

# 1 Amoebozoan testate amoebae illuminate the diversity of heterotrophs and the 2 complexity of ecosystems throughout geological time

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## 40 Abstract

41 Heterotrophic protists are vital in Earth's ecosystems, influencing carbon and nutrient  
42 cycles and occupying key positions in food webs as microbial predators. Fossils and  
43 molecular data suggest the emergence of predatory microeukaryotes and the  
44 transition to a eukaryote-rich marine environment by 800 million years ago (Ma).  
45 Neoproterozoic vase-shaped microfossils (VSMs) linked to Arcellinida testate  
46 amoebae represent the oldest evidence of heterotrophic microeukaryotes. This study  
47 explores the phylogenetic relationship and divergence times of modern Arcellinida  
48 and related taxa using a relaxed molecular clock approach. We estimate the origin of  
49 nodes leading to extant members of the Arcellinida Order to have happened during  
50 the latest Mesoproterozoic and Neoproterozoic (1054 - 661 Ma), while the  
51 divergence of extant infraorders postdates the Silurian. Our results demonstrate that  
52 at least one major heterotrophic eukaryote lineage originated during the  
53 Neoproterozoic. A putative radiation of eukaryotic groups (e.g. Arcellinida) during the  
54 early-Neoproterozoic sustained by favorable ecological and environmental conditions  
55 may have contributed to eukaryotic life endurance during the Cryogenian severe ice  
56 ages. Moreover, we infer that Arcellinida most likely already inhabited terrestrial  
57 habitats during the Neoproterozoic, coexisting with terrestrial Fungi and green algae,  
58 before land plant radiation. The most recent extant Arcellinida groups diverged  
59 during the Silurian Period, alongside other taxa within Fungi and flowering plants.  
60 These findings shed light on heterotrophic microeukaryotes' evolutionary history and  
61 ecological significance in Earth's ecosystems, using testate amoebae as a proxy.

62

## 63 Significance Statement

64 Arcellinida shelled amoebae are heterotrophic microbial eukaryotes with an  
65 extensive Neoproterozoic fossil record represented by the vase-shaped microfossils  
66 (VSMs), a diverse group that is abundant and widespread in late Tonian rocks  
67 (VSMs). Here we combined phylogenomic sampling and the fossil record to generate  
68 time-calibrated trees. Our results illuminate key events in the history of life, including:  
69 i) the Tonian origin of extant microbial eukaryote lineages; ii) a speculative proposed  
70 radiation of eukaryotes before the Cryogenian, "Tonian revolution"; iii) the  
71 establishment of complex terrestrial habitats before the Cryogenian; iv) a post-

72 Silurian divergence of modern Arcellinida sub-clades in terrestrial (including  
73 freshwater) habitats. Our results provide valuable insights into the evolution of life  
74 throughout geological time and are congruent with recent discoveries regarding the  
75 early diversification of eukaryotes, including the Precambrian history of eukaryotic  
76 protosteroids.

77

## 78 **Introduction**

79 Heterotrophic microbial eukaryotes play a crucial ecosystem role by contributing to  
80 the carbon and nutrient cycles (1, 2). These organisms, capable of phagocytosis, act  
81 as predators on bacterial and eukaryotic communities, playing a significant role in  
82 complex food webs supported by primary producers (1). Additionally, predation is an  
83 evolutionary innovation that likely contributed to the diversification of eukaryotes (3).  
84 The Last Eukaryotic Common Ancestor (LECA) was heterotrophic and capable of  
85 phagocytosis. However, the timing and specific conditions under which diverse  
86 lineages of heterotrophic microeukaryotes have proliferated in Earth's ecosystems  
87 remain unclear (4-6).

88 Evidence from fossils, biomarkers, geochemical proxies, genomic data, and  
89 molecular clocks indicate that eukaryotes first originated during the Stenian (1200-  
90 1000 Ma) and Tonian periods (1000-720 Ma) (7-11). This led to a transition from a  
91 prokaryotic- to a eukaryotic-rich marine environment (6, 12-14), by 800 Ma, likely  
92 triggered by increased phosphorus, nitrate, and silica availability (14-17). From  
93 around this time, Neoproterozoic vase-shaped microfossils (VSMs) represent the  
94 remnants of an early eukaryotic divergence event. Organisms represented by VSMs  
95 are generally thought to have lived in marine environments, although a terrestrial  
96 habitat for these organisms is also plausible (18-20). The well-preserved nature of  
97 VSMs has allowed for detailed investigation and comparisons of their morphology to  
98 modern eukaryotic groups. These investigations support the current interpretation of  
99 a large fraction of VSMs being members of the stem or crown groups of testate  
100 amoeba order Arcellinida, due to both morphological affinities and congruence with  
101 molecular phylogenetic reconstructions (19, 21, 22). Other VSMs, such as  
102 *Melicerion poikilon*, had suggested affinities to Euglyphida, a distantly-related,  
103 convergent rhizarian clade of testate amoebae (18, 19). However, morphological  
104 evidence, in this case, is tentative, and the suggestion for a Euglyphida affinity is  
105 currently incongruent with molecular phylogenetic reconstructions, as Euglyphida is

106 part of a clade of cercozoan filose amoebae, which appears to be much younger,  
107 around 292 Ma (23). Arcellinida is a diverse lineage of extant heterotrophic  
108 microeukaryotes within the Amoebozoa, found in terrestrial and freshwater  
109 environments (19-22). Since many VSMs have been recognized as Arcellinida, they  
110 are accepted to represent the oldest and most diverse fossil evidence of  
111 heterotrophic microeukaryotes (19-20, 24,25). Elucidating the origin and evolutionary  
112 history of Arcellinida (and derived lineages) in light of their microfossil record is  
113 pivotal to illuminate the early evolution and possible radiation of heterotrophic  
114 microbial eukaryotes, and serve as a proxy to infer the complexity of Earth's  
115 ecosystems over geological time (19-20, 25, 26).

116 Recent efforts of sampling diverse amoebozoan testate amoebae in a  
117 phylogenomic framework have resolved their deep phylogenetic relationships (21,  
118 27). Amoebozoa is home to at least two testate amoebae groups: Arcellinida and  
119 Corycidia. Arcellinida is a diverse order represented by lineages that build hard  
120 extracellular shells, with the potential to generate exceptionally preserved fossilized  
121 remains (19, 21). Corycidia is a recently established subclade of Amoebozoa  
122 represented by the lineages of testate amoebae that produce flexible shells and do  
123 not branch within Arcellinida (27). Despite recent advances, many lineages still  
124 remain unsampled (21).

125 In addition to expanding the diversity of sampled Arcellinida at the genome  
126 level, the potential of a highly resolved phylogeny to provide insight into timing the  
127 Arcellinida origin and divergence of sub-clades has not been explored (22, 25).  
128 VSMs can be interpreted either as stem or crown Arcellinida. In either case, these  
129 fossils can be used to calibrate a phylogenetic tree and estimate the divergence time  
130 of lineages both within Arcellinida, as well as closely related amoebozoans. These  
131 diverse VSMs found in sedimentary deposits around the world have been  
132 continuously investigated, and their stratigraphy refined over time, enabling us to  
133 constrain these fossils' ages (19-20, 28-36). In this context, combining the VSMs and  
134 phylogenetic tree calibration opens up avenues to time the evolution of Arcellinida  
135 and closely related groups.

136 Here, we investigate the origin and divergence times of Arcellinida and closely  
137 related amoebozoan taxa using phylogenomics and a relaxed molecular clock  
138 approach. We expanded the taxonomic sampling for amoebozoan testate amoebae,  
139 including 14 taxa that lacked precise placement, to produce a new phylogenomic

140 dataset (utilizing 226 genes). We considered the diverse record of VSMs and  
141 Metazoa fossils to time calibrate this well-resolved deep phylogenomic tree. Different  
142 calibration strategies and molecular clock models support the divergence of extant  
143 Arcellinida lineages during the latest Mesoproterozoic and early to mid-  
144 Neoproterozoic, between 1060 and 661 Ma. We thus corroborate the origin of a  
145 major eukaryotic group by the Neoproterozoic, including a recorded establishment of  
146 heterotrophy predating the Cryogenian Period. Overall, using amoebozoan testate  
147 amoebae as a proxy, we provide insights into the evolution of microbial eukaryotes  
148 and Earth's early ecosystems.

149

## 150 **Results**

151

### 152 ***A resolved tree of amoebozoan testate amoebae***

153 We constructed a concatenated supermatrix using 57 taxa and 226 genes (70,428  
154 amino acid sites) using the PhyloFisher v. 1.2.11 package (37, 38). We include novel  
155 data for 14 testate amoebae taxa, obtained through single-cell or whole-culture  
156 transcriptomics (**Fig. 1; Dataset S01, Tables S1 - S4**). The remaining testate  
157 amoebae and sister-group taxa were sampled based on previously available  
158 genomes and transcriptomes (**Dataset S01, Table S1**). The resulting phylogenomic  
159 tree recovers a monophyletic Arcellinida, with three well-defined suborders  
160 (Phryganellina, Organoconcha, and Glutinoconcha) and five infraorders within  
161 Glutinoconcha (**Fig. 2; Appendix S01, Figs. S1 and S2**). The Corycidia clade is  
162 also recovered with full support, with two families, Trichosidae and Amphizonelliidae  
163 fam. nov. (**Fig. 2**). Nearly all nodes of the tree are fully (=100%) or highly (>92%)  
164 supported by Maximum Likelihood nonparametric Real Bootstraps (MLRB), except  
165 for a single node within Sphaerothecina clade that has lower support (MLRB = 76%;  
166 **Fig. 2**). We have also produced single-gene reconstructions using SSU rDNA and  
167 Cytochrome Oxidase subunit I (COI) (**Dataset S01, Tables S5-S6**), which are the  
168 genes traditionally used to reconstruct relationships in Arcellinida. These analyses  
169 present broader taxon sampling but failed to recover most of the deeper  
170 relationships in Arcellinida (**Appendix S01, Figs. S3-S4**)

171

### 172 ***Time-calibrated tree of amoebozoan testate amoebae***

173 For estimating divergence times in testate amoebae evolution and closely related  
174 taxa, we expanded our phylogenomic supermatrix to consider a representative  
175 sampling of Amorphea, including Amoebozoa, Fungi, Metazoa, and their protistan  
176 relatives (**Dataset S01, Table S7**). For a comprehensive approach, taking into  
177 account the alternative interpretations of the VSM record, we implemented four  
178 different calibration strategies: i. calibration of nodes within Metazoa, excluding the  
179 VSM record to calibrate amoebozoan nodes; ii. calibration of nodes within Metazoa  
180 and calibration of Glutinoconcha+Organoconcha and Glutinoconcha nodes,  
181 considering VSMs as derived crown Arcellinida; iii. calibration of nodes within  
182 Metazoa and calibration of the Arcellinida node, considering VSMs as basal crown  
183 Arcellinida; iv. calibration of nodes within Metazoa and calibration of  
184 Arcellinida+Euamoebida node, considering VSMs as stem Arcellinida (**Dataset S01**,  
185 **Table S8 - S10**). To implement these four fossil calibration strategies, we performed  
186 a total of 36 experiments considering three different distributions (i.e., Uniform,  
187 Skew-Normal, or Truncated-Cauchy short-tail) under an uncorrelated or  
188 autocorrelated relaxed clock model with either a drift parameter of  $\alpha = 2$  and  $\beta = 2$  or  
189  $\alpha = 1$  and  $\beta = 10$ . We ran each experiment in two independent MCMC chains to  
190 check for convergence, which was achieved for all analyses (**Appendix S01, Fig.**  
191 **S5**).

192 Comparing all the calibration strategies and experiments, we observed overall  
193 similar inferred times with the uncorrelated clock model (median = 591 - 531 Ma and  
194 mean = 627 - 549 Ma; **Dataset S01, Table S11 and Appendix S01, Figs. S6 - S43**)  
195 and the autocorrelated clock model (median = 592 - 531 Ma and mean = 671 - 557  
196 Ma; **Dataset S01, Table S11**). Regarding implemented distributions, overall the  
197 uniform distribution inferred the youngest ages, while Truncated-Cauchy inferred  
198 slightly older ages (**Dataset S01, Table S11**). The drift parameter (i.e.,  $\alpha = 2$  and  $\beta =$   
199 2 vs.  $\alpha = 1$  and  $\beta = 10$ ) had virtually no impact independent of the clock model,  
200 distribution, and calibration strategy (**Dataset S01, Table S11**). The estimated times  
201 for the Arcellinida node by excluding VSMs from the calibration (mean = 911 - 734  
202 Ma; 95% highest probability density confidence interval (HPD CI) = 1060 - 605) and  
203 by including VSMs in the calibration (mean = 930 - 746 Ma; 95% HPD CI = 1054 -  
204 661 Ma) are highly congruent. This agrees with the current interpretation that VSMs  
205 represent fossil remains of Arcellinida, supporting their use to calibrate amoebozoan  
206 nodes. Aiming for a comprehensive approach, the time estimation results shown and

207 discussed hereafter focus on the full range of times estimated based on the  
208 calibration strategies that included the VSMs, the three distributions, and the drift  
209 parameter of  $\alpha = 2$  and  $\beta = 2$  under the uncorrelated or autocorrelated relaxed clock  
210 model.

211 The molecular clock analyses inferred a mean time for the root of Amorphea  
212 to be between 1640 and 1393 Ma (95% HPD CI = 1843 - 1088 Ma; **Fig. 3, Dataset**  
213 **S01, Table S11, and Appendix S01, Fig. S7 - S43**). For Metazoa, the mean time  
214 estimated ranged between 835 and 734 Ma (95% HPD CI = 872 - 673 Ma; **Fig. 3,**  
215 **Dataset S01, Table S11, and Appendix S01, Fig. S7 - S43**). The mean estimated  
216 for the origin of Amoebozoa ranged from 1607 - 1298 Ma (95% HPD CI = 1795 -  
217 1045 Ma; **Fig. 3, Dataset S01, Table S11, and Appendix S01, Fig. S7 - S43**). The  
218 Arcellinida node is constrained within the mean 930 and 746 Ma (95% HPD CI =  
219 1054 - 661 Ma; **Fig. 3, Dataset S01, Table S11, and Appendix S01, Fig. S7 - S43**),  
220 estimating an origin for Arcellinida during the latest Mesoproterozoic and  
221 Neoproterozoic (**Fig. 4A**). For the early divergence time of Arcellinida subclades, the  
222 estimated times suggest that the split between the Organoconcha and  
223 Glutinoconcha branches occurred during the Neoproterozoic (mean = 855 - 679 Ma;  
224 95% HPD CI = 969 - 600 Ma; **Figs. 3 - 4B**). Regarding the suborders of Arcellinida,  
225 we inferred a mean origin for Phryganellida between 534 - 265 Ma (95% HPD CI =  
226 661 - 175 Ma; **Figs. 3 - 4C**), for Organoconcha 735 - 550 Ma (95% HPD CI = 839 -  
227 463 Ma; **Figs. 3 - 4D**), and for Glutinoconcha between 790 - 621 Ma (95% HPD CI =  
228 897 - 539 Ma; **Figs. 3 - 4E**). For the deeper nodes of Glutinoconcha, the most  
229 sampled suborder of Arcellinida, the estimated divergence times ranged between  
230 Cryogenian and Carboniferous (mean = 643 - 393 Ma; 95% HPD CI = 705 - 335;  
231 **Figs. 3 - 4F - H**). The inferred ages for Glutinoconcha infraorders are relatively more  
232 widespread depending on the calibration strategy when compared to other nodes.  
233 Treating VSMs as derived crown taxa constrains the infraorders origin between  
234 Ediacaran and early Cretaceous (mean = 422 - 221 Ma; 95% HPD CI = 575 - 123  
235 Ma; **Figs. 3 - 4I - M**) while considering VSMs as basal crown or stem Arcellinida  
236 estimate their origin mostly between Silurian and early Cretaceous (mean = 339 -  
237 172 Ma; 95% HPD CI = 444 - 122 Ma; **Figs. 3 - 4I - M**). It is worth noting that only  
238 the calibration using a uniform autocorrelated clock model and VSMs as derived  
239 crown arcellinids inferred ages as old as the Ediacaran for the Glutinoconcha  
240 infraorders. All other distribution-clock models consistently led to ages constrained

241 within the Paleozoic. Most inferred ages for the nodes representing the origin of the  
242 modern extant genera and species of Arcellinida postdate the Silurian (mean = 416 -  
243 125 Ma; **Dataset S01, Table S11**). Besides the testate amoebae, the inferred mean  
244 times for the other orders and major groups of Amoebozoa we sampled ranged from  
245 585 Ma to 1288 Ma, placing their origin mostly during the Neoproterozoic (**Dataset**  
246 **S01, Table S11**). The results and time-calibrated trees for all experiments are  
247 present in the supplemental material (**Dataset S01, Table S11 and Appendix S01,**  
248 **Fig. S6 - S43**).

249

## 250 **Ancestral habitat reconstruction of Arcellinida**

251 We performed statistical analyses on the ancestral habitat of key hypothetical  
252 ancestors within Arcellinida considering alternative scenarios of a terrestrial or  
253 marine origin for the crown group (**Dataset S01, Table S12 and Appendix S01, Fig.**  
254 **S44**). The unrestrained reconstruction inferred a terrestrial habitat (100% probability)  
255 for all nodes within Arcellinida. The ancestral reconstruction that sets the fixed value  
256 of a marine state on the last common ancestor of modern Arcellinida inferred a high  
257 probability of terrestrial habitat for all nodes within Arcellinida (>93%), implying at  
258 least two independent transition events (2TE) from marine to terrestrial habitats (**Fig.**  
259 **5**). The ancestral reconstruction that sets the fixed value of a marine state on the last  
260 common ancestor of both the Arcellinida and the Organoconcha+Glutinoconcha  
261 clades inferred with high probability (>88%) a terrestrial habitat for the hypothetical  
262 ancestors of Phryganellina, Organoconcha, and Glutinoconcha, implying at least  
263 three independent transition events (3TE) from marine to terrestrial habitats (**Fig. 5**).

264

## 265 **Discussion**

266

### 267 ***A resolved tree of amoebozoan testate amoebae***

268 The phylogenomic dataset constructed in this study improves several aspects of the  
269 previously available amoebozoan testate amoebae dataset (21, 27). Through the  
270 PhyloFisher pipeline, we were able to construct a curated phylogenomic matrix that  
271 is free of paralogs and contamination (**Appendix S01, SI1**). Moreover, the matrix  
272 constructed is an accessible and easy-to-update dataset, since newly sequenced  
273 transcriptomes can be easily added through PhyloFisher to further expand the  
274 taxonomic sampling of amoebozoan testate amoebae in a phylogenomic approach.

275 In general terms, the phylogenomic tree obtained here is consistent with the  
276 previously published phylogenomic tree for amoebozoan testate amoebae (**Fig. 2**;  
277 21, 27). By recovering all major groups of the Arcellinida and Corycidia clades with  
278 full support, in accordance with previous reconstructions, we corroborate the  
279 robustness of the backbone of the amoebozoan testate amoebae tree. The  
280 phylogenomic analyses enable us to place previously unsampled taxa within  
281 Arcellinida suborders, supporting the taxonomic actions regarding *Heleopera lucida*  
282 comb. nov. and the genus *Microcorycia* (**Appendix S01, SI2 - SI3**). Moreover, the  
283 phylogenomic tree identifies taxa that will need future taxonomic revision based on  
284 further morphological and molecular studies, those being *Diffugia* cf. *capreolata* and  
285 the genera *Cyclopyxis* and *Phryganella*. Detailed discussion on the placement of  
286 newly sequenced amoebozoan testate amoebae is presented in **Appendix S01, SI2**  
287 - **SI3**.

288

### 289 **Robust Amorphea time-calibrated trees using VSMs**

290 Our time-calibrated trees are congruent with several molecular clocks that have  
291 sampled the diversity of eukaryotes (**Fig. 3, Dataset S01, Table S11, and Appendix**  
292 **S01, Fig. S6 - S43**). The estimated ages for the root of our tree fall within the range  
293 inferred by a recent molecular timescale for eukaryotes (95% confidence interval =  
294 2177 - 1356 Ma for Amorphea; 39). Similarly, the estimated ages for the origin of  
295 other major clades align with previous studies, including Obazoa (95% HPD CI =  
296 2305 - 1526 Ma), opisthokonts (95% HPD CI = 2019 - 1051 Ma), and animals (95%  
297 HPD CI = 833 - 680 Ma) (39-42). The inferred times place the origin of Arcellinida  
298 during the latest Mesoproterozoic and early to mid-Neoproterozoic, most likely during  
299 the Tonian Period and no later than the Cryogenian (**Fig. 4, Dataset S01, Table**  
300 **S11, and Appendix S01, Fig. S6 - S43**). It is worth noting that by considering  
301 different calibration strategies to account for alternative plausible interpretations of  
302 VSMs and the unavoidable fossil record uncertainties, the times we estimated for  
303 each node have wide ranges. However, independently of the calibration strategy,  
304 distribution, and clock model, the Arcellinida origin is mostly placed within the  
305 Neoproterozoic, and all estimated mean times suggest a Tonian origin. Notably, the  
306 time-calibrated trees we generated without using the VSMs as calibration data  
307 inferred times congruent with our analyses considering the Tonian VSMs. This  
308 demonstrates that inferred ages are not a fossil calibration bias and serves as further

309 corroboration for the interpretation of VSMs as fossil members of, at the very least,  
310 the stem Arcellinida group (**Fig. 4, Dataset S01, Table S11**). Also, the ages we  
311 inferred for the origin of modern groups (i.e., genera) are highly congruent with the  
312 recent Arcellinida fossil record which postdates the Carboniferous Period and  
313 preserves fossils assigned to genera like *Arcella*, *Diffugia*, and *Centropyxis* (**Dataset**  
314 **S01, Table S11**; 43-45). Collectively, the consistency of the estimated times, and  
315 their congruence with previous molecular clocks and with the fossil record, support  
316 that including VSMs to calibrate a comprehensive phylogenomic sampling of  
317 Amorphea leads to robust results.

318

### 319 **The Neoproterozoic diversification of heterotrophic microbial eukaryotes**

320 Our time-calibrated trees reveal that Arcellinida originated most likely during the  
321 Neoproterozoic, with most inferred times and all of the means falling within the  
322 Tonian Period (**Fig. 4 A - B**). The mean estimated times for other major groups of  
323 Amoebozoa also indicate a Tonian origin. Similarly, previous molecular clock  
324 analyses have indicated the divergence of multiple heterotrophic eukaryotes during  
325 this period (40, 41). However, their diversity has been challenging to examine due to  
326 the lack of a fossil record for these organisms. Nevertheless, the estimated  
327 Neoproterozoic origin for the Arcellinida crown group and the diversity of Tonian  
328 VSMs, currently represented by 14 morphologically diverse genera (19-20, 25, 31),  
329 suggest that by the Tonian Period, Earth's ecosystems had witnessed the origin of  
330 modern heterotrophic microbial eukaryotes.

331 An inferred Tonian origin for diverse heterotrophic microbial eukaryotes is  
332 congruent with proposals of ecosystem establishment during the Neoproterozoic,  
333 stemming from various disciplines. Previous studies speculate the existence of a  
334 “Tonian revolution”, based on evidence from biomarkers, fossil records, and  
335 molecular data that infer a marked transition from prokaryotic- to eukaryotic-rich  
336 ecosystems by 800 Ma (6, 11-14, 46). This transition was likely facilitated by factors  
337 such as increased availability of nitrate, phosphorus, silica, and reduced toxicity,  
338 which provided a favorable condition linked to the documented diversification of  
339 eukaryotic phototrophs at that time (14-17). In turn, the establishment of a  
340 community of phototrophs may have served as a favorable condition for the  
341 diversification of heterotrophs. Fossil and geochemical evidence suggest that  
342 photosynthetic biological mats contributed to the establishment of Oxygen Oases

343 during the Tonian, likely triggered by a higher capacity of oxygen productivity of  
344 eukaryotic phototrophs (7, 10, 47, 48). These oases probably represented an  
345 increase in aerobic conditions, and food availability, that were permissive to the  
346 survival and proliferation of heterotrophs like Arcellinida. Consequently, a combined  
347 interpretation of the geochemical and fossil records indicates that complex  
348 ecosystems were established by the Neoproterozoic.

349 VSMs serve as a unique testimony of the putative “Tonian revolution” on  
350 eukaryotic diversification and ecosystems established during the Neoproterozoic.  
351 The VSMs have been found in rocks characterized by organic-rich sediments  
352 corroborating the close association of heterotrophic eukaryotes with microbial mats  
353 (19). Also, the organisms represented by these VSMs likely preyed on both the  
354 bacterial and eukaryotic communities, similar to extant Arcellinida (19, 49, 50).  
355 Culture observations have demonstrated diverse strategies of extant Arcellinida to  
356 prey on various organisms, including diatoms, fungi, and nematodes (51-54).  
357 Moreover, several VSMs exhibit holes on their shells interpreted as predation marks,  
358 suggesting they also served as prey to other heterotrophs (3, 55). Predation has  
359 been interpreted as one of the triggers for eukaryotic diversification, including for  
360 animals, and VSMs are among the oldest records of this evolutionary innovation (3,  
361 56-58). Thus, while most of the microbial eukaryotes did not leave fossils, VSMs  
362 stand out as a robust fossil record highlighting the rise of predation and increase of  
363 food web complexity on Earth’s ecosystems no later than the Tonian Period.

364 Our time-calibrated trees suggest that the divergence of some modern  
365 Arcellinida lineages happened during the Neoproterozoic. The inferred times for the  
366 split and origin of the Organoconcha and Glutinoconcha suborders mostly fall within  
367 the Neoproterozoic between the Tonian and Ediacaran Periods, with only some  
368 estimated times suggesting an early-Paleozoic origin for Organoconcha. Specifically,  
369 the ages predicted for the Organoconcha and Glutinoconcha split either predate or  
370 overlap with the Cryogenian Period and its glaciations. It is worth noting that  
371 Phryganellina is currently the least genomically sampled Arcellinida group,  
372 represented only by two genera, thus it is difficult to assess when this suborder  
373 originated. Altogether, the origin and early divergence of the Arcellinida crown group,  
374 and the diverse VSMs record, imply the capability of Neoproterozoic ecosystems not  
375 only to sustain heterotrophic eukaryote groups but also to allow their diversification.

376

377 **A possible Pre-Cryogenian Eukaryotic Diversification and Survival during**  
378 **Earth's most severe Ice Ages**

379 The Sturtian and Marinoan glaciation periods witnessed extensive ice coverage  
380 across the planet, with glaciers extending into tropical regions (59-61). The survival  
381 of life during Cryogenian glaciations can be explained by the presence of refugia and  
382 the adoption of dormancy strategies (62-69). The presence of cyst-like structures  
383 identified inside VSMs serves as direct evidence that these organisms were already  
384 capable of entering dormancy stages, similar to extant Arcellinida (70). Additionally,  
385 recent discoveries have revealed that Tonian VSM taxa persisted into the  
386 Cryogenian Period, providing fossil records that showcase the diversity of  
387 heterotrophic eukaryotes during glaciation periods (69). The inferred times for the  
388 split and origin of Glutinoconcha and Organoconcha suggest they possibly originated  
389 during Cryogenian, indicating not only survival but also divergence of novel modern  
390 eukaryotic taxa during Cryogenian glaciations (**Fig. 4 D - E**). Consequently, the VSM  
391 record and the timing of early Arcellinida evolution enable speculation about a  
392 possible radiation of heterotrophic eukaryotic life shortly before and during the  
393 Cryogenian Period. This, coupled with a capacity for dormancy and the exploitation  
394 of habitat refugia, may explain the endurance of life during Earth's most severe ice  
395 age.

396

397 **Timing of terrestrial conquest by Arcellinida**

398 Currently, while it is largely suggested that the organisms represented by Tonian  
399 VSMs lived in shallow marine environments, a terrestrial habitat cannot be ruled out.  
400 To date, VSMs have been reported from Tonian sedimentary deposits described as  
401 fully or partially marine (18-19, 25). However, although scarcely discussed in the  
402 literature, it is plausible to hypothesize that the organisms represented by the VSMs  
403 may have lived in terrestrial environments and were deposited in marine sediments  
404 through a number of possible mechanisms, including: surface runoff, river discharge,  
405 wind blowing, or supratidal spillover. In any case, these organisms ultimately  
406 fossilized in a marine setting (71). Consequently, considering the alternative  
407 interpretations of VSMs' natural habitat and their affinity to Arcellinida (i.e., stem or  
408 crown Arcellinida), different scenarios can be explored regarding Arcellinida's  
409 conquest of terrestrial habitat.

410 Our ancestral habitat reconstructions indicate three alternative scenarios, a  
411 terrestrial origin for Arcellinida, a marine origin with two independent transition  
412 events (2TE) from marine to terrestrial habitats, and a marine origin with three  
413 independent transition events (3TE) (**Fig. 5 and Appendix S01, Fig. S44**). Although  
414 the reconstruction of a terrestrial origin is statistically superior to the other scenarios  
415 (Likelihood Ratio Test - LRT), this was already expected since all extant Arcellinida  
416 are terrestrial/freshwater inhabiting, leading to the reconstruction of a terrestrial  
417 ancestor (i.e. a possible systemic bias). However, interpreting VSMs as stem or  
418 crown Arcellinida and enforcing a marine origin for the Arcellinida stem and early  
419 crown groups, 2TE or 3TE are plausible and statistically equivalent based on LRT, in  
420 accordance with previous hypotheses (18-21, 25, 72). Multiple transition events are  
421 biologically plausible: Arcellinida-related amoebozoan lineages are often represented  
422 by both marine and terrestrial species, even within the same genera (e.g., *Vannella*,  
423 *Mayorella*, and *Trichamoeba*).

424 Coupling the reconstructed scenarios with the timing of Arcellinida origin and  
425 early divergence of its sub-clades, we infer that many Arcellinida probably inhabited  
426 terrestrial environments already in the Neoproterozoic, no later than the Ediacaran  
427 Period. Even if we consider the latest transition scenario reconstructed (3TE) the  
428 inferred times place the terrestrialization event of Glutinoconcha and Organoconcha  
429 most likely between the Tonian and Ediacaran periods (**Fig. 5**). The time of  
430 diversification of life in terrestrial habitats has been traditionally discussed based on  
431 the time of divergence of land plants (embryophytes), which is constrained within a  
432 Paleozoic diversification (73, 74). However, recent inferences based on  
433 phylogenomic reconstructions and molecular clocks have suggested that modern  
434 eukaryotic lineages, like Fungi and green algae, diverged on land no later than  
435 Cryogenian (73, 74). Congruently, our estimated times for Arcellinida  
436 terrestrialization are constrained within the Neoproterozoic. These findings suggest a  
437 Neoproterozoic establishment of relatively complex terrestrial ecosystems inhabited  
438 by diverse organisms, including phototrophic (green algae), absorptive heterotrophic  
439 (Fungi), and phagotrophic heterotrophic protists.

440 The inferred divergence of Arcellinida sub-clades in terrestrial habitats, well  
441 represented by Glutinoconcha (currently the best-sampled suborder), is also  
442 congruent to the diversification timing estimated for other eukaryotic groups. The  
443 Glutinoconcha infraorders' split is mostly constrained between the late-

444 Neoproterozoic and mid-Paleozoic (Devonian Period). The radiation of Fungi and the  
445 diversification of extant land plants are estimated to the same window (ca. 480 Ma)  
446 (75). Subsequently, the origin of extant Arcellinida groups, represented by the origin  
447 of all Glutinoconcha infraorders, is mostly constrained between the Silurian and  
448 Cretaceous. This is contemporaneous with the documented Late Paleozoic  
449 diversification of seed plants and saprotrophic mushrooms (75). Similarly, the  
450 estimated time for the divergence of Arcellinida genera, mostly post-dating early  
451 Mesozoic, is congruent to the radiation of diverse groups of Fungi and land plants,  
452 including pine trees (Pinaceae) and flowering plants (Angiosperm) (75). Altogether,  
453 congruences between the timing of the origin of various eukaryotic groups suggest  
454 an integrated and synchronous diversification of life in terrestrial habitats, enabling  
455 speculation about a possible radiation of Arcellinida in this time period. However, this  
456 claim requires explicit testing and corroboration via well-sampled studies of  
457 background diversification rates using fossils.

458

## 459 **Conclusions**

460 Timing the origin of modern Arcellinida testate amoebae and the divergence times of  
461 their subclades illuminate the evolution of heterotrophic microbial eukaryotes in  
462 geological time. To estimate this timing, we expanded the phylogenomic sampling of  
463 amoebozoan testate amoebae and generated robust time-calibrated Amorphea trees  
464 based on both Arcellinida and Metazoa fossil records. The estimated times for the  
465 origin of Arcellinida and other amoebozoans, mostly constrained within the Tonian  
466 Period, are congruent with the previously speculated “Tonian revolution” for a  
467 diversification of eukaryotes in this Period. This consistency suggests that Earth’s  
468 ecosystems had witnessed the divergence of both phototrophic and heterotrophic  
469 eukaryotic lineages, including Arcellinida, during the Neoproterozoic, no later than  
470 the Tonian Period. A putative radiation of eukaryotes before the Cryogenian Period,  
471 coupled with the exploitation of refugia habitats and dormancy strategy, may have  
472 contributed to their endurance during Earth’s most severe ice ages. Although the  
473 ancestral habitat of Arcellinida and the possibility of transition between environments  
474 (marine vs. terrestrial) remain contentious, considering the plausible alternative  
475 scenarios we infer that Arcellinida were most likely already inhabiting terrestrial  
476 habitats between the Tonian and Ediacaran Periods. Together with the previously  
477 suggested diversification of Fungi and green algae on land during the Cryogenian

478 Period, the inferred time for terrestrial Arcellinida is congruent with a Neoproterozoic  
479 establishment of relatively complex ecosystems composed of phototrophic (green  
480 algae), absorptive heterotrophic (Fungi), and phagotrophic heterotrophic eukaryotes,  
481 preceding the diversification of land plants. Similarly, the estimated post-Silurian  
482 origin of modern Arcellinida (i.e., infraorders) suggests a contemporaneity to the  
483 diversification of other groups, including diverse Fungi and land plants. Ultimately,  
484 we suggest the Arcellinida testate amoebae are a key group to further explore the  
485 diversification of heterotrophic microbial eukaryotes and the establishment of  
486 ecosystems starting in the Neoproterozoic.

487

## 488 **Material and Methods**

489

### 490 **Sampling, RNA extraction, and sequencing**

491 We generated transcriptomes for 14 previously genomically unsampled amoebozoan  
492 testate amoeba species through mRNA extraction from either monoclonal cultures or  
493 single-cells isolated from natural samples (**Dataset S01, Table S1**). We synthesized  
494 cDNA from RNA extractions using an adaptation of the Smart-seq2 protocol (77). We  
495 prepared our cDNA libraries for sequencing on the Illumina platform using a  
496 NexteraXT DNA Library Preparation Kit (Illumina) following the manufacturer's  
497 recommended protocol. Libraries were then pooled and sequenced (**Appendix S01,**  
498 **SI1; Dataset S01, Table S1**).

499

### 500 **Trimming, Transcriptome Assembly, and Quality Assessment**

501 We trimmed primers, adaptors, and low-quality bases from raw Illumina reads using  
502 Trimmomatic v. 0.36 (77). We then assembled the surviving reads with Trinity v.  
503 2.12.0 (78). We predicted amino acid sequences (proteomes) from the assembled  
504 transcriptomes using Transdecoder v. 5.5.0. Finally, we assessed the completeness  
505 of all newly sequenced transcriptomes using BUSCO v. 5.3.2 (79) (**Dataset S01,**  
506 **Table S1**). Further details on trimming, transcriptome assembly, and quality  
507 assessment are presented in **Appendix S01, SI1**.

508

### 509 **Phylogenomic dataset construction**

510 We constructed our amoebozoan phylogenomic dataset using the database and  
511 tools available in PhyloFisher v. 1.2.11 (37) following the steps outlined at

512 <https://thebrownlab.github.io/phylofisher-pages/detailed-example-workflow> and in  
513 Jones et al. (38). Our final concatenated matrix used in subsequent phylogenetic  
514 analyses consisted of 226 genes (70,428 amino acid sites) and 57 amoebozoan taxa  
515 (**Dataset S01, Table S4**). From each individual ortholog that was concatenated to  
516 create the aforementioned matrix, we constructed single ortholog trees to be used as  
517 input for coalescent-based phylogenomic analyses. Further details on our approach  
518 for phylogenomic dataset construction are presented in **Appendix S01, SI1**.

519

### 520 **Phylogenomic analyses**

521 We performed maximum likelihood phylogenetic reconstruction using our final matrix  
522 with IQ-TREE2 v. 2.0-rc (80). We initially inferred a tree from our matrix under the  
523 LG+C20+G4 model. We used the resulting tree as a guide tree to infer a Posterior  
524 Means Site Frequency model (81) using the ML model LG+C60+G4+PMSF in IQ-  
525 TREE2. We assessed the topological support for the resulting tree by 100 Real  
526 nonparametric Bootstrap replicates in IQ-TREE (IQ-TREE v. 2.1.2 COVID-edition)  
527 using the PMSF model. The topological support values inferred from MLRB were  
528 mapped onto the ML tree using RAxML v. 8.2.12 (82) using the “-f b” option. We  
529 carried out coalescent-based phylogenomic analyses with ASTRAL-III v. 5.7.3 (83).  
530 Statistical support for our ASTRAL-III analysis was assessed via local posterior  
531 probability values.

532

### 533 **Bayesian molecular dating**

#### 534 *Dataset and Topology*

535 Utilizing previously identified orthologs already present in the publicly available  
536 PhyloFisher database, we expanded our amoebozoan phylogenomic dataset to  
537 include a representative sampling of Amorphea. Amorphea is the eukaryotic clade  
538 composed of Amoebozoa, Fungi, Animals, and some other unicellular lineages.  
539 These ortholog sequences were aligned, trimmed, and concatenated as described  
540 above (*Phylogenomic dataset construction section*). Our final expanded dataset  
541 used in the subsequent phylogenetic reconstruction and for the molecular dating  
542 analysis consisted of 230 genes (73,467 amino acid sites) and 96 taxa (**Dataset**  
543 **S01, Table S7**). Maximum likelihood phylogenetic analysis was performed as  
544 described above (*Phylogenomic analyses section*).

545

546 *Fossil calibrations*

547 As external calibration information, we considered fossils to calibrate five internal  
548 nodes of the Metazoa clade and explored three different strategies to calibrate  
549 amoebozoan nodes (**Dataset S01, Tables S8 - S10**). We strictly derived the fossil  
550 calibration for Metazoa from dos Reis et al. (42) and Benton et al., (84), which have  
551 carefully evaluated the diversity of fossils available and calibration strategies for the  
552 Metazoa lineage. We derived the fossil calibration for Arcellinida based on previous  
553 analyses and interpretations of the morphological relationship between VSMs and  
554 extant Arcellinida (19, 21, 22). Currently, VSM's can be interpreted either as: i) the  
555 fossil record of stem Arcellinida; ii) basal crown Arcellinida closely related to  
556 Arcellinida common ancestor, or; iii) derived crown Arcellinida (19, 21, 22).  
557 Specifically, interpretations of Bayesian and maximum likelihood ancestral  
558 reconstructions of Arcellinida shell morphology suggest a morphological congruence  
559 between the VSM *Melanocyrillum* to the Glutinoconcha+Organonconcha hypothetical  
560 ancestor and between the VSM *Cyliocyrillum* to Glutinoconcha hypothetical  
561 ancestor, thus suggesting they may represent derived crown Arcellinida (21).  
562 Consequently, three different calibration strategies can be implemented from these  
563 alternative interpretations. VSM's can be considered to calibrate: i)  
564 Glutinoconcha+Organonconcha and Glutinoconcha nodes; ii) calibrate the Arcellinida  
565 node, or; iii) calibrate the node shared between Arcellinida and its closest sister  
566 group, the amoebozoan order Euamoebida. Aiming for a comprehensive approach  
567 we considered these three strategies and generated comparable time tree  
568 estimations. To constrain the VSMs ages, we considered the literature which has  
569 described these microfossils and refined their stratigraphical distribution (**Dataset**  
570 **S01, Tables S9**; 36). Since molecular dating considers the fossil information as  
571 statistical distributions, and different distributions may impact the time estimation  
572 differently, we followed dos Reis et al. (42) strategy and used a total of three different  
573 distributions to represent the calibrations derived from the fossil record: i. uniform; ii.  
574 skew-normal; and iii. truncated-cauchy short-tail. Full details on our approach for  
575 fossil calibration are presented in **Appendix S01, SI1**.

576

577 *Divergence time estimation*

578 We performed Bayesian molecular dating with the MCMCTree program,  
579 implemented within the PAML package (**Dataset S01, Table S10**; 85). We estimated

580 a mean substitution rate of  $0.03135 \text{ replacement site}^{-1} 10^{-8} \text{ Myr}^{-1}$  for our dataset with  
581 IQ-TREE2 v. 2.0.6 (80) phylogenetic dating under the LG+G model. Within the PAML  
582 package, we set the overall substitution rate ('rgene\_gamma =  $\alpha$ ,  $\beta$ ' parameter) as a  
583 gamma-Dirichlet prior following dos Reis et al. (42), with  $\alpha = 2$  and  $\beta = 63.78$ . This  
584 substitution rate was implemented for all dating experiments. For the rate drift  
585 parameter ('sigma<sup>2</sup>\_gamma =  $\alpha$ ,  $\beta$ '), we independently implemented two alternatives,  
586  $\alpha = 2$  and  $\beta = 2$  or  $\alpha = 1$  and  $\beta = 10$ . Similarly, we considered both uncorrelated and  
587 autocorrelated relaxed clock models. For all experiments, we analyzed the data  
588 under the LG+G model as a single partition, constrained the root age between 1.6 -  
589 3.2 Ga, and considered a uniform birth-death tree prior and 100 million years as one  
590 time unit. We performed a total of 36 experiments, considering three distributions  
591 (i.e., uniform, skew-normal, and truncated-Cauchy short-tail), varying the rate drift  
592 parameter (i.e.,  $\alpha = 2$  and  $\beta = 2$  or  $\alpha = 1$  and  $\beta = 10$ ) and clock model (i.e.,  
593 uncorrelated and autocorrelated). For each experiment, we ran two independent  
594 MCMC chains to verify convergence, discarding the first 2,000 iterations as burn-in  
595 and considering the following 20 million generations. To check the influence of fossil  
596 calibrations using Neoproterozoic VSMs on the estimated dates, we performed  
597 experiments calibrating only the nodes within the Animal clade, applying Uniform and  
598 Skew-Normal calibration strategies, under an uncorrelated or autocorrelated relaxed  
599 clock model with a drift parameter of  $\alpha = 2$  and  $\beta = 2$  or  $\alpha = 1$  and  $\beta = 10$ , following  
600 the same approach described above, as detailed in **Appendix S01, SI1**.

601

## 602 ***Ancestral reconstruction of Arcellinida habitat***

603 We applied a maximum likelihood (ML) method, implemented in BayesTraits v. 4.0.1  
604 (86), to statistically reconstruct the ancestral habitat states of Arcellinida and  
605 compare evolutionary scenarios. Currently, the diverse Tonian VSM record has been  
606 documented from environments described as fully or partially marine, suggesting the  
607 organisms represented by these fossils inhabited marine environments (19, 25).  
608 However, we cannot rule out the possibility of a terrestrial habitat for these  
609 organisms, since their dead remains could have been transported from terrestrial to  
610 marine environments where they fossilized. To explore this issue, we combine the  
611 fossil evidence and the phylogenomic reconstruction with branch lengths to  
612 reconstruct the potential ancestral habitat states (i.e., marine habitat vs. terrestrial  
613 habitat) of key Arcellinida clades through the BayesTraits MultiState method (86).

614 Specifically, we used the 100 topologies with branch lengths obtained for the  
615 phylogenomic Real Bootstrap topological support assessment (see *Phylogenomic*  
616 *analyses section*) and implemented four different ancestral reconstruction analyses:  
617 i. ancestral reconstruction without fossilizing (assigning a fixed ancestral state value)  
618 nodes; ii. ancestral reconstruction fossilizing Arcellinida node as terrestrial, which  
619 interprets the organisms represented by VSMs as terrestrial; iii. ancestral  
620 reconstruction fossilizing Arcellinida node as marine, which interprets the organisms  
621 represented by VSMs as marine; iv. ancestral reconstruction fossilizing Arcellinida  
622 and Organoconcha+Glutinoconcha nodes as marine, which interprets the organisms  
623 represented by VSMs as derived crown Arcellinida that lived in marine habitat  
624 (**DatasetS01, Table S12 and Appendix S01, Fig. S44**). To compare the  
625 reconstructed scenarios, we applied a likelihood ratio test (LRT), which we  
626 considered as significant a difference of  $LRT \geq 2$  (87).

627

## 628 **Data Availability**

629 Raw sequencing files are deposited at the NCBI SRA repository under the Bioproject  
630 PRJNA1032600. Phylogenomic supermatrix, single gene marker datasets, and input  
631 information for the molecular clock and ancestral reconstruction analyses are  
632 presented in DatasetS01. All molecular data associated with this manuscript are  
633 available on FigShare (10.6084/m9.figshare.25749276). This includes transcriptome  
634 assemblies, predicted proteomes, alignments (trimmed and untrimmed), as well as  
635 phylogenetic trees.

636

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649

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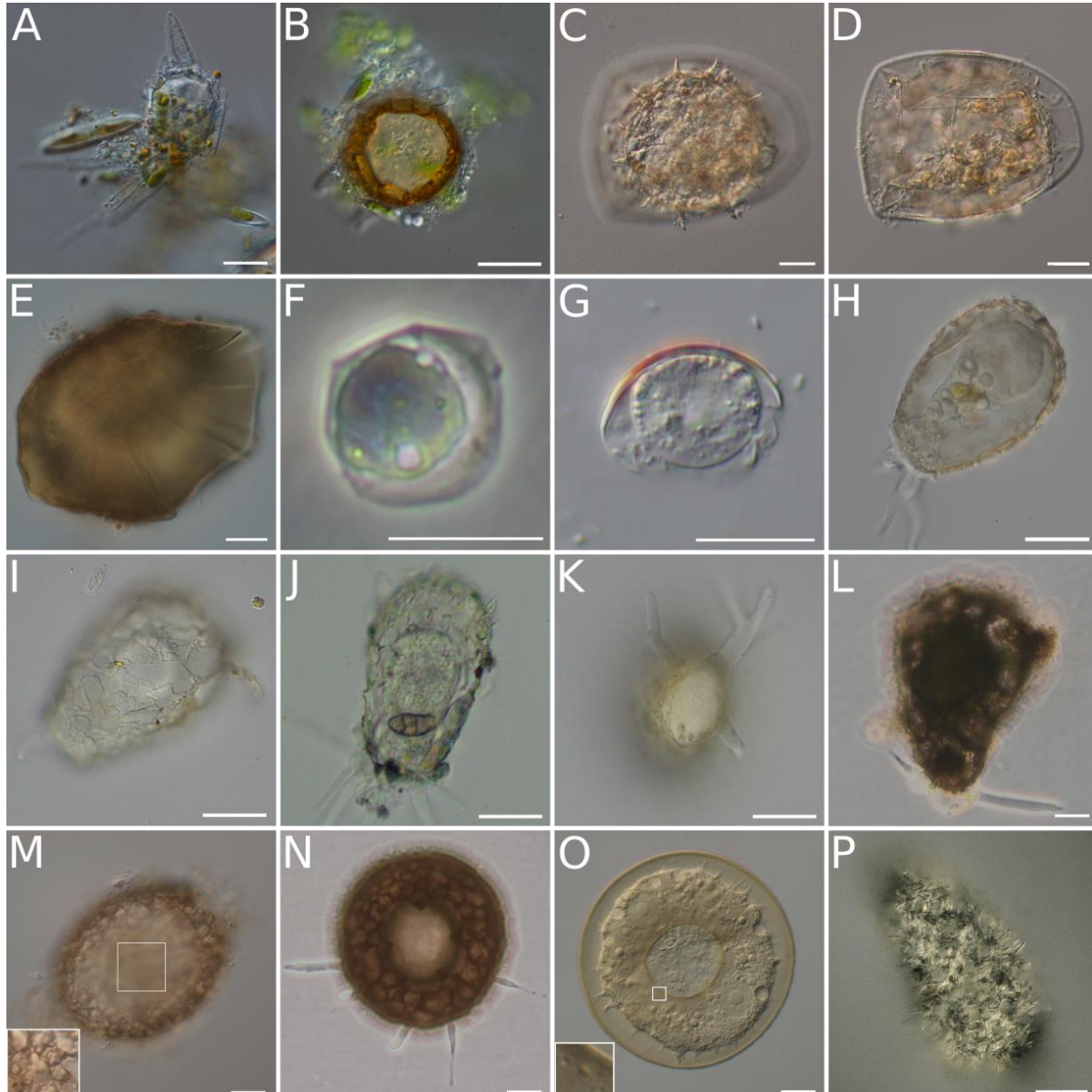
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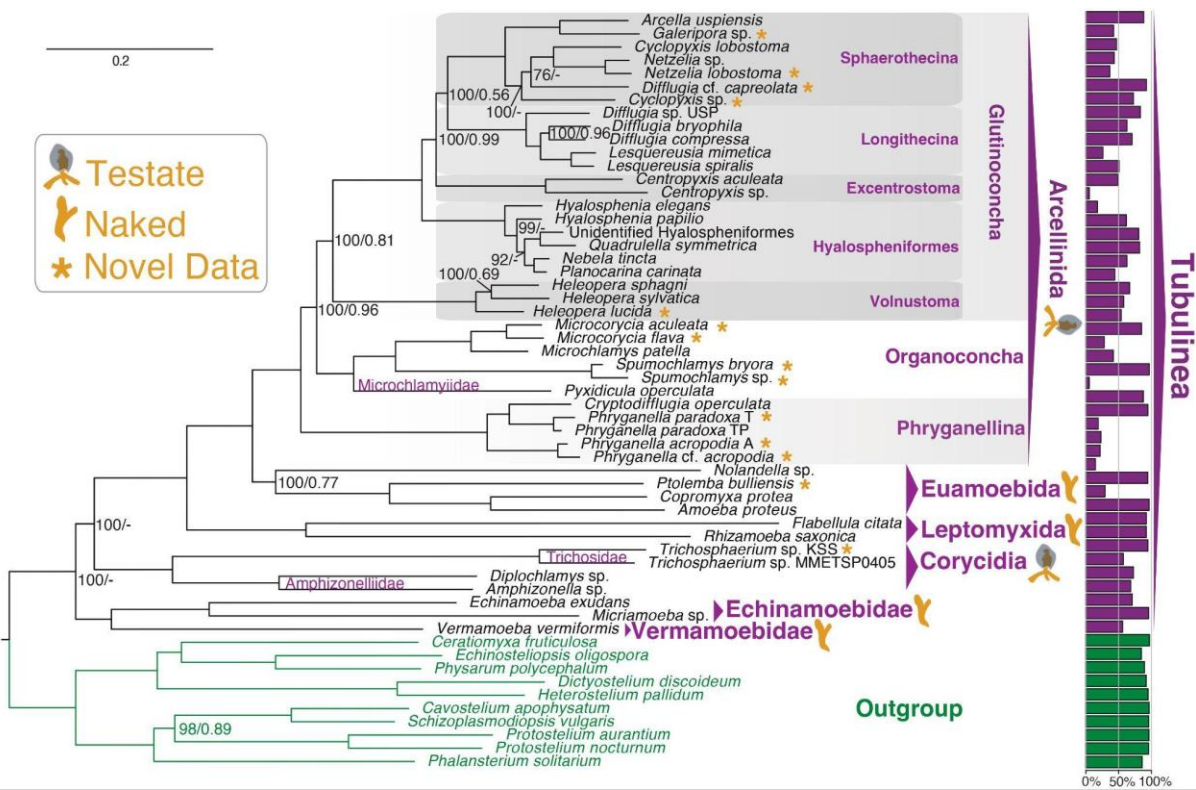
884 **Figures**



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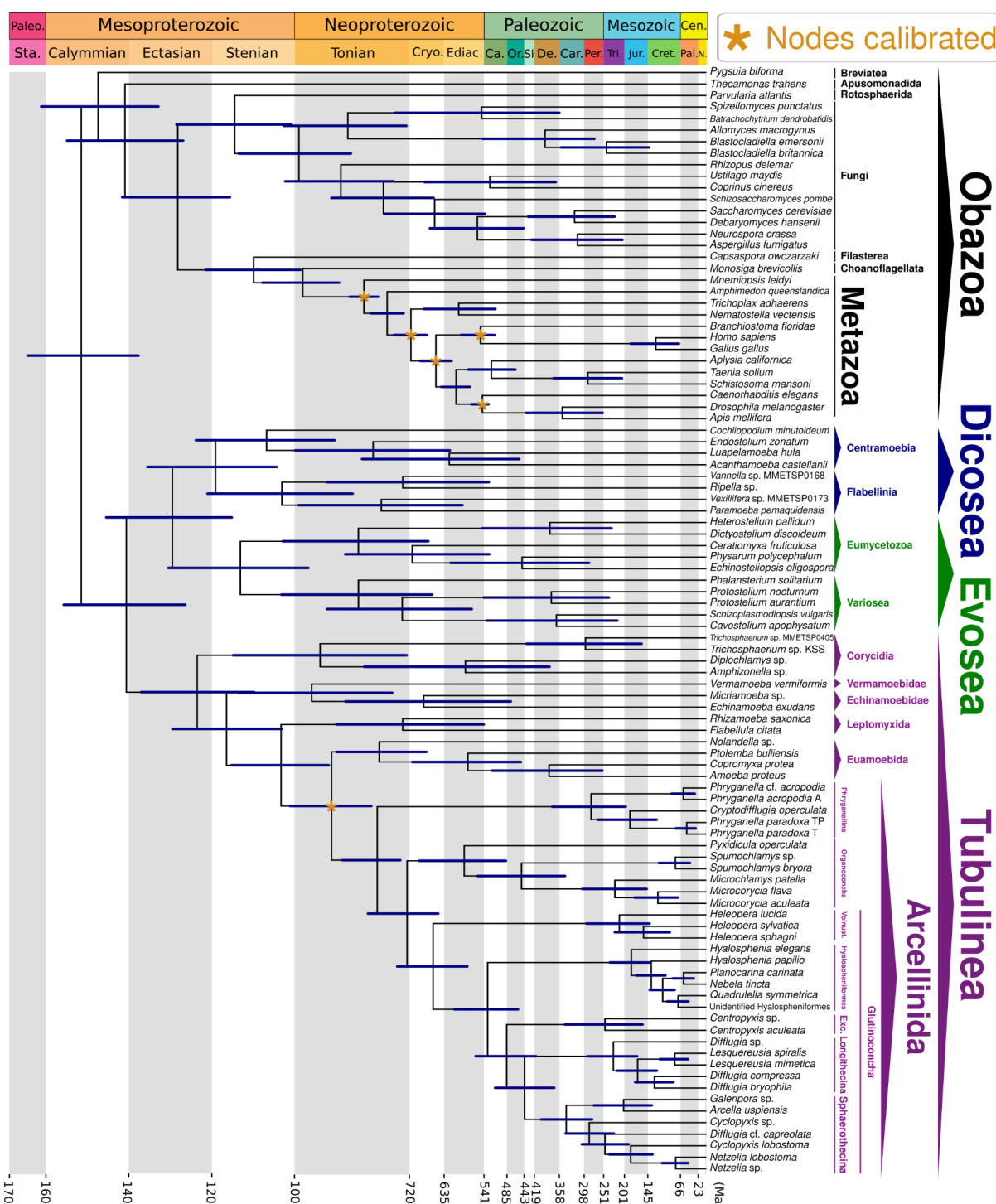
886 **Figure 1. Sampled testate amoebae — Arcellinida and Corycidia.** The pictured  
887 organisms were photodocumented prior to molecular processing and represent  
888 individuals or cultures from which the transcriptomic data was obtained. The scale  
889 bars represent 20  $\mu\text{m}$ , except when specified. **A.** *Phryganella paradoxa* T, lateral  
890 view. **B.** *Phryganella acropodia* A, apertural view. **C - D.** *Microcorycia aculeata*,  
891 dorsal view (C) and apertural view (D); **E.** *Microcorycia flava*, dorsal view focusing on  
892 the flexible part of the shell; **F.** *Spumochlamys* sp., dorsal view; **G.** *Spumochlamys*  
893 *bryora*, lateral view; **H - K.** *Heleopera lucida* comb. nov. (previously *Diffugia lucida*),  
894 lateral view focusing on the cell within the shell (H), lateral view focusing on the shell  
895 (I - J), and apertural view focusing on the compressed aspect of the shell (K); **L.**  
896 *Diffugia* cf. *capreolata*, lateral view, scale bar 40  $\mu\text{m}$ ; **M.** *Netzelia lobostoma*, lateral  
897 view, white square focusing on details of the shell; **N.** *Cyclopyxis* sp., apertural view,  
898 scale bar 40  $\mu\text{m}$ ; **O.** *Galeripora* sp., apertural view, white square focusing on the  
899 pores which surround the shell aperture; **P.** *Trichosphaerium* sp. KSS. Measured

900 morphometric characteristics of the newly sequenced testate amoebae taxa are  
901 present on **Dataset S01, Table S2**.



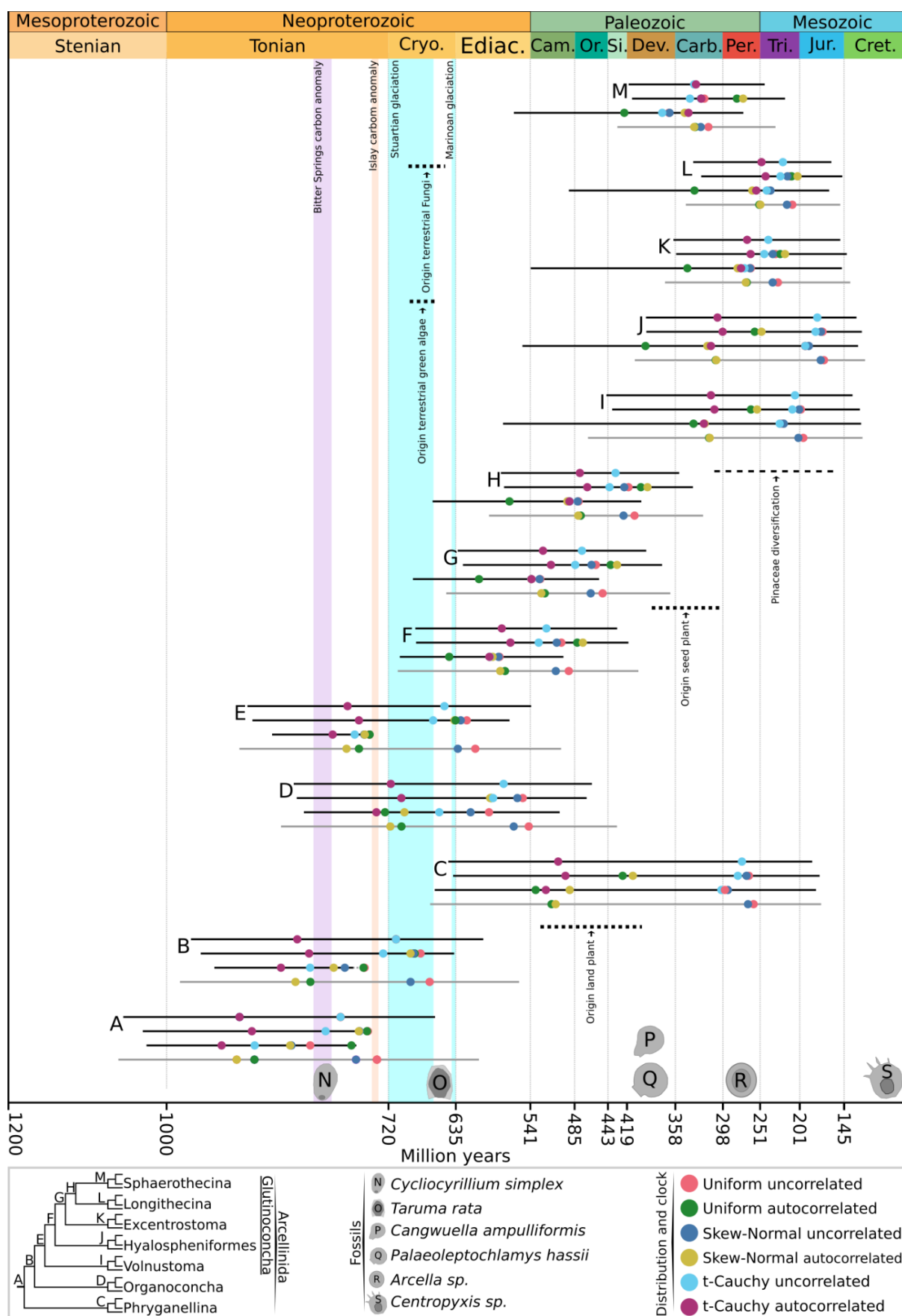
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903 **Figure 2. The tree of amoebozoan testate amoebae.** 226 genes (70,428 amino  
904 acid sites) phylogeny of amoebozoan testate amoebae rooted with Evosea  
905 (Amoebozoa). The tree was initially built using IQ-TREE2 v. 2.0-rc1 under the  
906 LG+C20+G4 model of protein evolution and further used to infer a Posterior Means  
907 Site Frequency model using the ML model LG+C60+G4+PMSF. Topological support  
908 was assessed by 100 Maximum Likelihood Real Bootstrap Replicates (MLRB) and  
909 local posterior probability values (LPP) calculated using ASTRAL-III v. 5.7.3, and are  
910 shown in the format (MLRB/LPP).

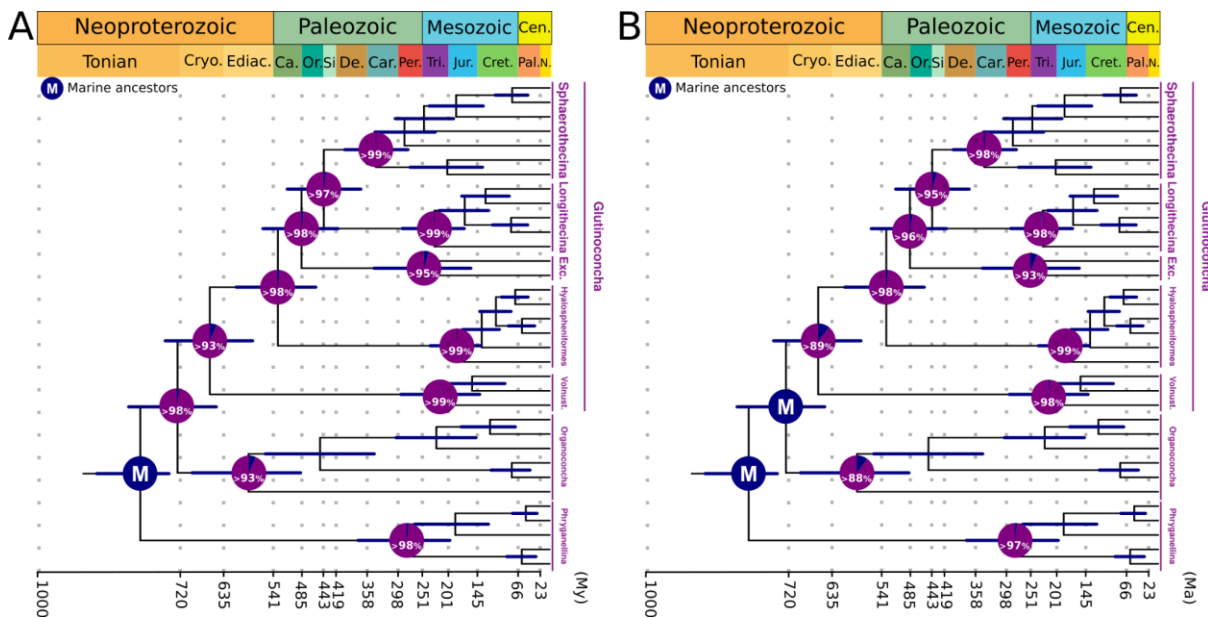


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912 **Figure 3. Amorphea time-calibrated tree inferred under autocorrelated relaxed**  
913 **clock model, applying a truncated-Cauchy distribution for node calibration and**  
914 **drift parameter of  $\alpha = 2$  and  $\beta = 2$ , considering VSMs as stem Arcellinida to**  
915 **calibrate Arcellinida+Euamoebida node.** Bars at nodes are 95% highest  
916 probability density confidence intervals (HPD CI). Asterisks indicate the nodes  
917 calibrated based on external fossil information. Out of the 36 time-calibrated trees  
918 generated, we display here the one representing the analysis that estimated the  
919 youngest minimum 95% HPD CI for the two earliest nodes of Arcellinida, thus  
920 showing the youngest age estimated for the origin of nodes leading to extant  
921 members of the Arcellinida Order. The results and time-calibrated trees for all

922 experiments are present in the supplemental material (**Dataset S01, Table S11 and**  
923 **Appendix S01, Fig. S6 - S43**). Abbreviations: Exc.- Excentrotoma; Vonust. -  
924 Volnustoma; Paleo. - Paleoproterozoic; Sta. - Statherian; Cryo. - Cryogenian; Ediac.  
925 - Ediacaran; Ca. - Cambrian; Or. Ordovician; Si - Silurian; De. - Devonian; Car.  
926 Carboniferous; Per. - Permian; Tri. - Triassic; Jur. - Jurassic; Cret. - Cretaceous;  
927 Cen. - Cenozoic; Pal. - Paleogene; N. - Neogene; My - Million Years.

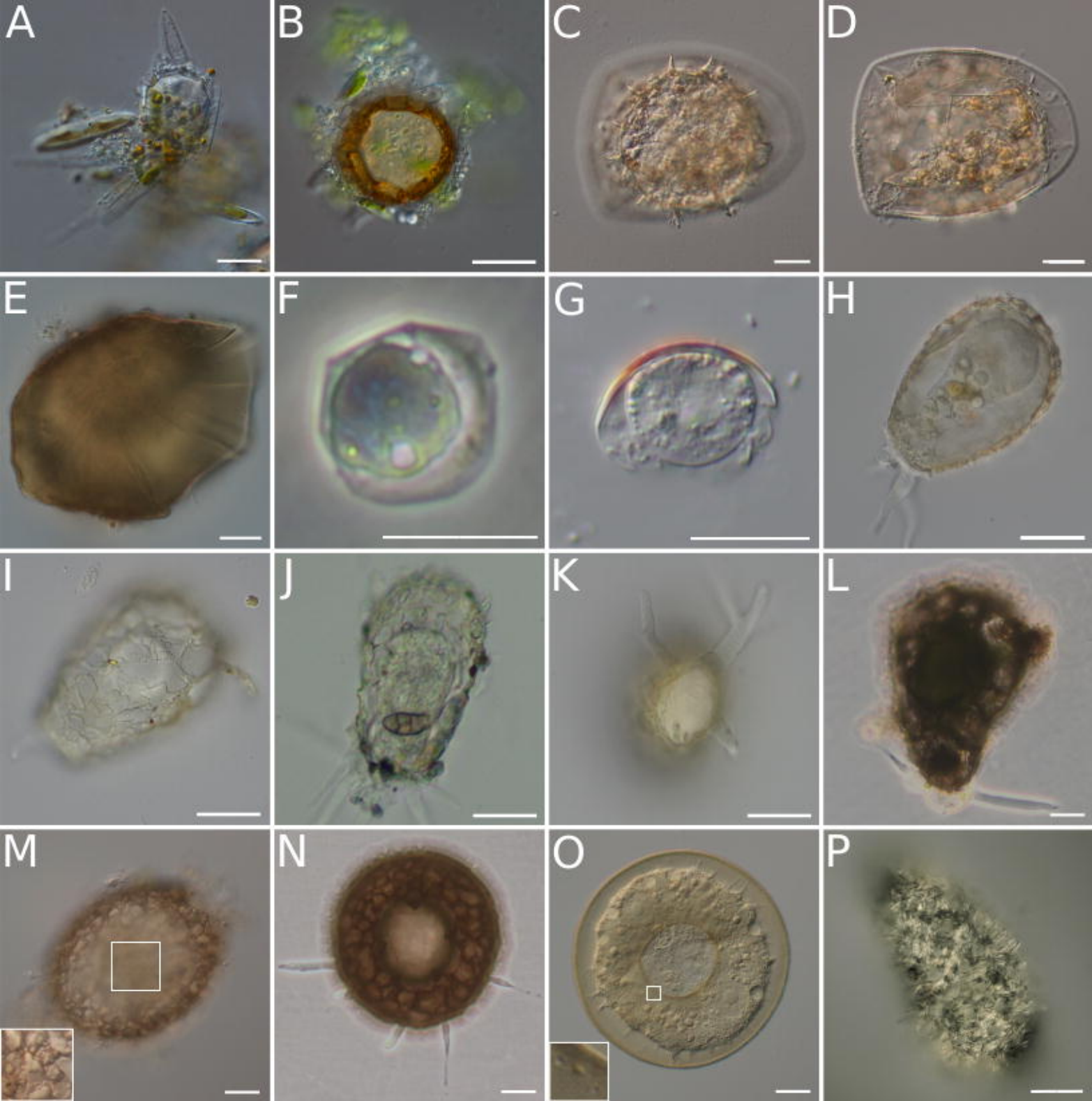


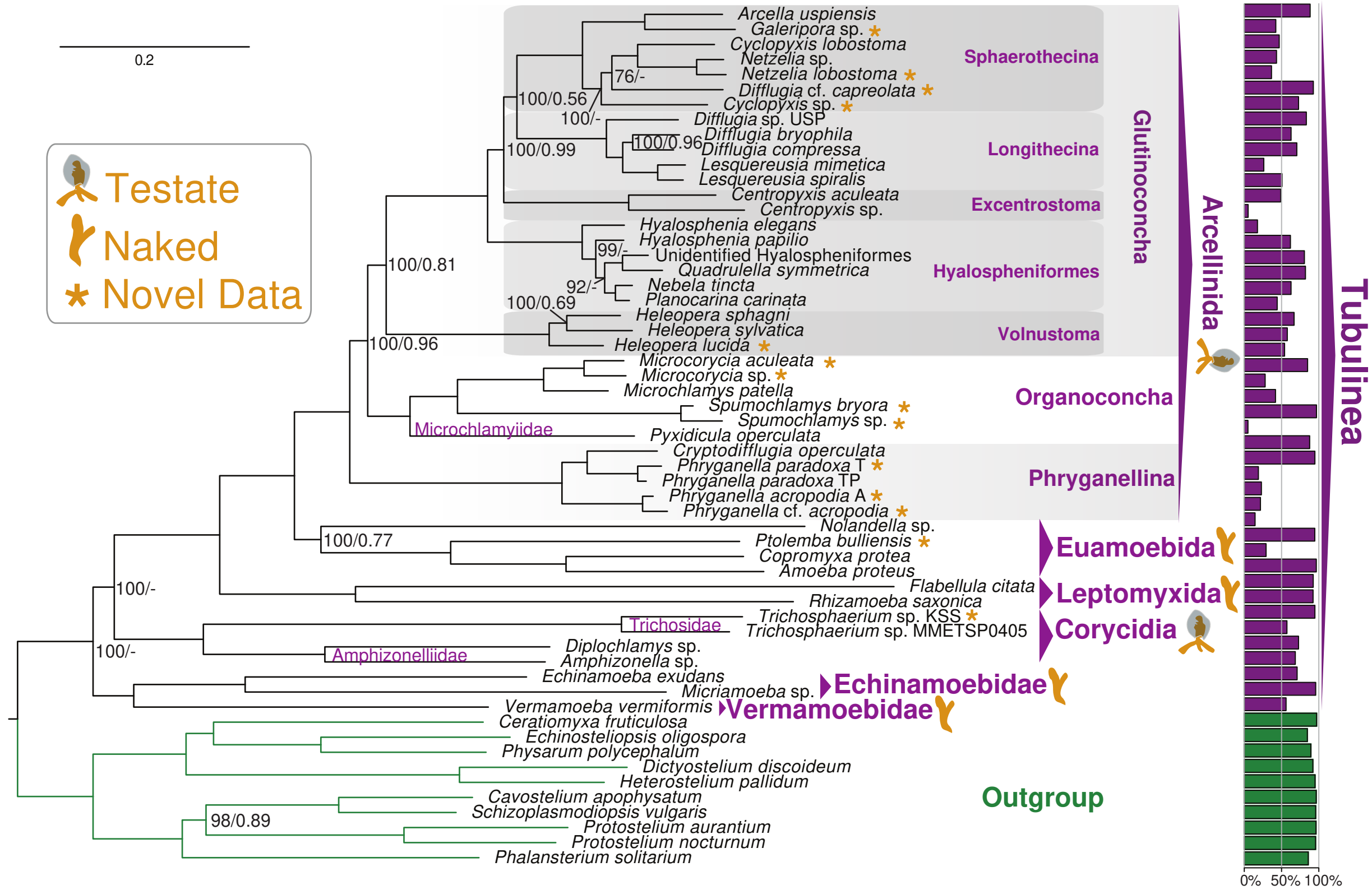
932 suggested by previous studies (horizontal dotted lines; 75). Displayed for each node  
933 are four bars representing, from bottom to top, the calibration strategy not  
934 considering VSM record to calibrate amoebozoan nodes (gray bar), the calibration  
935 strategy considering VSMs as derived crown Arcellinida, calibration strategy  
936 considering VSMs as basal crown Arcellinida, and calibration strategy considering  
937 VSMs as stem Arcellinida. The bars represent the combination of all 95% highest  
938 probability density confidence intervals estimated by each distribution-clock model  
939 considered and the colored dots represent the mean estimated time by each  
940 distribution-clock model. The results and time-calibrated trees for all experiments are  
941 present in the supplemental material (**Dataset S01, Table S11 and Appendix S01**,  
942 **Fig. S6 - S43**).

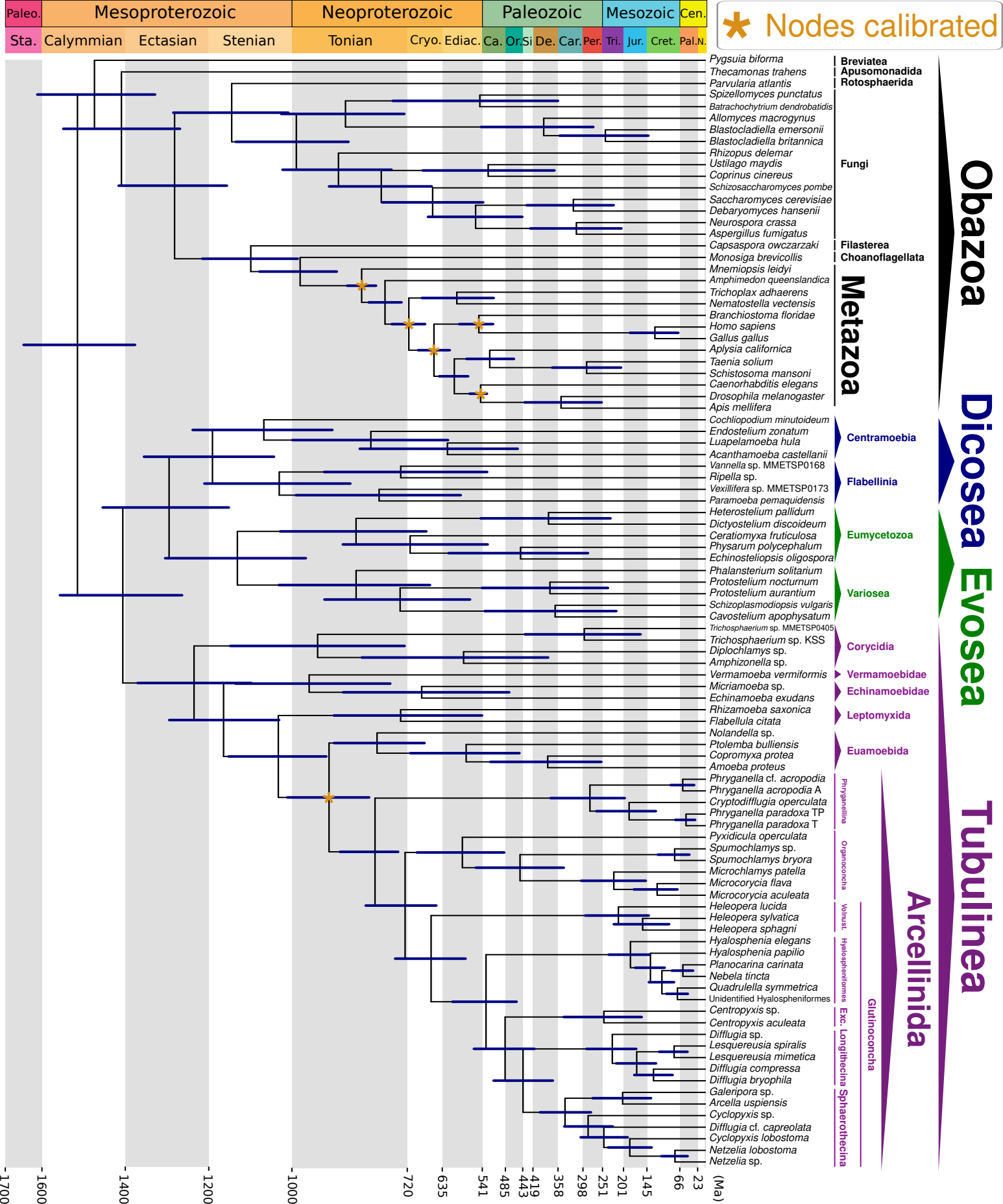


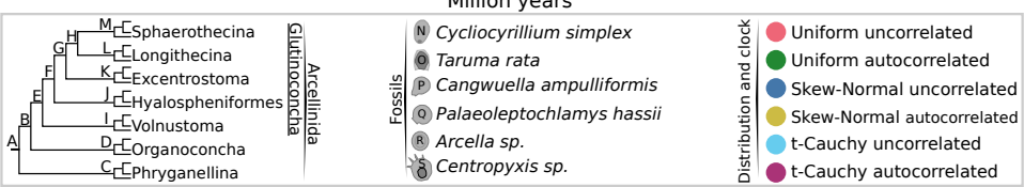
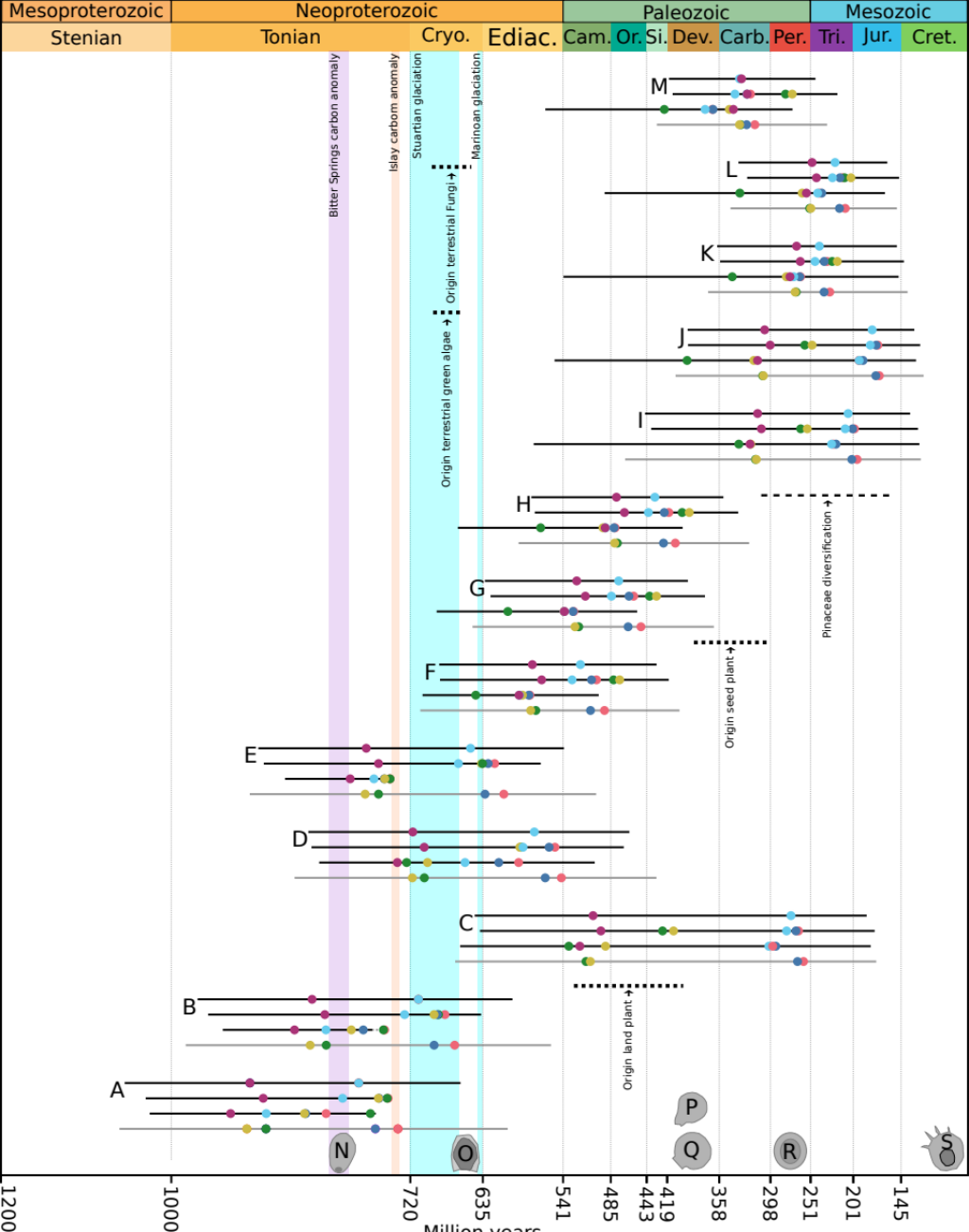
**Figure 5. Ancestral reconstructions of Arcellinida habitats using BayesTraits.**

Pie charts at each node indicate the mean probabilities of a hypothetical marine ancestor (blue) or a hypothetical terrestrial ancestor (purple). **A.** Reconstruction considering Arcellinida ancestor as marine, implying at least two independent transition events (2TE). **B.** Reconstruction considering Arcellinida and Organococoncha+Glutinoconcha ancestors as marine, implying at least three independent transition events (3TE). Bars at nodes are 95% highest probability density confidence intervals estimated by the calibration strategy using an autocorrelated relaxed clock model, applying a truncated-Cauchy distribution for node calibration and drift parameter of  $\alpha = 2$  and  $\beta = 2$ , considering VSMs as stem Arcellinida to calibrate Arcellinida+Euamoebida node. The complete results are shown on **Dataset S01, Table S12 and Appendix S01, Fig. S44**.

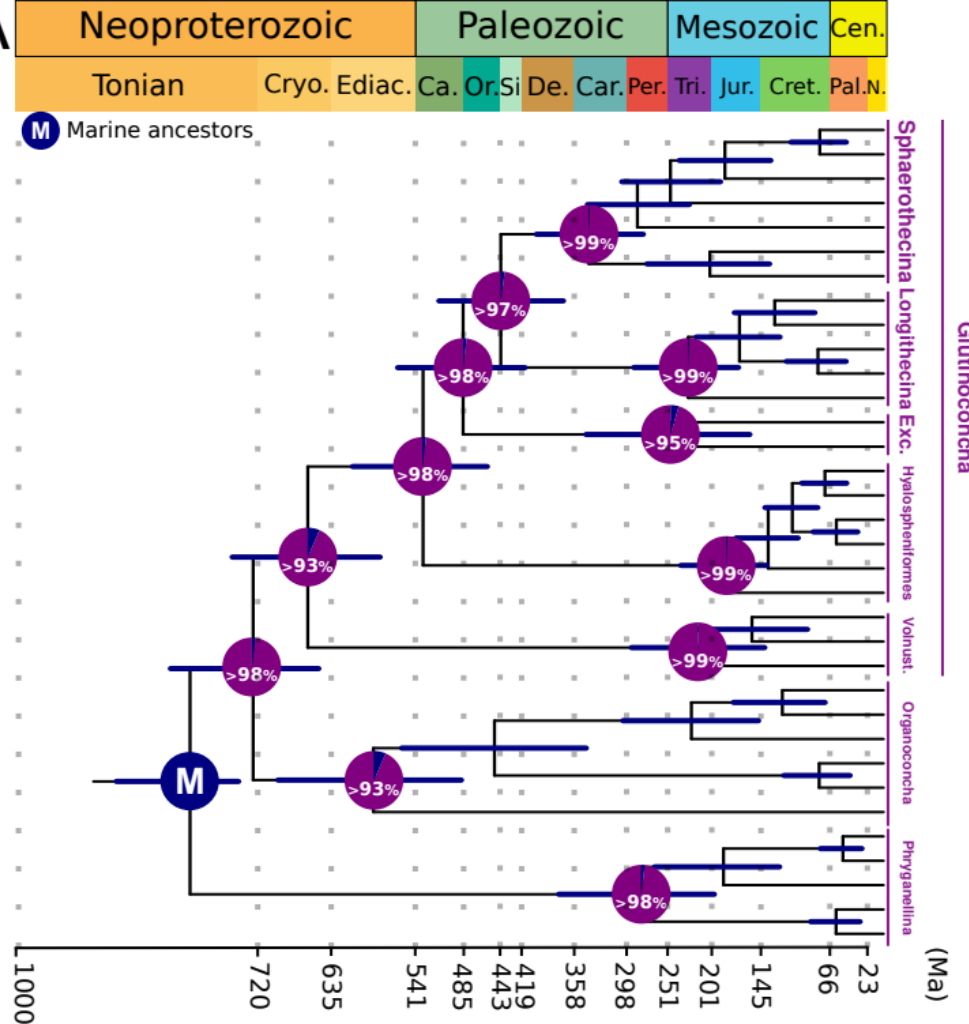








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