Fig. 3) in fossils, as has been suggested for *Steropodon* (see Introduction). Based on the information from the virtual reconstructions and comparisons with new evidence available for stem fossil monotremes [29, 30] and *Hadrocodium* [25, 26, 28], we re-coded the states of the postdentary trough and Meckel's groove for several taxa (see Results) in two morphological datasets [21, 22]. Each of these matrices has a different taxonomic focus, but together they cover all of the relevant, known lineages for evolution of the DMME. We also used these μCT scans (Fig. 3) to identify any developmental evidence for the presence of a postdentary trough in extant monotremes.

Phylogenetic analyses
For both datasets [21, 22], we explored the ancestral condition of the two osteological characters that have mainly defined the debate on single versus multiple origins of the DMME: the postdentary trough and Meckel's groove (see Introduction). We used Bayesian inference within MrBayes 3.2.4 [41, 42] to infer the phylogeny using each recoded dataset, and ancestral state reconstruction for the postdentary trough and Meckel's groove characters. Two independent analyses were run on three Markov chains for 5,000,000 generations, with trees sampled every 5000 generations for the second dataset [21], and for 10,000,000 generations, with trees sampled every 10,000 generations for the first dataset [22]. These chain lengths ensured clade frequency convergence between runs and estimated sample sizes for substitution parameters were >100 (using Tracer v1.5 [43]).

Both matrices were analysed with five partitions: 1. mandibulodental, 2. postcranial, 3. cranial characters, 4. Meckel's groove, and 5. the postdentary trough. The Mk v + gamma evolutionary model allows for different evolutionary rates among characters, and was employed for the multi-character partitions. The Mk v model was used for Meckel's groove and the postdentary trough, given that rates across sites (e.g. gamma) models are redundant for single character partitions.

Although the first dataset [22] includes 114 taxa, we only included 84 of these in our analyses. We excluded highly incomplete taxa (<12.5% of the characters; e.g. *Albertatherium*, *Aegialodont*) because they potentially bias branch length estimation [44], and hence phylogeny and ancestral state reconstruction. It is important that rate models for middle ear and Meckel's groove evolution reflect evolvability during the Mesozoic. So in addition, the 25 modern placentals and marsupials were also excluded, because canalisation in these groups during the Cenozoic has apparently limited the plasticity of these traits and their inclusion would artifactually deflate inferred evolutionary rates for tracing their history [45]. Many other taxa are also <50% complete, but are less concerning because they are largely complete for the anatomical region partitions in which they are included. Similarly, for the second matrix [21] we excluded 3 taxa, *Drescheratherium* (low completeness), the modern *Erinaceus*, and *Didelphis*. 
Alternative ancestral state reconstructions for each of the matrices were performed to cover different phylogenetic possibilities that stem primarily from arguments over molar cusp pattern homologies (e.g. [24, 46]), which are correlated among many characters, and therefore, could strongly mislead phylogeny if they were incorrectly assigned. Of particular importance are controversial molar cusp homologies between australosphenidans, boreosphenidans and the pseudotribosphenic shuotheriids, and between multituberculdates and haramiyidans. We note also that the evolutionary rate inferred for cheek tooth characters is 2.5-fold higher than among other anatomical partitions, and high evolutionary rates increase phylogenetic signal erosion, and hence, susceptibility to non-phylogenetic biases, such as functional correlations [47]. To check for phylogenetic anomalies we ran both datasets [21, 22] without cheek tooth characters (Fig. 4d; Additional file 1: Figure S1 and S2). For the first dataset, [22], the “pseudotribosphenic” shuotheriids fell outside crown mammals and placement of the enigmatic, fossorial Fruitafossor was clarified as grouping closer to therians than to monotrems (Fig. 4d; Additional file 1: Figure S2). The second dataset [21] did not include Fruitafossor or the pseudotribosphenic shuotheriids (purple) was clarified as grouping closer to therians and multituberculdates (e.g. [25]). For the second dataset [21], which does not include the multicuspate taxa, two analyses were performed: the first allows the unstable Kuehneotherium to fall freely within the phylogeny, while the second forces it outside Mammalia, in agreement with our analysis in which cheek teeth characters were excluded.

In recognition of these findings and phylogenetic uncertainties, the ancestral state reconstructions were performed on the full set of characters with (1) Shuotheriidae allowed to fall freely within the phylogeny (e.g. with australosphenidans), (2) constraining Shuotheriidae to fall outside Mammalia, and (3) constraining monophyly of the multituberculdate haramiyidans and multituberculdates (e.g. [25]). For the second dataset [21], which does not include the multicuspate taxa, two analyses were performed: the first allows the unstable Kuehneotherium to fall freely within the phylogeny, while the second forces it outside Mammalia, in agreement with our analysis in which cheek teeth characters were excluded.

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Availability of data and materials
The information supporting the conclusions of this article are included within the article and its additional files.

Authors’ contributions
Conceived and designed the experiments: HER-C, MWP, VW. Performed the experiments: HER-C, MWP. Analysed the data: HER-C, MWP, VW. Wrote the paper: HER-C, MWP, VW, SW. All authors gave final approval for publication.

Competing interest
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
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Additional file

Additional file 1: Figure S1. Bayesian inference phylogeny using Rougier et al.’s [21] dataset when cheek tooth characters were excluded. This analysis clarified the placement of the early “symmetrodont” Kuehneotherium outside the mammalian crown. Figure S2. Bayesian inference phylogeny for Luo et al.’s [22] dataset, but with cheek tooth characters excluded. The “pseudotribosphenic” shuotheriids (purple) fell outside crown mammals and the placement of the enigmatic, fossorial Fruitafossor (yellow) was clarified as grouping closer to therians than to monotrems (green). Table S1. Posterior probabilities. Description: A. Shuotheriidae allowed to group with australosphenidans. B. Constraining Shuotheriidae to fall outside Mammalia. C. Constraining monophyly of the multicuspate haramiyidans and multituberculdates. D. Forces Kuehneotherium outside Mammalia. E. Allows the unstable Kuehneotherium to fall freely in the phylogeny. 1: Alternative analyses including Aukstribosphenos and Sterapodon with postdentary trough coded as absent. (DOCX 231 kb)
possess an internal mandibular trough for

Asfaltomylos patagonicus

Haramiyavia

and Teinolophos trusleri

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