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# An estimate of the deepest branches of the tree of life from ancient vertically-evolving genes

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## 18 Abstract

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20 Core gene phylogenies provide a window into early evolution, but different gene sets and 21 analytical methods have yielded substantially different views of the tree of life. Trees inferred 22 from a small set of universal core genes have typically supported a long branch separating the archaeal and bacterial domains. By contrast, recent analyses of a broader set of non-23 24 ribosomal genes have suggested that Archaea may not be very divergent from Bacteria, and 25 that estimates of inter-domain distance are inflated due to accelerated evolution of ribosomal 26 proteins along the inter-domain branch. Resolving this debate is key to determining the 27 diversity of the archaeal and bacterial domains, the shape of the tree of life, and our 28 understanding of the early course of cellular evolution. Here, we investigate the evolutionary 29 history of the marker genes key to the debate. We show that estimates of a reduced Archaea-30 Bacteria (AB) branch length result from inter-domain gene transfers and hidden paralogy in 31 the expanded marker gene set, which act to artifactually diminish the genetic distance between the two domains. By contrast, analysis of a broad range of manually curated marker gene 32 33 datasets from a sample of 700 Archaea and Bacteria reveal that current methods likely 34 underestimate the AB branch length due to substitutional saturation and poor model fit; that 35 the best-performing phylogenetic markers tend to support longer inter-domain branch lengths; and that the AB branch lengths of ribosomal and non-ribosomal marker genes are statistically 36 37 indistinguishable. A phylogeny of prokaryotes inferred from the 27 highest-ranked marker 38 genes, including ribosomal and non-ribosomal markers, supported a long AB branch, 39 recovered a clade of DPANN at the base of the Archaea, and placed CPR within Bacteria as 40 the sister group to the Chloroflexota.

### 41 Introduction

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Much remains unknown about the earliest period of cellular evolution and the deepest 43 44 divergences in the tree of life. Phylogenies encompassing both Archaea and Bacteria have 45 been inferred from a "universal core" set of 16-56 genes encoding proteins involved in 46 translation and other aspects of the genetic information processing machinery(Ciccarelli et al., 47 2006; Fournier and Gogarten, 2010; Harris et al., 2003; Hug et al., 2016; Mukheriee et al., 48 2017; Petitjean et al., 2014; Ramulu et al., 2014; Raymann et al., 2015; Theobald, 2010; 49 Williams et al., 2020). These genes are thought to predominantly evolve vertically and are thus best-suited for reconstructing the tree of life (Ciccarelli et al., 2006; Creevey et al., 2011; 50 51 Ramulu et al., 2014; Theobald, 2010). In these analyses, the branch separating Archaea from 52 Bacteria (hereafter, the AB branch) is often the longest internal branch in the tree(Cox et al., 53 2008; Gogarten et al., 1989; Hug et al., 2016; Iwabe et al., 1989; Pühler et al., 1989; Williams 54 et al., 2020). In molecular phylogenetics, branch lengths are usually measured in expected 55 numbers of substitutions per site, with a long branch corresponding to a greater degree of 56 genetic change. Long branches can therefore result from high evolutionary rates, long periods of absolute time, or a combination of the two. If a sufficient number of fossils are available to 57 58 calibrate them then molecular clock models can, in principle, disentangle the contributions of 59 these effects. However, only very few fossil calibrations (Sugitani et al., 2015) are currently 60 available that are old enough to calibrate early divergences (Betts et al., 2018; Horita and 61 Berndt, 1999; Lepland et al., 2002; van Zuilen et al., 2002), and as a result, the ages and 62 evolutionary rates of the deepest branches of the tree, and estimates of the true biodiversity 63 of the archaeal and bacterial domains, remain highly uncertain.

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65 Recently, Zhu et al. (Zhu et al., 2019) inferred a phylogeny from 381 genes distributed across Archaea and Bacteria using the supertree method ASTRAL (Mirarab et al., 2014). In addition 66 67 to a large increase in the number of genes compared to other universal marker sets, the 68 functional profile of these markers comprises not only proteins involved in information 69 processing but also proteins affiliated with most other functional COG categories, including 70 metabolic processes (Table S1). The genetic distance (branch length) between the domains 71 (Zhu et al., 2019) was estimated from a concatenation of the same marker genes, resulting in 72 a much shorter AB branch length than observed with the core universal markers (Hug et al., 73 2016; Williams et al., 2020). These analyses were consistent with the hypothesis (Petitjean et 74 al., 2014; Zhu et al., 2019) that the apparent deep divergence of Archaea and Bacteria might 75 be the result of an accelerated evolutionary rate of genes encoding translational and in 76 particular ribosomal proteins along the AB branch as compared to other genes. Interestingly, 77 the same observation was made previously using a smaller set of 38 non-ribosomal marker 78 proteins (Petitjean et al., 2014), although the difference in AB branch length between 79 ribosomal and non-ribosomal markers in that analysis was reported to be substantially lower 80 (roughly two-fold, compared to roughly ten-fold for the 381 protein set (Petitjean et al., 2014; 81 Zhu et al., 2019).

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A higher evolutionary rate of ribosomal genes might result from the accumulation of compensatory substitutions at the interaction surfaces among the protein subunits of the ribosome (Petitjean et al., 2014; Valas and Bourne, 2011), or as a compensatory response to the addition or removal of ribosomal subunits early in evolution (Petitjean et al., 2014). 87 Alternatively, differences in the inferred AB branch length might result from varying rates or 88 patterns of evolution between the traditional core genes (Spang et al., 2015; Williams et al., 89 2020) and the expanded set (Zhu et al., 2019). Substitutional saturation (multiple substitutions 90 at the same site (Jeffroy et al., 2006)) and across-site compositional heterogeneity can both 91 impact the inference of tree topologies and branch lengths (Foster, 2004; Lartillot et al., 2007; Lartillot and Philippe, 2004; Quang et al., 2008; Wang et al., 2008). These difficulties are 92 93 particularly significant for ancient divergences (Gouy et al., 2015). Failure to model site-94 specific amino acid preferences has previously been shown to lead to under-estimation of the 95 AB branch length due to a failure to detect convergent changes (Tourasse and Gouy, 1999; Williams et al., 2020), although the published analysis of the 381 marker set did not find 96 97 evidence of a substantial impact of these features on the tree as a whole (Zhu et al., 2019). 98 Those analyses also identified phylogenetic incongruence among the 381 markers, but did 99 not determine the underlying cause (Zhu et al., 2019).

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101 This recent work (Zhu et al., 2019) raises two important issues regarding the inference of the 102 universal tree: first, that estimates of the genetic distance between Archaea and Bacteria from 103 classic "core genes" may not be representative of ancient genomes as a whole, and second, 104 that there may be many more suitable genes to investigate early evolutionary history than 105 generally recognized, providing an opportunity to improve the precision and accuracy of deep 106 phylogenies. Here, we investigate these issues in order to determine why different marker sets 107 support different Archaea-Bacteria branch lengths. First, we examine the evolutionary history of the 381 gene marker set (hereafter, the expanded marker gene set) and identify several 108 109 features of these genes, including instances of inter-domain gene transfers and mixed 110 paralogy, that may contribute to the inference of a shorter AB branch length in supertree and 111 concatenation analyses. Then, we re-evaluate the marker gene sets used in a range of 112 previous analyses to determine how these and other factors, including substitutional saturation and model fit, contribute to inter-domain branch length estimations and the shape of the 113 114 universal tree. Finally, we identify a subset of marker genes least affected by these issues, 115 and use these to estimate an updated tree of the primary domains of life and the genetic 116 distance between Archaea and Bacteria.

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## 117 Results and Discussion

# Gene transfers and hidden paralogy obscure the genetic distance between Archaea and Bacteria

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#### 121 Genes from the expanded marker set are not widely distributed in Archaea

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123 The 381 gene set was derived from a larger set of 400 genes used to estimate the phylogenetic 124 placement of new lineages as part of the PhyloPhIAn method (Segata et al., 2013). Perhaps 125 reflecting the focus on bacteria in the original application, the phylogenetic distribution of the 126 381 marker genes in the expanded set varies substantially (Table S1), with many being poorly represented in Archaea. Indeed 25% of the published gene trees (https://biocore.github.io/wol/ 127 128 (Zhu et al., 2019)) contain less than 0.5% archaeal homologues, with 21 (5%) and 69 (18%) 129 of these trees including no or less than 10 archaeal homologues, respectively. For the 130 remaining 75% of the gene trees, archaeal homologs comprise 0.5%-13.4% of the dataset. 131 While there are many more sequenced bacteria than archaea, 63% of the gene trees 132 possessed genes from less than half of the 669 archaeal genomes included in the analysis, 133 whereas only 22% of the gene trees possessed fewer than half of the total number of 9906 134 sampled bacterial genomes. These distributions suggest that many of these genes are not 135 broadly present in both domains, and that some might be specific to Bacteria.

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# Conflicting evolutionary histories of individual marker genes and the inferred species tree

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141 In the focal analysis of the 381 gene set, the tree topology was inferred using the supertree 142 method ASTRAL (Mirarab et al., 2014), with branch lengths inferred on this fixed tree from a 143 marker gene concatenation (Zhu et al., 2019). The topology inferred from this expanded 144 marker set (Zhu et al., 2019) is similar to published trees (Castelle and Banfield, 2018; Hug et 145 al., 2016) and recovers Archaea and Bacteria as reciprocally monophyletic domains, albeit 146 with a shorter AB branch than in earlier analyses. However, the individual gene trees (Zhu et 147 al., 2019) disagree regarding domain monophyly: Archaea and Bacteria are recovered as 148 reciprocally monophyletic groups in only 24 of the 381 published (Zhu et al., 2019) maximum 149 likelihood (ML) gene trees of the expanded marker set (Table S1).

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151 Since single gene trees often fail to strongly resolve ancient relationships, we used 152 approximately-unbiased (AU) tests (Shimodaira, 2002) to evaluate whether the failure to 153 recover domain monophyly in the published ML trees is statistically supported. For 154 computational tractability, we performed these analyses on a 1000-species subsample of the 155 full 10,575-species dataset that was compiled in the original study (Zhu et al., 2019). For 79 156 of the 381 genes, we could not perform the test because the gene was not found on any of the 74 archaeal genomes present in the 1000-species subsample. For the remaining 302 157 158 genes, domain monophyly was rejected (p < 0.05) for 232 out of 302 (76.8%) genes. As a 159 comparison, we performed the same test on several smaller marker sets used previously to 160 infer a tree of life (Coleman et al., 2021; Petitjean et al., 2014; Williams et al., 2020); none of bioRxiv preprint doi: https://doi.org/10.1101/2021.01.19.427276; this version posted May 7, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY 4.0 International license.

161 the markers in those sets rejected reciprocal domain monophyly (p > 0.05 for all genes, Figure 162 1(a)). In what follows, we refer to four published marker gene sets as: the Expanded set (381 genes (Zhu et al., 2019)), the Core set (49 genes (Williams et al., 2020), encoding ribosomal 163 164 proteins and other conserved information-processing functions; itself a consensus set of several earlier studies (Da Cunha et al., 2017; Spang et al., 2015; Williams et al., 2012)), the 165 166 Non-ribosomal set (38 genes, broadly distributed and explicitly selected to avoid genes 167 encoding ribosomal proteins (Petitjean et al., 2014)), and the Bacterial set (29 genes used in 168 a recent analysis of bacterial phylogeny (Coleman et al., 2021)).

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170 To investigate why 232 of the marker genes rejected the reciprocal monophyly of Archaea and 171 Bacteria, we returned to the full dataset (Zhu et al., 2019), annotated each sequence in each 172 marker gene family by assigning proteins to KOs, Pfams, and Interpro domains, among others 173 (Table S1, see Methods for details) and manually inspected the tree topologies (Table S1). 174 This revealed that the major cause of domain polyphyly observed in gene trees was inter-175 domain gene transfer (in 357 out of 381 gene trees (93.7%)) and mixing of sequences from 176 distinct paralogous families (in 246 out of 381 gene trees (64.6%)). For instance, marker 177 genes encoding ABC-type transporters (p0131, p0151, p0159, p0174, p0181, p0287, p0306, (i.e. p0000, p0011, p0020, p0091, p0094, p0202), 178 po0364), tRNA synthetases 179 aminotransferases and dehydratases (i.e. p0073/4-aminobutyrate aminotransferase; 180 p0093/3-isopropylmalate dehydratase) often comprised a mixture of paralogues.

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182 Together, these analyses indicate that the evolutionary histories of the individual markers of 183 the expanded set differ from each other and from the species tree. Zhu et al. acknowledged 184 (Zhu et al., 2019) the varying levels of congruence between the marker phylogenies and the 185 species tree, but did not investigate the underlying causes. Our analyses establish the basis 186 for these disagreements in terms of gene transfers and the mixing of orthologues and paralogues within and between domains. Concatenation is based on the assumption that all 187 188 of the genes in the supermatrix evolve on the same underlying tree; genes with different gene 189 tree topologies violate this assumption and should not be concatenated because the 190 topological differences among sites are not modelled, and so the impact on inferred branch 191 lengths is difficult to predict. In practice, it is often difficult to be certain that all of the markers 192 in a concatenate share the same gene tree topology, and the analysis proceeds on the 193 hypothesis that a small proportion of discordant genes are not expected to seriously impact 194 the inferred tree. However, the concatenated tree inferred from the expanded marker set 195 differs from previous trees in that the genetic distance between Bacteria and Archaea is greatly 196 reduced, such that the AB branch length appears comparable to distances among bacterial 197 phyla (Zhu et al., 2019). Because an accurate estimate of the AB branch length has a major 198 bearing on unanswered questions regarding the root of the universal tree (Gouy et al., 2015), 199 we next evaluated the impact of the conflicting gene histories within the expanded marker set 200 on inferred AB branch length.

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# 202The inferred branch length between Archaea and Bacteria is artifactually shortened by203inter-domain gene transfer and hidden paralogy

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To investigate the impact of gene transfers and mixed paralogy on the AB branch length inferred by gene concatenations (Zhu et al., 2019), we compared branch lengths estimated from markers that rejected (AU < 0.05) or did not reject (AU > 0.05) the reciprocal monophyly of Bacteria and Archaea in the 381 marker set (Figure 1(a)). To estimate AB branch lengths

209 for genes in which the domains were not monophyletic in the ML tree, we first performed a 210 constrained ML search to find the best gene tree that was consistent with domain monophyly for each family under the LG+G4+F model in IQ-TREE 2 (Minh et al., 2020). While it may 211 212 seem strained to estimate the length of a branch that does not appear in the ML tree, we 213 reasoned that this approach would provide insight into the contribution of these genes to the 214 AB branch length in the concatenation, in which they conflict with the overall topology. AB 215 branch lengths were significantly ( $P = 2.159 \times 10^{-12}$ , Wilcoxon rank sum test) shorter for markers 216 that rejected domain monophyly (Figure 1(a); <0.05: mean AB branch length in expected 217 substitutions/site 0.0130, >0.05: mean AB branch length 0.559). This result suggests that 218 inter-domain gene transfers reduce the AB branch length when included in a concatenation. 219 This behaviour might result from marker gene transfers reducing the number of fixed 220 differences between the domains, so that the AB branch length in a tree in which Archaea and 221 Bacteria are constrained to be reciprocally monophyletic will tend towards 0 as the number of 222 transfers increases. Consistent with this hypothesis, we observed that  $\Delta LL$ , the difference in 223 log likelihood between the constrained ML tree and ML gene tree (used here as a proxy for 224 gene verticality), correlates negatively with AB branch length (Figure 1(b)). Furthermore, AB 225 branch length decreased as increasing numbers of low-verticality markers were added to the 226 concatenate (Figure 1(c)). Taken together, these results indicate that the inclusion of genes 227 that do not support the reciprocal monophyly of Archaea and Bacteria in the universal 228 concatenate reduces the estimated AB branch length by homogenizing the genetic diversity 229 of the two domains.



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Figure 1: Expanded set genes in which Archaea and Bacteria are not monophyletic support a shorter AB branch. (a) Expanded set genes that reject domain monophyly (p < 0.05, AU test) support significantly shorter AB branch lengths when constrained to follow a domain monophyletic tree ( $p = 2.159 \times 10-12$ , Wilcoxon rank-sum test). None of the marker genes from several other published analyses reject domain monophyly (p > 0.05, AU test) for all genes tested. (b) Marker gene verticality ( $\Delta LL$ , see below) for the expanded gene set normalized by alignment length correlates negatively with the length of the AB branch between Archaea and Bacteria (R2=0.03998, p = 0.0004731). (c) Concatenations of 20-100 markers of the expanded set markers ranked by marker gene verticality ( $\Delta LL$ ) show the same trend, with a reduction in AB branch length as markers with a greater  $\Delta LL$  are added to the concatenate.  $\Delta LL$  is the difference between the log likelihood of the ML gene family tree under a free topology search and the log likelihood of the best tree constrained to obey domain monophyly. The trendline is estimated using LOESS regression.

# The age of the last universal common ancestor (LUCA) inferred from strict clocks does not predict marker gene quality

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233 Reliable age estimates using molecular clock methods require calibrations, but few 234 calibrations exist for the deeper branches of the tree of life. Zhu et al. (Zhu et al., 2019) argued 235 that the expanded marker set is useful for deep phylogeny because estimates of the age of 236 the last universal common ancestor (LUCA) obtained by fitting molecular clocks to their 237 dataset are in agreement with the geological record: a root (LUCA) age of 3.6-4.2 Ga was 238 inferred from the entire 381-gene dataset, consistent with the earliest fossil evidence for life 239 (Betts et al., 2018; Sugitani et al., 2015), whereas estimates from ribosomal markers alone supported a root age of 7 Ga. This age might be considered implausible because it is much 240 241 older than the age of the Earth and Solar System (with the moon-forming impact occurring 242 ~4.51 Ga (Barboni et al., 2017; Hanan and Tilton, 1987)). However, the palaeobiological 243 plausibility of the age estimate from the 381 gene set does not, in itself, constitute evidence of 244 marker gene suitability. In the original analyses, the age of LUCA was estimated using a 245 maximum likelihood approach, as well as a Bayesian molecular clock with a strict clock 246 (assuming a constant evolutionary rate) or a relaxed clock with a single calibration. A strict 247 clock model does not permit changes in evolutionary rate through time or across branches, 248 and so a longer AB branch will lead to an older inferred LUCA age. Likewise, a relaxed clock 249 model with a single calibration may fail to distinguish molecular distances and geological time. 250 Given that the short AB branch in the expanded gene set results, in part, from phylogenetic incongruence among markers, we evaluated the age of LUCA inferred from the subset of the 251 252 expanded gene set least affected by these issues. To do so, we analysed the top 5% of gene 253 families according to their  $\Delta LL$  score (a set of 20 genes, which includes only 1 ribosomal 254 protein) under the same clock model parameters as the original dataset (Figure 2). This 255 analysis resulted in a significantly more ancient age estimate for LUCA (5.5-6.5 Ga), and 256 trimming the alignment to remove poorly-aligning regions resulted in a still older estimate 257 (6.34-6.89 Ga), approaching that of the ribosomal genes (7.46-8.03 Ga). These analyses 258 suggest that, for these data and calibrations, the inferred age for LUCA is not a reliable 259 indicator of marker quality, because analyses using the subset of the data least affected by 260 incongruence more clearly reveals the underlying limitations of strict clock analyses (and 261 indeed relaxed clocks with few calibrations) for dating ancient divergences. In principle, more 262 reliable estimates of LUCA's age might be obtained by using more calibrations. However, 263 unambiguous calibrations remain elusive, particularly for the root and other deep branches of 264 the tree. Despite advances in molecular clock methodology, such calibrations represent the 265 only way to reliably capture the relationship between genetic distance and divergence time. 266



Figure 2. The inferred age of LUCA is not a reliable indicator of marker quality. Posterior node age estimates from Bayesian molecular clock analyses of 1) Conserved sites as estimated previously (Zhu et al., 2019); 2) Random sites (Zhu et al., 2019) 3) Ribosomal genes (Zhu et al., 2019) 4) The top 5% of marker gene families according to their  $\Delta LL$  score (including only 1 ribosomal protein) and 5) The same top 5% of marker genes trimmed using BMGE(Criscuolo and Gribaldo, 2010) to remove highly variable sites. In each case, a strict molecular clock was applied, with the age of the Cyanobacteria-Melainabacteria split constrained between 2.5 and 2.6 Ga.

# Phylogeny of Archaea and Bacteria using ancient vertically evolving genes

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#### 270 Finding ancient vertically-evolving genes

272 To estimate the AB branch length and the phylogeny of prokaryotes using a dataset that 273 resolves some of the issues identified above, we performed a meta-analysis of several 274 previous studies to identify a consensus set of vertically-evolving marker genes. We identified 275 unique markers from these analyses by reference to the COG ontology (Dombrowski et al., 276 2020; Galperin et al., 2019), extracted homologous sequences from a representative sample 277 of 350 archaeal and 350 bacterial genomes, and performed iterative phylogenetics and 278 manual curation to obtain a set of 54 markers that recovered archaeal and bacterial monophyly 279 (see Methods). Subsequently, we ranked these 54 genes by the extent to which they 280 recovered established within-domain relationships using the split score, a criterion described 281 previously (Dombrowski et al., 2020) (see Methods) yielding a final set of 27 markers that were used for inferring an updated universal species tree (see below). Marker genes that better 282 283 resolved relationships within each domain also supported a longer AB branch length (Figure 284 3).

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# Distributions of AB branch lengths for ribosomal and non-ribosomal marker genes are similar

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289 Traditional universal marker sets include many ribosomal proteins (Ciccarelli et al., 2006; 290 Fournier and Gogarten, 2010; Harris et al., 2003; Hug et al., 2016; Williams et al., 2020). If 291 ribosomal proteins experienced accelerated evolution during the divergence of Archaea and 292 Bacteria, this might lead to the inference of an artifactually long AB branch length (Petitjean et 293 al., 2014; Zhu et al., 2019). To investigate this, we plotted the inter-domain branch lengths for 294 the 38 and 16 ribosomal and non-ribosomal genes, respectively, comprising the 54 marker 295 genes set. We found no evidence that there was a longer AB branch associated with ribosomal 296 markers (Figure 4; mean AB branch length for ribosomal proteins 1.35, mean for non-297 ribosomal 2.25). Prior to manual curation, non-ribosomal markers had a greater number of 298 HGTs and cases of mixed paralogy. In particular, for the original set of 95 markers, 62% of 299 the non-ribosomal markers and 21% of the ribosomal markers were not monophyletic, 300 respectively. These values were 69% and 29% for the 54 markers, and 50% and 33% for the 301 27 markers. These results imply that manual curation of marker genes is important for deep 302 phylogenetic analyses, particularly when using non-ribosomal markers.

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*Figure 3. Better phylogenetic markers have longer AB branches.* The plot shows the relationship between split score (a lower split score denotes better recovery of established within-domain relationships, see Methods) and AB branch length (in expected number of substitutions/site) for the 54 highest-ranked marker genes. Marker genes with higher split scores (that split monophyletic groups into multiple subclades) have shorter AB branch lengths (P = 0.0311, r = 0.294). Split scores of ribosomal and non-ribosomal markers were statistically indistinguishable (P = 0.828, Figure S3).

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Figure 4. Among vertically-evolving marker genes, ribosomal genes do not have a longer AB branch length. The plot shows functional classification of markers against AB branch length using the 54 most vertically evolving markers. We did not see a significant (P = 0.619, Wilcoxon rank sum test) difference between AB branch lengths for ribosomal and non-ribosomal genes.

# 304 Substitutional saturation and poor model fit contribute to underestimation of AB branch 305 length

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307 For the top 50% of marker genes as determined by split scores (27 genes), we performed an 308 additional round of single gene tree inference and manual review to identify and remove 309 remaining sequences which had evidence of HGT or represented distant paralogs. The 310 resulting single gene trees are provided in the Data Supplement 311 (10.6084/m9.figshare.13395470). To evaluate the relationship between site evolutionary rate 312 and AB branch length, we created two concatenations: fastest sites (comprising sites with 313 highest probability of being in the fastest Gamma rate category; 868 sites) and slowest sites 314 (sites with highest probability of being in the slowest Gamma rate category, 1604 sites) and 315 compared relative branch lengths inferred from the entire concatenate using IQ-TREE 2 to 316 infer site-specific rates (Figure 5). As expected, total tree length is shorter from the slow-317 evolving sites, but the relative AB branch length is longer (1.2 substitutions/site, or ~2% of 318 total tree length, compared to 2.6 substitutions/site, or ~0.04% total tree length for the fastest-319 evolving sites). This result suggests that, at fast-evolving sites, some changes along the AB 320 branch have been overwritten by later events in evolution --- that is, that substitutional 321 saturation leads to an underestimate of the AB branch length.

322

323 Another factor that has been shown to lead to underestimation of genetic distance on deep 324 branches is a failure to adequately model the site-specific features of sequence evolution 325 (Lartillot and Philippe, 2004; Schrempf et al., 2020; Wang et al., 2018; Williams et al., 2020). 326 Amino acid preferences vary across the sites of a sequence alignment, due to variation in the 327 underlying functional constraints (Lartillot and Philippe, 2004; Quang et al., 2008; Wang et al., 328 2008). The consequence is that, at many alignment sites, only a subset of the twenty possible 329 amino acids are tolerated by selection. Standard substitution models, such as LG+G4+F, are 330 site-homogeneous, and approximate the composition of all sites using the average 331 composition across the entire alignment. Such models underestimate the rate of evolution at 332 highly constrained sites because they do not account for the high number of multiple 333 substitutions that occur at such sites. The effect is that site-homogeneous models 334 underestimate branch lengths when fit to site-heterogeneous data. Site-heterogeneous 335 models have been developed that account for site-specific amino acid preferences, and these 336 generally show improved fit to real protein sequence data (reviewed in (Williams et al., 2021)). 337 To evaluate the impact of substitution model fit for these data, we fit a range of models to the 338 full concatenation, assessing model fit using the Bayesian information criterion (BIC) in IQ-339 TREE 2. The AB branch length inferred under the best-fit model, the site-heterogeneous 340 LG+C60+G4+F model, was 2.52 substitutions/site, ~1.7-fold greater than the branch length 341 inferred from the site-homogeneous LG+G4+F model (1.45 substitutions/site). Thus, 342 substitution model fit has a major effect on the estimated length of the AB branch, with better-343 fitting models supporting a longer branch length (Table 1). The same trends are evident when 344 better-fitting site-heterogeneous models are used to analyse the dataset of Zhu et al.: 345 considering only the top 5% of genes by  $\Delta LL$  score, the AB branch length is 1.2 under 346 LG+G4+F, but increases to 2.4 under the best-fitting LG+C60+G4+F model (Figure S2). 347



**Figure 5.** Slow- and fast-evolving sites support different shapes for the universal tree. (i) Tree of Archaea and Bacteria inferred from a concatenation of 27 core genes; (ii) Tree inferred from the fastestevolving sites; (iii) Tree inferred from the slowest-evolving sites. To facilitate comparison of relative diversity, scale bars are provided separately for each panel. Slow-evolving sites support a relatively long inter-domain branch and less diversity within the domains (that is, shorter between-taxa branch lengths within domains). This suggests that substitution saturation (overwriting of earlier changes) may reduce the relative length of the AB branch at fast-evolving sites and genes. bioRxiv preprint doi: https://doi.org/10.1101/2021.01.19.427276; this version posted May 7, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY 4.0 International license.

34	8
34	9

Substitution model	BIC	AB branch length
LG+G4+F	5935950.053	1.449090256
LG+C20+G4+F	5783903.997	2.139350118
LG+C40+G4+F	5756823.360	2.469702112
LG+C60+G4+F	5746886.292	2.517828771

351

350 Table 1. The inferred AB length from a concatenation of the top 27 markers using a simple model versus models which account for site compositional heterogeneity. Using 352 better fitting models, i.e models which allow for across-site compositional heterogeneity, a 353 longer AB branch is inferred.

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#### A phylogeny of Archaea and Bacteria inferred from 27 vertically-evolving marker genes 356

357 The topology of our phylogeny of the primary domains of life (Figure 6) is consistent with recent 358 single-domain trees inferred for Archaea and Bacteria independently (Coleman et al., 2021; 359 Dombrowski et al., 2020; Williams et al., 2017), although the deep relationships within Bacteria 360 are only poorly resolved, with the exception of the monophyly of Gracilicutes (Figure 6). A 361 recent analysis suggested that, among extant lineages, the metabolisms of Clostridia, 362 Deltaproteobacteria, Actinobacteria and some Aquificae might best preserve the metabolism 363 of the last bacterial common ancestor (Xavier et al., 2021). Assuming a universal root between 364 Archaea and Bacteria (Dagan et al., 2010; Gogarten et al., 1989; Iwabe et al., 1989), none of 365 these groups branch near the bacterial root in our analysis (Figure 6). This is consistent with 366 previous work (Castelle and Banfield, 2018; Hug et al., 2016; Parks et al., 2017; Raymann et 367 al., 2015) including the inference of an updated and rooted bacteria phylogeny (Coleman et 368 al., 2021). Notably, our analysis placed the Candidate Radiation (CPR) (Brown et al., 2015) 369 as a sister lineage to Chloroflexi (Chloroflexota) rather than as a deep-branching bacterial 370 superphylum. While this contrasts with initial trees suggesting that CPR may represent an 371 early diverging sister lineage of all other Bacteria (Brown et al., 2015; Castelle and Banfield, 372 2018; Hug et al., 2016), our finding is consistent with a recent analysis that recovered CPR 373 within the Terrabacteria (Coleman et al., 2021). Together, these analyses suggest that the 374 deep-branching position of CPR in some trees was a result of long branch attraction, a 375 possibility that has been raised previously (Hug et al., 2016; Méheust et al., 2019).

376

377 The deep branches of the archaeal subtree are well-resolved in the ML tree and recover clades 378 of DPANN (albeit at 51% bootstrap support), Asgard (100% bootstrap support), and TACK 379 Archaea (75% bootstrap support), in agreement with a range of previous studies (Dombrowski 380 et al., 2020; Guy and Ettema, 2011; Raymann et al., 2015; Williams et al., 2017). We also find 381 support for the placement of Methanonatronarchaeia (Sorokin et al., 2017) distant to 382 Halobacteria within the Methanotecta, in agreement with recent analyses and suggesting their 383 initial placement with Halobacteria (Sorokin et al., 2017) may be an artifact of compositional 384 attraction (Aouad et al., 2019; Dombrowski et al., 2020; Martijn et al., 2020). Notably, the 385 Hadesarchaea (92% bootstrap support) and a clade comprising Theionarchaea, 386 Methanofastidiosa, and Thermococcales (92% bootstrap support) branch basal to the TACK and Asgard Archaea, respectively, in our analysis, rather than with other Euryarchaeota.
These positions have been previously reported (Adam et al., 2017; Raymann et al., 2015;
Williams et al., 2017), though the extent of euryarchaeotal paraphyly and the lineages involved
has varied among analyses.

391

392 A broader observation from our analysis is that the phylogenetic diversity of the archaeal and bacterial domains, measured as substitutions per site in this consensus set of vertically-393 394 evolving marker genes, appears to be similar (Figure 5(i); the mean root to tip distance for 395 archaea: 2.38, for bacteria: 2.41, the range of root to tip distances for archaea: 1.79-3.01, for bacteria: 1.70-3.17). Considering only the slowest-evolving category of sites, branch lengths 396 397 within Archaea are actually longer than within Bacteria (Figure 5(iii)). This result differs from 398 some published trees (Hug et al., 2016; Zhu et al., 2019) in which the phylogenetic diversity 399 of Bacteria has appeared to be significantly greater than that of Archaea. By contrast to those 400 earlier studies, we analysed a set of 350 genomes from each domain, an approach which may 401 tend to reduce the differences between them. While we had to significantly downsample the 402 sequenced diversity of Bacteria, our sampling nonetheless included representatives from all 403 known major lineages of both domains, and so might be expected to recover a difference in 404 diversity, if present. Our analyses and a number of previous studies (Hug et al., 2016; Parks 405 et al., 2018; Petitjean et al., 2014; Zhu et al., 2019) indicate that the choice of marker genes 406 has a profound impact on the apparent phylogenetic diversity of prokaryotic groups; for 407 instance, in the proportion of bacterial diversity composed of CPR (Hug et al., 2016; Parks et 408 al., 2017). Our results demonstrate that slow and fast-evolving sites from the same set of 409 marker genes support different tree shapes and branch lengths; it therefore seems possible 410 that between-dataset differences are due, at least in part, to evolutionary rate variation within 411 and between marker genes.



**Figure 6:** A phylogeny of Archaea and Bacteria inferred from a concatenation of 27 marker genes. Consistent with some recent studies (Dombrowski et al., 2020; Guy and Ettema, 2011; Raymann et al., 2015; Williams et al., 2017), we recovered the DPANN, TACK and Asgard Archaea as monophyletic groups. Although the deep branches within Bacteria are poorly resolved, we recovered a sister group relationship between CPR and Chloroflexota, consistent with a recent report (Coleman et al., 2021). The tree was inferred using the best-fitting LG+C60+G4+F model in IQ-TREE 2 (Minh et al., 2020). Branch lengths are proportional to the expected number of substitutions per site. Support values are ultrafast (UFBoot2) bootstraps (Hoang et al., 2018). Numbers in parenthesis refer to number of taxa within each collapsed clade. Please note that collapsed taxa in the Archaea and Bacteria roughly correspond to order- and phylum-level lineages, respectively.

#### 412 Conclusion

#### 413

414 Core gene phylogenies provide a window into the earliest period of archaeal and bacterial 415 evolution. Concatenation is useful for pooling signal across individual genes, but topology and 416 branch length estimates from concatenations only reflect the underlying tree of life if the 417 individual genes share the same evolutionary history. Our analysis of published datasets 418 (Coleman et al., 2021; Petitjean et al., 2014; Williams et al., 2020; Zhu et al., 2019) indicates 419 that incongruence among marker genes resulting from inter-domain gene transfer and hidden 420 paralogy can lead to an under-estimate of the inter-domain branch length. We performed a re-421 analysis of marker genes from a range of published analyses, manually curated datasets to 422 identify and remove transferred genes, and estimated an updated phylogeny of Archaea and 423 Bacteria. Considering only this manually curated consensus marker gene dataset, we found 424 no evidence that ribosomal markers overestimate stem length; since they appear to be 425 transferred less frequently than other genes, our analysis affirms that ribosomal proteins are 426 useful markers for deep phylogeny. In general, better markers, regardless of functional 427 category, support a longer AB branch length. A phylogeny inferred from the 27 best-performing 428 markers was consistent with some recent work on early prokaryotic evolution, resolving the 429 major clades within Archaea and nesting the CPR within Terrabacteria. Our analyses suggest 430 that both the true Archaea-Bacteria branch length (Figure 7), and the phylogenetic diversity of 431 Archaea, may be underestimated by even the best current models, a finding that is consistent 432 with a root for the tree of life between the two prokaryotic domains.

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Figure 7. The impact of marker gene choice, phylogenetic congruence, alignment trimming, and substitution model fit on estimates of the Archaea-Bacteria branch length. Analysis using a sitehomogeneous model (LG+G4+F) on the complete 381-gene expanded set results in an AB branch substantially shorter than previous estimates. Removing the genes most seriously affected by inter-domain gene transfer, trimming poorly-aligned sites using BMGE (Criscuolo and Gribaldo, 2010), and using the bestfitting site-heterogeneous model available (LG+C60+G4+F) substantially increase the estimated AB length, such that it is comparable with published estimates from the "core" set (Williams et al., 2020) and the consensus set of 27 markers identified in the present study. Branch lengths measured in expected number of substitutions/site.

### 433 Methods

#### 434 **Data**

435 We downloaded individual alignments (Zhu 2019) the from et al.. (https://github.com/biocore/wol/tree/master/data/), along with the genome metadata and the 436 437 individual newick files. We checked each published tree for domain monophyly, and also 438 performed approximately unbiased (AU) (Shimodaira, 2002) tests to assess support for 439 domain monophyly on the underlying sequence alignments using IQ-TREE 2 (Minh et al., 440 2020). The phylogenetic analyses were carried out using the 'reduced' subset of 1000 taxa outlined by the authors (Zhu et al., 2019), for computational tractability. These markers were 441 442 also trimmed according to the protocol in the original paper (Zhu et al., 2019), i.e sites with 443 >90% gaps were removed, followed by removal of sequences with >66% gaps.

We also downloaded the Williams et al. (Williams et al., 2020) ("core"), Petitjean et al. (Petitjean et al., 2014) ("non-ribosomal") and Coleman et al. (Coleman et al., 2021) ("bacterial") datasets from their original publications.

#### 447

#### 448 Annotations

449 Proteins used for phylogenetic analyses by Zhu et al. (Zhu et al., 2019), were annotated to 450 investigate the selection of sequences comprising each of the marker gene families. To this 451 end, we downloaded the protein sequences provided by the authors from the following 452 repository: https://github.com/biocore/wol/tree/master/data/alignments/genes. To obtain 453 reliable annotations, we analysed all sequences per gene family using several published 454 databases, including the arCOGs (version from 2014) (Seemann, 2014), KOs from the KEGG 455 Automatic Annotation Server (KAAS; downloaded April 2019) (Aramaki et al., 2020), the Pfam database (Release 31.0)(Bateman et al., 2004), the TIGRFAM database (Release 15.0) (Haft 456 457 et al., 2003), the Carbohydrate-Active enZymes (CAZy) database (downloaded from dbCAN2 458 in September 2019)(Cantarel et al., 2009), the MEROPs database (Release 12.0) (Rawlings 459 et al., 2016), (Saier et al., 2006), the hydrogenase database (HydDB; downloaded in 460 November 2018) (Søndergaard et al., 2016), the NCBI- non-redundant (nr) database 461 (downloaded in November 2018), and the NCBI COGs database (version from 2020). 462 Additionally, all proteins were scanned for protein domains using InterProScan (v5.29-68.0; 463 settings: --iprlookup --goterms) (Jones et al., 2014).

464

465 Individual database searches were conducted as follows: arCOGs were assigned using PSI-BLAST v2.7.1+ (settings: -evalue 1e-4 -show\_gis -outfmt 6 -max\_target\_seqs 1000 -dbsize 466 467 10000000 -comp based stats F -seg no) (Altschul et al., 1997). KOs (settings: -E 1e-5), 468 PFAMs (settings: -E 1e-10), TIGRFAMs (settings: -E 1e-20) and CAZymes (settings: -E 1e-469 20) were identified in all archaeal genomes using hmmsearch v3.1b2(Finn et al., 2011). The 470 MEROPsand HydDB databases were searched using BLASTp v2.7.1 (settings: -outfmt 6, -471 evalue 1e-20). Protein sequences were searched against the NCBI\_nr database using 472 DIAMOND v0.9.22.123 (settings: -more-sensitive -e-value 1e-5 -seq 100 -no-self-hits -473 taxonmap prot.accession2taxid.gz) (Buchfink et al., 2015). For all database searches the best 474 hit for each protein was selected based on the highest e-value and bitscore and all results are 475 summarized in the Data Supplement Table, 476 Annotation Tables/0 Annotation tables full/All Zhu marker annotations 16-12477 2020.tsv.zip. For InterProScan we report multiple hits corresponding to the individual domains
478 of a protein using a custom script (parse\_IPRdomains\_vs2\_GO\_2.py).

479

Assigned sequence annotations were summarized and all distinct KOs and Pfams were collected and counted for each marker gene. KOs and Pfams with their corresponding descriptions were mapped to the marker gene file downloaded from the repository: <u>https://github.com/biocore/wol/blob/master/data/markers/metadata.xlsx</u> and used in summarization of the 381 marker gene protein trees (Table S1).

485

486 For manual inspection of single marker gene trees, KO and Pfam annotations were mapped to the tips of the published marker protein trees, downloaded from the repository: 487 488 https://github.com/biocore/wol/tree/master/data/trees/genes. Briefly, the Genome ID, Pfam, 489 Pfam description, KO, KO description, and NCBI Taxonomy string were collected from each 490 marker gene annotation table and were used to generate mapping files unique to each marker 491 which links the Genome ID to the annotation information phyloaeny. aene 492 (GenomeID|Domain|Pfam|Pfam Description|KO|KO Description). An in-house perl script 493 replace tree names.pl

(https://github.com/ndombrowski/Phylogeny\_tutorial/tree/main/Input\_files/5\_required\_Scripts
 ) was used to append the summarized protein annotations to the corresponding tips in each
 marker gene tree. Annotated marker gene phylogenies were manually inspected using the
 following criteria including: 1) retention of reciprocal domain monophyly (Archaea and
 Bacteria) and 2) for the presence or absence of potential paralogous families. Paralogous
 groups and misannotated families present in the gene trees were highlighted and violations of
 search criteria were recorded in Table S1.

#### 501 *Phylogenetic analyses*

#### 502 COG assignment for the Core, Non-Ribosomal, and Bacterial marker genes

503 First, all gene sequences in the three published marker sets (core, non-ribosomal, and 504 bacterial) were annotated using the NCBI COGs database (version from 2020). Sequences 505 were assigned a COG family using hmmsearch v3.3.2 (Finn et al., 2011) (settings: -E 1e-5) 506 and the best hit for each protein sequence was selected based on the highest e-value and bit 507 score. To assign the appropriate COG family for each marker gene, we quantified the 508 percentage distribution of all unique COGs per gene, and selected the family representing the 509 majority of sequences in each marker gene.

Accounting for overlap, this resulted in 95 unique COG families from the original 119 total marker genes across all three published datasets (Table S2). Orthologues corresponding to these 95 COG families were identified in the 700 genomes (350 Archaea, 350 Bacteria, Table S3) using hmmsearch v3.3.2 (settings: -E 1e-5). The reported BinID and protein accession were used to extract the sequences from the 700 genomes, which were used for subsequent phylogenetic analyses.

#### 516 Marker gene inspection and analysis

517 We aligned these 95 sequence sets using MAFFT-linsi (Katoh and Toh, 2008) and removed 518 poorly-aligned positions with BMGE (Criscuolo and Gribaldo, 2010). We inferred initial 519 maximum likelihood trees (LG+G4+F) for all 95 markers and mapped the KO and Pfam 520 domains and descriptions, inferred from annotation of the 700 genomes, to the corresponding 521 tips (see above). Manual inspection took into consideration monophyly of Archaea and 522 Bacteria and the presence of paralogs, and other signs of contamination (HGT, LBA). 523 Accordingly, single gene trees that failed to meet reciprocal domain monophyly were excluded. 524 and any instances of HGT, paralogous sequences, and LBA artefacts were manually removed 525 from the remaining trees resulting in 54 markers across the three published datasets that were 526 subject to subsequent phylogenetic analysis (LG+C20+G4+F) and further refinement (see 527 below).

#### 528

#### 529 Ranking markers based on split score

530 We applied an automated marker gene ranking procedure devised previously, the split score 531 (Dombrowski et al., 2020), to rank each of the 54 markers that satisfied reciprocal monophyly 532 based on the extent to which they recovered established phylum-, class- or, order-level 533 relationships within the archaeal and bacterial domains (Table S4).

534 The script quantifies the number of splits, or occurrences where a taxon fails to cluster within 535 its expected taxonomic lineage, across all gene phylogenies. Monophyly of archaeal and 536 bacterial lineages was assessed based on clades defined in Table S4. Briefly, we used 537 Cluster1 for Archaea in combination with Cluster0 (phylum) or Cluster3 (i.e. on class-level if defined and otherwise on phylum-level; Table S4) for Bacteria. We then ranked the marker 538 539 genes using the following split-score criteria: the number of splits per taxon and the splits 540 normalized to the species count. The percentage of split phylogenetic groups was used to 541 determine the highest ranking (top 50%) markers.

#### 542 Concatenation

543 Based on the split score ranking of the 54 marker genes (above), the top 50% (27 markers, 544 Table S4) marker genes were manually inspected using criteria as defined above, and 545 contaminating sequences were manually removed from the individual sequence files. 546 Following inspection, marker protein sequences were aligned using MAFFT-LINSI (Katoh and 547 Standley, 2013) and trimmed using BMGE (Criscuolo and Gribaldo, 2010). We concatenated 548 the 27 markers into a supermatrix, which was used to infer a maximum-likelihood tree (Figure 549 6, under LG+C60+G4+F), evolutionary rates (see below), and rate-category supermatrices 550 as well as to perform model performance tests (see below).

#### 551 Constraint analysis

552 We performed a maximum likelihood free topology search using IQ-TREE 2 (Minh et al., 2020) 553 under the LG+G4+F model, with 1000 bootstrap replicates on each of the markers from the 554 expanded, bacterial, core and non-ribosomal sets. We also performed a constrained analysis with the same model, in order to find the maximum likelihood tree in which Archaea and 555 Bacteria were reciprocally monophyletic. We then compared both trees using the 556 557 approximately unbiased (AU) Shimodaira (2002) test in IQ-TREE 2 (Minh et al., 2020) with 558 10,000 RELL (Shimodaira, 2002) bootstrap replicates. To evaluate the relationship between 559 marker gene verticality and AB branch length, we calculated the difference in log-likelihood 560 between the constrained and unconstrained trees in order to rank the genes from the 561 expanded marker set, made concatenates comprised of the top 20-100 (intervals of 5) of these

562 marker genes, and inferred the tree length under LG+C10+G4+F with 1000 bootstrap 563 replicates.

#### 564 Site and gene evolutionary rates

We inferred rates using the --rate option in IQ-TREE 2 (Minh et al., 2020) for both the 381 marker concatenation from Zhu (Zhu et al., 2019) and the top 5% of marker genes based on the results of difference in log-likelihood between the constrained tree and free-tree search in the constraint analysis (above). We also used this method to explore the differences in rates for the 27 marker set. We built concatenates for sites in the slowest and fastest rate categories, and inferred branch lengths from each of these concatenates using the tree inferred from the corresponding dataset as a fixed topology.

#### 572 Substitution model fit

573 Model fit tests were undertaken using the top 5% concatenate described above, with the 574 alignment being trimmed with BMGE 1.12 (Criscuolo and Gribaldo, 2010) with default settings 575 (BLOSUM62, entropy 0.5) for all of the analyses except the 'untrimmed' LG+G4+F run, other 576 trimmed alignment LG+G4+F, models on the were LG+R4+F and 577 LG+C10,20,30,40,50,60+G4+F, with 1000 bootstrap replicates. Model fitting was done using 578 ModelFinder (Kalyaanamoorthy et al., 2017) in IQ-TREE 2 (Minh et al., 2020). For the model 579 testing for the 27 concatenation, we performed a model finder analysis (-m MFP) including 580 additional models evolution, complex of (i.e. 581 LG+C60+G4+F,LG+C50+G4+F,LG+C40+G4+F,LG+C30+G4+F,LG+C20+G4+F,LG+C10+G 582 4+F,LG+G4+F,LG+R4+F) to the default, to find the best fitting model for the analysis. This 583 revealed that, according to AIC, BIC and cAIC, LG+C60+G4+F was the best fitting model. For 584 comparison. also performed analyses usina following models: we the 585 LG+G4+F,LG+C20+G4+F,LG+C40+G4+F (Table 1).

#### 586 Molecular clock analyses

587 Molecular clock analyses were devised to test the effect of genetic distance on the inferred 588 age of LUCA. Following the approach of Zhu et al. (Zhu et al., 2019), we subsampled the 589 alignment to 100 species. Five alternative alignments were analysed, representing conserved 590 sites across the entire alignment, randomly selected sites across the entire alignment, only 591 ribosomal marker genes, the top 5% of marker genes according to ΔLL and the top 5% of 592 marker genes further trimmed under default settings in BMGE 1.12 (Criscuolo and Gribaldo, 593 2010). Divergence time analyses were performed in MCMCTree (Yang, 2007) under a strict 594 clock model. We used the normal approximation approach, with branch lengths estimated in codeml under the LG+G4 model. In each case, a fixed tree topology was used alongside a 595 596 single calibration on the Cyanobacteria-Melainabacteria split. The calibration was modelled 597 as a uniform prior distribution between 2.5 and 2.6 Ga, with a 2.5% probability that either 598 bound could be exceeded. For each alignment, four independent MCMC chains were run for 599 2,000,000 generations to achieve convergence.

#### 600 Plotting

601 Statistical analyses were performed using R 4.0.4 (R Core Team, 2021), and data were plotted 602 with ggplot2 (Wickham, 2016).

### 603 Data and code availability

All of the data, including sequence alignments, trees, annotation files, and scripts associated
with this manuscript have been deposited in the FigShare repository at DOI:
10.6084/m9.figshare.13395470.

607

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