Genome beginnings: rooting the tree of life

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A rooted tree of life provides a framework to answer central questions about the evolution of life. Here we review progress on rooting the tree of life and introduce a new root of life obtained through the analysis of indels, insertions and deletions, found within paralogous gene sets. Through the analysis of indels in eight paralogous gene sets, the root is localized to the branch between the clade consisting of the Actinobacteria and the double-membrane (Gram-negative) prokaryotes and one consisting of the archaeabacteria and the firmicutes. This root provides a new perspective on the habitats of early life, including the evolution of methanogenesis, membranes and hyperthermophily, and the speciation of major prokaryotic taxa. Our analyses exclude methanogenesis as a primitive metabolism, in contrast to previous findings. They parsimoniously imply that the ether archaeabacterial lipids are not primitive and that the cenancestral prokaryotic population consisted of organisms enclosed by a single, ester-linked lipid membrane, covered by a peptidoglycan layer. These results explain the similarities previously noted by others between the lipid synthesis pathways in eubacteria and archaeabacteria. The new root also implies that the last common ancestor was not hyperthermophilic, although moderate thermophily cannot be excluded.

Keywords: tree of life; root; indels; eubacteria; firmicutes; archaeabacteria; eukaryotes

1. INTRODUCTION

Today, there is enormous interest in reconstructing the rooted tree of prokaryotic life, but there is little or no agreement about the topology of the net, the web, the ring, the tree or the non-tree of life (Hilario & Gogarten 1993; Doolittle 1999; Jain et al. 2003; Rivera & Lake 2004; Konstantinidis & Tiedje 2005; Dagan & Martin 2006; McInerney & Pisani 2007; Sorek et al. 2007), and there is even less agreement about the location of the root.

Rooting is important because rooted ‘trees’ provide a framework for answering fundamental questions about the evolution of life. An accurate representation of a rooted tree, or graph, of life’s history allows one to test theories for novel innovations in the evolution of life. With rooting information, it becomes possible to relate genetic, biochemical, ultrastructural and behavioural innovations to geological, paleontological and climatological events, thereby allowing one to trace the interdependent histories of the Earth and its microbiota and to test theories for the order of appearance of novel biological innovations.

Rooted trees allow central assumptions to be tested, such as: did the cenancestor, the last common population ancestral to all extant life, live in a hot environment? Many think so, but others argue for moderate temperatures (Miller & Bada 1988), and even the concept of a knowable cenancestor (Gogarten et al. 2002) has been questioned. Did the archaeabacteria evolve in a hot environment? Again many think so, but the guanine–cytosine (GC) compositions of ribosomal RNAs suggest that the most recent common ancestor was not an extreme thermophile (Galtier et al. 1999). Did carbon heterotrophy evolve before autotrophy, or vice versa? Did prokaryotes with single membranes precede (Gupta 1998) or follow (Cavalier-Smith 2002) prokaryotes with double membranes? Did early organisms respire, or was substrate level phosphorylation used, and was methanogenesis early or late (Russell & Hall 1997; Grassineau et al. 2001; Bada & Lazcano 2002; Martin & Russell 2003; Purdy et al. 2003; Chistoserdova et al. 2004; Russell & Martin 2004; Ferry & House 2006; Gribaldo & Brochier-Armanet 2006)? Are phylogenetic signals sufficiently well preserved in eukaryotic genomes to permit the founding prokaryotic partners in eukaryogenesis to be identified, as some have claimed (Lake 1988; Woese 2002), or has all signal been overwritten by the passage of time and subsequent gene transfers (GTs)? All of these questions are controversial, and obtaining a workable representation of prokaryotic evolution may help answer them.
2. FOUR ROOTED TREES OF LIFE

Four different rooted trees of life are illustrated in figure 1. The root, or cenancestor (Fitch & Upper 1987), of the traditional tree based on the pioneering work of Gogarten et al. (1989), Iwabe et al. (1989) and their collaborators, is placed between the archaebacteria and the bacteria, as shown in figure 1a. The non-tree, tree of life (Doolittle 1999), in figure 1b, illustrates the extensive lateral/horizontal GTs that may have erased most, if not all, of the phylogenetic signal from the early evolution of life on Earth, making it unlikely that the topology of the tree, and even less likely that the root, can be determined. A tree rooted within the double-membrane prokaryotes, based on transition analyses (Cavalier-Smith 2006), is illustrated in figure 1c. Figure 1d illustrates the indel-based rooted tree/ring of life recently developed in a series of papers from our laboratory (Lake et al. 2008).

As the variety of differing rooted trees suggests, determining roots is a complex problem that is sensitive to various sources of error. Here we discuss some of the difficulties encountered in rooting the tree of life and describe some recent progress made in this area.

Based on the proximity of the archaebacteria to the traditional root (figure 1a), it is thought that some archaeabacterial metabolisms and adaptations, such as methanogenesis and hyperthermophily, may be indicators of the energy sources used and the environments that were present at the time of the cenancestor. Thus there is uncertainty about whether the archaebacteria are phylogenetically ancient, or whether they simply evolved rapidly after diverging from eubacterial ancestors. Furthermore, the most deeply branching bacterial phylum, the Aquificae, is adjacent to the root, adding support for a hyperthermophilic cenancestor. Since a hyperthermophilic cenancestor is inconsistent with molecular stability arguments (Miller & Lazcano 1995) and with correlations between phylogenetic analyses and ribosomal RNA GC compositions (Galtier et al. 1999), this suggests that life might have started at mesophilic temperatures. These findings call the traditional root of the tree of life into question and make it important to examine other types of rooting data.

3. LONG BRANCH ATTRACTION

There is also a growing awareness that the traditional root might be an artefact of phylogenetic reconstruction resulting from long branch attraction (Felsenstein 1978; Lake 1991) and other sequence analysis artefacts (Philippe & Forterre 1999; Zhaxybayeva et al. 2005). Because the root is located at the bottom of the tree, the phylogenetic signal generated by this earliest bifurcation has had the longest time to decay. This makes the location of the root extremely sensitive to artefacts of phylogenetic reconstruction, arguably making the
root of the tree of life one of the most difficult phylogenetic signals to reconstruct. As the name ‘long branch attraction’ implies, this artefact can erroneously connect the longest branches in trees together. Thus when one uses paralogous gene sets to locate the root between the sets, long branch attraction can connect the fast-evolving, i.e. the longest-branched, taxa in each of the sets and thereby incorrectly root the tree (Lake 1991; Philippe & Forterre 1999).

One way to reduce the harmful effects of long branch attraction is to use phylogenetic characters that evolve more slowly than the nucleotide and amino acid sequences traditionally used. Potentially useful slower evolving characters include using the absences/presences of genes and the absences/presences of indels (insertions and deletions within genes). Indels are named indels because it is not always possible to determine whether they are inserts or deletions, and calling them indels acknowledges this inherent uncertainty. Since it is known that indels can persist over long time scales without changing length or position, as evidenced by the fact that numerous amino acid replacements occur within indels, they are potentially less affected by long branch attraction and other artefacts. Hence, in principle, they permit one to delve deeper into the rooted tree of life.

4. GENE TRANSFERS
Lateral/horizontal GTs represent a second confounding artefact. When genes are transferred between taxa, they misplace branches. There are three principal mechanisms that facilitate GT in prokaryotes, reviewed in Syvanen & Kado (1998). Genes can be transferred by means of transformation, conjugation and transduction. Transformation is the process whereby prokaryotes take up free DNA from their surroundings. Conjugation, also known as bacterial sex, occurs when an organism builds a tube-like structure known as the pilus, joins it to its ‘mate’ and transfers a plasmid through the tube. This process can transfer genes between very different taxa. Most astounding is the demonstration of Escherichia coli conjugating with the eukaryote Saccharomyces cerevisiae (Heinemann & Sprague 1989). Finally, transduction is a process for moving genes from one prokaryotic species to another via viruses (Hendrix et al. 1999).

GTs directly affect the calculation of the root by moving branches, thereby turning trees into tangles of webs, or nets, and making prokaryotic evolution less clonal and more population driven (Doolittle 1999; Gogarten et al. 1999; Ochman et al. 2000; Zhaxybayeva & Gogarten 2004; Lake et al. 2005) (also see the articles by Gogarten et al., Dagan & Martin, Beiko & Ragan and Sorensen et al. this volume). However, the effects of GT can be reduced by the careful choice of taxa used for analyses.

5. THE DIVERSITY OF LIFE ON EARTH
Because of the pervasiveness of GTs, one cannot realistically expect to reconstruct the root of a perfectly bifurcating, high-resolution tree. However, since GTs are less likely to occur between phylogenetically and functionally distant groups (Jain et al. 1999; Jain 2003), their effects may be reduced by choosing taxonomic groups that are phylogenetically well separated, thereby minimizing intergroup gene/indel transfers. These phylogenetically well-separated and relatively homogeneous prokaryotic taxa (Skophammer et al. 2006, 2007; Lake et al. 2007) are: the archaeabacteria, the Bacilli and relatives, the Clostridia and relatives, the Actinobacteria and the double-membrane, Gram-negative, prokaryotes (Cyanobacteria, Proteobacteria, Spirochaetes, Chlorobi, Chloroflexi and 17 additional phyla) (for more detailed definitions of these groups, including some recent taxonomic changes, see Ohno et al. (2000), Garrity & Holt (2001), Wu et al. (2005) and Lake et al. (2008)). Together these five prokaryotic super-taxa, plus the eukaryotes, include all-known life (Boone & Castenholz 2001).

The Actinobacteria, characterized by high GC genomic compositions, are morphologically diverse and include many human pathogens, including those that cause leprosy and tuberculosis. The archaeabacteria, a presumably paraphyletic taxon (Gouy & Li 1989; Archibald 2008; Cox et al. 2008), contain extreme halophiles, methanogens, hyperthermophiles and other unique phenotypes. The Clostridia and the Bacilli are firmicutes, characterized by their low GC compositions, although not exclusively (Ueda et al. 2004). The Clostridia are unique among the single-membrane prokaryotes for including photosynthetic as well as fermenting organisms. The double-membrane prokaryotes constitute an inordinately speciose, probably primitively photosynthetic, taxon. Some of their novelty is associated with the double-membrane system that surrounds them and encloses the periplasmic space.

6. A ROOTING EXAMPLE
The Actinobacteria are an extremely diverse prokaryotic taxon and have properties that suggested that this taxon might contain the root of life. They are among the most morphologically diverse prokaryotes and are widely distributed in both terrestrial and aquatic ecosystems (Embley & Stackebrandt 1994). Actinobacteria employ varied metabolic mechanisms, although no photosynthetic members are known. They are primarily chemoheterotrophs, which either respire or ferment. Their oxygen tolerances vary from strictly aerobic, to facultatively anaerobic, to microaerophilic, or strictly anaerobic. In addition to using some unique biochemical pathways not found in other prokaryotes, they also synthesize many macromolecules absent from other organisms, such as unique cell wall peptidoglycans (Gokhale et al. 2007). Given their diverse morphological and biochemical repertoires (Embley & Stackebrandt 1994; Boone & Castenholz 2001; Garrity & Holt 2001), properties that might indicate a deep placement in the tree of life, we were anxious to learn whether the root of life was contained within the Actinobacteria.

The GyrA protein contains an indel (Gao & Gupta 2005) that is ideal for determining whether the root is
Table 1. Summary of the GyrA/ParC indel. A summary of GyrA and ParC alignments within the NGSSG/GPDFPT region, corresponding to *Escherichia coli* residues 167–217 in the outgroup ParC sequence.

| GyrA       | paralogous ParC sequences shown at the bottom of the table, and is absent in GyrA protein sequences from all other prokaryotic taxa. In the top of the table, and is absent in GyrA protein sequences from all known GyrA proteins from Actinobacteria, shown at
| A-Actinobacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| D-Proteobacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| D-Cyanobacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| F-Firmicutes | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| R-Archaebacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |

| ParC       | sequences preceded by an A, D, F or R correspond to sequences from all known GyrA proteins from Actinobacteria, shown at the top of the table, and is absent in GyrA protein sequences from all other prokaryotic taxa. In the paralogous ParC sequences shown at the bottom of the table, the indel is absent in all sequences (Servin et al. 2008).
| A-Actinobacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| D-Proteobacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| D-Cyanobacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| F-Firmicutes | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |
| R-Archaebacteria | NGSSGIAVGMATNIPPHNLGELDIADLVIDNPAELE-----LSELME1IPGDFTP |

The process of indel rooting (Rivera & Lake 1992; Baldauf & Palmer 1993; Gupta 1998) is illustrated in figure 2 using the GyrA/ParC indels from table 1. The top two taxa in the trees, shown by shading at the left and the right of the figure, correspond to the actinobacterial taxa shown at the top of table 1. The sequences corresponding to the actinobacterial sequences, also shown by shading, are at the top of the centre panel and contain the four amino acid insert. The bottom two-thirds of the GyrA (orthologue 1) sequences and all of the ParC (orthologue 2) sequences lack the four amino acid insert and are not shaded. The tree on the left side of the figure is rooted outside the actinobacterial clade, as illustrated. This root requires only a single change (an insertion at the top of the tree) to produce the observed indel pattern. By contrast, the tree on the right side of the figure is rooted within the actinobacterial clade and requires two changes. Four amino acids must be inserted somewhere between the orthologue 1 and the orthologue 2 sequences, and this sequence must be subsequently deleted on the branch leading from the Actinobacteria to the remainder of the orthologue 1 tree. We conclude most parsimoniously that the root of the tree of life cannot be located within the shaded regions corresponding to the Actinobacteria. In this way, analyses of indels in orthologous/paralogous gene pairs can be used to exclude the root from regions of the tree.

In earlier studies (Rivera & Lake 1992; Gupta & Singh 1994; Philippe et al. 1999), it was assumed that incomplete gene sets could not be used for rooting, since ubiquitous gene sets were required for sequence-based rooting (Gogarten et al. 1989; Iwabe 1989; Brown & Doolittle 1995; Boucher et al. 2003; Zhaxybayeva et al. 2005). We now know that ubiquitous genes are not necessarily required for indel rooting since indel- and sequence-based rooting studies use different types of information. Although the ParC sequence is missing from the archaeabacteria, this complication was not considered in the simplified analysis in figure 2. However, when insertions and deletions of *genes* and insertions and deletions of *indels* were simultaneously analysed, it was shown that the GyrA/ParC data exclude the root within the Actinobacteria (Servin et al. 2008). In fact, one frequently finds indel sets that exclude some roots, even though genes are missing from some taxa.

Recently, solutions have been found for some technical problems that previously prevented the analyses of indel data. For example, in the past, it was generally thought that a single indel could not be statistically
significant. Now, however, methods have been developed for determining the statistical significance of indels by analysing the large amounts of sequence data that are available at the margins of indels. The statistical significance of an indel is a function of the length of the indel (longer indels are better), the proportion of exceptions to the indel pattern (fewer exceptions are better) and the evolutionary histories of the flanking sequences (Lake et al. 2008). Solutions have also been found for other problems related to indel rooting. For example, improved methods for identifying paralogous gene sets are now available (Skophammer et al. 2007), further increasing the number of useful indels.

Having workable solutions to some of the technical problems associated with indels rooting has greatly increased the number of indel sets available for rooting studies. In the following sections, we discuss the current state of indel-based rooting studies and some of the resulting biological implications.

7. A NEW ROOT

To date, indel-based rooting has excluded the root from the tree/graph of life, except for the site marked ‘root’ in the rooted tree and graph in figure 3. The prokaryotic tree of life is shown in figure 3a, and the graph of life, including the eukaryotes, is shown in figure 3b. The figure summarizes the results of eight indel analyses that exclude the root from the tree/ring of life consisting of the Actinobacteria, A; the double-membrane prokaryotes, D; the firmicutes, F; the archaeabacteria, R and the eukaryotes, K. The excluded regions are indicated by the surrounding balloons, and the names of the indel-containing proteins that exclude these roots are given within the balloons. All eight indels significantly, \( p > 0.95 \), exclude their respective clades.

In addition to providing a new root, indel-based rooting provides a non-tree-based view of evolution. For example, the tree of life becomes a graph of life

**Figure 2.** The process of indel rooting illustrated for two alternative rootings. In the centre of the figure, the two top sequences within orthologue 1 contain an insertion (shaded), whereas the bottom two-thirds of orthologue 1 and all of orthologue 2 lack the insertion. The trees on the left and the right sides of the figure represent two different rooted trees that relate orthologues 1 and 2. The tree on the right is rooted through the shaded region corresponding to those sequences that contain the insert, and the tree on the left is rooted outside the shaded region. The right tree is less parsimonious than the left tree, indicating that the root of the tree cannot lie within the shaded region.

**Figure 3.** (a) A summary of the new root of the tree of life and (b) for the ring of life. The relevant four taxa representing known prokaryotic diversity are the double-membrane prokaryotes (D), the firmicutes (F), the Actinobacteria (A) and the archaeabacteria (R). The eukaryotes (K) are present in the ring of life (b), and the Bacilli (B) and the Clostridia (C) form a paraphyletic grouping within the ring. The regions from which the root is excluded are circled and labelled with the name(s) of the relevant indel(s) that exclude(s) them. The dots present on the distal portions of the leaves represent the last common ancestors of each crown group. For reference, the root based on ground-breaking analyses of anciently duplicated gene paralogues (Gogarten et al. 1989; Iwabe 1989), marked by an ‘X’, is located between the archaeabacteria and (b) the Archaea contain extreme halophiles, methanogens, hyperthermophiles and other unique phenotypes, bacteria, and the root based on transition analysis (Cavalier-Smith 2006), marked by an ‘X’, is within the double-membrane prokaryotes.
when the eukaryotes are included because indel-based analyses provide data, directed quartets (Lake 2008), that allow one to distinguish between trees and graphs. To understand how this is possible, consider the phylogenetic relationships between eukaryotes and archaeobacteria and the eukaryotes and the double-membrane prokaryotes. Based on indels present in protein synthesis initiation factor IF2 and in protein synthesis elongation factor EFG, the eukaryotes and the archaeobacteria are sister taxa, as shown by the relevant excluded region in figure 3b. By contrast, based on the indel present in heat shock protein Hsp70, the eukaryotes and the double-membrane prokaryotes are sister taxa (figure 3b). Since trees do not allow a taxon to be simultaneously the sister taxon to two different taxa, i.e. the archaeobacteria and the double-membrane prokaryotes, these results can no longer be represented by a tree. In other words, it is impossible for a tree to join the eukaryotes to the archaeobacteria and simultaneously to the double-membrane prokaryotes. This relationship can only be represented by a ring, as shown in figure 3b. This ability of indel-based rootings to distinguish graphs from trees may become important in future studies designed to reconstruct a low-resolution graph of life.

As previously discussed, an important property of indel-based rooting is that indels can sometimes exclude roots, even though genes may be missing from some taxa. For example, the Hsp70/MreB sequences that exclude the root from the double membrane/eukaryotic clade shown in figure 3b were suitable for analysis, despite the possibility that some of these genes had been transferred from eubacteria to the eukaryotes and directly argue against the root based on transition analyses shown in figure 3b (Cavalier-Smith 2006). Again this fits well with other data and is consistent with the derivation of double-membrane prokaryotes from simpler single-membrane prokaryotes. The double-membrane arrangement is known to greatly complicate many processes that are much simpler in single-membrane prokaryotes. For example, complex design changes are needed to accommodate transport across the double-membrane arrangement. Specifically, special ABC transporters differing considerably from those present in single-membrane prokaryotes are required to facilitate the uptake of vitam in B12 in double-membrane prokaryotes ( Locher et al. 2002), and the process of flagellar assembly is considerably more complex in double-membrane prokaryotes than in single-membrane prokaryotes. In double-membrane prokaryotes, the process requires the construction of novel flagellar rings, the L and P ring assemblies, in addition to the M and S rings found in the cytoplasmic membrane of single- and double-membrane prokaryotes (Macnab 2003).

Parsimonious reasoning applied to this root also indicates that members of the cenancestral population were enclosed by ester-linked lipid membranes and surrounded by a peptidoglycan layer, because the Actinobacteria and the double-membrane prokaryotes on the left side of the root in figure 3b and the Clostridia and the Bacilli on the right side of the root in figure 3b, all share these character states. The location of the root also implies that during the transition from the root to the archaeobacteria, through the Clostridia and the Bacilli (figure 3b), the peptidoglycan layer was lost and the ester-linked membrane lipids were replaced with ether-linked lipid membranes. If this occurred,
then it is possible that remnants of this transition still exist in the lipid biosynthetic pathways found in the firmicutes today.

This possibility prompted us to re-examine the taxonomic distributions of genes coding for archaebacterial lipid pathways. With the appearance of the first archaebacterial and firmicute genomes, detailed phylogenetic comparisons of lipid biosynthetic pathways pointed towards some novel lipid-based archaebacterial–firmicute connections (Lange et al. 2000; Smit & Mushegian 2000; Boucher et al. 2003, 2004; Daiyasu et al. 2005). These simply did not make sense in light of the then current ideas concerning the uniqueness and the early evolution of the archaebacteria. However, those genomic findings make sense when viewed in the light of the new rooted tree of life. The findings reported in those prior genomic analyses suggest that parts of the mevalonate (MVA) synthesis pathway, central to archaebacterial lipid synthesis, may have been present in the common population immediately ancestral to the Bacilli and the archaebacteria. In addition, they imply that one of the enzymes thought to be responsible for the unique archaebacterial sn-1 stereochemistry was present in this ancestral population.

Many bacilli have complete, or nearly complete, MVA lipid synthesis pathways, similar to those found in the archaebacteria (Smit & Mushegian 2000; Boucher et al. 2004), whereas most eubacteria use the DXPS lipid synthesis pathway (Lange et al. 2000). In detailed phylogenetic analyses of the genes present in the prokaryotic MVA pathways (Boucher et al. 2004), few double-membrane prokaryotes contain MVA genes. Furthermore, the MVA genes that are present in double-membrane prokaryotes are non-parsimoniously distributed throughout the trees in no particular phylogenetic order, suggesting that they arose from multiple GTs. In contrast, the Bacilli form a single clade in most MVA gene trees (Boucher et al. 2004). The phylogenetic distribution of MVA biosynthetic genes within the Bacilli parsimoniously suggests that many genes in the pathway were vertically inherited from the last common ancestral population, relating the Bacilli and the archaebacteria.

Similar evidence suggests that at least one antecedent of the enzymes contributing the unique sn-1 stereochemistry existed in the ancestral population common to the Bacilli and the archaebacteria. Geranylgeranylglyceryl phosphate (GGGP) synthase, a terminal member of the MVA pathway, is one of the enzymes thought to be responsible for the unique sn-1 stereochemistry of the archaebacterial glycerol phosphate backbone (Boucher et al. 2004). GGGP synthase appears to have been vertically inherited from the ancestral population common to the Bacilli and the archaebacteria, since, except for a single Cytophaga species (Boucher et al. 2004), only the Bacilli and the archaebacteria contain genes for GGGP synthase. Furthermore, the Bacilli and the archaebacteria are not intermixed in the GGGP synthase tree, as would be expected if extensive GT from either group had been the source of GGGP synthase. Instead, they are resolved into their respective groups. Although the GGGP synthase genes cannot polarize the root, they are consistent with our indel evidence, excluding the root from the firmicute/archaebacterial clade. We parsimoniously infer that the ancestral population contained a nearly complete archaebacterial-like MVA pathway and one of the genes necessary for producing the unique archaebacterial lipid backbone stereochemistry. Thus this new root offers a simple explanation for a previously puzzling observation.

Almost no proposal regarding the evolution of life has provided more thought provoking discussion than the possibility of a hyperthermal origin of life. The new root discussed here does not support the proposition that the cenancenal population was hyperthermophilic, because the taxa adjacent to the root are not hyperthermophilic. The Actinobacteria, adjacent to the left side of the root, are primitively mesophilic, and the Clostridia on the right of the root are not hyperthermophilic (figure 3b). However, neither a moderately thermophilic nor a mesophilic cenancensator can be excluded for this root, since two recently discovered thermophilic Clostridia occupy a position near the root in concatenated protein sequence trees (Wu et al. 2005). Symbiobacterium thermophilum (Ohno et al. 2000) grows optimally at 60°C, and Carboxydothemus hydrogenoformans (Wu et al. 2005) grows optimally at 78°C. Thus this new root parsimoniously places constraints on the growth temperature of the cenancensator. These results, together with other evidence and arguments for mesophilic origins (Miller & Lazcano 1995; Gautier et al. 1999; Philippe & Forterre 1999), make lower temperature hydrothermal sites such as the Lost City field (Russell & Martin 2004; Kelly et al. 2005), increasingly attractive for the cenancenal evolution of life. We hope that this new root will form a basis for synthesizing the rapidly growing database of genomic- and Earth science-based information becoming available on the early evolution of life.

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REFERENCES


