Phylogenomics provides robust support for a two-domains tree of life

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Hypotheses about the origin of eukaryotic cells are classically framed within the context of a universal ‘tree of life’ based on conserved core genes. Vigorous ongoing debate about eukaryote origins is based on assertions that the topology of the tree of life depends on the taxa included and the choice and quality of genomic data analysed. Here we have reanalysed the evidence underpinning those claims and apply more data to the question by using supertree and coalescent methods to interrogate >3,000 gene families in archaea and eukaryotes. We find that eukaryotes consistently originate from within the archaea in a two-domains tree when due consideration is given to the fit between model and data. Our analyses support a close relationship between eukaryotes and Asgard archaea and identify the Heimdallarchaeota as the current best candidate for the closest archaeal relatives of the eukaryotic nuclear lineage.

Current hypotheses about eukaryotic origins generally propose at least two partners in that process: a bacterial endosymbiont that became the mitochondrion and a host cell for that endosymbiosis.1–4 The identity of the host has been informed by analyses of conserved genes for the transcription and translation machinery that are considered essential for cellular life.5–7 Traditionally, the host was considered to be a eukaryote on the basis of ribosomal RNA trees in either unrooted8 or rooted form.9 In these trees, archaea, bacteria and eukarya form three separate primary domains, with the rooted version suggesting that archaea and eukarya are more closely related to each other than to bacteria.9 A criticism of these three-domains (3D) trees is that they were constructed using overly simple phylogenetic models.10–12 Phylogenetic analyses using models that better fit features of the data,13–15, coupled with an expanded sampling of prokaryotic diversity,16–18, have supported a two-domains (2D) tree consistent with the eocyte hypothesis whereby the eukaryotic nuclear lineage—the host for the mitochondrial endosymbiont—originated from within the archaea (reviewed in refs. 19,20). The 2D tree has gained increasing traction in the field,10–12 particularly with the discovery of the Asgard archaea19,20. The Asgard archaea branch together with eukaryotes in phylogenetic trees and their genomes encode homologues of eukaryotic signature proteins—proteins which underpin the defining cellular structures of eukaryotes, and which were previously thought21 to be unique to eukaryotes. However, the discoveries and analyses that support the 2D tree have been criticized from a variety of perspectives.

It has been suggested22–24 that the close relationship between eukaryotes and Asgard archaea in 2D trees25,26 is due to eukaryotic contamination of Asgard metagenomes combined with phylogenetic artefacts caused by the choice of genes analysed and the inclusion of fast-evolving archaea in tree reconstructions;22–24 see also the comment25 and response26 to those analyses. The phenomenon of long-branch attraction due to the presence of fast-evolving sequences is a well-known artefact in phylogenetic analyses26–28. Indeed, it has previously been suggested that it is the 3D tree, rather than the 2D tree, that is an artefact of long-branch attraction29–31, both because analyses under better-fitting models have recovered a 2D tree but also because the 3D topology is one in which the two longest branches in the tree of life—the stems leading to bacteria and to eukaryotes—are grouped together. Nevertheless, when putative fast-evolving sequences were removed, Forterre and colleagues22–24 recovered a monophyletic archaea within a 3D tree, whether analysing 35 core genes, a particular subset of six genes or RNA polymerases alone. Claims that the 2D tree is a product of unbalanced taxonomic sampling and inclusion of fast-evolving sequences have also been made by others32.

In a more general criticism it has been suggested30–33 that protein sequences do not harbour sufficient signal to resolve the 2D/3D debate due to mutual saturation (but see refs. 11,12). One suggested solution is to analyse conserved structural motifs (folds) in proteins rather than primary sequence data31,33,34. Three-dimensional structures are thought to be more highly conserved than primary sequences. It has therefore been suggested that they should provide a more reliable indicator of ancient relationships, although it is not yet clear how best to analyse fold data for this purpose. Published unrooted trees based upon analyses of protein folds have recovered archaea, bacteria and eukaryotes as separate groups34,35, a result that is consistent with the 3D but not the 2D tree. Analyses of protein folds have recently been extended to use non-stationary models to infer a rooted tree of life31. In these analyses the inferred root separated cellular life into prokaryotes (archaea plus bacteria, termed akaryotes) and eukaryotes31,33. This tree is incompatible with the idea that archaea and eukaryotes share closer common ancestry, and recapitulates the hypothesis36 that the deepest division in cellular life is between prokaryotes and eukaryotes.

In this paper, we have evaluated the analyses and data that have led to conflicting hypotheses of relationships between the major groups of cellular life and for the position of the eukaryotic nuclear lineage. We have also performed phylogenomic analyses using
the best-available supermatrix, supertree, and coalescent methods on an expanded sample of genes and taxa, to further explore the deep structure of the tree of life and the relationship between archaea and eukaryotes.

Results and discussion

Analysis of core genes consistently supports two primary domains, not three. It has recently been argued\textsuperscript{22–24} that the 2D tree is an artefact of data and taxon sampling and that resolution of those issues provides support for a 3D tree. The molecular data at the core of this debate had first been used\textsuperscript{22} to support a 2D tree in which eukaryotes clustered within archaea as the closest relatives of the Asgard archaea. The original dataset\textsuperscript{19} comprised a concatenation of 36 ‘universal’ genes for 104 taxa. In the initial critique, it was claimed that the close relationship reported\textsuperscript{19} between Asgard archaea and eukaryotes was caused by the inclusion in the dataset of a contaminated elongation factor 2 (EF2) gene for *Lokiarchaeum* sample Lok13 (ref. \textsuperscript{24}; now *Heimdallarchaeota*)\textsuperscript{25} and by the inclusion of fast-evolving archaeal lineages in the analysis. However, recent data suggest that the EF2 gene of *Heimdallarchaeota* is not contaminated with eukaryotic sequences because similar EF2 sequences have been found in additional *Heimdallarchaeota* metagenome-assembled genomes prepared from different environmental DNA samples in different laboratories\textsuperscript{26,27}.

The claim\textsuperscript{22–24} that the presence of fast-evolving sequences might be affecting the topology recovered could be seen as a reasonable challenge, since long-branch attraction can influence the tree topology recovered. A problem for this specific critique\textsuperscript{26}, however, is that no single, clear and consistent criterion was used to identify the ‘fast-evolving’ sequences that were removed from the original dataset\textsuperscript{19} to recover the 3D tree. Long-branched archaea might result from either a fast evolutionary rate or a long period of time and these possibilities are difficult to distinguish a priori. Moreover, the historical papers\textsuperscript{38,39} cited\textsuperscript{22} as providing topological evidence that some sequences are fast-evolving used site- and time-homogeneous phylogenetic models (models in which the process of evolution is constant over the sites of the alignment and branches of the tree) which often fit data poorly. To investigate further, we ranked all of the taxa in the original dataset\textsuperscript{19} according to their root-to-tip distances for each species. This is equal to the summed branch length (expressed as expected number of substitutions per site) from the root of the tree (rooted between bacteria and archaea) to the relevant tip. We calculated distributions and 95% credibility intervals (Supplementary Table 1) for each of these root-to-tip distances from the samples drawn during a Markov Chain Monte Carlo (MCMC) analysis under the best-fitting (see below) CAT + GTR + G4 model in PhyloBayes, to perform Bayesian relative rates tests (Supplementary Table 1).

The 23 taxa previously identified as fast-evolving sequences are not the 23 taxa with the longest root-to-tip distances. Meanwhile some of the taxa chosen for exclusion ("Parvarchaeum, Micrarchaeum, Nanoarchaeum Nst1, Nanosalinarum and Korarchaeum") are indeed relatively long-branching, others ("Iainarchaeum, Nanoarchaeum G17 and Aenigmaarchaeon") are in the bottom half of the branch length distribution and many of the longest-branching archaea (including the *Thaumarchaeota*) were retained. Nevertheless, analysis\textsuperscript{22} of the reduced dataset did recover a 3D tree, raising the question of why this result was obtained. In the following analyses we have followed the recent renaming\textsuperscript{22} of the three ‘Loki’ metagenome-assembled genomes originally analysed as *Lokiarchaeum* sp. GC14_75 (formerly Lok1), *Heimdallarchaeota* archaeon LC_2 (Lok2) and *Heimdallarchaeota* archaeon LC_3 (Lok3).

The published 3D tree\textsuperscript{22} was recovered from the 35-gene concatenated dataset under the LG + G4 + F model\textsuperscript{20} in PhyML 3.1 (ref. \textsuperscript{41}), with moderate support (76% bootstrap) for monophyletic archaea (fig. 5b in ref. \textsuperscript{22}). In repeating this analysis, we noted that although PhyML returned a 3D tree, analysis of the same alignment under the same substitution model (LG + G4 + F) with IQ-TREE 1.6.2 (ref. \textsuperscript{42}) and RAxML 8.2.4 (ref. \textsuperscript{43}), two other maximum-likelihood phylogeny packages, instead yielded a 2D tree where Heimdallarchaeota and *Lokiarchaeum* were together the sister group to eukaryotes, with a better likelihood score (Supplementary Fig. 1 and Supplementary Table 2). To investigate further, we computed the log likelihoods of the 2D and 3D trees in all three packages, keeping the alignment and model constant (Supplementary Table 2). All three implementations accord the 2D tree a higher likelihood than the 3D tree (lnl approximately equal to −684701.2, compared to −684716.1 for the 3D tree). It thus appears that the recovery of a 3D tree reflects a failure of PhyML to find the more likely 2D tree, rather than to the removal of problematic sequences. The differences between the likelihoods are not significant according to an approximately unbiased test (AU = 0.229 for the 3D tree, 0.771 for the 2D), meaning that analysis of the 35-gene dataset under LG + G4 + F is equivocal with respect to the 2D and 3D trees; contrary to previous claims\textsuperscript{22}, analysis of the 35-gene concatenation under the LG + G4 + F model provides no unambiguous evidence to prefer the 3D tree.

A number of newer models accommodate particular features of empirical data better than the LG + G4 + F, so we investigated which trees were produced from the 35-gene dataset using these models. We addressed three issues in particular: among-site compositional heterogeneity due to site-specific biochemical constraints\textsuperscript{44}, changing composition in different lineages over time\textsuperscript{45}, and variations in site- and lineage-specific evolutionary rates (heterochronous evolution)\textsuperscript{46}.

The CAT + GTR + G4 model\textsuperscript{45,47} is an extension to the standard GTR model that allows compositions to vary across sites. Analysis of the 35-gene dataset using this model produced a 2D tree where eukaryotes group with Heimdallarchaeota and *Lokiarchaeum* with maximal support (Fig. 1). It was previously reported\textsuperscript{22} that convergence in Bayesian analyses is a problem for this dataset using the CAT + GTR + G4 (ref. \textsuperscript{45}) model. In our analyses, we achieved good convergence between chains as assessed both by comparison of split frequencies and, for the continuous parameters of the model, means and effective sample sizes (Supplementary Table 4). As an additional check, we also carried out maximum-likelihood analyses using the LG + C60 + G4 + F model, which improves on the LG + G4 + F model by modelling site-specific compositional heterogeneity using a mixture of 60 composition categories. This model fits the data much better than the LG + G4 + F according to the Bayesian Information Criterion (BIC; Supplementary Table 3) and, like CAT + GTR + G4, it recovered a 2D tree with high bootstrap support (Supplementary Fig. 1c). The 3D tree (AU = 0.036) could also be rejected at P < 0.05 using an AU test, based on the LG + C60 + G4 + F model and the 35-gene alignment.

Bayesian posterior predictive simulations\textsuperscript{48} provide a tool for evaluating the adequacy of models, by testing whether data simulated under a model is similar to the empirical data. Figure 2 plots the 2D tree (inferred under CAT + GTR + G4) and the 3D tree (inferred under LG + G4 + F in PhyML) on the same scale (Fig. 2a), revealing that—from the same alignment—CAT + GTR + G4 infers that many more substitutions have occurred in the core gene set during the evolutionary history of life. Model fit tests (Fig. 2b and Supplementary Table 4) indicate that LG + G4 + F provided a much poorer fit to the data (larger Z-scores) than CAT + GTR + G4 in terms of across-site compositional heterogeneity (Z = 64.2 for LG + G4 + F and Z = 6.9 for CAT + GTR + G4) and therefore systematically underestimated the probability of convergent substitutions (Z = 19.7 for LG + G4 + F and Z = 7.62 for CAT + GTR + G4). These differences arise because LG + G4 + F assumes that amino acid frequencies are the same at all sites, whereas in empirical data sets different sites have different compositions, arising from distinct biochemical and selective constraints. Since this means the effective number of amino acids per site is in reality lower than that
predicted by LG + G4 + F, the probability of parallel convergence to the same amino acid in independent lineages is higher (Supplementary Table 5). CAT + GTR + G4 accounts for this across-site variation by incorporating site-specific compositions and is therefore less prone to underestimating rates of convergent substitution. This is important because the longest branches in both the 2D and 3D trees are the lineages leading to the bacteria and eukaryotes. The lesser ability of LG + G4 + F to detect convergent substitutions along these branches may favour inference of a 3D tree. While CAT + GTR + G4 provides a better fit than LG + G4 + F, neither model completely fits the composition of the data ($P = 0$ for all tests; Supplementary Table 5). As a further data exploration step, we recoded the amino acid alignment into four categories (Supplementary Table 5). As a further data exploration step, we recoded the amino acid alignment into four categories, for all tests; Supplementary Table 5). As a further data exploration step, we recoded the amino acid alignment into four categories (Supplementary Table 5). As a further data exploration step, we recoded the amino acid alignment into four categories. We used the GHOST model of IQ-Tree to analyse the 35-gene dataset. While CAT + GTR + G4 gave a 2D tree with maximal posterior support (PP = 1; Supplementary Fig. 3), as did a partitioned analysis using the best-fitting homogeneous model for each of the 12 gene partitions (LG + G4 + F in all cases; Supplementary Fig. 3; PP = 1). We also inferred a phylogeny from the entire 35-gene dataset under the branch-heterogeneous node-discrete compositional heterogeneity (NDCH) model, which explicitly incorporates changing sequence compositions across the tree. NDCH2 is an extension of the NDCH model; it has a separate composition vector for each tree node and is constrained via a sampled concentration parameter of a Dirichlet prior. Thus, the model adjusts to the level of across-branch compositional heterogeneity in the data during the MCMC analysis. For reasons of computational tractability, this analysis could only be run on the SR4-recoded version of the 35-gene alignment. NDCH2 obtained adequate model fit with respect to across-branch compositional heterogeneity ($P = 0.7838$) and recovered a 2D tree with Heimdallarchaea as the sister group to eukaryotes (PP = 0.85; Supplementary Fig. 2).

A failure to account for heterotachy or rates of molecular evolution that are both site- and branch-specific, has been posited as a potential issue for phylogenomic analyses of ancient core genes. We used the GHOST model of IQ-Tree to analyse the 35-gene alignment. GHOST is an edge-unlinked mixture model in which the sites of the alignment evolve along a shared tree topology but are fit by a finite mixture of GTR exchangeabilities, sequence compositions and branch lengths. We fit a four-component mixture model to both the original amino acid alignment (LG + G4 + F components) and the SR4-recoded version (GTR + F components). The resulting trees were a weakly supported (amino acids; 58% bootstrap.

Fig. 1 | The 35-gene matrix of Da Cunha et al. favours a 2D tree using the best-fitting models in both maximum-likelihood and Bayesian analyses. Note the eukaryotes (green) group with the sampled Asgard archaea (pink) with maximum posterior support. Bacteria are in grey, TACK archaea in orange, Euryarchaeota in purple. This is a consensus tree inferred under the CAT + GTR + G4 model in PhyloBayes-MPI; branch lengths are proportional to the expected number of substitutions per site, as indicated by the scale bar. A 2D topology was obtained under a variety of other models in maximum-likelihood analyses (LG + G4 + F, LG + PMSF + G4, LG + G4 + F; Supplementary Fig. 1) and also with four-state Susko–Roger recoding under the CAT + GTR + G4 and NDCH2 models (Supplementary Fig. 2).
Fig. 2 | Evidence that the 3D tree is an artefact of long-branch attraction. a, Da Cunha et al.22 analysed a dataset of 35 core protein-coding genes under the LG + G4 + F model and obtained a 3D tree; the better-fitting (Supplementary Table 4) CAT + GTR + G4 model recovers a 2D tree. b, Posterior predictive tests indicate that CAT + GTR + G4 performs significantly better than LG + G4 + F in capturing the site-specific evolutionary constraints reflected by lower biochemical diversity approaching that of the empirical data. This results in more realistic estimates of substitutional saturation and convergence found in the data. The longest branches on both the 3D and 2D trees in a are the stems leading to the bacteria and eukaryotes (in yellow and green, respectively). CAT + GTR + G4 identifies many more convergent substitutions on these branches than does LG + G4 + F, as can be seen by comparing the branch lengths in a. This failure to detect convergent substitutions under LG + G4 + F has the effect of drawing the bacterial and eukaryotic branches together because convergences are mistaken for homologies (synapomorphies), resulting in a 3D tree. Bootstrap support (a) and Bayesian posterior probability (b) are indicated for the key nodes defining the 3D and 2D trees. Asgard refers to a clade of Heimdallarchaeota and Lokiarchaeum. Plotting these trees to the same scale (in terms of substitutions per site) illustrates major differences in these analyses. The 3D/LG + G4 + F analysis suggests that, on average, 30.77 changes have taken place per site; the 2D/CAT + GTR + G4 analysis suggests that 47.4 changes per site have occurred. This difference amounts to ~128,511 additional substitutions in total inferred under the CAT + GTR + G4 model.
support for eukaryotes plus Heimdallarchaeota and Lokiarchaeum) or strongly supported (recoded data; 95% bootstrap support for eukaryotes and Heimdallarchaeota) 2D tree (Supplementary Fig. 5).

In summary, all of our analyses of the 35-gene alignment using better models recovered a 2D tree in which eukaryotes are either the sister group of Heimdallarchaeota and Lokiarchaeum or Heimdallarchaeota alone, rather than the 3D tree which the data has previously been claimed\(^\text{23}\) to support.

**Do some core genes have different histories?** Based upon AU tests under the LG + G4 + F model for individual genes in the 35-gene dataset, it was suggested\(^\text{22}\) that the 35-gene dataset contains two subsets of genes with different evolutionary histories: a larger set supporting the 2D tree and a smaller set supporting the 3D tree. We used the better-fitting CAT + GTR + G4 model to analyse a concatenated dataset of the six genes that significantly favoured the 3D tree under LG + G4 + F and we also analysed a four-state recoded version of the same alignment. Analysis of the original amino acids recovered a moderately supported 3D tree, while analysis of the recoded alignment recovered a weakly supported 2D tree (Supplementary Fig. 4); posterior predictive simulations indicated that model fit was improved by SR4 recoding (Supplementary Table 7), suggesting that support for the 3D tree from these six genes under LG + G4 + F may be due to model misspecification.

It has also been suggested that phylogenetic analyses of RNA polymerase subunits\(^\text{24}\) provide robust support for a 3D tree. By contrast, other\(^\text{29}\) analyses of RNA polymerase subunits have already suggested that better-fitting models prefer a 2D tree. We evaluated the fit of both models, LG + G4 + F and CAT + GTR + G4, used\(^\text{22}\) to recover a 3D tree from RNA polymerase subunits, using posterior predictive simulations (Supplementary Discussion) and found that both models provide an adequate fit to the data (Supplementary Table 8). Model fit was improved following SR4 recoding (Supplementary Table 8) and this analysis recovered a weakly supported and poorly resolved 2D tree (Supplementary Fig. 6).

Expanded gene and taxon sampling supports a clade of eukaryotes and Asgard archaea. We took advantage of the recent dramatic improvements in genomic and transcriptomic sampling of free-living bacteria, archaea and microbial eukaryotes to assemble a dataset of 125 species, including 53 eukaryotes, 39 archaea (including an expanded set of Asgard metagenome-assembled genomes\(^\text{20}\) representing two new groups, Odinarchaeota and Thorarchaeota) and 33 bacteria, on the principle that improved sampling can sometimes help to resolve difficult phylogenetic problems\(^\text{25}\). We used free-living representatives of eukaryotic groups to avoid the well-documented problems for tree reconstruction caused by sequences from parasitic eukaryotes\(^\text{26}\). Our sampling of archaea and bacteria was also expanded to include representatives from the large number of uncultivated lineages that have recently been identified by single-cell genomics and metagenomics\(^\text{33,36,37}\).

To further investigate the claim\(^\text{25}\) that the tree inferred depends on the choice of universal marker genes, we used the Orthologous MAnrix (OMA\(^\text{38}\)) algorithm to identify single-copy orthologues de novo on the 125 genome set. Benchmarks\(^\text{39}\) indicate that OMA is conservative, in that it returns a relatively low number of orthologues but that these orthologues perform better than other methods at recovering the species tree. Combining OMA analysis with manual filtering to remove EF2 and genes of endosymbiotic origin (see Methods), we identified 21 broadly conserved marker genes found in at least half of our set of bacteria, archaea and eukaryotes, and 43 genes encoded by at least half of the archaea and eukaryotes (see Methods). We concatenated the 21 genes conserved in all three domains and inferred a tree under CAT + GTR + G4 (Fig. 3a). rooting on the branch separating bacteria and archaea resulted in a 2D tree, in which eukaryotes form a maximally supported clade with Asgard archaea (Fig. 3a); within Asgards, the closest relatives of eukaryotes was recovered as the Heimdallarchaeota, although with only modest support (PP = 0.79).

We next analysed the expanded set of genes conserved between archaea and eukaryotes, placing the root outside the TACK/Asgard/eukaryote clade as suggested by the previous analysis including bacteria. The consensus tree under CAT + GTR + G4 (Fig. 3b) resolves a clade of eukaryotes and Heimdallarchaeota with maximal posterior support; within that clade, eukaryotes group with one Heimdallarchaeota metagenome bin (LC3) with high (PP = 0.95) support.

Given ongoing debates about the impact of even single genes within concatenated datasets, we investigated in detail the overlap between the 35-gene set, the 21 genes selected by OMA and a 29-gene set used in some previous analyses\(^\text{11,12,60,61}\) (Supplementary Table 10). After removing EF2, seven genes are found in all three sets; 27 in at least two of the three and 50 genes in total are present in at least one of the datasets. We obtained the orthologues for the 50-gene families from the 125 species dataset and inferred trees using the best-fit maximum-likelihood model in IQ-Tree on the 7-, 27- and 50-gene concatenations (Supplementary Fig. 8). We also expanded species sampling for the 35 genes to compare with the analyses described above. Analysis under the best-fitting maximum-likelihood model for all four concatenates resulted in a 2D tree, with either all Asgards (the 7- and 35-gene datasets) or Heimdallarchaeota (27- and 50-gene datasets) as sister to eukaryotes with moderate (7-gene set) to high (the other sets) bootstrap support. These results indicate that there is a congruent signal for a 2D tree, and a relationship between eukaryotes and Asgard archaea, that is robust to moderate differences in the choice of marker genes. The results of all our concatenation analyses are summarized in Supplementary Table 11.

**Supertree and multispecies coalescent methods support the 2D tree.** Concatenation allows phylogenetic signal to be pooled and permits the use of complex, parameter-rich substitution models but its assumptions are problematic in the context of microbial evolution. In particular, concatenation requires that all of the genes share a common phylogeny\(^\text{62,63}\), an assumption that is difficult to test because trees inferred from individual genes are often poorly supported. Some incongruence between single-gene trees can be attributed to stochastic error or model misspecification\(^\text{14}\) but genuinely
different evolutionary histories for different genes can arise from incomplete lineage sorting, gene duplication and loss and horizontal gene transfer. We therefore investigated alternative methods for integrating phylogenetic signal from multigene datasets that account for gene tree incongruence in different ways. The probabilistic supertree method of Steel and Rodrigo (SR2008) and the
When data were simulated under the KVR model, the method model to recover the root of the tree from protein fold datasets. division within cellular life and rejecting both the 2D and 3D trees. as the sister lineage to the three Heimdallarchaeota metagenome method and ASTRAL placed eukaryotes within the Asgard archaea our dataset (of these 3,199 gene families, 479 included at least one broadly conserved across the species of interest, we analysed a set classification of proteins families against a set of bacterial, archaeal and eukaryotes as sister taxa. These results suggest that there is a split presence–absence (SPA) method, are supertree methods that model differences between gene trees as stochastic noise; ASTRAL is a supertree method that is consistent under the multispecies coalescent. These methods have their own assumptions and limitations but these are distinct from—and provide a useful contrast to—concatenation. As these methods do not require genes to be broadly conserved across the species of interest, we analysed a set of 3,199 single-copy orthologues found in at least four of the taxa in our dataset (of these 3,199 gene families, 479 included at least one archaeon and one eukaryote; see Supplementary Table 12 for the taxonomic distribution and phylogenetic relationships supported by the individual trees).

All of these analyses resolved a 2D tree including a clade of eukaryotes and Asgard archaea with high to maximal support (Supplementary Figs. 9–10). Supertrees inferred under the SPA method and ASTRAL placed eukaryotes within the Asgard archaea as the sister lineage to the three Heimdallarchaeota metagenome bins (Supplementary Figs. 9–10), while the SR2008 supertree recovered eukaryotes and Asgard archaea as monophyletic sister lineages (Supplementary Fig. 10). To compare these supertrees independently of their models and assumptions, we calculated the summed quartet distances between the set of input trees and each supertree: the total number of quartets (subtrees of four leaves) that differ between the input trees and each supertree (Table 1). The tree with the best score by this metric was the model-based MSC ASTRAL analysis which, like the SPA supertree, recovered Heimdallarchaeota and eukaryotes as sister taxa. These results suggest that there is a congruent genome-wide signal for a specific relationship between eukaryotes and the Heimdallarchaeota, and that the 2D tree does not appear to be an artefact of concatenation.

Is there support from protein folds for a root between prokaryotes and eukaryotes? Debates about the 2D and 3D trees have typically assumed that the root of the tree lies on the branch separating bacteria and archaea or within the bacteria. Recently, a non-stationary model of binary character evolution (the KVR model) was used to infer a rooted tree of life from a matrix of protein fold presence–absence data. Fold presence and absence were quantified by searching Hidden Markov Models (HMMs) corresponding to structural classification of proteins families against a set of bacterial, archaeal and eukaryotic genomes. The inferred trees are intrinsically rooted because the model is non-stationary: in this model there is one composition (probability of protein fold presence) at the root of the tree and a second composition elsewhere. These analyses recovered a root between prokaryotes and eukaryotes, suggesting this is the primary division within cellular life and rejecting both the 2D and 3D trees.

We performed simulations to evaluate the ability of the KVR model to recover the root of the tree from protein fold datasets. When data were simulated under the KVR model, the method recovered the true root of the simulation tree as might be expected. However, when protein fold compositions were allowed to vary over the tree, something which is observed in the empirical data, the model fails to find the true root. Under these conditions, KVR finds a root on one of the branches with atypical sequence composition (see Supplementary Discussion). In the empirical data matrix, the eukaryotes encode significantly more protein folds than either bacteria or archaea (median of 871 folds per eukaryotic genome, compared to 521 for archaea and 615 for bacteria; \( P < 10^{-4} \) for the eukaryote–archaea and eukaryote–bacteria comparisons, \( P = 0.000278 \) comparing bacteria and archaea; \( n = 47 \) eukaryotes, 47 bacteria and 47 archaea, Wilcoxon rank-sum tests) but their higher compositions are in the minority because the matrix contains an equal number of genomes from each of the three domains. Thus, the inferred root between prokaryotes and eukaryotes may result from the model’s bias in placing the root on a branch with atypical composition; in simulations, the root inference can be controlled by varying which composition among tips—high or low—is in the majority (Supplementary Discussion). These results agree with recent work in suggesting that non-reversible models may provide reliable rooting information when the assumptions of the model are met but that root inferences are sensitive to model misspecification. The KVR model is only one of the many possible non-stationary and non-homogeneous models and does not appear to be well-suited to these data. Models that better describe the process by which fold (or sequence) compositions change through time and across the tree—or indeed those that make use of other sources of time information—may perform better for rooting deep phylogenies. How best to root ancient radiations remains an open question and method development is still at an early stage. A key challenge will be the development of methods that account for the heterogeneity of the evolutionary process across the data and through evolutionary time (across the branches of the tree).

A potentially bigger problem than model misspecification for the published analyses is their assumption that the entire protein fold set evolves on a single underlying tree. This assumption is unlikely to be realistic because of the different histories generated by widespread horizontal gene transfer and, in eukaryotes, by endosymbiotic gene transfer from the bacterial progenitors of mitochondria and plastids. The assumption of a single underlying tree to explain fold distributions also means that, despite claims to the contrary, the published analyses cannot be used to reject the 2D tree because, as generally formulated, it seeks to explain the inheritance of only a subset of the genes on cellular genomes.

To evaluate whether the protein folds in the published matrix share a common evolutionary tree, we inferred single-gene phylogenies for each fold (Supplementary Discussion). Although weakly supported, these trees are consistent with there being extensive disagreement between single fold-based topologies: only 22 of the protein folds supported the monophyly of eukaryotes and none recovered all three domains as potentially monophyletic groups, even though this was the consensus topology obtained from analysis of the complete matrix. The trees contained signals for sister-group relationships between eukaryotes and alphaproteobacteria (the most frequent sister group among the protein folds shared between eukaryotes and bacteria) and for a relationship between eukaryotes and the TACKL archaea. These analyses are consistent with endosymbiotic theory and the ideas that underpin the 2D tree, namely that eukaryotes contain a mixture of genes from the archaean host cell and the bacterial endosymbiont that became the mitochondrion (Supplementary Discussion).

Conclusions

Identifying the tree that best depicts the relationships between the major groups of life is important for understanding eukaryotic origins and the evolution of the complexity that distinguishes
eukaryotic cells. It has recently been asserted that the tree recovered depends upon the species investigated and the choice and quality of the molecular data analysed12,20. In the present study we have investigated the datasets used to underpin these claims and find no compelling evidence to support them. Analyses using better-fitting phylogenetic models consistently recovered a 2D tree11,12,13,15,16,17,19,20 wherein eukaryotes are most closely related to members of the recently discovered Asgard archaea. These results are also supported by additional analyses of expanded concatenations and increased species sampling, and from large-scale genome-wide datasets analysed using supertree and coalescence methods.

We also investigated support from analyses of whole-genome protein folds for a rooted universal tree in which the deepest division is between prokaryotes and eukaryotes. Taken at face-value this tree would reject the 2D and 3D trees that are the focus of robust discussion in the current literature9–20. However, while protein structure is a useful guide to identifying homology when primary sequence similarity is weak, how best to analyse fold data to resolve deep phylogenetic relationships is still not clear. Published analyses11 do not account for the varied evolutionary histories of individual folds due to endosymbiosis and gene transfer, and our simulations suggest that root inference under existing models is unreliable and affected by variation in the abundance and distribution of folds across genomes. At present, the best-supported root is on the branch separating bacteria and archaea11,13,14,20 or among the bacteria11,12, and the hypothesis that eukaryotes are younger than prokaryotes is supported by a range of phylogenetic, cell biological8 and palaeontological11,12,16–20 evidence.

Our analyses and published trees11,16,20 imply that the eukaryotic nuclear lineage evolved from within the archaea. They provide robust phylogenomic support for a clade of eukaryotes and Asgard archaea, and identify the Heimdallarchaeota as the best candidate among sampled lineages11,13,14,20 for a sister group to eukaryotes. This sister-group relationship will no doubt change with further sampling of the potentially vast archaeal diversity in nature still to be discovered. The prize will be ever more reliable inferences of the features that were in place in the last common ancestor of both groups and an improved evidence-based understanding of the building blocks that underpinned the transition from prokaryotic to eukaryotic cells.

Methods

Sequence alignment. For the reanalyses of the Da Cunha et al. and Spang et al. datasets, alignments were obtained from the supplementary material of Da Cunha et al.21 and the EF2 gene removed according to the coordinates provided; the alignments from Spang et al.14 were generously provided by the authors. OMA 2.1.1 (ref. 13) was used to identify putative single-copy orthologues among a dataset of 92 eukaryotic, archaean and bacterial genomes. For putative orthologues present in at least half of the sampled species, single-gene trees were inferred for each candidate under the LG+G+F model in IQ-TREE and the trees were manually inspected to filter out eukaryotic genes that were acquired from the mitochondrial or plastid endosymbionts. We also performed a BLASTP screen to identify organellar genes that might have been missed via the tree inspection approach. This procedure resulted in the retrieval of 43 single-copy orthologues shared between archaea and eukaryotes and 21 genes shared among all three domains, that were used for concatenation-based phylogenomic analyses. For all OMA gene families found in at least four species, we used a BLASTP-based screen to identify and filter out eukaryotic gene families of bacterial origin, resulting in 3,261 gene families in four or more species that are either eukaryotic-specific inventions or shared between eukaryotes and archaea. For the comparisons of core gene sets, an iterative process of manual comparisons, similarity searches and tree building was used to identify common and distinct markers in the published sets, identify seed sequences for each marker in the genomes of Dictyostelium discoideum, Sulfolobus solfataricus and Escherichia coli strain K12, and build HMMs for each marker using the existing datasets. We used domain-specific HMM searches in HMMER3 (ref. 12) followed by the reciprocal best-hit criterion against our domain-specific reference genomes to identify candidate orthologues, followed by gene tree inference and manual curation to assemble final marker sets. Sequences were aligned using the L-INS-i mode in Mafft 7 (ref. 12) and poorly aligning regions identified and removed using the BLOSUM30 model in RMGE 1.12 (ref. 12).

Phylogenetics. Maximum-likelihood analyses were performed using IQ-TREE 1.6.2 (ref. 13) and bootstrap supports were computed using UFBoot2 (ref. 13), except where indicated in the main text. Model fitting was carried out using the M property in IQ-TREE, adding the empirical site profile models (C20–C60) to the default candidate model set. Bayesian phylogenies were inferred under the CAT+GTR+G4 model in PhyloBayes-MP1 1.8 (ref. 14), using the bpcomp and tracceomp programmes to monitor convergence of two MCMC chains for each analysis. Posterior predictive simulations were performed using readbp_mpi in PhyloBayes. Tests for across-branch compositional heterogeneity were performed in p4 (ref. 15); we inferred maximum-likelihood gene trees for each of the 35 genes in the concatenation, then simulated data for each gene under the LG+G+F model. A Chi-square statistic reflecting compositional heterogeneity was calculated on the original and simulated datasets and the values from the simulated data were used as a null distribution with which to evaluate the test statistic from the original data.

Supertrees. Supertrees were inferred from the maximum-likelihood phylogenies for each single gene, with substitution models chosen as described above. MRP, SR2008 and SPA supertrees were inferred using p4 (ref. 16). Multispecies coalescent trees were inferred using ASTRAL-III (ref. 17).

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The data associated with our analyses are available in the FigShare repository at https://doi.org/10.1084/nmfigshare.8950859.v2.

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**Author contributions**

All authors contributed to the conception and design of the project and to the interpretation of results. T.A.W., C.J.C., P.G.F. and G.J.S. performed analyses. T.A.W. and T.M.E. wrote the manuscript, with input from all authors.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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- No software was used.

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- OMA was used for orthologue inference. Mafft was used for sequence alignment, BMGE for identifying and removing poorly-aligned sites. IQ-Tree 1.6.2 was used for inference of maximum likelihood trees (with RAxML 8 and PhyML 3.1 where indicated). PhyloBayes-MPI 1.8 and p4 1.2.0 were used for Bayesian analyses.

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Sampling strategy: Taxa were subsampled from the larger diversity of each of the domains so as to maximize the representation of known diversity within a total dataset size for which fitting of the best available phylogenetic models is tractable.

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