The origin of the eukaryotic cell can be regarded as one of the hallmarks in the history of life on our planet. Yet, despite its evolutionary significance, the emergence of eukaryotes remains poorly understood (Embley and Martin 2006). At the cellular level, the gap between prokaryotes (Bacteria and Archaea) and Eukarya is immense, with the latter cell types being exceedingly compartmentalized with organelles and cell structures such as mitochondria, peroxisomes, Golgi complex, and endoplasmic reticulum, and the central nexus of the eukaryotic cell, the nucleus. Although cellular compartmentalization is also observed in several prokaryotic lineages (e.g., in Cyanobacteria, Planctomycetes, and Crenarchaeota), these compartments do not seem evolutionarily related to those observed in eukaryotes (e.g., see McInerney et al. 2011). The absence of such missing links, or intermediate stages of eukaryogenesis, significantly hampers the delineation of more sophisticated models for the emergence of the eukaryotic cell (Martijn and Ettema 2013).

The lack of understanding of how eukaryotes emerged echoes back in another central dilemma. How are the domains of cellular life related to each other? Ever since Carl Woese and George Fox (1977) discovered the “third domain of life” in the late 1970s, the Archaea, followed by their subsequent proposal for the classical three domains of life model (Fig. 1A) (Woese 1987; Woese et al. 1990), scientists have had heated debates on their evolutionary affiliation. In particular, they have failed to reach consensus regarding the placement of the eukaryotic branch in the tree of life. Early attempts
Figure 1. Overview of scenarios for the origin of the eukaryotic cell. Schematic depiction of the classical three-domain tree of life (A) and a tree that supports fusion hypotheses in which the eukaryotic nuclear lineage evolved from within the archaeal radiation (B). Of the latter category, a number of hypotheses have been proposed that can be classified as amitochondriate fusion scenarios (i.e., the fusion event leads to a mitochondrion-lacking proto-eukaryotic lineage). (Legend continues on following page.)
to resolve the placement of eukaryotes in the tree of life, including these of Carl Woese himself, placed eukaryotes as a sister clade with respect to the Archaea (Fig. 1A), whereas others proposed an alternative topology in which eukaryotes grouped within the Archaea (Fig. 1B), as a sister to the Crenarchaeota (“eocytes”) (Henderson et al. 1984; Lake et al. 1984).

The phylogenetic incongruence regarding the placement of eukaryotes in the tree of life has triggered the emergence of a wide variety of evolutionary models to explain the origin of eukaryotes, each of which having its own unique angle to it. Whereas some of these models are essentially compatible with the classical three domains view of life, such as the Archezoa hypothesis (Cavalier-Smith 1989), other models propose an autonomous origin of eukaryotes, such as the Neomuran hypothesis (Cavalier-Smith 2002a,b), which proposes that Archaea and Eukaryotes emerged from a common ancestor related to Gram-positive bacteria, and the PVC hypothesis, which suggests that eukaryotes evolved from an ancestor of the PVC superphylum (Devos and Reynaud 2010; Forterre and Gribaldo 2010; Forterre 2011; Reynaud and Devos 2011). Yet another class of hypotheses imply an important role for viruses in the prokaryote-to-eukaryote transition (Bell 2001; Takemura 2001; Forterre 2005, 2006), and some suggest that Bacteria, Archaea, and eukaryotes have evolved from a complex, eukaryote-like ancestor (Kurland et al. 2006). However, in-depth analyses of the genomic data that became available during the last decade of the previous century deemed many of these hypotheses implausible. Importantly, these analyses pointed out two other important issues. First, they confirmed that Archaea had highly distinct gene repertoires, compatible with them evolving independently from the bacterial domain (Bult et al. 1996; Brown and Doolittle 1997; Olsen and Woese 1997; Rivera et al. 1998; Makarova et al. 1999; Ettema et al. 2005), and second, they pointed out that eukaryotic genomes are chimeric in nature, comprising two distinct gene sets (Rivera et al. 1998); genes for information storage and processing are Archaea related (see, for example, Yutin et al. 2008), and genes for metabolic or “operational” processes are mostly bacterial in nature. These observations suggest that at the root of the eukaryotic domain of life, some sort of fusion event has taken place that involved an archaeal partner, and one or more bacterial partners (Koonin 2010).

The idea of a cellular fusion event at the basis of the eukaryotic lineage is not new. During the late 1980s, Zillig and colleagues interpreted the myriad of characters shared between Archaea, Bacteria, and Eukarya as evidence of chimerism.
in the eukaryotic domain of life (Zillig et al. 1985, 1989a,b). The fusion hypothesis postulated by Zillig and coworkers fitted in nicely with the Serial Endosymbiosis Theory (repopularized by Lynn Margulis), which placed fusion events as a central avenue in cellular evolution, in particular, that of eukaryotes (Sagan 1967). Altogether, both concepts stood at the basis of a new wave of hypotheses for the origin of the eukaryotic cell in which fusion events played a central role. Although the identity, nature, and amount of fusion partners differed markedly between different hypotheses, we can roughly enumerate two classes based on whether the end product of the fusion represents an amitochondriate (mitochondrion-lacking) (Fig. 1C) or a mitochondriate (mitochondrion-bearing) (Fig. 1D) cellular entity. The former “amitochondriate category” is compliant with the Archezoa theory (Cavalier-Smith 1989), and other “Woesean” models, in the sense that they implement the origin of the eukaryotic cell and mitochondrion as two independent evolutionary events. Several studies have suggested, however, that these two events most likely co-occurred, based on several lines of evidence. For example, we now know that previously assumed amitochondriate eukaryotes do harbor remnants of the mitochondrion past in the form of hydrogenosomes, mitosomes, and possibly other mitochondria-like organelles. In addition, phylogenetic analyses have argued against the existence of a deeply rooting Archezoan clade (see Embley 2006 for a detailed argument).

Significant heterogeneity exists among the fusion models with respect to the envisioned archaeal fusion partner (Fig. 1C,D). Most of these models, especially those that were delineated before the genomic age, are based on biological (metabolic or cytological) considerations. For example, several models implement a syntrophic interaction between a methanogenic archaeon and hydrogen-producing bacterial partner (e.g., the hydrogen and other syntrophic hypotheses; see Fig. 1C,D). Yet another group of models envision that the nucleus emerged from an archaeal endosymbiont (so-called endokaryotic models) (Martin 2005), such as the eocyte (Lake 1988) and pyrococcus-gammaproteobacteria fusion models (Fig. 1C) (Horiike et al. 2004).

Obviously, obtaining information regarding the identity and nature of the archaeal fusion partner holds the key to solving the evolutionary puzzle of the origin of the eukaryotic cell. Yet, the plethora of phylogenetic studies that has tried to resolve this conundrum has thus far failed to reach consensus, prompting some to even speak of a “phylogenomic impasse” (Gribaldo et al. 2010). Clearly, the phylogenomic approaches taken to resolve these deep evolutionary relationships have to be performed with great care as they are prone to all sorts of biases and artifacts (Delsuc et al. 2005; Gribaldo et al. 2010). Some of those, such as compositional bias and varying evolutionary rates (heterotachy), are inherent in the nature of biological sequence data. Yet, apart from biological issues, the outcome of phylogenomic studies are strongly influenced by data selection (genes and taxa), site filtering, choice of sequence alignment and phylogenetic algorithms, evolutionary model, etc. (see Delsuc et al. 2005 for an overview). Delsuc et al. (2005) hit the nail on the head by summarizing these issues wittily: “garbage in, garbage out.” Therefore, the outcome of the earliest genome-scale phylogenetic studies, in which such issues were not yet properly addressed, should be regarded in light of such artifacts. With this in mind, it is interesting to note that the first phylogenomic studies that aimed to depict the tree of life generally supported the iconic “Woesean” three-domains tree (e.g., Snel et al. 1999; Tekaia et al. 1999; Ciccarelli et al. 2006). However, recent studies that analyzed more extensive genomic data sets with more sophisticated phylogenetic models and algorithms embedded the eukaryotic branch within the archaeal domain, creating a paraphyletic archaeal domain. More specifically, they seem to support a scenario in which eukaryotes emerged from the “TACK superphylum” (Cox et al. 2008; Foster et al. 2009; Guy and Ettema 2011; Williams et al. 2012), which, apart from Cren-, Kor-, and Thaumarchaeota, also comprises the recently proposed Aigarchaeota phylum (Nunoura et al. 2011). These latter findings thus lend support to an extended version of the eocyte hypothesis.
(Lake 1988), or the recently proposed phagocytosing archaeon theory (PhAT), which poses that eukaryotes evolved from a phagocytic TACK archaeon (Martijn and Ettema 2013). It should, however, be noted that whereas the eocyte hypothesis represents a so-called endokaryotic fusion model (Lake 1988; Martin 2005) with an amitochondriate, nucleated eukaryote as result, the PhAT hypothesis, as an extension of previous scenarios (e.g., Yutin et al. 2009), proposes a cellular fusion that results in a mitochondrion-containing but nucleus-lacking cell (Fig. 1). Apart from phylogenetic studies, the evolutionary affiliation between eukaryotes and the TACK superphylum is seemingly supported by the recent identification of several previously presumed eukaryotic signature proteins (ESPs) in genomes of TACK Archaea (Guy and Ettema 2011). These ESPs are generally involved in pivotal processes in eukaryotes and, among others, include bona fide archaeal orthologs of actin (Yutin et al. 2009; Bernander et al. 2011; Ettema et al. 2011), tubulin (Yutin and Koonin 2012), ESCRT proteins (Lindás et al. 2008; Ettema and Bernander 2009; Makarova et al. 2010), as well as several proteins that are involved in transcription and translation (Guy and Ettema 2011).

Here, we present a reassessment of the phylogenetic affiliation between the archaeal domain and eukaryotes by performing phylogenomic analyses in which we have investigated the impact of gene and taxon selection and compositional bias removal while using a variety of phylogenetic algorithms and evolutionary models. Our analyses indicate that a broadened taxon sampling and increased gene selection unequivocally support a scenario in which the eukaryotic branch emerged from within or at the base of the TACK superphylum.

RESULTS AND DISCUSSION

An Unbiased Phylogenomics Approach to Study the Evolutionary Affiliation between Archaea and Eukaryotes

A dual strategy was designed to investigate the evolutionary affiliation between Archaea and eukaryotes. One analysis included a phylogenomic analysis of highly conserved, universal protein coding genes, and a second analysis focused on the phylogenetic analysis of 16S/18S and 23S/28S ribosomal DNA (rDNA) genes (Fig. 2). Because it was anticipated that the eukaryotic lineage might branch within the archaeal domain, care was taken to include an unbiased sample of all main archaeal groups in all data sets that were analyzed. The latter was achieved by clustering concatenated protein sequences comprising 55 panorthologs present in 120 archaeal species using data extracted from the arCOG database (Fig. 2; see supplemental Fig. 1 online) (Wolf et al. 2012). Using a clustering cutoff of 70% sequence identity, a set of 58 taxa was obtained that represented all major archaeal groups. This set was completed by 10 bacterial and 10 eukaryotic genomes, as well as eight recently published archaeal genomes, including the genome of the Geoarchaeote NAG1, which was proposed to represent a novel archaeal phylum (Kozubal et al. 2013) (see supplemental Table 1 online for a detailed list). Assignment of proteins encoded by each of these genomes to the arCOG clusters was achieved by performing PSI-BLAST (position-specific iterative basic local alignment search tool) analysis followed by phylogenetic evaluation of each updated arCOG to remove paralogous sequences, as well as genes that were obviously horizontally transferred (see the Material and Methods section for details). Eventually, a data set was obtained that comprised 67 protein clusters (see supplemental Table 2 online), each of which contained at least 52 archaeal sequences, as well as at least three bacterial and three eukaryotic protein sequences. This data set represented the starting point of a number of analyses that aimed to gain insight into the phylogenetic position of eukaryotes in the tree of life, focusing on effects of gene selection, taxon selection, and compositional bias. In parallel, 16S/18S and 23S/28S rDNA gene sequences were obtained from the same taxa present in the protein data set, followed by addition of 16S and 23S sequences from further archaeal taxa representing deeply diverging lineages. These sequences were also used to infer the phylogenetic position of eukaryotes in the rDNA-based tree of life and
Figure 2. Flowcharts of the data selection processes. Numbers in blue outside boxes represent the number of sequences or organisms included. Numbers in red represent the number of genes or clusters. Abbreviations in green show the phylogeny algorithms applied to the data set: ML, maximum-likelihood (RAxML); BP, BLG, and BGTR, Bayesian under CAT-Poisson, CAT-LG, and CAT-GTR model, respectively (Phylobayes). (A) Protein concatenated data sets. (B) Ribosomal RNA genes phylogeny. (C) Archaeal diversity tree.
investigate any potential taxon sampling effects on this placement (see Fig. 2).

**Robust Support for the Archaeal TACK Superphylum**

Recent studies have reported that the eukaryotic lineage emerged from within the so-called TACK superphylum, comprising the Thaum-, Aig-, Cren-, and Korarchaeota (Guy and Ettema 2011; Williams et al. 2012). To reevaluate these findings, we decided to first investigate whether the existence of this proposed superphylum is supported by means of phylogenetic analysis of the concatenated protein and rDNA data sets. Bayesian and maximum-likelihood (ML) analyses were performed on the data set of 67 conserved proteins clusters (see above and Fig. 2), but leaving out eukaryotic sequences, hence, comprising sequences up to 66 archaeal and 10 bacterial taxa. Likewise, Bayesian and ML analyses were performed on concatenated data sets of archaeal and bacterial 16S and 23S rDNA sequences. Bayesian and ML analyses of the concatenated protein data set, as well as the concatenated rDNA data set, provided very strong support of the existence of the TACK superphylum with BS = 100 and PP = 1.00 for both data sets (Table 1; see supplemental Fig. 2 online).

The strong support obtained here agrees with previous phylogenetic analysis of 26 conserved protein-coding genes (Guy and Ettema 2011) and universally conserved ribosomal protein sequences (Wolf et al. 2012; Yutin et al. 2012). Moreover, the branching order within TACK observed in the present analysis corresponds with the branching order observed in both studies, with Korarchaeota representing the deepest branch, and Aig- and Thaumararchaeota representing sister-phyla, subseeded by Crenarchaeota—(((Thaum,Aig)Cren)Kor). It should be noted that the analysis by Yutin et al. (2012) did not include sequence data from Aigarchaeota.

**Phylogenomic Analysis of Concatenated Protein and rDNA Data Sets Supports the Affiliation between TACK and Eukaryotes**

Having established strong phylogenetic support for the existence of a TACK superphylum in the archaeal domain, we sought to address the phylogenetic relationship between this group of Ar-
We performed Bayesian and ML analyses on the same data set of 67 conserved proteins, as well as on the concatenated data set of 16S and 23S rDNA sequences, but now including protein, 18S, and 28S data from up to 10 eukaryotic taxa. The results of the protein data set analyses provide strong support for a phylogenetic affiliation between the TACK superphylum and eukaryotes, with bootstrap support (BS) of 85 and posterior probability (PP) of 1.00 (Table 1; see supplemental Fig. 3C,D online). The analysis of the concatenated 16S/18S and 23S/28S rDNA data sets also supported the TACK-eukaryotes affiliation, albeit with slightly lower support values in the ML analysis (BS = 79, PP = 1.00) (Table 1; see supplemental Fig. 4A online; Fig. 4B). The same analyses performed on a larger data set (162 clusters) (see supplemental Table 2 online) in which more missing data was tolerated (presence of at least 90% of archaean sequences and one eukaryotic sequence) yielded similar results, with the TACK superphylum supported with BS = 74 and PP = 1.00 (Table 1; see supplemental Fig. 3A,B online). Altogether, these results corroborate those of a recent study performed by Williams et al. (2012), which rejected the three-domain tree of life by providing strong support of a phylogenetic affiliation of TACK and eukaryotes in both concatenated protein and rDNA gene data sets.

The Phylogenetic Affiliation between TACK and Eukaryotes Is Improved by Removal of Discordant Protein Clusters

To mitigate any conflicting effect of horizontal gene transfers or fast-evolving eukaryotic sequences on the inferred TACK-eukaryotes relationship, protein clusters displaying the most divergent phylogenetic signals were removed using a discordance filtering procedure similar to the one used by Williams et al. (2010). After ranking all 67 protein clusters based on their discordance score (Fig. 3A), different fractions of the most discordant clusters were gradually removed and the remaining data sets were analyzed using an ML analysis. Removal of the most discordant clusters resulted in an increased support for the TACK-eukaryotes affiliation (red circles in Fig. 3B), and any remaining support for the three-domain tree topology (green crosses in Fig. 3B) was completely diminished after removal of 15% of most discordant clusters. These results indicate that the discordant clusters mainly were supporting the three-domain tree topology, which might point at the presence of fast-evolving eukaryotic sequences and traces of chimerism in composite microbial genomes that resided in the protein clusters (see supplemental Fig. 5 online).

Removing the 15% most discordant protein clusters further substantiated the support for the eukaryotic clade to nest within the TACK superphylum, in addition to the ML analysis (BS = 100), Bayesian phylogenies run under three different models (CAT-Poisson, CAT-GTR, and CAT-LG) unanimously confirm it with posterior probabilities of 1.00. Interestingly, removing discordant protein clusters also increasingly associated eukaryotes with Korarchaeota (blue triangles, Fig. 3B). Stepwise removal of the most discordant clusters caused the BS for this affiliation to increase from 85 (no removal) to 100 (15% removal), and robust support was observed even when removing up to half of all data (BS = 94) (see Fig. 3B and supplemental Fig. 6 online). Given that this affiliation has also previously been observed by Williams et al. (2012, see their Fig. 2A), we decided to analyze the data set in which 15% of the most discordant clusters were removed using ML and Bayesian methods under the CAT-Poisson, CAT-LG, and CAT-GTR models. Intriguingly, only CAT-LG failed to retrieve the eukaryotes-Korarchaeota affiliation with significant support (PP = 0.50) (see supplemental Fig. 3H online). The ML analysis very strongly supported (BS = 100) (see supplemental Fig. 3E online) the eukaryotes-Korarchaeota affiliation, and the Bayesian analyses of this data set with the CAT-Poisson and CAT-GTR models, which are regarded to be significantly more robust against long-branch attraction artifacts compared to all other models (Lartillot et al. 2009), confidently retrieved this grouping as well (PP of 0.99 and 1.00 for CAT-Poisson and CAT-GTR, respectively) (see Fig. 4A, Table 1,
and supplemental Fig. 3F,G online). The eukaryotes-Korarchaeota clade is also supported in the ML analyses of both larger data sets (BS = 74 and 85 for the 162 and 67 protein data sets, respectively) (see supplemental Fig. 3A,C online), but not in the Bayesian analyses performed on the same set in which eukaryotes are a sister clade to the TACK superphylum (PP = 1.00 and 0.92 for the 162 and 67 protein data sets, respectively) (see supplemental Fig.
Figure 4. Bayesian trees obtained from concatenated amino acid sequence alignments (A) and the concatenated sequences of the small and large ribosomal subunits (SSU and LSU, respectively) (B). In A, the tree was obtained from the alignment of the 85% least-divergent proteins (data set discFilter15p), running four chains of PhyloBayes under a CAT-Poisson model. In B, the sequences for both ribosomal subunits were concatenated and a tree was obtained with MrBayes. The tree was rooted with Archaea only. The eukaryotes-Korarchaeota affiliation will remain stable with the inclusion of more genomic data from Korarchaeota and related archaean species whenever this becomes available.
The Phylogenetic Affiliation between TACK and Eukaryotes Is Not Caused by Compositional Bias

Compositional bias in biological sequence data has been shown to be a major source of incorrect inference of phylogenetic relationships (Delsuc et al. 2005). To assess whether such bias is also affecting the abovementioned TACK-eukaryotes relation and nested affiliation of eukaryotes with Korarchaeota within TACK, we performed an analysis in which we gradually removed sites that were potentially compositionally biased. To do so, a χ² filter was used that determines the relative contribution to the global amino acid composition heterogeneity for each site in a given alignment (Viklund et al. 2012). ML analyses of protein data sets in which those sites that contributed most to the composition heterogeneity were removed in a stepwise manner revealed that both the TACK-eukaryotes affiliation, as well as the nested affiliation of eukaryotes with Korarchaeota, remained strongly supported, even when removing sites more than half a standard deviation away from the average amino acid heterogeneity, that is, in total 28.4% of the most heterogeneous sites of the alignment (Fig. 3C; see supplemental Fig. 7 online). These findings indicate that the phylogenetic signal underlying the observed TACK-eukaryotes affiliation is not caused by compositional bias.

Taxon Sampling Effects Do Not Affect the Affiliation of the TACK Superphylum and Eukaryotes

In addition to the availability of informative characters in phylogenetic analyses, empirical studies underline the impact that taxon sampling can exert on the reliability of phylogenetic inference (Townsend and Lopez-Giraldez 2010). Archaeal taxa are generally sampled sparsely with respect to amount, diversity, and available genomic sequences and, moreover, several deeply rooting phyla comprise of only a single representative (e.g., Korarchaeota, Aigarchaeota, and Nanoarchaeota). Therefore, taxon-sampling artifacts might have a significant impact on the phylogenomic reconstruction of the archaeal species tree itself and placement of the eukaryotic root, in particular. To assess such effects on the placement of eukaryotes in the archaeal tree in concatenated protein and rDNA data sets, we decided to perform phylogenetic analysis on data sets from which predefined (combinations of) archaeal clades were omitted (Table 2).

The analyses of the concatenated protein data sets using ML methods indicated that removal of individual archaeal TACK phyla, as well as removal of any permutations thereof, had only very little affect on the TACK-eukaryotes affiliation. Even if the data set included only a single TACK phylum, all bootstrap values were above 70 (Table 2; see supplemental Fig. 8 online). Interestingly, in the case in which only a single phylum was removed, only removal of Korarchaeota (represented by only a single taxon) decreased support for the TACK-eukaryotes affiliation (BS = 92 vs. BS = 100 for separate removal of Thaum-, Aig-, or Crenarchaeota). The latter observation could indicate that the affiliation between Korarchaeota and eukaryotes that was observed before (see above) is genuine rather than the result of an artifact.

Support for TACK superphylum remains strong in phylogenetic analyses of concatenated rDNA sequences with a combination of excluded taxa within the TACK-eukaryotes affiliation (Table 2; see supplemental Fig. 9 online). In trees in which a single TACK phylum was removed, BS was equal or higher than 70 except for those from which Thaumarchaeota was removed. In trees where two different clades were removed (CK, AK, AC, TC, TK, or TA; for abbreviations, see Table 2), TACK + E support was above 70 in three of them (AC, TC, and TK), and in the rest of the combinations, the placement of the eukaryotic root could not be confidently established (Table 2).

In light of the results discussed above and those presented in Table 2, the effect of taxon sampling in terms of removal of any combination of TACK phyla on the phylogenetic affiliation between TACK and eukaryotes seems to be minor, and none of the analyses of concatenated protein and rDNA data sets retrieved significant support for the three-domains tree of life scenario.

The Archaeal Legacy of Eukaryotes

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CONCLUDING REMARKS

The origin of the eukaryotic cell represents an enigmatic evolutionary puzzle—a puzzle that is currently lacking too many pieces to fully appreciate the process of eukaryogenesis. Phylogenomic and comparative genomic analyses of genomic sequence data that has recently become available have already started to unveil key pieces of this puzzle. Several independent phylogenomic studies (Cox et al. 2008; Foster et al. 2009; Guy and Ettema 2011; Kelly et al. 2011; Williams et al. 2012) have now reported results that refute the three-domains tree of life, supporting scenarios in which the eukaryotic lineage evolved from a fusion event that involved a “garden variety” archaeon (Koonin 2010) as a source for the eukaryotic nuclear lineage.

In the present study, we have reevaluated the phylogenetic affiliation between Archaea and eukaryotes, focusing on effects of marker (gene) selection, taxon sampling, and compositional bias removal. The results in our study unequivocally support a phylogenetic affiliation between eukaryotes and the archaeal TACK superphylum, underscoring the conclusions drawn in recent studies (Guy and Ettema 2011; Williams et al. 2012), and providing further support for fusion-like scenarios at the root of the eukaryotic lineage. Moreover, the apparent affiliation with the TACK superphylum provides evidence against fusion scenarios that involve archaeal members of the Euryarchaeota, such as the hydrogen and syntrophic hypotheses, which involve a methanogenic fusion partner, and the sulfur syntrophy hypothesis and serial endosymbiotic theory, which envision a Thermoplasma-like fusion partner (Fig. 1C,D).

Yet, our data is compatible with an extended version of the eocyte theory (Williams et al. 2012) or the recently proposed PhAT (Martijn and Ettema 2013), which both propose TACK-affiliated archaeal fusion partners. It should be

Table 2. Effect of taxon removal on the position and phylogeny of the TACK superphylum and eukaryotes

<table>
<thead>
<tr>
<th>Removed taxa</th>
<th>Remaining taxa</th>
<th>TACK + E support</th>
<th>Topology inside TACK</th>
<th>See supplemental figures online</th>
<th>SSU + LSU</th>
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All phylogenies were inferred with RAxML under PROTCATLG, running 100 bootstraps. T, Thaumarchaeota; A, Aigarchaeota; C, Crenarchaeota; K, Korarchaeota; E, eukaryotes; B, bacteria; NA, not applicable; SSU, small ribosomal subunits; LSU, large ribosomal subunits.

aTopology inside the TACK + E superphylum using a pseudo-Newick format.

bThese two values represent support for the TACK superphylum, without eukaryotes.
noted that, whereas the eocyte hypothesis represents a so-called endokaryotic fusion model (Lake 1988; Martin 2005) with an amitochondriate, nucleated eukaryote as result (Fig. 1C), the PhA T hypothesis proposes a cellular fusion that results in a mitochondrion-containing but nucleus-lacking cell (Fig. 1D).

In several of our analyses we retrieved support for a eukaryotes-Korarchaeota sister relationship (also observed in Williams et al. 2012). Although we cannot rule out that this affiliation is artificial, for example, the result of poor taxon sampling (Korarchaeota are represented only by a single, deeply rooting taxon), it could also indicate that the eukaryotic nuclear lineage evolved from an archaean lineage that is distantly related to Korarchaeota and which remains to be identified. Possibly, this lineage could belong to the genomically unexplored lineages that are part of the TACK superphylum (Fig. 5), such as archaeal clades with enigmatic names as Deep Sea Archaea Group, Marine Hydrothermal Vent Group, and Ancient Archaea Group (Fig. 5). The genomic exploration of these archaean lineages, as well as those that thus far have remained undetected in the microbial biosphere, represents a unique opportunity to identify some key pieces of the puzzle of eukaryotic origins, which will hopefully allow us to gain a more profound insight into the evolutionary transition from prokaryotic to eukaryotic life on our planet.

**MATERIALS AND METHODS**

A detailed description of the materials and methods is available in the supplemental material online.
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The Archaeal Legacy of Eukaryotes: A Phylogenomic Perspective

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