

# Mitochondria, bioenergetics and the origins of eukaryotic complexity

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## Abstract

Eukaryogenesis is thought to be one of the major events in the history of life on Earth, as the appearance of eukaryotes is often considered a prerequisite for the eventual evolution of complex multicellularity. Accordingly, figuring out how exactly it happened has generated considerable interest over the decades, as has the more general question of what factors enabled/drove complexification once eukaryotes had evolved. Phylogenomic studies over the last few decades have helped tremendously with the clarification of the phylogenetic relationships of eukaryotes, which are now understood to be the product of an endosymbiosis between an archaeal host and an  $\alpha$ -proteobacterial symbiont. However, debate continues regarding the cellular complexity of the host, the role the protomitochondrion played in the process and the factors that steered eukaryote evolution in the direction of further increases in complexity. The hypothesis that the mitochondrion provided an enormous boost in available “power per gene” relative to prokaryotes and this allowed cellular and genomic complexification to proceed has been advanced over the last few years, has become quite popular, and has generated significant controversy. Here, we examine this hypothesis in the light of our current understanding of the *process* of eukaryogenesis, discuss why it is not necessary to explain the origin of complex eukaryotic features, and consider alternative explanations for their appearance.

Cellular life on Earth is divided in two fundamentally different categories in terms of organization – prokaryotes and eukaryotes. While a few examples of simple multicellularity can be found in prokaryotes [2–4, 81], multicellularity has evolved on a large number of occasions in eukaryotes [5, 6], and complex multicellular organisms can only be found among them, including metazoans and human beings, which are usually perceived, deservedly or not, as the pinnacle of life’s complexity. For these reasons, the appearance of an eukaryote-level cellular organization is typically seen as a prerequisite for the evolution of “truly complex” and eventually sentient organisms, even if it is not possible to strictly prove the validity of such a claim. Eukaryogenesis is therefore often viewed as one of the key evolutionary transitions in the history of life on our planet, and has as a result attracted an immense amount of attention from theoretical and organismal biologists over the decades.

A robust scenario for eukaryogenesis needs to explain both the mechanistic details of how the process unfolded and the evolutionary forces that drove and shaped it. On a broader level, there is also the question of why it was eukaryotes to which the path towards greater and greater complexification was open, in contrast to the case of prokaryotes,

which never progressed beyond very simple multicellularity. That is a question that may turn out to be in principle unanswerable, as it involves speculations regarding what did not happen in prokaryotes and why that might have been, but it is also one that will nevertheless persist in the minds of researchers interested in the subject. even more so given that examples of many of the hallmark features of the eukaryote cell can in fact be found in some form in at least some prokaryotes. Yet currently we have evidence for only one transition to an eukaryote-level of genomic and organismal complexity in the whole history of life on Earth.

Eukaryotes are distinguished from prokaryotes by numerous traits, among them the enclosure of genetic material in a double membraned nuclear compartment and its organization into multiple linear chromosomes consisting of histone-based nucleosomal chromatin, the existence of an intracellular membrane network, with sophisticated mechanisms for internal membrane transport between compartments, the ancestral presence of mitochondria (sometimes reduced to mitochondria-related organelles, or MROs, and lost completely on only one known occasion [7]), the presence of a complex cytoskeleton, mitotic cell division, generally larger cellular size, and an overall higher tendency

to contain/tolerate endosymbionts. However, examples of many of these features can be found in prokaryotes too [8], such as cells with internal membrane-bound compartments, in some cases even enclosing the nucleoid [9–13], giant size [14–17], multiple chromosomes/replicons [18], linear chromosomes [19], presence of endosymbionts [20], and others. Yet these are generally isolated examples, and the full complement of eukaryotic features is indeed unique to eukaryotes.

## The chimeric origin of eukaryotes

Given the long history of heated debates surrounding eukaryogenesis and its very distant from us timing (ensuring that little solid evidence is available to constrain theories), it is no surprise that there is an enormous variety of hypotheses regarding how it happened. Our goal is not to comprehensively review all of them, but to specifically examine the role of the mitochondrion in the process. We will therefore primarily focus on the consensus view of how eukaryotes evolved that has been emerging in recent years [21–31], together with some of the prior hypotheses most relevant to the discussion.

It is now almost universally accepted that the mitochondrion originated as a result of an endosymbiotic event involving a member of the  $\alpha$ -proteobacterial clade. The first endosymbiotic proposals were advanced already back in the early 20th century by Mereschkowski regarding the origin of chloroplasts [32, 33]; that the mitochondrion also originated that way became a popular view in the 1960s thanks to the endosymbiotic theory advanced by Lynn Margulis [34]. The relationship between  $\alpha$ -proteobacteria and mitochondria was later confirmed by early sequencing studies [35–37], and has been robustly reproduced in subsequent phylogenomic studies [38–40], although the exact positioning of mitochondria relative to the extant  $\alpha$ -proteobacterial diversity is still a subject of debate [41–44].

The next major step in piecing together the puzzle of eukaryote origins came from the discovery by Carl Woese, during pioneering rRNA sequencing efforts in the 1970s, that archaea constitute an independent domain of cellular life [45]. Based on early phylogenetic analyses, a three-domain system was proposed, in which bacteria, archaea, and eukaryotes form distinct domains, with eukaryotes being more closely related to archaea than to bacteria [45, 46]. A relationship between archaea and eukaryotes was also supported early on by observations of commonalities between the information processing machineries in the two groups, such as RNA polymerases and other components of the transcriptional and translation apparatuses [47, 48], the presence of nucleosomal histone proteins in archaea but not in bacteria [49–52], and others.

Already in the 1980s, the eocyte hypothesis was proposed as an alternative of the three-domain system, initially based on phylogenetic analysis of ribosomal structure and later supported by studies of individual highly conserved

genes [53–57]. It has eukaryotes emerging from within archaea rather than being a sister lineage that diverged early from a common ancestor of both.

The eocyte view was bolstered by early phylogenomics studies, once whole-genome sequences for a sufficiently large sampling of microbial diversity became available [22–24, 26, 27, 29–31], although it was for a long time difficult to establish the exact phylogenetic relationship between the various diverse archaeal lineages and eukaryotes. Increasingly many aspects of cellular biology thought to be specific to eukaryotes were over the years also found to be present in at least some archaeal lineages [58–60], suggesting a rather complex eukaryote ancestor. However, these genes are overall dispersed around the archaeal tree and not all found in a single lineage [61, 62].

A major breakthrough was made in the last few years through the discovery of first, the Lokiarchaeota archaeal lineage [63], and then the broader Asgard archaea grouping [64–66] (combining Lokiarchaeota and the related Thor-, Odin- and Heimdallarchaeota clades). There are some caveats to be mentioned when discussing Asgard archaea, first, that the interpretation of a direct relationship between them and eukaryotes has been challenged [67–69], and, second, and more important, that they have not yet been isolated, cultured and observed directly, but are instead only known from metagenomic sequence assemblies meaning that we cannot be absolutely certain about their cellular organization. Nevertheless, these archaeal lineages do appear to be the most eukaryote-like archaea based on the gene content of their genomes, with the presence of numerous genes homologous to components of the eukaryote membrane dynamics and trafficking machinery being of particular interest.

The emerging consensus view therefore is that eukaryotes arose from within the archaea. Still open, however are (among many others) the following questions:

1. When exactly in the process of eukaryogenesis did the mitochondrial endosymbiosis become established?
2. What was the level of cellular organization of the host?
3. What evolutionary forces drove the evolution of the hallmarks of the eukaryotic cell, in particular its distinctly complex relative to prokaryotes features
4. What influence did the mitochondrion have on this process?

As the mitochondrion itself is one of these hallmark complex features of eukaryotes, there are three broad classes of possibilities for the relative timing of its acquisition in the context of eukaryogenesis (Figure 1):

1. The host was already a fully fledged but amitochondriate eukaryotic cell that then acquired an  $\alpha$ -proteobacterial endosymbiont, which in turn evolved into the modern mitochondrion

2. The host was a complex archaeon (more complex than any known so far in some detail extant archaeal lineage)
3. The endosymbiotic event coincided with and perhaps triggered the process of eukaryogenesis.

The first possibility has been traditionally referred to as the “Archezoan” scenario, and enjoyed widespread popularity in the 1980s and 1990s [71, 72] based on the observation that numerous eukaryotes lack mitochondria and that they tended to group together as early diverging branches in early phylogenies. However, such topologies eventually turned out to be long-branch attraction artifacts while upon detailed examination, each of the examples of amitochondriate eukaryotes turned out to be a case of secondary reduction of mitochondria into MROs, such as hydrogenosomes and mitosomes [24, 73–79]. Thus no primitively amitochondriate eukaryotes are at present known (of note, recently an example of a completely amitochondriate eukaryote, *Monoecocomonoides* sp., was reported [7], but it is still clearly a product of secondary loss as it belongs to the oxymonads and not to a new early branching eukaryote lineage), the Archezoan scenario has been largely rejected, and it is inferred that the Last Common Eukaryote Ancestor (LECA) had fully functioning mitochondria.

Deciding between the remaining two classes of hypotheses is considerably more challenging, at least in part because the concept of a “complex archaeon” is not strictly defined. There have been proposals that the archaeal host was already capable of phagocytosis (“phagocytosing archaeon theory” [80], and phylogenomic analyses have been published suggesting a late acquisition of the mitochondrion [82]. However, the latter have been seriously challenged on technical grounds [83, 84], and the former hypothesis seems unlikely to be true. The gene content of Asgard archaea genomes and recent biochemical results showing that they possess functional profilins capable of regulating the polymerization of actin filaments [85] have provided support for eukaryote-like cytoskeleton and membrane dynamics being present in these organisms. However, bona fide phagocytosis might have in fact evolved independently in eukaryotes on multiple occasions (based on the observation that very few of the relevant components are conserved across all eukaryotes, unlike what is observed for most ancestral to all eukaryotes features [86]), and it is likely that for phagocytosis to evolve, it is in fact a requirement that a functioning mitochondrion is present in the cell (see further discussion below). Such a view is supported by recently published analyses of the Asgard archaea genomes from the perspective of the phagocytosis machinery as found in various eukaryotes, which predict that Asgard archaea are not capable of phagocytosis [87].

Undoubtedly, future direct studies of the cell biology of Asgard archaea, once they are isolated and cultured, plus the possible finding of even more eukaryote-like lineages will shed a lot more light on the state of the eukaryotic ancestor.

It is still unlikely, however, that the archaeal host that gave rise to protoeukaryotes approached the ground state of complexity for eukaryotes. LECA seems to already have possessed a long list of complex features shared between all eukaryotes: endomembrane system, meiosis, nuclear pores, nucleosomal chromatin, linear chromosomes with telomeres, spliceosome and spliceosomal introns, small nucleolar RNPs, and others [88]. We have to then answer the question how and why these features, as well as additional complexifications restricted to certain branches of the eukaryotes, such as massive increases in genome size and gene number as well as multicellularity, arose.

Our focus for the remainder of the current discussion will be on the role that the endosymbiont played in the process of eukaryogenesis and how it might have influenced the evolution of eukaryote complexity.

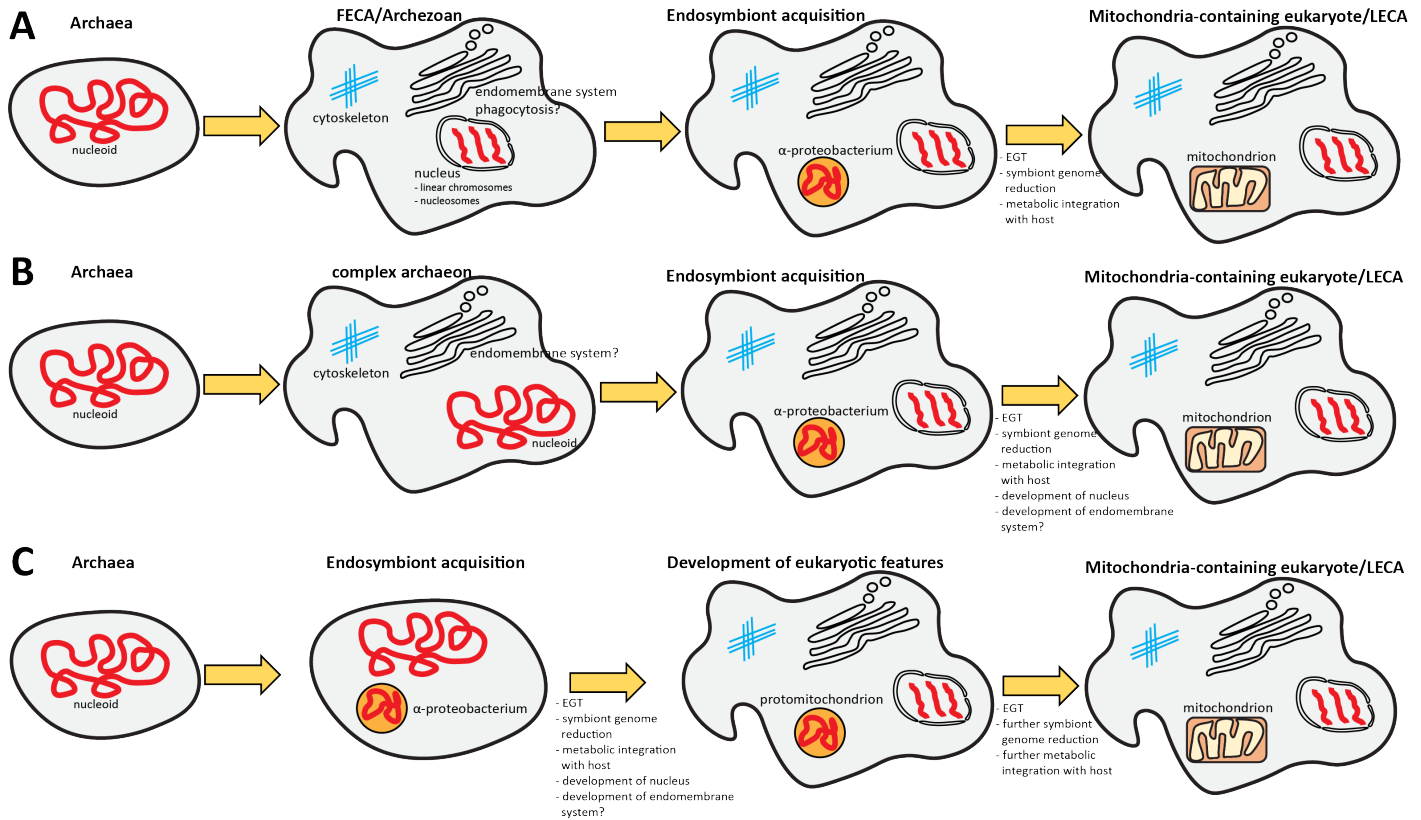
## Energy and mitochondria

What prompted us to write this review is an argument that has been for several years going on in the literature regarding the bioenergetic influence that the mitochondrion had on the process of eukaryogenesis [8, 89–103]. **It has generated both much confusion in need of clarification and an unnecessary amount of vitriol unrelated to the purely scientific issues. It is these scientific questions that we want to focus on here at some length.**

The modern mitochondrion (when not secondarily reduced) functions as a dedicated cellular organelle in which aerobic respiration occurs (among some additional functions). It is true that aerobic respiration provides much more energy to the cell than anaerobic alternatives, therefore the naïve explanation that the energetic boost provided by the mitochondrion enabled the later complexification of eukaryotes sounds very appealing on the surface. However, the proper comparison is not between aerobic eukaryotes and anaerobic prokaryotes, but between aerobic eukaryotes and aerobic prokaryotes, and quantitative support of any evolutionary scenario is much preferable to mere verbal arguments.

A more sophisticated version of the position that the bioenergetic effects of the mitochondrion were decisive for the emergence of eukaryote complexity was formulated by Lane and Martin in 2010 [89] and in later publications [8, 90] (hereafter referred to as “power-per-gene” and “bioenergetic hypothesis”). Key to Lane and Martin’s argument is the concept of “power per gene”, i.e. how much power the cell generates for each nuclear gene; the claim is that the existence of numerous organelles entirely dedicated to energy production increases the energy availability per gene, enabling expansion of genome size and gene content and of organizational complexification [90]:

The situation is more pronounced in terms of gene number. An average bacterial genome (such as *E. coli*) contains nearly 5,000 genes, compared with some 20,000 in an average protist, such as *Euglena*



**Figure 1: Different scenarios for eukaryogenesis and the relative placement and timing of symbiont acquisition, symbiont reduction and host complexification.** (A) Archezoan scenario. An archaeal ancestor develops into an organism with most typical eukaryotic features with the exception of the presence of mitochondria. An  $\alpha$ -proteobacterial endosymbiont is acquired. As a result of endosymbiotic gene transfer (EGT), endosymbiont genome reduction, and metabolic integration with the host, the  $\alpha$ -proteobacterial endosymbiont/protomitochondrion eventually turns into the highly reduced mitochondrion of the Last Eukaryotic Common Ancestor (LECA). (B) The complex archaeal ancestor scenario. An archaeal ancestor develops certain eukaryote-like features, i.e. some combination of a complex cytoskeleton, an endomembrane system. An  $\alpha$ -proteobacterial endosymbiont is acquired. EGT, endosymbiont genome reduction, and metabolic integration with the host follow resulting in the highly reduced mitochondrion of the Last Eukaryotic Common Ancestor (LECA). In parallel, the nucleus evolves, possibly triggered by the introduction of the endosymbiont [70]. (C) The protomitochondrion as the trigger of eukaryogenesis. The  $\alpha$ -proteobacterial endosymbiont is acquired by a relatively simple archaeal host. Eukaryotic features develop in parallel with the reduction of the endosymbiont into the LECA-state mitochondrion, and likely as a more or less direct consequence of its presence.

ranging up to 40,000 in *Paramecium* [62,63,70,71]. At a metabolic rate of 0.49 pW per cell, a bacterium with 5000 genes has only 0.1 fW per gene. Smaller bacteria, with around 2,500 genes, have a power of 0.2 fW per gene. Larger bacteria, with around 10,000 genes are surely close to a lower functional limit, with a power of just 0.05 fW per gene. In contrast, at a power of 2,286 pW per cell and 20,000 genes, an average protist has about 115 fW per gene, over 1000-fold more energy per gene than an average bacterium, and more than 2000-fold more than a large bacterium.

mitochondrial genome is highly reduced and compact. The ancestral  $\alpha$ -proteobacterial genome most likely contained a few thousand genes packed into several megabases of genomic space, as its modern relatives do. While some plant mitochondria (e.g. those of the *Silene* genus) have secondarily ballooned to multimegabases sizes [104], most mitochondrial genomes are only a few tens of kilobases long and they all contain only a few dozen protein coding genes at most [105]; in almost all cases, those genes are a subset of the  $\sim 66$  genes found in the mitochondrial genomes of the jakobids [106], which are the least reduced in terms of gene content mitogenomes known so far [107]. This extreme reduction is due to a combination of gene loss and endosymbiotic gene transfer (EGT) to the nuclear genome. The ques-

A critical part of Lane and Martin's argument is that the

tion why mitochondrial genomes persist at all rather than having been transferred to the nucleus altogether has long been pondered. Given that the retained genes are mostly components of the redox systems in the mitochondrial inner membrane, the best explanation proposed until now seems to be that direct and rapid regulation of the expression of these genes is necessary (“colocation for redox regulation of gene expression”, or “CoRR” hypothesis; [108]), necessitating the retention of a mitochondrial genome physically close to the location of electron transport chains (ETCs).

Lane and Martin suggest that it is not possible for a prokaryote cell to develop an analogous system of dedicated, membrane-separated organelle with its own small compact genome “servicing” its ETCs. Of course, the thylakoids of cyanobacteria are a membrane-bound ETC-containing organelle, and other prokaryotes also have developed analogous structures (e.g. the chromatophores of *Rhodobacter sphaeroides* [13]), but these do not contain their own genomes. It is therefore proposed that the only path towards increased size for a prokaryote is whole-genome polyploidy as high-copy number plasmids servicing the ETC have never been observed and are claimed to be an impossibility. However, that does not result in high energy availability per gene [90]:

Bacteria and archaea respire over their plasma membrane and thus are subject to surface-area-to-volume constraints. Increasing their linear dimension 25-fold increases their surface area 625-fold, and volume 15,000-fold. ATP synthesis could therefore increase 625-fold, but such an increase would require a 625-fold increase in the number of respiratory proteins, ATP synthase enzymes, and all other molecular machinery needed to transcribe and translate the genes. Transcription from a single bacterial genome could hardly be increased 625-fold. We must then scale up the number of genomes accordingly. To a superficial first approximation, scaling up ATP synthesis 625-fold would require haploid genome number to increase by 625-fold. Energy per gene would remain unchanged. If we take internal volume into consideration, the same principles apply. Protein synthesis could not increase 15,000-fold from a single genome, but if the number of haploid genomes were increased 15,000-fold, energy per gene would fall by 25-fold. So scaling up a bacterium to mean eukaryotic volume would cut energy per gene by  $5000 \times 25 = 125,000$ -fold [90]. All else being equal, a eukaryotic-sized bacterium should have about five orders of magnitude less energy per gene than the eukaryote [89, 90].

These numbers may seem absurd but are supported by the few examples of giant bacteria known, notably *Epulopiscium* and *Thiomargarita*. These cells are larger than most protists and show extreme polyploidy. *Epulopiscium* has tens of thousands of copies of its complete genome [17], whereas

*Thiomargarita* has around 15,000 copies [89, 109]. In both cases, the genomes are placed next to the plasma membrane, and the internal volume of the cell is metabolically inactive (being a giant vacuole in the case of *Thiomargarita*). Thus, despite their giant size, their energy per gene is exactly equivalent to that of *E. coli*, as predicted by the scaling argument above. In comparison, eukaryotic protists such as *Euglena* and *Amoeba proteus* have orders of magnitude more energy per gene.

The Lane and Martin bioenergetic argument has become quite popular in recent years (the original publication has been cited  $\geq 600$  times and we have seen the overall idea referred to as authoritative in numerous papers).

However, evolutionary hypotheses should ideally meet higher epistemic standards than the mere statement of an argument in order to be widely accepted as true. **A plausible description of the evolutionary process and the evolutionary forces in operation need to be provided.**

**We heavily stress that the bioenergetic hypothesis and the “power-per-gene” notion it introduces do not meet these criteria, and are difficult to connect to any known mechanisms for how evolution actually occurs. Much of what follows ignores that larger issue and examines the arguments for and against the bioenergetic hypothesis as if it was a much more reasonable proposition than it actually is.**

**We also stress that a minimum requirement of a set of evolutionary explanations for how different aspects of cell biology evolved is that the different explanations are consistent with each other, and that throughout what follows we will mostly evaluate the Lane and Martin’s bioenergetic relative to other hypotheses regarding eukaryogenesis advanced by Lane and Martin themselves in recent years.**

Back in 2015, as part of a larger long-term endeavor aiming at understanding cellular bioenergetics and its impact on evolution, we carried out an analysis of the biochemical costs of replicating, transcribing and translating genes [92]. We made several observations that relate to the bioenergetic hypothesis as stated by Lane and Martin. First, no discontinuity between prokaryotes and single-celled eukaryotes is observed in the metabolic scaling of energy requirements and cell size, **meaning that eukaryotes are not in fact endowed with greater bioenergetic capacity than equivalently sized aerobic prokaryotes would be.** Second, the evolution of expanded genomes in eukaryotes can be readily explained simply as a consequence of their larger physical size and lower effective population size ( $N_e$ ), without having to invoke any bioenergetic impacts by the mitochondrion.

To briefly summarize, the replication, transcription, and translation of a gene impose metabolic costs on the cell, which are a function of gene and protein length and expression levels. This cost results in a negative fitness effect  $s_c = s_{DNA} + s_{RNA} + s_{PRO}$  of magnitude on the order of the ratio between the absolute cost of maintenance of the gene and the total energy budget of the cell. Then the total fitness effect of the gene is  $s_n = s_a - s_c$ , where  $s_a$  is the direct

selective advantage conferred by the functionality provided by the gene. In order for the gene to be actively selected for or against,  $s_n$  has to exceed the drift threshold ( $1/N_e$  for haploid and  $1/2N_e$  for diploid organisms). As it turns out after calculating the biochemical costs of replication, transcription and translation, accounting for the total energy budget and the total abundance of transcripts and proteins in different types of cells, the fractional metabolic cost of genes decreases with increasing cell size, and combined with the well known fact that physically larger organisms tend to have lower effective population sizes, it appears that DNA is under selection at the metabolic level in prokaryotes, but is largely free to accumulate in large-sized eukaryotic cells.

It thus follows that even aside from the fact that the mitochondrion does not appear to have provided a major such advantage, a bioenergetic boost triggered by the presence of mitochondrion in ancient eukaryotes is not necessary to explain the complexification of their genomes.

In a later study [99], we examined in more detail the energetic and biosynthetic capacity of prokaryotes and eukaryotes as well as the cost of building cellular membranes using similar approaches. We found that, contrary to expectations, eukaryotes do not in fact have a higher ATP production capability than prokaryotes when normalized for cell size, i.e. the ATP synthesis capacity of a hypothetical cell in which all ETC complexes reside in the cell membrane is not substantially different from that of a cell of comparable size in which ETCs are located inside mitochondria. We also did not find a discontinuity in the scaling with cell size of the number of ribosomes per cell. Finally, it turns out that the addition of internal membranes does in fact impose significant metabolic costs on cells due to the high ATP cost of synthesizing lipids.

These two studies have been commented on in a negative light by Martin, Lane and colleagues on a fairly large number of occasions [96–98, 100, 101, 103], critiques that have included some rather incorrect statements. For example, in [96] we read that:

For example, population geneticists contend that mitochondria had nothing whatsoever to do with eukaryote origin (Lynch and Marinov 2015).

Another excerpt, this time from [101]:

In 2016, many if not most authors publishing on the topic of eukaryogenesis still see no crucial role either for symbiosis or for mitochondria at eukaryote origin (Gray, 2014; Keeling et al., 2015; Booth and Doolittle, 2015; Forterre, 2011; Baum, 2015; Dacks et al., 2016; Lynch and Marinov, 2015, 2016; Koonin, 2015; Martijn and Ettema, 2013). For some, “luck” is preferable to endosymbiosis as an evolutionary mechanism (Keeling et al., 2015; Booth and Doolittle, 2015).

And finally, some quotes from [98]:

The energetic and symbiogenic argument that mitochondria really made a difference at eukaryote origin (54) prompted Booth and Doolittle (15) to argue from a philosophical standpoint that mitochondria had no impact on evolution, to which an exchange (99, 100) was published. It also prompted Lynch and Marinov to argue from a population genetic standpoint that mitochondria had no impact on eukaryote evolution and that “*variation in the power of random genetic drift has played a central role in the historical diversification of genome and possibly cellular architecture across the tree of life*” (16)

[...]

Lynch usually argues that nothing other than non-adaptive processes or  $N_e$  impacted any aspect of the evolutionary process, in any organism group.

[...]

Lynch and Marinov’s (16, 17, 102) proposition is that mitochondria are irrelevant to eukaryote origin and that nothing more than point mutations and the power of population bottlenecks are needed to transform a prokaryote into a eukaryote.

As a general comment, the argument that “nothing other than nonadaptive processes or  $N_e$  impacted any aspect of the evolutionary process” has never been made; the point has always been that all evolutionary forces, one of which is genetic drift, have to be taken into account in quantitative manner when analyzing evolutionary processes, that neutral evolution is the null hypothesis to be rejected, and that when such an analysis is carried out, it turns out that selection is invoked as the default explanation in too many cases where it need not be [110–112]. **Another important point is that greatly increased complexity is generally maladaptive, first, because it results in more fragile and likely to break systems, and second, because from an evolutionary point of view it is much preferable to be a prokaryote with enormous absolute and effective population sizes than to be a highly vulnerable to extinction megafaunal vertebrate with very low absolute and effective population sizes.**

Specifically with respect to mitochondria and eukaryogenesis, the statement that “the mitochondrion had nothing to do with it” is also widely inaccurate. It is the much more specific claim that the bioenergetic impact of the mitochondrion was crucial for the origin and complexification of eukaryotes that seems doubtful to us. **This is a conclusion we initially reached based on the two studies referred to above, but it is also supported by several other lines of reasoning, to be expanded on below.**

We note that for an evolutionary explanation to be a solid and reliable one, it needs to provide an account not just of the effects that the trait in question has today but also of the *process* of evolution and of the impact the trait had during that process. A trait might be having a certain effect at a given point in time but it need not be the case

that this effect played a causal selective role in the process of evolution up to that point.

The mitochondrion is indeed the power station of modern eukaryotic cells, and it does indeed provide a large “power per (nuclear) gene” at the cost of a relatively small investment in genetic real estate. However, as we will show, even if we accept the premise that “power-per-gene” is an evolutionarily meaningful concept, it is far from warranted to conclude that these effects played a causal role in the evolution of what we recognize as core eukaryotic features or even that it had these effects when they were evolving in the very distant geologic past.

## The progression and impact of endosymbiosis

The first important consideration to take into account is that endosymbiosis is a process, and a complex one. After (and possibly even during) initial establishment of endosymbiosis, the following key interrelated aspects of that process need to unfold for an endosymbiont to become an “organelle”:

1. Biochemical integration between host and endosymbiont, involving mechanisms for transport of metabolites across membranes, has to evolve
2. Extensive EGT has to happen from the endosymbiont genome to the host genome.
3. Mechanisms for targeting and import of proteins into the endosymbiont have to develop.

The two key endosymbiotic events in the history of the planet are first, the acquisition of the mitochondrion by the protoeukaryote, and second, the acquisition of a cyanobacterial endosymbiont, later turned into a plastid, by the ancestor of the modern Archeplastida lineage. Both of these events happened in the very distant past and we have very scarce information on which to rely on for figuring out the details of the process.

However, we do have plenty of examples of more recent endosymbiotic events, from which we can obtain an idea of how analogous processes can unfold.

The original archeplastid cyanobacterial symbiosis took a cyanobacterial genome containing several thousand genes down to  $\leq 250$  genes (in red algae) and even fewer  $\leq 100$  genes in green algae and land plants. While that original event happened more than a billion years ago, unlike the situation with the mitochondrion, whose origin is, as far as we can tell, a singular event, we do have a recent example of a second primary endosymbiotic event between an eukaryote and cyanobacteria, that of the rhizarian *Paulinella*. The chromatophore of *Paulinella chromatophora* is estimated to have been acquired 60–100 Mya [113]. In that time, its genome has been reduced to a size of 1.02 Mbp containing 867 protein coding genes [114], and up to 450 now nuclear

encoded genes are trafficked back into the chromatophore [115, 116].

Another well known example of what is often referred to (correctly or not) as “organelle-formation-in-progress” are the spheroid bodies in rhopalodiacean diatoms, also of cyanobacterial origin. Notably, this is a very recent event, dating to the Miocene some 12 Mya [117]. The sequenced spheroid body genome of *Epithemia turgida* is 2.79 Mbp in size and contains 1,720 protein coding genes [118], while that of *Rhopalodia gibberula* is 3.02 Mbp in size and contains 1,671 protein coding genes [119].

A wide variety of bacterial endosymbionts are found in insects, usually exhibiting a great extent of genome reduction. The smallest endosymbiont genome known to date is that of *Nasua deltocephalinicola*, found in the phloem-feeding insect *Macrosteles quadrilineatus*, which is only 112 kbp in size [120]. It is thought to descend from a colonization event that happened  $\sim 250$  Mya or earlier [121].

These and other examples show that while the speed of genome reduction and EGT probably varies between groups, it still takes on the order of hundreds of millions of years for an endosymbiont genome to be reduced to less than 100 Kb. Of course, it is in principle possible that this happened quicker in the case of the protoeukaryote, but a counter argument can be made that modern eukaryotes are all descendants of organisms that have already at least once successfully managed to integrate an endosymbiont, thus there are numerous components of the protein targeting and compound import/export machineries that could be reused/adapted in the case of further endosymbiotic events, a luxury that the protoeukaryote did not have. In any case, it is highly doubtful that the protomitochondrion had its genome reduced extremely quickly on a geological time scale. The fact that the Jakobids, which place a lower bound on the mitogenome complexity of LECA, have such a fairly large complement of genes also argues against such a conclusion. We have no idea how long it took to go from FECA (the first eukaryotic common ancestor) to LECA, but it is likely that it was a substantial amount of time given the sheer number of unique to yet shared between all eukaryotes complex features of their cell biology.

This has some significant consequences for the Lane and Martin hypothesis, a key feature of which is that the small size of the mitochondrial genome and its complete dedication to energy production provides a much larger “power per nuclear” gene relative to what prokaryotes have to live with. The early endosymbiont was not reduced, and it most likely took a long time for it to be reduced to the point where such an effect could become significant to have a major impact on evolution.

## The impact of the (proto)mitochondrion versus the role of mature mitochondria

### *The nature of the initial endosymbiotic relationship*

An even more serious problem is that the original endosymbiont quite likely was not even a mitochondrion to begin with but had a rather different kind of relationship with the host.

The diversity of proposals on how and why exactly that relationship was established is large and difficult to summarize in a short amount of space [122]. We will restrict ourselves to pointing out that key to answering this question are the accurate estimation of the timing of the establishment of endosymbiosis and good understanding of the geochemical conditions on Earth at that time and thereafter. The original endosymbiotic theory paper by Lynn Margulis [34] placed much emphasis on the role of oxygen, envisioning a symbiotic relationship between an anaerobic host and an aerobic endosymbiont in the context of the Great Oxidation Event (GOE; the appearance of oxygen in Earth's atmosphere as a result of the activity of photosynthetic cyanobacteria  $\sim 2.5$  Gya); more recent papers have also developed the idea that the acquisition of the mitochondrion was beneficial because it conferred oxygen tolerance to the host [75, 123, 124]. However, the fossil record [125, 126] and molecular clocks [127] have not so far suggested a timing of eukaryogenesis immediately coinciding with the GOE, and it is also significant that for the rest of the Precambrian oxygen concentrations remained low ( $\leq 1\%$ ) compared to conditions in the Phanerozoic. Thus it is not clear to what extent there was a causal relationship between the GOE and eukaryogenesis. In addition, the oxygen tolerance hypothesis has serious issues given that mitochondria themselves are a major source of reactive oxygen species in modern cells [8, 128].

A popular scenario for how the endosymbiosis was established is the hydrogen hypothesis first stated by Martin and Müller in 1998 [129]. According to the hydrogen hypothesis, the archaeal host had a hydrogen-dependent methanogenic metabolism utilizing hydrogen and  $\text{CO}_2$  while the endosymbiont was facultatively anaerobic capable of generating hydrogen and  $\text{CO}_2$  as products of anaerobic respiration. The hydrogen hypothesis will be tested to some extent once the metabolism of Asgard archaea and relatives is better understood. But it need not be correct for the more general point that it illustrates: under the hydrogen hypothesis the endosymbiont did not initially function as a mitochondrion but had a very different in nature metabolic relationship with the host, and this is a general feature of many proposals for how the endosymbiosis was originally established.

The bioenergetic “power per gene” hypothesis does not apply at all for the period between the acquisition of the endosymbiont and its conversion into a true mitochondrion, and, as discussed above, some of its key arguments also do not hold until the combination of EGT and loss of non-essential genes sufficiently reduced the size of its genome.

This process could well have taken tens and hundreds of millions of years. But during all that time the endosymbiont impacted the host in numerous ways having little to do with **the bioenergetic contribution it might have made to the host, and even less with that of the yet-to-evolve mitochondrion**. In fact, many of the proposed scenarios for how some of specific complex aspects of eukaryotic cell biology evolved feature exactly that sort of impacts. We note that we do not necessarily agree with all of the proposals listed below, but they nevertheless do illustrate how hallmark eukaryote traits could have evolved without having to invoke the bioenergetic hypothesis as formulated by Lane and Martin **(we also again stress that most of these hypotheses have been originally advanced by Martin and collaborators themselves)**.

### *The origin of the endomembrane system*

While membrane invaginations and even membrane-bound compartments are not unknown in prokaryotes, the complex endomembrane system consisting of an endoplasmic reticulum, Golgi apparatus and a variety of endomembrane vesicles (lysosomes, peroxisomes, autophagosomes, and others) is, as far as we know and ignoring any potential huge surprises lurking in uncultured Asgard and related archaea, unique to eukaryotes. How did it evolve? As with most other aspects of eukaryogenesis, there are numerous scenarios and we cannot be certain about the validity of any of them. Here we will briefly consider the one proposed by Gould, Gard and Martin a couple years ago [130].

It is based on the observation that prokaryotes produce extracellular membrane vesicle called outer membrane vesicles, or OMVs [131–133]. Both bacteria and archaea have been observed to generate OMVs but important to the OMV hypothesis are the ones budding off bacterial membranes. Another relevant observation is that mitochondrial membranes also generate vesicles, called mitochondria-derived vesicles, or MDVs [133–136]. Finally, a well known general dichotomy between bacteria and archaea concerns the composition of their membranes. Archaeal lipids contain isoprenoid hydrocarbon side chains and glycerol-1-phosphate (G1P), connected via an ether linkage. In contrast, bacterial lipids are based on an ester linkage between glycerol-3-phosphate (G3P) and fatty acids. Eukaryote membranes are all of the bacterial type, thus if the eukaryote ancestor was an archaeon the transition from archaeal to bacterial lipids has to be explained (although it is to be noted that a recent phylogenomic analyses suggests that Lokiarchaeota together with some Euryarchaeota groups lack G1P synthesis capacity and instead have the capacity to synthesize G3P-based “chimeric lipids” [137]). The OMV hypothesis ties these observations together by postulating that an outward flux of OMVs from the bacterial endosymbiont gave rise to the endomembrane system in protoeukaryotes.

**We are not entirely convinced of the validity of this hypothesis, autogenous models for the origin of the endomem-**

brane system (e.g. [138]) may turn out to be more in line with what actually happened, but this is Martin's own latest model. We just note that the process as outlined could have well unfolded and proceeded quite far along the path towards the modern eukaryotic state long before the  $\alpha$ -proteobacterial endosymbiont was reduced to a true mitochondrion so that the bioenergetic hypothesis could even apply. However, the endosymbiosis event itself obviously played a major causative role (under the OMV hypothesis).

### *The evolution of phagocytosis*

Phagocytosis has been central to eukaryogenesis hypothesis for much of the history of the subject as many have assumed that a phagocytosing host is a prerequisite for the establishment of endosymbiosis [139, 140] (the "failed predation" hypothesis). There are several problems with such a view. First, bacterial endosymbionts inside other bacteria [20, 141–143] and even inside mitochondria [144] are known, meaning that phagocytosis by the host is not necessary for endosymbiosis (and on a broader speculative level, it might be argued that a rare and improbable event fits better with the singular nature of the prokaryote-to-eukaryote evolutionary transition).

Second, and more important, phagocytosis is most likely incompatible with a cellular membrane that also contains ETCs. The reason for this is that lysosomes in modern eukaryotes maintain a highly acidic pH, which is necessary for the proper functioning of the digestive enzymes contained in them. But ATP synthases generate ATP by transporting protons along the electrochemical gradient generated by electron transport, and if ATP synthases were to end up in the lysosomal membrane as a result of a phagocytosis event, the acidic pH in the lysosomal lumen would be very quickly dissipated. A similar line of reasoning was advanced by Martin and colleagues in 2017 [145].

Interestingly, this argument for why the archaeal host could not have been phagocytosing can be turned around to explain why eukaryotes did evolve phagocytosis. Once the endosymbiont/protomitochondrion (it need not have fully evolved into the LECA mitochondrial state) had taken over most ATP production in the cell, there was no longer a need to maintain ETC in the external cell membrane, and the cell was now free to evolve phagocytosis.

This was a major influence that the endosymbiont had on the (proto)eukaryote cell but, again, it was at best an indirect effect of its bioenergetic impact. Predation is known in prokaryotes (e.g. *Bdellovibrio* [146]), but it involves the predatory cell invading the prey. Phagocytosis is of a fundamentally different nature, with the predator engulfing the prey. Its appearance likely had significant ramifications for the subsequent evolution of eukaryotes, e.g. by enabling increased cell size and driving the evolution of defense mechanisms, such as rigid and armored cell walls, as a means of protection against predation.

### *The origin of spliceosomal introns*

A radical difference between the organization of eukaryotic and prokaryotic genes is the presence of spliceosomal introns in the former. Some eukaryotes have very few or almost no introns, but multiple phylogenetic reconstructions of the evolution of intron positions across the whole eukaryotic tree of life unequivocally point to the conclusion that LECA had an intron-dense genome as numerous introns in modern eukaryotes can be traced back to it [147–150]. Spliceosomal introns are spliced by the spliceosome, a rather complex biochemical machinery consisting of upwards of a hundred proteins plus several small nuclear RNAs (snRNAs), a state that too appears to be ancestral to extant eukaryotes [151].

The origin of spliceosomal introns is one of the pieces of the eukaryogenesis puzzle that we are most certain about. The core of the spliceosome is actually a ribozyme as it is the snRNAs that catalyze the splicing reaction [152], and the overall structure that the snRNAs adopt is highly reminiscent of the conformation of the self-splicing Group II introns [153], which are found in bacteria and the organelles of various eukaryotes [154]. Thus it is highly likely that the spliceosome evolved from such self-splicing Group II introns. Remarkably, Group II introns appear to be ancestrally absent from archaeal genomes, thus the endosymbiont giving rise to the mitochondrion plays a central role in the working hypothesis for how they evolved into the spliceosome [70, 155]. Originally the archaeal host had no Group II introns, but once the endosymbiosis was established, Group II introns that were present in the  $\alpha$ -proteobacterial endosymbiont began colonizing the host genome through EGT. The degeneration of these self-splicing introns through mutations made it necessary to evolve mechanisms to splice them in *trans*; the snRNAs evolved from fragments of Group II introns to take over that role, and with time the protein components of the spliceosome were added around them.

Given that EGT could have started, and probably indeed did start as soon as the endosymbiont was acquired, this example of a major impact that it had on the host is also almost completely independent from any bioenergetic effect it might have also had once it became a mitochondrion.

### *The origin of the nucleus*

Yet this line of thinking can be extended even further, and this is what was done by Martin and Koonin about a decade ago [70, 155], when they proposed that the invasion by Group II introns is what drove the evolution of the nucleus. The argument goes as follows. In prokaryotes transcription and translation are directly coupled in time and space because there is no physical separation between the two processes, while in eukaryotes the nuclear membrane compartmentalizes them. However, removal of spliceosomal introns is relatively slow, and therefore abundant spliceosomal introns would impose a major fitness cost in terms of mistranslated proteins in the absence of decoupling be-

tween transcription and translation. This is posited to have provided selective pressure for the evolution of the nucleus.

There are significant question marks regarding how exactly an organism so fundamentally flawed might have survived at all for sufficiently long to evolve a nuclear membrane, but this is beside the point. The important thing to note here is that when it comes to the specifics of the models that exist in the literature for how such a major discontinuity between prokaryotes and eukaryotes as the nucleus could have evolved, the bioenergetic effect of the mitochondrion is once again absent from the picture, and the process as proposed could have unfolded entirely within the time period from endosymbiont acquisition to its conversion into a mitochondrion with a reduced genome.

### *The evolution of sex*

EGT has also been postulated to have driven the evolution of sex, another feature that seems to have been present in LECA [156], for example, by Lane in 2011. **The hypothesis is that the invasion of mitochondrial DNA into the host genome through EGT greatly elevated the rate at which genes are inactivated by deleterious mutations (for example, through the insertion of Group II introns with diminished self-splicing activity). The protoeukaryote is proposed to have offset these fitness effects by cell fusion and masking these mutations by intact copies of these genes from other cells.** This scenario was formulated in the context of a host cell that has been freed of restriction of genome size by a fully formed mitochondrion, but there is really little reason why it would not apply from the onset of endosymbiosis and EGT. In addition, as we noted before [92, 99], the mitochondrion did not in fact provide the cell with a quantum boost in energy availability compared to a prokaryote of similar size.

### *The complexity of the archaeal host and its implications*

Finally, we come back to the fact that the recent genomic analyses of Asgard archaea [63–66, 157] as well as biochemical experiments using purified expressed proteins from their genomes [85] hint that these cells might possess fairly sophisticated cytoskeletons and membrane trafficking machineries. How complex exactly the organization of the immediate eukaryote ancestor was remains to be constrained by direct studies of these and, hopefully, additional yet to be discovered archaeal lineages. What is important to note is that the more complex that ancestor turns out to be, the less likely the bioenergetic “power-per-gene” hypothesis is to be true, as all of that complexification has to have developed before the mitochondrial endosymbiosis had even occurred.

## Summary and conclusions

### **The status of the bioenergetic hypothesis**

The key observations based on which we can evaluate the bioenergetic hypothesis are the following:

1. Accounting for cell size, there is no discontinuity between prokaryotes and eukaryotes in terms of energy budget.
2. The mitochondrion does not appear to provide aerobic cells with a major advantage in terms of energy availability relative to comparatively-sized aerobic cells without mitochondria.
3. As a result of their greater size and population genetic environment characterized by low  $N_e$ , the path towards passive genomic expansions is open to eukaryotes but not to
4. Many of the hallmark features of eukaryote could have evolved in the absence of a fully fledged mitochondrion, as a result of the impacts of having an endosymbiont yet to become a mitochondrion.
5. The endosymbiont likely had not become a true mitochondrion for a lengthy period of time following its establishment in FECA, allowing for such evolution to take place before the bioenergetic impact of the mitochondrion was present.

We can evaluate the bioenergetic hypothesis in the light of three broad eukaryogenesis scenarios outlined in Figure 1.

It is obviously incompatible with the Archezoan hypothesis, as in that sequence of events all major eukaryotic features are already present prior to the endosymbiosis taking place. Importantly, the mitochondrion plays little role in the development of shared eukaryote complexity under that hypothesis. However, the Archezoan scenario appears to be false as far as we can tell.

Complex archaeon scenarios are not incompatible with the bioenergetic hypothesis, but they nevertheless do argue against it to an extent, the more strongly so the more complex the archaeal host and the later the acquisition of the mitochondria turn out to be. In such sequences of events non-trivial cellular complexity would have evolved even without a mitochondrion, and also possibly even under anaerobic conditions.

But even the scenario of eukaryogenesis coinciding entirely with the establishment of the mitochondrial endosymbiosis does not support the bioenergetic hypothesis, because, as we saw above, complexification could have been triggered by the presence of the endosymbiont, independent of its bioenergetic effects and likely before they even took shape. However, the mitochondrion still plays an absolutely critical role in the process under that scenario, as it is these impacts (impacts on the host’s genome through EGT, possible impacts on the development of an endomembrane system, indirect impacts by enabling the evolution of

features such as phagocytosis, and others) that drove much of the process of eukaryogenesis. Thus equating skepticism of the bioenergetic hypothesis with claiming that the mitochondrion had no role in the evolution of eukaryotes is a rather gross misrepresentation.

### How and why eukaryotic complexity evolved

As has been demonstrated extensively previously [110–112, 158–160], many of the complex features of eukaryote genomic organization, such as the larger size of eukaryote genomes, the presence of introns, the accumulation of transposable elements and intergenic space, the baroque complexity of gene regulation in multicellular organisms (many of the intermediate steps that lead from simple to more complex gene regulatory networks are non-adaptive and are thus more easily tolerated in organisms with lower  $N_e$  [160]), etc., most of them often thought of as adaptive improvements over the simple eukaryote state, can in fact be readily and much more parsimoniously explained by the inability of natural selection to get rid of slightly deleterious features in a population genetic environment characterized by low effective population size. Our bioenergetic analyses [92, 99] provided further support for that view, by identifying differences between the situation in prokaryotes and eukaryotes with respect to the fitness impact that the presence of additional DNA and its expression has on the cell.

The key variable that emerges from this analyses is cellular size. Larger cells are more tolerant to the presence of additional DNA (cases of massive whole-genome polyploidization aside) because of how much smaller the fractional cost of its maintenance is, and also because physical size is generally inversely correlated with  $N_e$ . Increased tolerance towards the addition of genes and the expansion of noncoding space can then lead to increases in the complexity of the gene repertoire of the cell and of the regulatory networks governing the expression of these genes. The latter, by virtue of the multiple alternative cell states they can generate, are probably a prerequisite for the emergence of complex multicellularity.

The reasons why the tolerance towards (or, alternatively, inability to purge) extra DNA due to the combination of large cellular size and low  $N_e$  is conducive to complexification are severalfold. At the gene level, complexification occurs through the de novo origin of new genes from intergenic space, the acquisition of new genes through horizontal gene transfer, or, perhaps most importantly, through subfunctionalization or neofunctionalization of duplicate copies of existing genes. The maintenance of intergenic space imposes metabolic costs on the cell proportional to its length, which, as discussed above, are sufficiently large to be subject to selection against in prokaryotes, plus it represents an increase in the size of the mutational target in the cell, also proportional to its length and also potentially sufficiently large to be subject to selection in lineages with very large  $N_e$ . A similar line of reasoning applies to genes acquired through HGT, which is dependent on the interplay

between the magnitude of their positive fitness effect to the cell and the negative fitness effect of the metabolic cost of maintaining and expressing them. Finally, theoretical analyses point to subfunctionalization becoming an increasingly more probable mechanism for the preservation of duplicate genes with decreasing population size [161, 162].

At the level of gene regulation, complexification occurs through the expansions of the set of regulatory elements (REs) controlling the expression of each gene (such as enhancers and insulators). The typical human gene is associated with as many as a dozen distally located such elements [163], and it is often the case that different enhancers are utilized to drive the expression of a gene in different cell types and tissues. This is in marked contrast with prokaryotes and many unicellular eukaryotes where gene regulation is mostly promoter proximal. The proliferation of REs is of key importance to the evolution of new cell types, and in turn, of organismal complexity, as it allows the same set of genes to be used in different combinations in novel contexts. However, this is not necessarily an adaptive process, as discussed in detail previously [160]. In short and in a most simplified form, addition of regulatory embellishments increases the size of the target for degenerative mutations, thus it has a negative fitness effect proportional to the mutation rate  $\mu$ , and it also imposes a metabolic cost through its maintenance. Larger cells and population genetic environments characterized by lower  $N_e$  are therefore more conducive to increases in regulatory complexity.

We can thus understand the evolution of eukaryotes and complex life on planet Earth on the broadest level as driven by an evolutionary ratchet towards increased complexity emerging passively from the neutral fixation of otherwise nonadaptive genomic changes in lineages with long-term population genetic environments permissive to it. In contrast, prokaryotes remain relatively simple and streamlined due to their small size and very high  $N_e$ , which efficiently eliminate the genomic alterations that might otherwise have led to significantly increased complexity from their populations. The process goes as follows: increases in cell size, decreases in  $N_e$  and changes in genomic organization in early eukaryotes make it easier to acquire and tolerate additional DNA. The emergence of features such as phagocytosis and a complex cytoskeletal organization might enable and further drive cell size increases. The increased complexity of gene composition and gene regulation eventually leads to the appearance of simple multicellularity, which in turn lowers  $N_e$  even further as multicellular organisms are typically physically larger than their unicellular relatives. The lowered  $N_e$  facilitates even further complexification of gene regulatory networks, enabling the evolution of yet larger and more complex organisms, which in turn have even lower  $N_e$ . This state need not be an inevitable product – the vast majority of eukaryote lineages are still unicellular [164] – but it is also true that such an evolutionary path is probably closed to populations of small-sized aerobic prokaryotes.

The mitochondrial endosymbiosis was the key event

that set this process in motion. Yet it most likely did so through its numerous other impacts on its host rather than through an increase in the available “power-per-gene” as proposed by Lane and Martin.

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