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Bayesian Morphological Clock versus Parsimony: An insight into the relationships and dispersal events of postvacuum Cricetidae (Rodentia, Mammalia)

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ABSTRACT

Establishing an evolutionary timeline is fundamental for tackling a great variety of topics in evolutionary biology, including the reconstruction of patterns of historical biogeography, coevolution and diversification. However, the tree of life is pruned by extinction and molecular data cannot be gathered for extinct lineages. Until recently methodological challenges have prevented the application of tip-dating Bayesian approaches in morphology-based fossil-only datasets. Herein, we present a morphological dataset for a group of cricetid rodents to which we apply an array of methods fairly new in palaeontology that can be used by palaeontologists for the analysis of entirely extinct clades. We compare the tree topologies obtained by traditional parsimony, time-calibrated and non-calibrated Bayesian inference phylogenetic approaches and calculate stratigraphic congruence indices for each. Bayesian tip-dated clock methods outperform parsimony in the case of our dataset, which includes...
highly homoplastic morphological characters. Regardless, all three topologies support the
monophyly of Megacricetodontinae, Democricetodontinae and Cricetodontinae. Dispersal
and speciation events inferred through Bayesian Binary Markov chain Monte Carlo and
biodiversity analyses provide evidence for a correlation between biogeographic events,
climatic changes and diversification in cricetids.

Keywords: Cricetidae, Miocene, Bayesian tip-dating, morphological clock, parsimony,
STRAP, palaeobiogeography, palaeoecology, palaeobiodiversity
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The fossil record is our only resource for providing a deep-time perspective of ecosystem processes and, therefore, for understanding the dynamics that have shaped our current biota. It is increasingly clear that fossil data are fundamental to infer species diversification, particularly extinction rates (Mitchell et al., 2019). Despite the fact that morphological phylogenetics have been increasingly marginalised in the last decades, morphological data remain the only available information to reconstruct evolutionary scenarios and reconcile the fossil record with molecular trees (Wright 2017; Lee and Palci 2015). The importance of morphological phylogenetics for dating rigorously the tree of life is now widely recognized and has been bolstered by recent methodological developments (Hunt and Slater 2016; Lee and Palci 2015; Wright 2017). Recent developments applying Bayesian methods using fossil taxa as tips, what is called the morphological clock, have been revitalizing the use of morphological data to elucidate the dynamics of evolution over time and across the tree of life (Varela et al. 2019; Simões et al. 2018, 2020a). While these analyses are usually carried out with data from both fossils and extant taxa, a very interesting application of this methodology is that it can be employed with data from extinct clades only, which adds to the palaeontologist’s toolbox another method of reconstructing the evolution. This makes it now possible to compare phylogenies of extinct taxa obtained by means of evolutionary models with those resulting from maximum parsimony, which is the most widely applied method for analysing morphological data.

Time provides palaeontologists with a unique perspective on phylogeny. A few methods to integrate stratigraphic data with parsimony analyses were already available in the 90’s (Wagner, 1995; Fisher, 1994). However, palaeontologists had to devote considerable time and effort in the process of calibrate them manually because of the lack of user-friendly software facilitating time-calibration of cladograms. Moreover, depending on the number of taxa included in the dataset palaeontologists have to infer the distribution of morphological
characters without the inclusion of temporal data, with the subsequent loss of information. Parsimony was until recently the only way for palaeontologists to analyse their morphological fossil datasets. However, since the introduction of Bayesian tip-dated phylogenetic methods, which were first applied with uniform tree prior (Ronquist et al. 2012) and then with fossilized birth-death (FBD) tree priors (Stadler 2010; Heath et al. 2014), the inclusion of stratigraphic data into phylogenetic analyses boomed. The development of tip-dating with FBD tree priors that allow fossil species to be included as terminal tips has been particularly useful for palaeontologists. Recent numerical methods such as PaleoTree (Bapst, 2012, 2014) or STRAP (Stratigraphic Tree Analysis for Palaeontology) (Bell and Lloyd 2014) allow phylogenies resulting from both parsimony and Bayesian analyses of fossil taxa to be dated. The package STRAP allows besides to assess their stratigraphic congruence (Bell and Lloyd 2014; O’Connor and Wills 2016; King and Beck 2019; King 2021). So, the development of all these comparative methods have promoted the revival of phylogenetic analysis incorporating stratigraphic data and the testing of different techniques of phylogenetic reconstruction using morphological data (Bell and Lloyd 2014; Sansom et al., 2018; King 2021). This provides palaeontologists with a golden opportunity to expand considerably their research toolkit.

Over the years, the development of numerical, parsimony and Bayesian methods to determine trees has resulted in an intense study of certain clades (Benton 2015). However, the most striking aspect of rodent palaeophylogenetics is the low ratio of phylogenetic hypotheses to species number. In fact, few comprehensive morphological phylogenetic analyses have been carried out for these mammals and none has been performed applying morphological clock methods. Nonetheless, rodents have an excellent fossil record and their dentition provides an ideal dataset to characterize phenotypic variation and calibrating divergence time analyses. The present work rests on cricetids, which are an important group of rodents because they include representatives involved in the first radiation of modern rodents in Eurasia during the Miocene.
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They probably gave rise to other important lineages, such as arvicolines (voles), cricetines (hamsters) and even murines (mice). The Miocene comprises the first phase in the development of modern terrestrial ecosystems. This is a time interval that witnessed enormous geographical and environmental changes and during which the mammal record experienced a major turnover and successive dispersal events. Cricetids possibly appeared in China in Middle Eocene times (Tong 1992). After an initial radiation in Central Asia, the group expanded in Western Europe just after the “Grande Coupure” (Stehlin, 1910), when the Turgaï Strait dried out and a land bridge between Europe and Asia was established. They prospered in Europe during the Oligocene but primitive cricetids disappeared before the arrival of the more derived, typical Miocene ones. The time interval devoid of cricetids has been called the “cricetid vacuum” by Daams and Freudenthal (1989). Post-vacuum cricetids are known worldwide and under various morphologies since Miocene times, but their evolutionary history is poorly known. The biogeographic relationships and phylogenetic affinities of cricetid species with one another need to be studied in a time-scaled phylogenetic framework. This work aims to clear up the phylogenetic relationships and estimate divergence times of postvacuum cricetids to elucidate their origin and the dispersal events they underwent by applying parsimony and Bayesian methods to an array of important extinct lineages of Miocene cricetids from Europe, the Middle East and Asia. Moreover, the different approaches we use are assessed on the basis of the stratigraphic coherence of the resulting topologies in order to arrive at an informed opinion on which method is most likely to yield the most accurate results in the phylogenetic analyses of palaeontological (morphological) dataset.

MATERIAL AND METHODS

Material

The systematic study presented below is based on the examination of original specimens listed in Supplementary Tab. S1 provided in the Supplementary Information available in the
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First, second and third lower molars are designated as m1, m2 and m3, respectively and first, second and third upper molars as M1, M2 and M3, respectively.

Maximum Parsimony Analyses

Implied weights maximum parsimony analysis—A total of 82 phylogenetically informative dental characters from 74 taxa have been coded. The data matrix (Supplementary file S1 available on Dryad) has been built using Mesquite 3.04 (Maddison & Maddison 2009) and the analysis ran in TNT v.1.5 (Goloboff et al. 2003) using the new technology search algorithms and the implied weighting algorithm (Supplementary file S2 available on Dryad) (Goloboff et al. 2008). New technology search algorithms are recommended for large datasets because they allow the sampling of trees from a broader spectrum of local optima (Goloboff et al. 1999). Tree searches have been carried out using 1,000 initial trees by random addition sequences with 100 iterations or rounds for each of the four NTS algorithms: sectorial search, ratchet, drift and tree fusing. The analysis has been performed with K = 10 and collapsing all branches with support = 0. K values larger than the default (3.0) are more accurate to perform analyses for large datasets (Goloboff et al. 2017). The number of suboptimal trees to be retained was set at 10 and the relative fit difference at 0.1. The final output trees (22 MPTs+suboptimal trees) have been filtered for all the most parsimonious trees (MPTs). A total of 3 MPT with a length of 487 steps, a Consistency Index (CI) of 0.261 and a Retention Index (RI) of 0.765 have been obtained and have been used to calculate the strict consensus tree (488 steps) (Supplementary Figure S1 available on Dryad). Character 1, length of the M1, has been treated as additive. Branch support have been estimated through two complementary indices: Bremer Support (Bremer 1994) and Relative Bremer Support (Goloboff and Farris 2001).
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Equal weights maximum parsimony analysis—The analysis has been also carried out with TNT using equal weights. Tree searches have been conducted as performed for the implied weighting parsimony analysis. The final output trees (440 MPTs+suboptimal trees) have been filtered for all the most parsimonious trees (MPTs). A total of 375 MPT with 479 steps, a Consistency Index (CI) of 0.265 and a Retention Index (RI) of 0.767 have been obtained and have been used to calculate the strict (530 steps) and majority (488 steps) consensus trees (Supplementary Figures S2 and S3 available on Dryad).

Bayesian inference analyses

Analyses have been carried out using Mr. Bayes v.3.2.6 (Ronquist et al. 2012) and the BEAST2 package (Bouckaert R., et al. (2014) using the CIPRES Science Gateway v.3.3 (Miller et al., 2010).

Non-clock Bayesian analysis—The analysis has been performed with Mr. Bayes v.3.2.6 (Ronquist et al. 2012). The morphological dataset (Supplementary file S1 available on Dryad) has been analysed with the MkV model (Lewis, 2001) under the $\gamma$ model, 30 million generations and four independent runs. Convergence of independent runs is assessed by an average standard deviation of split frequencies of 0.0148, by an average potential scale reduction factors (PSRF) of 1 for all parameters and an effective sample size (ESS) greater than 200 for each parameter.

Time-calibrated relaxed-clock Bayesian inference analysis—The analysis has been performed in BEAST2 package (Bouckaert R. et al. 2014) (Supplementary file S3 available on Dryad). The Mkv model (Lewis 2001) has been used, with a gamma distribution with four rate categories to account for rate variation across sites. The prior on the gamma shape parameter is an exponential distribution. Following Simões et al. (2020a), the morphological dataset has
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been analysed using the fossilized birth-death tree model with sampled ancestors (FBD-SA) (Stadler 2010), under relaxed-clock models, 30 million generations and four independent runs with the initial 30% of samples removed as “burn-in”. The relaxed-clock model is the independent $\gamma$ rate relaxed-clock model, which is a continuous uncorrelated relaxed-clock model using a gamma distribution to assess clock rate variation across lineages (Simões et al 2018). It is compatible with the fossilized birth–death tree model. The base clock rate has been given an informative prior that derived from the undated Bayesian inference analysis: the median value for tree height (TL) in substitutions from the entire posterior trees sample (TL=9.9825) divided by the age of the tree (37.6 Ma), which is based on the median of the distribution for the root prior. So, the prior is: 9.9825/37.6=0.265. In contrast to Mr. Bayes, BEAST 2 does not allow different modelling strategies for how extant taxa are sampled and assumes a random sampling strategy (Simões et al, 2020b), which is unimportant in studies that do not include extant species. Our calibrations have been based on tip dating, which accounts for the uncertainty in the placement of fossil taxa and avoids the issue of bound estimates for node-based age calibrations (Ronquist et al. 2016, Simões et al. 2018). It has been substantiated that sampling fossil ages instead of fixing them to the midpoint within their stratigraphic age range avoids biases in divergence time estimations (Barido-Sotani et al., 2019). Thus, the fossil ages used in this work for tip dating correspond to the uniform prior distributions on the age range of the stratigraphic occurrence of the fossils. The age of the root has been set with a soft lower bound. The minimum age of the root corresponds to the oldest age for the oldest fossil belonging to Eucricetodon and the maximum root has been set at 41.2 Ma (Middle/Late Eocene boundary), which is the maximum soft age for the clade Eucricetodontinae. Conditioning on rho has not been possible because the data set of the analysis only contains extinct taxa so we have conditioned on the root and on sampling (Supplementary file S3 available on Dryad). An effective sample size (ESS) greater than 200 for the main parameters assesses the convergence of independent runs.
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Stratigraphic congruence

To assess the stratigraphic congruence of the parsimony and Bayesian inference-derived phylogenies resulting from our analyses we used the package STRAP for R with the default number of permutations (1000) (Bell and Lloyd, 2014). We have obtained the following stratigraphic fit indices: MIG (minimum implied gap (Norell and Novacek, 1992; Norell et al. 1992)), SCI (Stratigraphy Consistency Index (Huelsenbeck 1994)), RCI (Relative Completeness Index (Benton and Storrs 1994)), GER (Gap Excess Ratio (Wills 1999)), MSM* (modified Manhattan Stratigraphic Measure (Pol, 2001)) and Wills’ modifications of GER (GERt and GER*, Wills et al. (2008)). MIG provides the sum of the branch lengths excluding tip durations (the sum of the ghost ranges), SCI provides the proportion of nodes that are stratigraphically congruent in a tree, RCI provides the measure of the extent of observed ranges of taxa and the sum of the ghost ranges, MSM* corresponds to MIG for the maximally stratigraphically consistent possible tree divided by the actual MIG and GER to MIG minus the best possible stratigraphic fit, scaled by the contrast between the best and worst fit values (Lloyd and Bell, 2014; Wright and Lloyd, 2020)). SCI, GER, GERt, and MSM* scale between 0.0 (least congruent) and 1.0 (most congruent) (O’Connor and Wills 2016; Wright and Lloyd, 2020). Additionally, the program provides the significance test for those measurements (p.SCI, p.RCI, p.GER, and p.MSM*) as well as a combined fit and significance test measure (GER*and GERt) (Bell and Lloyd 2014). The p-values indicate the probability of the null hypothesis, which indicates random tree topology. So, very small p-values will indicate a significantly good fit to stratigraphy (Bell and Lloyd 2014).

Biogeographic Inference using Reconstruct Ancestral State in Phylogenies

In order to infer historical biogeography we have applied RASP 4.2 (Yu et al. 2015), a method to reconstruct ancestral geographical distributions using a combination of phylogenetic and geographical information. Bayesian Binary Markov chain Monte Carlo (BBM) method for
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ancestral state has been performed on the Maximum Clade Credibility Tree (MCCT) in RASP 4.2 (Yu et al. 2015). BBM calculates the probabilities of ancestral ranges using the probabilities of each unit area generated by MrBayes (Yu et al. 2015). The Markov chain Monte Carlo (MCMC) chains were run for five million generations and ten independent runs. The state was sampled every 100 generations. Fixed JC + G (Jukes-Cantor + Gamma) were used for BBM analysis. The distribution range of all species of cricetids included in this work has been divided into eight geographic areas, which have been carefully chosen according to dissimilarities in their faunal composition that provided evidence for palaeobiogeographic or palaeoenvironmental differences between them. They correspond to: A (southwestern Europe), B (Central Europe), C (Greece), D (Anatolia-Caucasus), E (west central Asia), F (Arabian Peninsula), G (southern Asia) and H (east central Asia).

Biodiversity rates

The combination of cladistic and biodiversity analyses highlights the phenomena of speciation, extinction, and diversity changes in a given group over time (Stigall 2010, López-Antoñanzas et al. 2015). This allows establishing the timing of biodiversity crises and, thus, deducing possible causes.

Per-capita rates for speciation ($\hat{q}$), extinction ($\hat{p}$), and diversity change ($d$) have been calculated (Supplementary Tab. S2 available on Dryad) following the equations given by Foote (2000) according to which:

\[
\hat{p} = -\frac{\ln(N_{bt}/(N_{bt} + N_{ft}))}{\Delta t}
\]

\[
\hat{q} = -\frac{\ln(N_{bt}/(N_{bt} + N_{bL}))}{\Delta t}
\]

\[
d = \hat{p} - \hat{q}
\]

where $N_{bt}$ indicates the number of species that cross both the upper and lower interval boundaries, $N_{ft}$ the number of species that originate within the interval and cross over the
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upper interval boundary and NbL the number of species that cross the lower interval boundary,
but become extinct during the interval and $\Delta t$ the duration of the interval $t_1-t_0$.

Rates of biodiversity change (R), speciation (S), and extinction (E) have been calculated
(Supplementary Tab. S2 available on Dryad) with the following equations (Stigall 2010):

$$R = \frac{\ln N_1 - \ln N_0}{\Delta t}$$

$$S = \frac{\ln (N_0 + o_0) - \ln N_0}{\Delta t}$$

$$E = \frac{\ln (N_0 + o_0) - \ln N_1}{\Delta t}$$

where $N_0$ is the initial number of species in a clade at time $t_0$, $N_1$ the number of species in a clade at time $t_1$, $o_0$ the number of speciation events during the interval $t_1-t_0$, and $\Delta t$ the duration of the interval $t_1-t_0$.

All rates have been calculated each 0.5 Ma using the phylogenetically corrected species ranges obtained from the MCCT. Values calculated from the first and last intervals have been excluded from the analysis to remove edge effects, following the criterion of Stigall (2010).

RESULTS AND DISCUSSION

Tree Topology

Maximum Parsimony and tip-dated and undated Bayesian trees (Figs. 1, 2, Supplementary Figs. S2-S4 available on Dryad) show three large clades that correspond to the rodent subfamilies Megacricetodontinae, Democricetodontinae and Cricetodontinae. Both in, the majority consensus tree resulting from equal weights maximum parsimony and undated Bayesian analysis (Supplementary Figures S3 and S4 available on Dryad) show Megacricetodontinae and Cricetodontinae more closely related to one another than either one is to Democricetodontinae. However, some studies have evidenced that downweighting characters according to their homoplasy (applying implied weighting) improves morphological data sets, particularly those that are highly homoplasmatic, and produces more resolved and accurate trees than standard equal weights (Goloboff et al., 2008; Smith, 2019).
As a well-recognized problem when working on fossil rodents is the supposedly high homoplasy of dental characters, discussion on parsimony results will be based on the topology of the tree resulting from the implied weighting analysis. Both implied weighting parsimony and tip-dated Bayesian analyses (Figs. 1 and 2) supported Cricetodontinae as the sister group to the clade consisting of Megacricetodontinae plus Democricetodontinae, which are sister clades of each other. These results place Democricetodontinae and Megacricetodontinae closer to one another than either is to the Cricetodontinae, which is in line with the phylogenetic hypothesis proposed by Flynn (2009).

Implied weights maximum parsimony—The topology of the tree shows three main clades: Megacricetodontinae, Democricetodontinae and Cricetodontinae.

The calibration of the strict consensus tree (Fig.1) shows that Democricetodontinae (node 83) originated circa 23.7 Ma. They share two non-exclusive synapomorphies (Supplementary Tab. S3, Supplementary Figure S1 (for numbers of each node in the tree) available on Dryad), which are the presence of a double protoloph on M2 (29(0→1)) and the metacone absent or included in crest on M3 (50(0→2)). All the taxa included in this clade also share the exclusive synapomorphy of having an m1 with anteroconid in lingual position (60(0→1)).

Megacricetodontinae (node 138) originated approximately 23.7 Ma. They share three exclusive synapomorphies. The first one consists in having a divided anterocone with the lingual anterocone smaller than the labial one on M1 (5(0→2)). The second one rests in having m3 quite reduced (with an Lm1/Lm3 ratio between 1.4-1.6 mm) (67(0→2)) and the third one in having the entoconid small but still distinct on m3. The taxa included in this clade (except for Megacricetodon andrewsi) also share the non-exclusive synapomorphy of having M3 with usually incomplete central atoll (45(0→1)). Megacricetodontinae comprise two clades. The
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first one regroups the species originating from node 137 (Shamalina tuberculata (Punjabemys
downsii, Sindemys shewanensis)). It is supported by two non-exclusive synapomorphies: the
presence of a small enterostyle isolated from the protocone and the lack of a well-developed
anterolophid on m3 ((18(0→1); 73(0→1)). The second clade consists in all the species that
originate from node 145 (Aktaumys dzhungaricus and more derived megacricetodontines).

They share the non-exclusive synapomorphies of having on M1 the lingual anteroloph weak or
absent (10(0→1)), a long mesoloph (19(1→0)) and the posteroloph that continues beyond the
point where it meets the metalophule (23(0→1)), and in having the protoloph slightly anteriorly
directed on M3 (41(0→1)).

Cricetodontinae are the species originating from node 91. They share the two exclusive
synapomorphies of having the anterocone divided in two parts of similar size on M1 (5(0→1))
and a distinct anterior ectoloph (12(0→1)). They are also sustained by five non-exclusive
synapomorphies: a long posteroloph that continues beyond the point where it meets the
metalophule (23(0→1)), a large single anteroconid on m1 (54(0→1)) and the lingual
anterolophid weak or absent on the lower molars (59(0→2), 68(0→1), 73(0→1)).

The split of Cricetodontinae from the sister clades Democricetodontinae plus
Megacricetodontinae is set up circa 23.8 Ma. The basalmost taxa within Cricetodontinae are
Cricetodon versteegi and, one node up, C. goklerensis, both from the Early Miocene of Turkey.
The remaining taxa are split into two very asymmetrical branches. One is represented by
plesiomorphic species of Cricetodon from the Early Miocene of Turkey (C. fikreti + C.
trallesensis), whereas the other form a much larger group that includes all the remaining species
of the subfamily. At the base of this clade a polytomy involving Cricetodon fengi,,
Mixocricetodon dehmi plus Cricetodon sonidensis, and a clade comprising all the more derived
species. The latter splits into two lineages. The smaller one includes the species originating
from node 103 with C. wanhei as most basal and C. aliveriensis plus C. kasapligili as most
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derived and *C. tobieni* in an intermediate position. This clade is supported by the non-exclusive
and unambiguous synapomorphy of having a complete central atoll on M3 (a reversal). This
character is only shared by the most plesiomorphic species of *Cricetodon* (*C. verstegi*) and
most of *Democricetodon* spp. The larger branch is constituted by a succession of lineages, the
most basal of which is that of the most plesiomorphic true European species, *C. meini* plus *C.
aureus*. One node up *C. orientalis* from the Middle Miocene of China branches and then *C.
soriae* from the Middle Miocene of Europe. The more derived species divide first in a clade of
Middle Miocene European species of *Cricetodon* (*C. jotae, C. jumaensis, C. albanensis, C.
bolliegeri, C. nievei, C. engesseri*). They share the two non-exclusive synapomorphies of
having the lingual anteroloph of M2 poorly developed (26(0→1)) and M2 elongated
(28(0→1)). The remaining species, originating from node 93, consist in a succession of
European (*C. sansaniensis, C. hungaricus, C. caucasicus*) and Turkish (*C. pasalarensis*) taxa
that splits (node 114) to give rise to *Byzantinia* and *Hispanomys* during the Middle Miocene.
The bifurcation in which *Hispanomys* originated (node 113) includes the late Middle Miocene
species of ‘*Cricetodon*’ from Central Europe (‘*C.* klariankae, and ‘*C.* venczeli’), which should
be considered as belonging to *Hispanomys*. Similarly, the bifurcation from which *Byzantinia*
evolved (node 116) contains the Middle Miocene ‘*Cricetodon*’ *fandli, ‘*C.* candirensis* and ‘*C.*
cariensis* basally, which may be reinterpreted, according to this analysis, as plesiomorphic
representatives of *Byzantinia*.

*Tip-dated Bayesian analysis*—The divergence between Cricetodontinae and
Democricetodontinae + Megacricetodon is set up at approximately 28.2 Ma, and that between
the sister clades Democricetodontinae and Megacricetodontinae circa 25.9 Ma.

Democricetodontinae: The topology of the tree (Fig. 2) shows two sister clades. The first
includes two sister species: *Democricetodon sui* from the Early Miocene of China (21.9-21.16
Ma), which is the oldest record of *Democricetodon* in Asia (Maridet et al., 2011), and *Primus*
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*microps* from the Early Miocene of Pakistan (circa 21-23 Ma) (Bruijn et al. 1981). The second one shows at the base the two plesiomorphic species of *Democricetodon* (*D. anatolicus* and *D. doukasi*) from the Early Miocene of Turkey, which insert sequentially on the steam, leading to more derived *Democricetodon* spp. *Democricetodon franconicus*, from the Early Miocene of Aliveri (Greece), is the most basal species of this clade. It is followed successively by *D. mutilus* and *D. gracilis*, which reveal the first entrance of Democricetodontinae in the central-western European bioprovince (circa 18.27 Ma). *D. gracilis* is sister species to a clade including mostly Asian species, providing evidence for an early Middle Miocene migration of *Democricetodon* from Europe towards China at approximately 16.1 Ma.

Megacricetodontinae: The topology of the tree (Fig. 2) shows two main clades within this subfamily of rodents. The first one includes an array of Early Miocene species of disputed origin (*Vallaris zappai*, (*Sindemys shewanensis*, *Punjabemys downsi*)). *Punjabemys downsi* from the Early Miocene of Pakistan was originally considered a member of Megacricetodontinae (Lindsay, 1988) and then as belonging to the Myocricetodontinae by Wessels (2009) together with *Vallaris zappai* and *Sindemys shewanensis* from the Early Miocene of Turkey and Pakistan, respectively. According to the topology of the tree, these taxa are either the basalmost representatives of Megacricetodontinae or could as well be considered as their sister group. Be that as it may, they are sister group to the clade that includes the other megacricetodontines, the basalmost taxon of which is *Shamalina tuberculata* from the Early Miocene of Saudi Arabia. This result agrees with Lindsay (1994), who thought that *Shamalina* could have given rise to *Megacricetodon*, but oppose the suggestion of Wessels (2009), who included this taxon within the Myocricetodontinae. *Shamalina* may have led to two main lineages. One consists of the sister species *Aktaumys dzhungaricus* and *Megacricetodon beijiangensis* and points to a dispersal event from Saudi Arabia towards Kazakhstan and China circa 21.28 Ma. The other lineage, which is more speciose, shows at its base *Megacricetodon*
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*hellenicus* from the Early Miocene of Aliveri, Greece (Oliver and Peláez-Campomanes, 2014), which provides evidence for the first entrance of *Megacricetodon* into the Aegean-Anatolian region at approximately 19.5 Ma ago. One node further up two clades substantiates two independent migration events. One involved *Megacricetodon* aff. *collongensis* (circa 18 Ma), the most primitive species of the typical central European “*Megacricetodon bavaricus* group” proposed by Oliver and Peláez-Campomanes (2013), and the other *M. andrewsi* and *M. primitivus* at approximately 17 Ma. These results agree with Oliver and Peláez-Campomanes (2016), who inferred at least three migration events in Europe for early *Megacricetodon* forms on the basis of important morphological differences.

Cricetodontinae originated in Turkey at approximately 25 Ma (Fig. 2). The basalmost taxon is *C. versteegi* and *C. goklerensis* is one node up in the clade. Both taxa come from the lower Miocene of Turkey (de Bruijn et al. 1993; Joniak et al. 2017). *Cricetodon goklerensis* is sister species to a large group that includes all the remaining species of the subfamily, the basalmost clade of which includes the most plesiomorphic species of *Cricetodon*, mostly from Turkey and China, and documents the first entrance of *Cricetodon* from Turkey into China at circa 19.4 Ma. The split of this basal clade and the clade comprising more derived species of Cricetodontinae is set up at 22.3 Ma. Within the latter, the most basal taxa are represented by the Turkish Early Miocene species *C. fikreti* and *C. trallesensis*. *Cricetodon trallesensis* diverges from the main clade approximately 20 Ma ago and led to two clades. The first one includes practically all European Middle Miocene species of *Cricetodon* and reveals the first entrance of the Cricetodontinae into Europe circa 17.77 Ma. The presence inside this group of the Asiatic species *C. orientalis* provides evidence for a second dispersal of Cricetodontinae into China, this time from Europe, which took place approximately 16.8 Ma. The second clade comprises the most derived species of Cricetodontinae. A sequence of the three most basal taxa from the Middle Miocene of Central Europe and Turkey (*C. hungaricus, C. caucasicus, C.

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pasalarensis) insert sequentially on the steam, leading to two important lineages. One leads to
Hispanomys and includes the late Middle Miocene species of “Cricetodon” from Central
Europe (C. fandli, C. klariankae and C. venzclí), which should be considered to belong to
Hispanomys, according to this analysis. Similarly, the bifurcation from which Byzantinia
evolved contains, in a basal position, the Middle Miocene Cricetodon cariensis and C.
candirensis, which would be plesiomorphic representatives of Byzantinia.

Bayesian analyses versus Parsimony

Tree topologies of undated Bayesian and equal weights maximum parsimony searches are
more similar to each other than either is to the implied weighting parsimony and tip-dating
Bayesian trees. In the same way, the topologies of implied weighting parsimony and tip-dating
Bayesian trees are more similar to each other than either to the two other topologies. Despite
the fact that the topologies of implied weight parsimony and tip-dated Bayesian trees (Figs. 1,
2) are similar as far as the relationships between large clades are concerned, they differ in the
placement of some taxa amongst the lineages inside these clades. This issue holds particularly
true for the clades that are not comprehensively sampled (e.g., Democricetodontinae and
Megacricetodontinae vs Cricetodontinae) and those that are weakly supported. Because of the
differences recovered between the topologies resulting from different methods, stratigraphic
congruence indices have been calculated to assess how well each of these phylogenies fits with
the chronostratigraphy. Our results show high levels of stratigraphic congruence for all metrics
for parsimony (equally weights and implied weights) and both Bayesian (undated and tip-
dating) methods (Supplementary Tab. S4 available on Dryad). As expected, the maximum
credibility tree recovered by tip-dating has significantly higher stratigraphic congruence than
the majority consensus trees resulting from undated Bayesian and parsimony analyses
(Supplementary Tab. S4 available on Dryad). The majority consensus tree resulting from
implied weighting parsimony (similar to the strict consensus tree) has higher stratigraphic
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congruence than the majority consensus trees recovered from equal weights parsimony and undated Bayesian searches. Recent studies provided evidence that the inclusion of stratigraphic age data in tip-dating impacts the topology of the tree, particularly when the datasets include incompletely scored taxa or taxa having weak character support (King 2021). The differences we found between the topologies of the tip-dated Bayesian tree and the undated Bayesian and parsimony trees are probably due to the presence of highly homoplastic characters in our dataset. Taken into account that the topology of the undated phylogenies shows some anomalous results (e.g. derived position of the oldest democricetodontines) and that time-scaling these topologies (Fig. 1) results in improbable long ghost lineages for a large number taxa, we consider that the topology obtained via the tip-dating Bayesian analyses is most likely to be more accurate with our kind of morphological dataset (fossil rodent teeth). Therefore, the discussion on palaeogeography and biodiversity will be based on the Maximum Clade Credibility Tree (MCCT) resulting from the tip-dating Bayesian analysis (Fig. 2). Interestingly, Lee and Yates (2018) also found that tip-dating approaches were better able to elucidate the relationship of some crocodilian groups that showed extensive convergent adaptations.

Palaeobiogeographic framework

During the Miocene, Democricetodontinae, Megacricetodontinae and Cricetodontinae were successful and exhibited a wide distribution, ranging geographically from western Europe to eastern Asia. Their evolution is complex and involves indigenous speciation phenomena and numerous intercontinental dispersal events. According to our results most immigration events involved Turkey (Fig. 3). Democricetodontinae may have originated in Turkey approximately 24.8 Ma and entered Asia about 23.9 Ma. The results of this work also provides evidence of a probable Turkish origin for European Democricetodontinae with a first arrival in central and western Europe circa 18.3 Ma. The delayed entrance of this group into Europe was probably
due to the geographical isolation of the Dinarian-Anatolian Island (Neubauer et al. 2015). A
Middle Miocene dispersal of *Democricetodon* from Europe to China has been evidenced in this
work and set up around 16.1 Ma.

A more comprehensive phylogenetic analysis that includes the Myocricetodontinae is
needed to elucidate if *Vallaris zappai* and more derived taxa belong to the Megacricetodontinae
or are, actually, closer to the Myocricetodontinae. Without this information, two scenarios are
plausible. The first one points to a Turkish origin of Megacricetodontinae circa 23.3 Ma,
whereas in the second one they would originate from either Turkey or Saudi Arabia around
22.6 Ma. The eastwards dispersal to Kazakhstan and western China is set circa 21.3 Ma and
that to the Aegean-Anatolian area (where they are recorded in Aliveri and Kēsekoy) circa 19.5
Ma. Two independent dispersal events took place then from this area towards the West first
and then toward eastern China, and Central Europe at approximately 18 Ma. A subsequent
dispersal from western Asia to the Aegean-Anatolian area occurred approximately one million
years later (circa 17 Ma) (Fig. 3).

Cricetodontinae are supposed to have originated from Turkey at 25 Ma, where several
lineages differentiated, and then passed into Greece at the earliest at 20 Ma and to eastern Asia
around 19.4 Ma (Fig. 3). Greek cricetodontines from the Early Miocene of Aliveri are not
closely related to the European Middle Miocene cricetodontines but to the Turkish Early
Miocene ones. In fact, at this time, Aliveri may not have belonged to the European region but
to the Aegean-Anatolian one, which was only intermittently in contact with Europe. Most of
the European cricetodontines originated from an independent Turkish lineage that dispersed
into the south-western part of the continent circa 18.5 Ma. Two dispersal events towards central
Europe occurred at 16.4 Ma and 15.5 Ma and two additional ones towards the Aegean-
Anatolian area and western China are inferred at 16.8 Ma and 16.2 Ma. On the basis of our
analysis, it turns out that the Late Miocene cricetodontines are not closely related to the Middle
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Miocene European ones. In fact, one important dispersal event at about 16.4 Ma from Turkey towards Europe together with an independent evolution in Turkey led to a later development of two dynamic centres of speciation in southwestern and central Europe (Hispanomys) and Anatolia (Byzantinia).

Palaeoclimatic context

During the Oligocene and the Early Miocene, the East Antarctic Ice Sheet was relatively unstable with periods of growth and others of decline (Miller et al., 2020). At the Eocene/Oligocene Transition and the beginning of the Oligocene (33.9-32 Ma), a large deglaciation triggered successive sea level rises up to 50 m. On the contrary, sea-level drops were recorded in early Late Oligocene times, at approximately 28 Ma (Oi1), and at the beginning of the Miocene, circa 23 Ma (Mi1) (Miller et al., 2020). Interestingly, this sea-level fall at the Oligocene/Miocene boundary coincides with a severe restriction of the marine connections between the Indian Ocean and the Mediterranean Sea (Hüsing et al., 2009). This could have opened dispersal land routes that could have allowed the migration of primitive megacricetodontines, such as Shamalina tuberculata from Saudi Arabia towards Kazakhstan and western China (~23.5 Ma). The dispersal of Democricetodon from Turkey to Pakistan and China (~23.9 Ma) might have been elicited by the same phenomenon. Until the beginning of the Miocene Climatic Optimum (17-15 Ma), eustatic sea level changes indicates the presence of a moderate to large ice sheet in Antarctica (Miller et al. 2020). Low sea level that allowed dispersal of faunas are indicators of cool intervals, whereas higher sea levels represents warmer periods. Climate variations are expected to have played a major role in shaping the diversity of small mammals in general and rodents in particular. So, confronting the timing of shifts of diversification and climatic perturbations can provide lines of evidence regarding the influence of the latter on the former. The analysis of the shifts of diversification across the phylogeny of Cricetodontinae, the most comprehensive subclade in this work, is shown in Fig. 4.
Regrettably, the poor fossil record of Cricetodontinae in Turkey between 23 Ma and 19 Ma, where and when this group originated and started diversifying, does not allow a robust interpretation of our results during this interval of time. The MCO (~17-15 Ma) was a warm period that is mostly characterized by high sea level and reduced ice volume. Since a little before the Miocene Climatic Optimum (18.5 Ma) and nearly throughout all of it, the rodent community of cricetodontines experienced a period of stability and success, during which the biodiversity was slightly but constantly increasing (Fig. 4). This increase in biodiversity resulted from a continuous decrease in the extinction rate of these rodents together with high speciation rates that remained nearly constant. At approximately 16.0 Ma, an interruption of the Miocene Climatic Optimum is evinced by a sea level fall of ~40 m (Mi2) (Miller et al. 2020), which would have allowed the second dispersal of *Democricetodon* from westwards to eastwards. This cooling event (Mi2) seems to have promoted a drop in the speciation of Middle Miocene European cricetodontines, which was fairly high up to that point (Fig. 4), while their extinction rate continued to decrease. After the MCO and during the Middle Miocene Climatic Transition (~15-13 Ma), three events of cooling and sea-level fall at 14.8 (Mi2a), 13.8 (Mi3) and 12.8 Ma (Mi4) took place that culminated with the establishment of the “permanent East Antarctic Ice Sheet” (Miller et al. 2020). After the Mi2 event, speciation and extinction rates tended to decrease continuously. However, our results show high extinction and speciation rates that are coupled with the Miocene cooling events Mi3 and Mi4 (Fig. 4). The most prominent turnover is detected during the Mi3 event (circa 13.8 Ma). Interestingly, this event seems to have been particularly impactful in southern Europe, where most of the cricetodontines lived at this time. Actually, a sudden and important drop in temperatures was evidenced in Spain after analysing oxygen isotopes of mammal teeth enamel (Domingo et al. 2009, 2012). The evolutionary radiation of various endemic lineages of *Byzantinia*, in Anatolia, and *Hispanomys*, in Europe, took place during the Mi3 event (Fig. 4). In these areas, parallel lineages showed similar trends towards an increase of size, a reduction of length of the
third molars and the development of a complete backwards paracone spur (Rummel 1998).

This suggests a specialisation toward food that includes more abrasive and fibrous plants, indicating in turn increasingly open environments. When Byzantinia and Hispanomys flourished, their potential competitors such as the Middle Miocene cricetodontines were driven to extinction. At the beginning of the Late Miocene, at about 11.5 Ma (Mi5), another cooling event associated with an increase in the ice volume in Antarctica took place (Cook et al., 2008).

This event is linked to a faunal turnover within the Cricetodontinae in which a drop in biodiversity caused by elevated extinction rates (oldest taxa belonging to Byzantinia) was followed by a second radiation of the most derived species of Hispanomys and Byzantinia.

It is evident that climatic events impacted the evolutionary history of the Cricetodontines. However, given that some of the oldest taxa belonging to Byzantinia are not dated precisely, the high extinction and speciation rates that we found in Mi5 may be exaggerated, whereas those corresponding to Mi 4 might be minored.

CONCLUSION

Our analyses evidence that time-calibrated Bayesian searches yield trees that have higher stratigraphic congruence compared with trees from undated Bayesian and parsimony searches. Majority and strict consensus trees resulting from implied weights parsimony show the second best stratigraphic concordance values. They are followed by equal character weighting parsimony and non-clock Bayesian searches. So, according to our results, if stratigraphic congruence is taken as a proxy for phylogenetic accuracy, then time-calibrated Bayesian inference analysis (morphological clock) is probably the more accurate method for analysing morphological characters with a high degree of homoplasy (such as those of rodent teeth). However, more studies comparing morphological parsimony and time-calibrated and not time-calibrated Bayesian methods are needed to conclude which of these approaches deal
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better with the morphological characters provided by fossil rodents, which are mainly based
on the dentition and considered highly convergent.

The Miocene is marked by obvious manifestations of climatic changes, which turn out to
have been very influential in the evolution of animal life. Cricetids arrived at different times
in different areas around the Mediterranean and became extinct asynchronously as well. Our
results suggest that the Miocene cooling events, particularly Mi2, Mi3, Mi4 and Mi5 that
took place at 16 Ma, 13.8 Ma, 12.8 Ma and 11.5, respectively, impacted cricetid evolution by
promoting dispersal and triggering important origination/extinction events.

DATA AVAILABILITY

The data underlying this article are available in the article and from the Dryad Digital
Repository https://datadryad.org/stash/share/Dmda4GGW64PE8-c4fZiwP2VbpnTiTZpWm4PRSm5L20Q

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Figure 2. Time-calibrated relaxed-clock Bayesian inference analysis with morphological tip-dating using the fossilized birth–death tree model. Summary of the MCCT depicting the median divergence time estimates for different cricetid lineages against a geological time scale. Numbers at nodes indicate median estimates for the divergence times, and node bars indicate the 95% highest posterior density for divergence times.

Figure 3. Ancestral state distributions at each node of the MCCT of cricetid obtained by BBM analysis implemented in RASP against a geological time scale. Pie charts indicate probabilities of alternative ancestral ranges. The colour indicates possible ancestral ranges at different nodes. The schematic map shows the biogeographical areas used in this work and colours correspond to: ocher (southwestern Europe), purple (central Europe), dark red (Greece), orange (Anatolia-Caucasus), green (west central Asia), light blue (Arabian Peninsula), dark blue (southern Asia) and yellow (east-central Asia).

Figure 4. Instantaneous per-capita (top) and deterministic (bottom) rates for Cricetodontinae speciation (p, S), extinction (q, E), and biodiversity change (d, R) calculated using
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phylogenetically constrained species ranges. Miocene cooling events against geological time
are indicated as Mi.

432x307mm (300 x 300 DPI)
Figure 2. Time-calibrated relaxed-clock Bayesian inference analysis with morphological tip-dating using the fossilized birth–death tree model. Summary of the MCCT depicting the median divergence time estimates for different cricetid lineages against a geological time scale. Numbers at nodes indicate median estimates for the divergence times, and node bars indicate the 95% highest posterior density for divergence times.

274x385mm (300 x 300 DPI)
Figure 3. Ancestral state distributions at each node of the MCCT of cricetid obtained by BBM analysis implemented in RASP against a geological time scale. Pie charts indicate probabilities of alternative ancestral ranges. The colour indicates possible ancestral ranges at different nodes. The schematic map shows the biogeographical areas used in this work and colours correspond to: ocher (southwestern Europe), purple (central Europe), dark red (Greece), orange (Anatolia-Caucasus), green (west central Asia), light blue (Arabian Peninsula), dark blue (southern Asia) and yellow (east-central Asia).
Figure 4. Instantaneous per-capita (top) and deterministic (bottom) rates for Cricetodontinae speciation (p, S), extinction (q, E), and biodiversity change (d, R) calculated using phylogenetically constrained species ranges. Miocene cooling events against geological time are indicated as Mi.

189x205mm (300 x 300 DPI)