

A new view of the tree of life

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The tree of life is one of the most important organizing principles in biology¹. Gene surveys suggest the existence of an enormous number of branches², but even an approximation of the full scale of the tree has remained elusive. Recent depictions of the tree of life have focused either on the nature of deep evolutionary relationships^{3–5} or on the known, well-classified diversity of life with an emphasis on eukaryotes⁶. These approaches overlook the dramatic change in our understanding of life's diversity resulting from genomic sampling of previously unexamined environments. New methods to generate genome sequences illuminate the identity of organisms and their metabolic capacities, placing them in community and ecosystem contexts^{7,8}. Here, we use new genomic data from over 1,000 uncultivated and little known organisms, together with published sequences, to infer a dramatically expanded version of the tree of life, with Bacteria, Archaea and Eukarya included. The depiction is both a global overview and a snapshot of the diversity within each major lineage. The results reveal the dominance of bacterial diversification and underline the importance of organisms lacking isolated representatives, with substantial evolution concentrated in a major radiation of such organisms. This tree highlights major lineages currently underrepresented in biogeochemical models and identifies radiations that are probably important for future evolutionary analyses.

Early approaches to describe the tree of life distinguished organisms based on their physical characteristics and metabolic features. Molecular methods dramatically broadened the diversity that could be included in the tree because they circumvented the need for direct observation and experimentation by relying on sequenced genes as markers for lineages. Gene surveys, typically using the small subunit ribosomal RNA (SSU rRNA) gene, provided a remarkable and novel view of the biological world^{1,9,10}, but questions about the structure and extent of diversity remain. Organisms from novel lineages have eluded surveys, because many are invisible to these methods due to sequence divergence relative to the primers commonly used for gene amplification^{7,11}. Furthermore, unusual sequences, including those with unexpected insertions, may be discarded as artefacts⁷.

Whole genome reconstruction was first accomplished in 1995 (ref. 12), with a near-exponential increase in the number of draft genomes reported each subsequent year. There are 30,437 genomes from all three domains of life—Bacteria, Archaea and Eukarya—which are currently available in the Joint Genome Institute's Integrated Microbial Genomes database (accessed 24 September 2015).

Contributing to this expansion in genome numbers are single cell genomics¹³ and metagenomics studies. Metagenomics is a shotgun sequencing-based method in which DNA isolated directly from the environment is sequenced, and the reconstructed genome fragments are assigned to draft genomes¹⁴. New bioinformatics methods yield complete and near-complete genome sequences, without a reliance on cultivation or reference genomes^{7,15}. These genome- (rather than gene) based approaches provide information about metabolic potential and a variety of phylogenetically informative sequences that can be used to classify organisms¹⁶. Here, we have constructed a tree of life by making use of genomes from public databases and 1,011 newly reconstructed genomes that we recovered from a variety of environments (see Methods).

To render this tree of life, we aligned and concatenated a set of 16 ribosomal protein sequences from each organism. This approach yields a higher-resolution tree than is obtained from a single gene, such as the widely used 16S rRNA gene¹⁶. The use of ribosomal proteins avoids artefacts that would arise from phylogenies constructed using genes with unrelated functions and subject to different evolutionary processes. Another important advantage of the chosen ribosomal proteins is that they tend to be syntenic and co-located in a small genomic region in Bacteria and Archaea, reducing binning errors that could substantially perturb the geometry of the tree. Included in this tree is one representative per genus for all genera for which high-quality draft and complete genomes exist (3,083 organisms in total).

Despite the methodological challenges, we have included representatives of all three domains of life. Our primary focus relates to the status of Bacteria and Archaea, as these organisms have been most difficult to profile using macroscopic approaches, and substantial progress has been made recently with acquisition of new genome sequences^{7,8,13}. The placement of Eukarya relative to Bacteria and Archaea is controversial^{1,4,5,17,18}. Eukaryotes are believed to be evolutionary chimaeras that arose via endosymbiotic fusion, probably involving bacterial and archaeal cells¹⁹. Here, we do not attempt to confidently resolve the placement of the Eukarya. We position them using sequences of a subset of their nuclear-encoded ribosomal proteins, an approach that classifies them based on the inheritance of their information systems as opposed to lipid or other cellular structures⁵.

Figure 1 presents a new view of the tree of life. This is one of a relatively small number of three-domain trees constructed from molecular information so far, and the first comprehensive tree to

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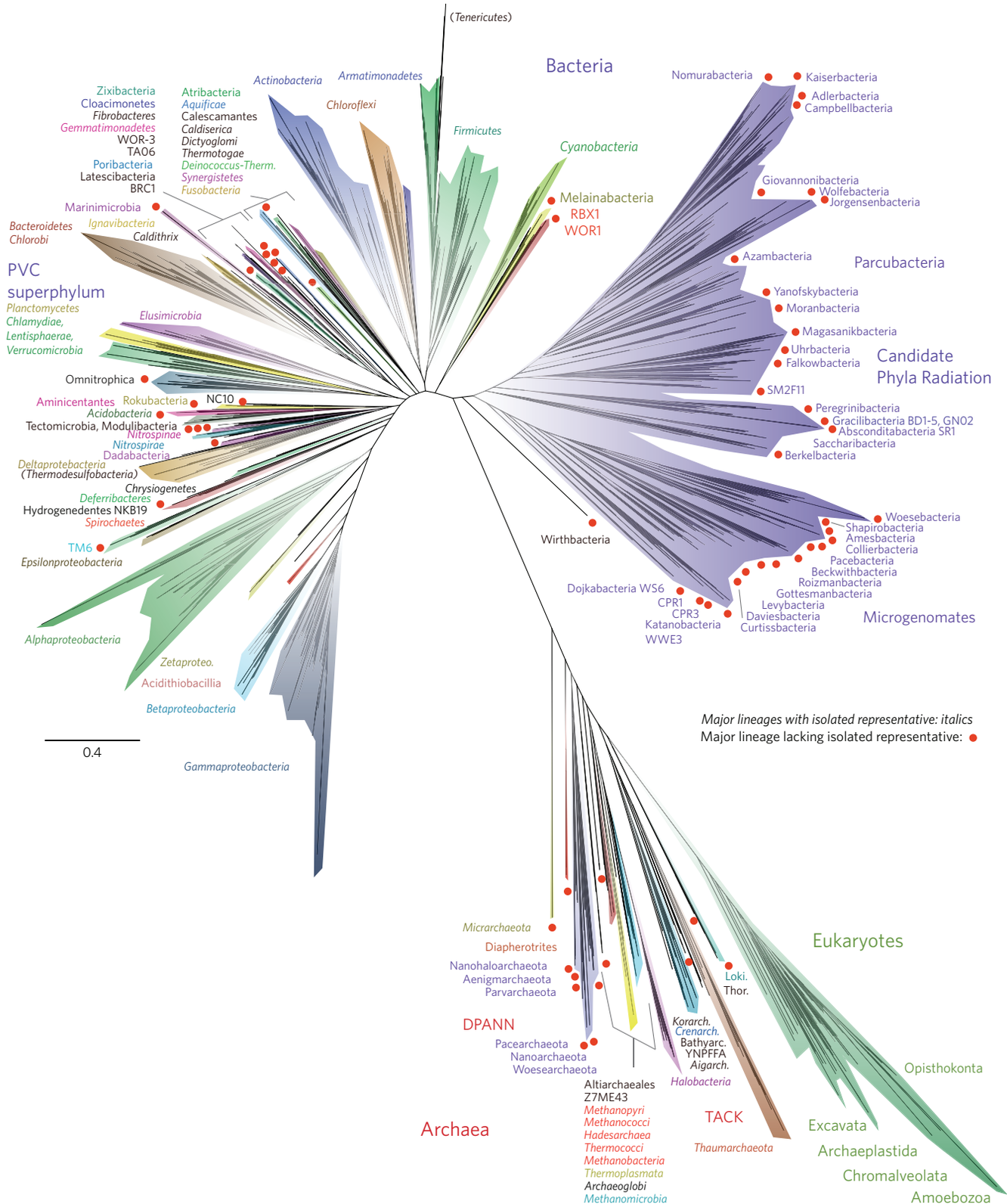


Figure 1 | A current view of the tree of life, encompassing the total diversity represented by sequenced genomes. The tree includes 92 named bacterial phyla, 26 archaeal phyla and all five of the Eukaryotic supergroups. Major lineages are assigned arbitrary colours and named, with well-characterized lineage names, in italics. Lineages lacking an isolated representative are highlighted with non-italicized names and red dots. For details on taxon sampling and tree inference, see Methods. The names Tenericutes and Thermodesulfobacteria are bracketed to indicate that these lineages branch within the Firmicutes and the Deltaproteobacteria, respectively. Eukaryotic supergroups are noted, but not otherwise delineated due to the low resolution of these lineages. The CPR phyla are assigned a single colour as they are composed entirely of organisms without isolated representatives, and are still in the process of definition at lower taxonomic levels. The complete ribosomal protein tree is available in rectangular format with full bootstrap values as Supplementary Fig. 1 and in Newick format in Supplementary Dataset 2.

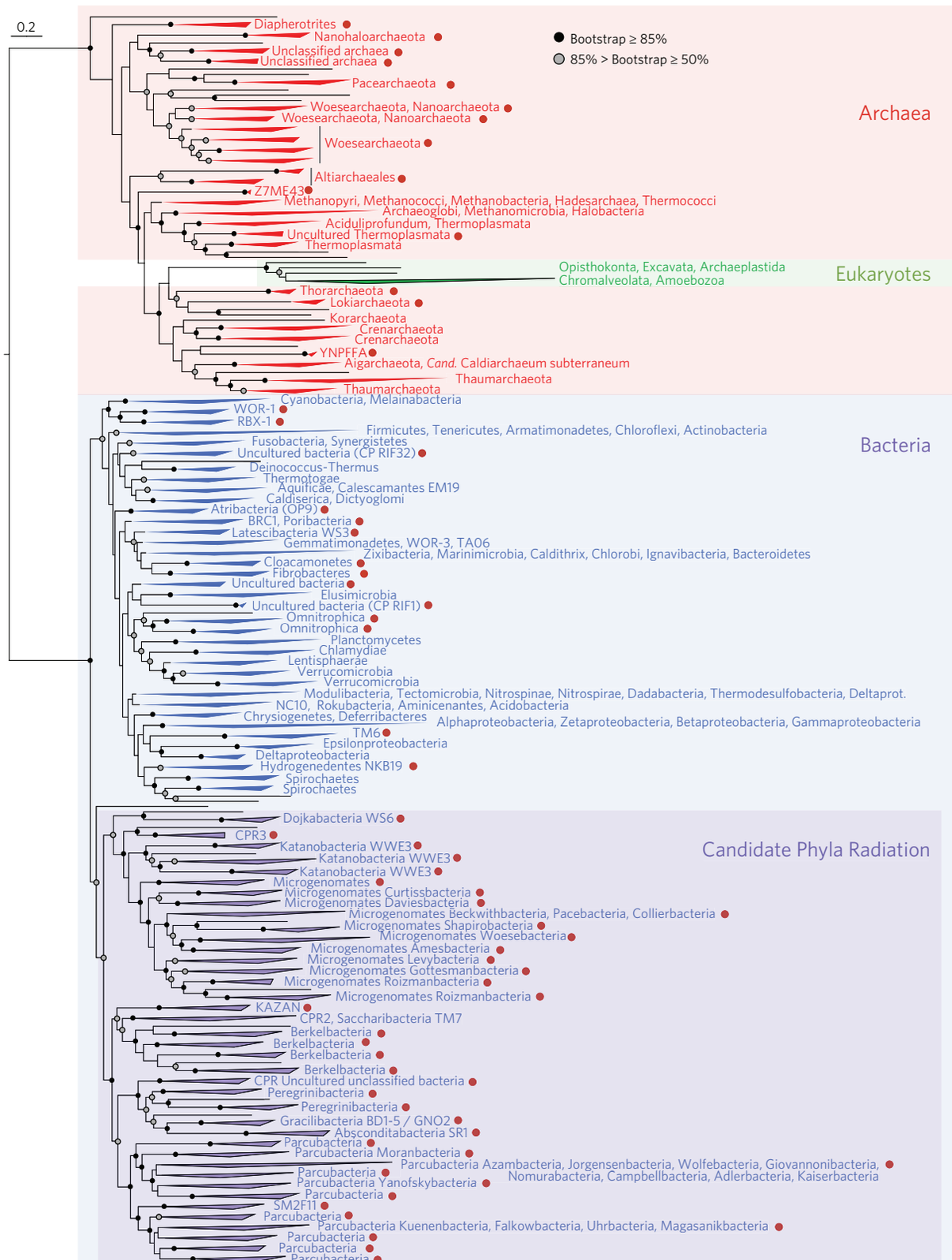


Figure 2 | A reformatted view of the tree in Fig. 1 in which each major lineage represents the same amount of evolutionary distance. The threshold for groups (coloured wedges) was an average branch length of <0.65 substitutions per site. Notably, some well-accepted phyla become single groups and others are split into multiple distinct groups. We undertook this analysis to provide perspective on the structure of the tree, and do not propose the resulting groups to have special taxonomic status. The massive scale of diversity in the CPR and the large fraction of major lineages that lack isolated representatives (red dots) are apparent from this analysis. Bootstrap support values are indicated by circles on nodes—black for support of 85% and above, grey for support from 50 to 84%. The complete ribosomal protein tree is available in rectangular format with full bootstrap values as Supplementary Fig. 1 and in Newick format in Supplementary Dataset 2.

be published since the development of genome-resolved metagenomics. We highlight all major lineages with genomic representation, most of which are phylum-level branches (see Supplementary Fig. 1 for full bootstrap support values). However, we separately identify the Classes of the Proteobacteria, because the phylum is not monophyletic (for example, the Deltaproteobacteria branch away from the other Proteobacteria, as previously reported^{2,20}).

The tree in Fig. 1 recapitulates expected organism groupings at most taxonomic levels and is largely congruent with the tree calculated using traditional SSU rRNA gene sequence information (Supplementary Fig. 2). The support values for taxonomic groups are strong at the Species through Class levels (>85%), with moderate-to-strong support for Phyla (>75% in most cases), but the branching order of the deepest branches cannot be confidently resolved (Supplementary Fig. 1). The lower support for deep branch placements is a consequence of our prioritization of taxon sampling over number of genes used for tree construction. As proposed recently, the Eukarya, a group that includes protists, fungi, plants and animals, branches within the Archaea, specifically within the TACK superphylum²¹ and sibling to the Lokiarchaeota²². Interestingly, this placement is not evident in the SSU rRNA tree, which has the three-domain topology proposed by Woese and co-workers in 1990¹ (Supplementary Fig. 2). The two-domain Eocyte tree and the three-domain tree are competing hypotheses for the origin of Eukarya⁵; further analyses to resolve these and other deep relationships will be strengthened with the availability of genomes for a greater diversity of organisms. Important advantages of the ribosomal protein tree compared with the SSU rRNA gene tree are that it includes organisms with incomplete or unavailable SSU rRNA gene sequences and more strongly resolves the deeper radiations. Ribosomal proteins have been shown to contain compositional biases across the three domains, driven by thermophilic, mesophilic and halophilic lifestyles as well as by a primitive genetic code²³. Continued expansion of the number of genome sequences for non-extremophile Archaea, such as the DPANN lineages^{8,13}, may allow clarification of these compositional biases.

A striking feature of this tree is the large number of major lineages without isolated representatives (red dots in Fig. 1). Many of these lineages are clustered together into discrete regions of the tree. Of particular note is the Candidate Phyla Radiation (CPR)⁷, highlighted in purple in Fig. 1. Based on information available from hundreds of genomes from genome-resolved metagenomics and single-cell genomics methods to date, all members have relatively small genomes and most have somewhat (if not highly) restricted metabolic capacities^{7,13,24}. Many are inferred (and some have been shown) to be symbionts^{7,25,26}. Thus far, all cells lack complete citric acid cycles and respiratory chains and most have limited or no ability to synthesize nucleotides and amino acids. It remains unclear whether these reduced metabolisms are a consequence of superphylum-wide loss of capacities or if these are inherited characteristics that hint at an early metabolic platform for life. If inherited, then adoption of symbiotic lifestyles may have been a later innovation by these organisms once more complex organisms appeared.

Figure 2 presents another perspective, where the major lineages of the tree are defined using evolutionary distance, so that the main groups become apparent without bias arising from historical naming conventions. This depiction uses the same inferred tree as in Fig. 1, but with groups defined on the basis of average branch length to the leaf taxa. We chose an average branch length that best recapitulated the current taxonomy (smaller values fragmented many currently accepted phyla and larger values collapsed accepted phyla into very few lineages, see Methods). Evident in Fig. 2 is the enormous extent of evolution that has occurred within the CPR. The diversity within the CPR could be a result of the early emergence of this group and/or a consequence of rapid evolution related to

symbiotic lifestyles. The CPR is early-emerging on the ribosomal protein tree (Fig. 1), but not in the SSU rRNA tree (Supplementary Fig. 2). Regardless of branching order, the CPR, in combination with other lineages that lack isolated representatives (red dots in Fig. 2), clearly comprises the majority of life's current diversity.

Domain Bacteria includes more major lineages of organisms than the other Domains. We do not attribute the smaller scope of the Archaea relative to Bacteria to sampling bias because metagenomics and single-cell genomics methods detect members of both domains equally well. Consistent with this view, Archaea are less prominent and less diverse in many ecosystems (for example, seawater²⁷, hydrothermal vents²⁸, the terrestrial subsurface¹⁵ and human-associated microbiomes²⁹). The lower apparent phylogenetic diversity of Eukarya is fully expected, based on their comparatively recent evolution.

The tree of life as we know it has dramatically expanded due to new genomic sampling of previously enigmatic or unknown microbial lineages. This depiction of the tree captures the current genomic sampling of life, illustrating the progress that has been made in the last two decades following the first published genome. What emerges from analysis of this tree is the depth of evolutionary history that is contained within the Bacteria, in part due to the CPR, which appears to subdivide the domain. Most importantly, the analysis highlights the large fraction of diversity that is currently only accessible via cultivation-independent genome-resolved approaches.

Methods

A data set comprehensively covering the three domains of life was generated using publicly available genomes from the Joint Genome Institute's IMG-M database (img.jgi.doe.gov), a previously developed data set of eukaryotic genome information³⁰, previously published genomes derived from metagenomic data sets^{7,8,31,32} and newly reconstructed genomes from current metagenome projects (see Supplementary Table 1 for NCBI accession numbers). From IMG-M, genomes were sampled such that a single representative for each defined genus was selected. For phyla and candidate phyla lacking full taxonomic definition, every member of the phylum was initially included. Subsequently, these radiations were sampled to an approximate genus level of divergence based on comparison with taxonomically described phyla, thus removing strain- and species-level overlaps. Finally, initial tree reconstructions identified aberrant long-branch attraction effects placing the Microsporidia, a group of parasitic fungi, with the Korarchaeota. The Microsporidia are known to contribute long branch attraction artefacts confounding placement of the Eukarya³³, and were subsequently removed from the analysis.

This study includes 1,011 organisms from lineages for which genomes were not previously available. The organisms were present in samples collected from a shallow aquifer system, a deep subsurface research site in Japan, a salt crust in the Atacama Desert, grassland meadow soil in northern California, a CO₂-rich geyser system, and two dolphin mouths. Genomes were reconstructed from metagenomes as described previously⁷. Genomes were only included if they were estimated to be >70% complete based on presence/absence of a suite of 51 single copy genes for Bacteria and 38 single copy genes for Archaea. Genomes were additionally required to have consistent nucleotide composition and coverage across scaffolds, as determined using the ggkbase binning software (ggkbase.berkeley.edu), and to show consistent placement across both SSU rRNA and concatenated ribosomal protein phylogenies. This contributed marker gene information for 1,011 newly sampled organisms, whose genomes were reconstructed for metabolic analyses to be published separately.

The concatenated ribosomal protein alignment was constructed as described previously¹⁶. In brief, the 16 ribosomal protein data sets (ribosomal proteins L2, L3, L4, L5, L6, L14, L16, L18, L22, L24, S3, S8, S10, S17 and S19) were aligned independently using MUSCLE v. 3.8.31 (ref. 34). Alignments were trimmed to remove ambiguously aligned C and N termini as well as columns composed of more than 95% gaps. Taxa were removed if their available sequence data represented less than 50% of the expected alignment columns (90% of taxa had more than 80% of the expected alignment columns). The 16 alignments were concatenated, forming a final alignment comprising 3,083 genomes and 2,596 amino-acid positions. A maximum likelihood tree was constructed using RAXML v. 8.1.24 (ref. 35), as implemented on the CIPRES web server³⁶, under the LG plus gamma model of evolution (PROTGAMMALG in the RAXML model section), and with the number of bootstraps automatically determined (MRE-based bootstrapping criterion). A total of 156 bootstrap replicates were conducted under the rapid bootstrapping algorithm, with 100 sampled to generate proportional support values. The full tree inference required 3,840 computational hours on the CIPRES supercomputer.

To construct Fig. 2, we collapsed branches based on an average branch length criterion. Average branch length calculations were implemented in the Interactive Tree of Life online interface³⁷ using the formula:

Average branch length = mean([root distance to tip] – [root distance to node]) for all tips connecting to a node.

We tested values between 0.25 and 0.75 at 0.05 intervals, and selected a final threshold of <0.65 based on generation of a similar number of major lineages as compared to the taxonomy-guided clustering view in Fig. 1. The taxonomy view identified 26 archaeal and 74 bacterial phylum-level lineages (counting the Microgenomates and Parcubacteria as single phyla each), whereas an average branch length of <0.65 resulted in 28 archaeal and 76 bacterial clades.

For a companion SSU rRNA tree, an alignment was generated from all SSU rRNA genes available from the genomes of the organisms included in the ribosomal protein data set. For organisms with multiple SSU rRNA genes, one representative gene was kept for the analysis, selected randomly. As genome sampling was confined to the genus level, we do not anticipate this selection process will have any impact on the resultant tree. All SSU rRNA genes longer than 600 bp were aligned using the SINA alignment algorithm through the SILVA web interface^{38,39}. The full alignment was stripped of columns containing 95% or more gaps, generating a final alignment containing 1,871 taxa and 1,947 alignment positions. A maximum likelihood tree was inferred as described for the concatenated ribosomal protein trees, with RAXML run using the GTRCAT model of evolution. The RAXML inference included the calculation of 300 bootstrap iterations (extended majority rules-based bootstrapping criterion), with 100 randomly sampled to determine support values.

To test the effect of site selection stringency on the inferred phylogenies, we stripped the alignments of columns containing up to 50% gaps (compared with the original trimming of 95% gaps). For the ribosomal protein alignment, this resulted in a 14% reduction in alignment length (to 2,232 positions) and a 44.6% reduction in computational time (~2,100 h). For the SSU rRNA gene alignment, stripping columns with 50% or greater gaps reduced the alignment by 24% (to 1,489 positions) and the computation time by 28%. In both cases, the topology of the tree with the best likelihood was not changed significantly. The ribosomal protein resolved a two-domain tree with the Eukarya sibling to the Lokiarchaeota, while the SSU rRNA tree depicts a three-domain tree. The position of the CPR as deep-branching on the ribosomal protein tree and within the Bacteria on the SSU rRNA tree was also consistent. The alignments and inferred trees under the more stringent gap stripping are available upon request.

Nomenclature. We have included names for two lineages for which we have previously published complete genomes⁴⁰. At the time of submission of the paper describing these genomes⁴⁰, the reviewer community was not uniformly open to naming lineages of uncultivated organisms based on such information. Given that this practice is now widely used, we re-propose the names for these phyla. Specifically, for WW3 we suggest the name Katanobacteria from the Hebrew 'katan', which means 'small', and for SR1 we suggest the name Absconditabacteria from the Latin 'Abscondo' meaning 'hidden', as in 'shrouded'.

Accession codes. NCBI and/or JGI IMG accession numbers for all genomes used in this study are listed in Supplementary Table 1. Additional ribosomal protein gene and 16S rRNA gene sequences used in this study have been deposited in Genbank under accession numbers KU868081–KU869521. The concatenated ribosomal protein and SSU rRNA alignments used for tree reconstruction are included as separate files in the Supplementary Information.

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Author contributions

L.A.H. and J.F.B. defined the research objective. L.A.H. generated data sets and conducted phylogenetic tree inferences. L.A.H., B.J.B. and J.F.B. conducted data analysis. L.A.H.,

B.J.B., K.A., C.T.B., A.J.P., C.J.C., C.N.B., A.W.H., Y.A., K.I., Y.S., N.D., D.A.R., K.M.F., R.A., B.C.T. and J.F.B. contributed to metagenome binning and genome analysis. L.A.H. and J.F.B. wrote the manuscript with input from all authors. All authors read and approved the final manuscript.

Additional information

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Competing interests

The authors declare no competing financial interests.



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