

The Syntrophy hypothesis for the origin of eukaryotes revisited

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20 The discovery of Asgard archaea, phylogenetically closer to eukaryotes than other archaea, together with 21 improved knowledge of microbial ecology impose new constraints on emerging models for the origin of the 22 eukaryotic cell (eukaryogenesis). Long-held views are metamorphosing in favor of symbiogenetic models 23 based on metabolic interactions between archaea and bacteria. These include the classical Searcy's and 24 hydrogen hypothesis, and the more recent Reverse Flow and Entangle-Engulf-Enslave (E³) models. Two 25 decades ago, we put forward the Syntrophy hypothesis for the origin of eukaryotes based on a tripartite 26 metabolic symbiosis involving a methanogenic archaeon (future nucleus), a fermentative myxobacterial-like 27 deltaproteobacterium (future eukaryotic cytoplasm) and a metabolically versatile methanotrophic 28 alphaproteobacterium (future mitochondrion). A refined version later proposed the evolution of the 29 endomembrane and nuclear membrane system by invagination of the deltaproteobacterial membrane. 30 Here, we adapt the Syntrophy hypothesis to contemporary knowledge, shifting from the original hydrogen 31 and methane-transfer-based symbiosis (HM-Syntrophy) to a tripartite hydrogen and sulfur-transfer-based 32 model (HS-Syntrophy). We propose a sensible ecological scenario for eukaryogenesis in which eukaryotes 33 originated in early Proterozoic microbial mats from the endosymbiosis of a hydrogen-producing Asgard 34 archaeon within a complex sulfate-reducing deltaproteobacterium. Mitochondria evolved from versatile, 35 facultatively aerobic, sulfide-oxidizing and, potentially, anoxygenic photosynthesizing, alphaproteobacterial 36 endosymbionts that recycled sulfur in the consortium. The HS-Syntrophy hypothesis accounts for 37 (endo)membrane, nucleus and metabolic evolution in a realistic ecological context. We compare and 38 contrast the HS-Syntrophy hypothesis to other models of eukaryogenesis, notably in terms of the mode and 39 tempo of eukaryotic trait evolution, and discuss several model predictions and how these can be tested. 40

41 Eukaryogenesis was a unique major evolutionary transition resulting in significant average cell complexity 42 increase. This foundational event led to an impressive radiation of morphologically diverse phyla, most of them unicellular (protists) but many including multicellular taxa such as animals, fungi, kelp and land 43 44 plants¹. Elusive for a long time, reconstructing a mechanistically plausible and ecologically realistic model 45 for the origin of eukaryotes appears now within reach thanks to recent advances in molecular 46 phylogenomic tools, genome-binning from metagenomes and a better knowledge of microbial diversity and function in natural ecosystems. Until recently, notwithstanding the generally accepted endosymbiotic 47 origin of mitochondria and chloroplasts, models proposing the symbiotic origin of eukaryotes directly 48 from bacterial and archaeal ancestors were largely dismissed^{2,3}. The prevailing view stated that an 49 50 independent proto-eukaryotic lineage sister to archaea evolved most eukaryotic features (complex cytoskeleton, endomembranes, nucleus, phagocytosis, etc.) before it engulfed the mitochondrial 51 alphaproteobacterial ancestor⁴⁻⁶. This view started to vacillate with the realization that truly primary 52 amitochondriate eukaryotes were not known⁷ and additionally deteriorated with phylogenomic trees 53 where eukaryotes branched within archaea, albeit without clear sister groups⁸. The discovery of Asgard 54 archaea, a phylogenetically deep-branching lineage sharing more and more similar genes with eukaryotes 55 than other archaea^{9,10}, has further fostered this paradigm shift on eukaryogenesis. Eukaryotes are no 56 longer on the same footing as archaea and bacteria as one of the original primary domains of life¹¹; they 57 are a third, but secondary, domain resulting from the evolutionary merging of specific archaeal and 58 59 bacterial linages^{2,6,12-14}. Moreover, current knowledge about Asgard general metabolic potential and preferred biotopes (mostly sediments and microbial mats, where intimate metabolic interactions are the 60 rule^{3,15}), realistically favor symbiogenetic models based on metabolic symbioses or syntrophies^{16,17}. This is 61 further supported by the syntrophic nature of the first cultured Asgard member, Candidatus 62 Prometheoarchaeum syntrophicum, an anaerobic organism able to grow in symbiosis with a sulfate-63 reducing deltaproteobacterium, a methanogenic archaeon or both¹⁸. Collectively, this strongly supports 64 cooperative models for the origin of the eukaryotic cell^{2,3,6,17} whereby higher complexity evolved from the 65 physical integration of prokaryotic cells combined with extensive gene and genome shuffling^{12,19-21}. 66

The first symbiogenetic models date back to more than 20 years ago. Among them, the more detailed 67 were the Serial Endosymbiosis Theory²²⁻²⁴, the Hydrogen hypothesis²⁵ and the Syntrophy hypothesis^{26,27}. 68 69 In the original Syntrophy hypothesis, we proposed that eukaryotes evolved from a tripartite metabolic 70 symbiosis based on i) interspecies hydrogen transfer from a fermenting deltaproteobacterial host to an endosymbiotic methanogenic archaeon and ii) methane recycling by a versatile methanotrophic, 71 facultative aerobic alphaproteobacterium²⁶ (Hydrogen-Methane –HM– Syntrophy). From an ecological 72 perspective, these metabolic interactions were reasonable, being widespread in anoxic and redox-73 transition settings². However, knowledge about archaeal diversity and metabolism was then much more 74 75 fragmentary than today, and the metabolic potential of uncultured lineages remained inaccessible. The 76 probable involvement of an Asgard archaeal relative in eukaryogenesis imposes new constraints, such 77 that realistic models need to take into account their metabolic potential and ecology. Accordingly, several 78 symbiogenetic models are currently being put forward. They differ on the metabolic interactions 79 proposed (Box 1) and, importantly, the tempo and mode of evolution of key eukaryotic traits (Box 2). 80 Here, we present an updated version of the Syntrophy hypothesis based on a tripartite metabolic 81 symbiosis involving interspecific hydrogen and sulfur-transfer (HS-Syntrophy) occurring in redox-82 transition ecosystems: a complex sulfate-reducing deltaproteobacterium (host), an endosymbiotic hydrogen-producing Asgard-like archaeon (future nucleus) and a metabolically versatile, facultatively
 aerobic, sulfide-oxidizing and potentially anoxygenic photosynthesizing, alphaproteobacterium (future

aerobic, sulfide-oxidizing and potentially anoxygenic photosynthesizing, alphaproteobacterium (future
 mitochondrion). We briefly discuss the evolution of the endomembrane system, the nucleus and the

86 genome¹⁹ in the framework of the HS-Syntrophy hypothesis. This model makes several predictions that

genome¹⁹ in the framework of the HS-Syntrophy hypothesis. This model makes several predictions that
differentiate it from alternative scenarios including, notably, the two-step origin of the nucleus (first, as

- distinct metabolic compartment before its consecration as major genetic reservoir and expression center)
- 89 and the bacterial origin of eukaryotic membranes and cytoplasm. We propose ways to specifically test
- 90 some aspects of different eukaryogenesis models and offer suggestions for future avenues of research.
- 91

92 The ecological context of the eukaryogenetic symbiosis

Despite the challenges associated to the interpretation of the earliest life traces, the oldest reliable 93 eukaryotic fossils can be dated back to at least 1.65 Ga²⁸. This imposes a minimal age for the origin of 94 95 eukaryotes that roughly agrees with the oldest boundaries of recent molecular dating estimates for the last eukaryotic common ancestor (LECA; 1.0-1.6 Ga)²⁹ and the eukaryotic radiation (<1.84 Ga)³⁰. At the 96 same time, LECA was complex, being endowed with mitochondria and resembling modern protists^{20,21}. 97 Although the alphaproteobacterial lineage that gave rise to the mitochondrion remains to be precisely 98 identified³¹, it is clear that the mitochondrial ancestor was aerobic³², but likely also possessed anaerobic 99 respiratory capacities (as in many modern protists). This implies that i) aerobic respiration had already 100 evolved in bacteria when the mitochondrial endosymbiosis occurred and ii) oxic or microaerophilic 101 102 conditions existed in the environment where the mitochondrial endosymbiosis took place or in its 103 immediate vicinity. Aerobic respiration possibly evolved (almost) in parallel to cyanobacterial oxygenic photosynthesis, which led to the oxygenation of the atmosphere, the Great Oxidation Event (GOE), some 104 2.4 Ga ago at the beginning of the Proterozoic (2.5-0.5 Ga)^{33,34}. Therefore, eukaryogenesis took place 105 between the GOE and the minimum age of the oldest unambiguous eukaryotic fossils²⁸. If some older, 106 more difficult to affiliate, fossils³⁵ are indeed eukaryotic, eukaryogenesis might have occurred during the 107 first three to five hundred million years after the GOE. 108

109 What did the Earth look like at that time? Before the GOE, the atmosphere and oceans were 110 essentially anoxic, which constrained existing biogeochemical cycles. The atmosphere rapidly oxygenated from 2.33 Ga but sulfate levels in oceans increased slower³³, limiting the biological S cycle^{36,37}. This means 111 that oceans were oxygen-poor during the early Proterozic, when eukaryotes evolved; the deep ocean 112 remained anoxic until the beginning of the Phanerozoic (500 Ma)³⁸⁻⁴⁰. If an aerobic mitochondrial 113 ancestor suggests oxygen availability at or near the environment where eukaryotes finally evolved, 114 115 current knowledge on Asgard archaea ecology and metabolism strongly suggests that the archaeon involved in eukaryogenesis, and hence the first eukaryogenetic steps, were strictly anaerobic. Asgard 116 archaea are mostly found in deep-sea sediments^{9,10,18,41} and microbial mats¹⁰, including thermophilic 117 ones^{10,42}. Thus, with the exception of some derived planktonic Heimdallarchaeota, which more recently 118 acquired the capacity to oxidize organics using nitrate or oxygen as terminal electron acceptors^{16,43}, the 119 vast majority of Asgard archaea thrive in anoxic environments, as their ancestors did, degrading organics 120 and producing or consuming hydrogen¹⁶. These observations argue in favor of redox transition 121 environments, where anoxic and oxic/microoxic zones are in close proximity, as favored ecosystems for 122 123 eukaryogenesis. Furthermore, since the early Proterozoic deep ocean was anoxic, it is more likely that 124 eukaryotes evolved in shallow sediments or microbial mats, where redox gradients established, like

today, from the oxygen-enriched surface where cyanobacterial oxygenic photosynthesis took place to theincreasingly anoxic layers below.

127 Phototrophic microbial mats are particularly interesting potential eukaryogenesis cradles. They were the Proterozoic 'forests', dominating shallow aquatic and terrestrial habitats, as abundant fossil 128 stromatolites (lithified microbial mats) show^{34,44,45}. These light- and redox-stratified microbial 129 communities are phylogenetically and metabolically diverse^{42,46,47}. Although microorganisms in modern 130 mats are different from their Proterozoic counterparts, core metabolic functions have been mostly 131 preserved across phyla⁴⁸ and at the ecosystem level, suggesting that functional shifts observed in mats 132 across redox gradients today reflect early metabolic transitions⁴⁹. Most primary production occurs in 133 upper layers, where light can penetrate, via photosynthetic carbon fixation. The upper cyanobacterial 134 135 oxygenic-photosynthesis layer is typically followed by a reddish layer dominated by oxygen-tolerant anoxygenic photosynthesizers (Alpha- and Gammaproteobacteria) and often an underlying green layer of 136 photosynthetic Chloroflexi and/or Chlorobi. Organic matter fixed in the upper mat layers is progressively 137 degraded in deeper, anoxic layers, by extremely diverse microbial communities^{42,50,51}. Two broad zones 138 can be distinguished in vertical anoxic profiles where, respectively, sulfate reduction and methanogenesis 139 dominate⁴⁶ (Fig. 1a). Here, like in anoxic sediments, the degradation of organic matter involves 140 syntrophy⁵², mostly implicating interspecies hydrogen (or, directly, electron⁵³) transfer. In anoxic 141 142 environments, pairs of electron donors and acceptors display low redox potential differences such that many energy-generating metabolic reactions can only proceed in the presence of syntrophic sinks⁵⁴. 143 144 Methanogenic archaea and sulfate-reducing bacteria (SRB) belonging to the Deltaproteobacteria are frequently engaged in syntrophy. Deltaproteobacteria are metabolically diverse and can use or produce 145 hydrogen or, directly, electrons⁵⁵, being frequently involved in interspecies hydrogen or electron 146 transfer⁵³. Many of them oxidize organic compounds with sulfate, but they can also be autotrophic⁵⁶ 147 (including in syntrophy⁵⁷), use other electron donors and acceptors (including metals, such as arsenic⁵⁸), 148 ferment or switch between metabolisms depending on the environmental conditions⁵⁹. 149 150 Deltaproteobacteria establish widespread syntrophies with archaea; with methanogens when acting as 151 hydrogen producers, with methanotrophic archaea when acting as hydrogen-consuming sulfatereducers⁶⁰. Mutualistic interactions between methanogens and deltaproteobacteria can be rapidly 152 selected, leading to specialized syntrophy^{61,62}. Deltaproteobacterial SRB also establish symbioses with 153 sulfide-oxidizing or other bacteria and eukaryotes^{2,59,63}. In addition to methanogens and SRB, a wide 154 155 variety of uncultured lineages occurs in anoxic sediment and microbial mat layers, where archaea thrive. Many of these archaea seem to be involved in cycling organics, particularly alkanes, being likely engaged 156 in syntrophies with hydrogen-scavengers^{49,52,64}. Indeed, the first cultured Asgard archaeon can grow by 157 degrading amino acids in syntrophy with either a sulfate-reducing deltaproteobacterium and/or a 158 159 methanogen¹⁸.

160

161 Eukaryogenetic syntrophies

162 Considering this historical and ecological context, we favor the idea that eukaryogenesis occurred in 163 microbial mats (or similarly stratified shallow sediments) with marked redox gradients (**Fig. 1a**), 164 potentially mildly warm. Oxygenic photosynthesis (and in close proximity, aerobic respiration) might have 165 first evolved in warm environments. Indeed, most deep-branching cyanobacteria are thermophilic^{65,66}. 166 Although universal molecular mechanisms to cope with reactive oxygen species exist^{67,68} and might have 167 been co-opted from antioxidant-prone compounds very early^{69,70}, oxygen toxicity would have been

advantageously relieved in thermophilic mats by its rapid release into the atmosphere (oxygen is poorly 168 169 soluble at high temperature). The original (HM) Syntrophy hypothesis postulated, on solid microbial 170 ecology grounds, the evolution of eukaryotes from well-known widespread symbioses between fermenting (hydrogen-producing), ancestrally SRB, deltaproteobacteria and methanogenic archaea. SRB 171 and methanogens, which compete for hydrogen, can readily evolve stable syntrophy in co-culture^{61,62}. We 172 additionally favored a myxobacterial-like deltaproteobacterium due to the similarities shared by these 173 complex social bacteria and eukaryotes^{19,26}. This symbiosis would have established at the sulfate-methane 174 transition zone but evolved upwards in the redox gradient, where an additional symbiosis formed with a 175 176 versatile methanotrophic alphaproteobacterium that scavenged the methane released by the primary 177 consortium. We cannot completely reject such tripartite metabolic symbiosis at the origin of eukaryotes since methanogenesis, originally thought exclusive of Euryarchaeota, occurs across archaeal phyla and 178 might have been ancestral to the archaeal domain⁷¹⁻⁷⁴. However, although methanogenesis might be 179 eventually discovered in Asgard archaea (some Asgard archaea do have methyl-coenzyme M reductases 180 probably involved in the reverse, anaerobic alkane oxidation, reaction⁴¹), current genomic comparisons 181 seem to exclude it from their ancestral metabolic capacities¹⁶. 182

183 In this context, we now favor a similar eukaryogenetic process but based on alternative, albeit 184 equally ecologically relevant, metabolic symbioses (Fig. 1). In our HS-Syntrophy hypothesis, we propose 185 that eukaryotes evolved from the syntrophic interaction of a sulfate-reducing (hydrogen/electronrequiring) deltaproteobacterium, possibly sharing some complex traits with myxobacteria, and a 186 187 hydrogen-producing Asgard-like archaeon. This deltaproteobacterium may have been metabolically 188 versatile or mixotrophic, but in symbiosis with the archaeon, it respired sulfate. This initial facultative 189 symbiosis was stabilized by the incorporation of the archaeon as endosymbiont (Fig. 1b-c). This 190 consortium likely established first in deeper anoxic layers and subsequently migrated upwards in the 191 redox gradient, where it established a second (initially facultative) symbiosis with a sulfide-oxidizing 192 alphaproteobacterium that acted as both, sulfide sink and sulfate donor for the Asgard-193 deltaproteobacterium consortium (Fig. 1b-c). Alternatively, the two facultative symbioses might have co-194 existed although, in this case, we favor a later obligatory endosymbiosis of the alphaproteobacterial ancestor¹⁹. This would be in line with genomic evidence suggesting a late mitochondrial symbiosis⁷⁵. 195 Given the dominance of H₂S-dependent anoxygenic photosynthesizing bacteria in microbial mats and 196 their interaction with SRB for sulfur cycling in upper layers^{76,77}, the versatile, facultatively aerobic 197 mitochondrial ancestor was likely also photosynthetic (or mixotrophic). Interestingly, the possibility that 198 199 mitochondrial cristae evolved from intracytoplasmic membranes typical of photosynthetic Alphaproteobacteria has been highlighted⁷⁸. This tripartite symbiotic consortium became definitely 200 201 stabilized when the alphaproteobacterium became an endosymbiont within the deltaproteobacterium 202 (Fig. 1d). In our view, the first eukaryotic common ancestor (FECA) is neither an archaeon nor a 203 bacterium, but the first obligatory symbiogenetic consortium. Strictly speaking, this would correspond to 204 the integrated symbiosis of the three partners that contributed to the final making of the eukaryotic cell 205 and genome. But the FECA stage could also be decoupled in time in two subsequent stages corresponding 206 to the integration of the Asgard archaeon within the deltaproteobacterium (FECA 1) and the acquisition 207 of the mitochondrial endosymbiont (FECA 2).

In our model, up to the FECA stage, the eukaryogenetic syntrophies were based on the samemetabolic exchange that occurred in the corresponding facultative symbioses (hydrogen between the

210 archaeon and the SRB; sulfide/sulfate between the SRB and SOB). However, the incorporation of the 211 mitochondrial ancestor as obligatory endosymbiont implied a radical change in the metabolism of the 212 whole consortium, constraining the outcome of the eukaryogenetic process. Because the mitochondrial 213 ancestor was also aerobic and could get a much higher energy yield by directly oxidizing organics, the consortium started to rely solely on aerobic respiration (Fig. 1e). This resulted in the loss of the less-214 efficient anaerobic archaeal metabolism and bacterial sulfate-reduction. At the same time, the proto-215 216 eukaryote migrated to the fully oxic layers of the mats, spreading on oxic surfaces and, upon the 217 development of motility mechanisms, colonizing the planktonic realm. Cellular changes that included the 218 development of an extensive endomembrane system (see below) led to the LECA stage (Fig. 1f).

219 The HS-Syntrophy model implies three prokaryotic partners that became integral part of the future 220 eukaryotic cell. However, other pre-eukaryogenetic symbioses might have occurred at the facultative syntrophy stage, eventually leaving historical traces in the form of transferred genes to the 221 222 eukaryogenetic symbiotic partners. One traditional criticism to symbiogenetic models proposing the 223 endosymbiosis of one prokaryote within another prokaryote is the absence of phagocytosis in prokaryotes^{2,4,6,79,80}. Mainstream models now accept a symbiogenetic origin of eukaryotes but only under 224 the premise that an endomembrane system, a developed cytoskeleton and phagocytosis evolved in the 225 archaeal ancestor prior to the engulfment of the mitochondrial ancestor^{6,81}. However, prokaryotes 226 227 harboring endosymbiotic prokaryotes are known and might be more frequent than currently thought. In 228 addition to the well-known cases of gammaproteobacterial symbionts within betaproteobacterial endosymbionts in mealybugs⁸² and rickettsiales in tick mitochondria⁸³, old electron microscopy studies⁸⁴⁻ 229 ⁸⁶ and more recent observations⁸⁷ suggest the potential occurrence of prokaryotic endosymbionts in 230 bacteria. Interestingly, a recent report of prey engulfment by planctomycetes⁸⁸ suggests that bona fide 231 bacterial phagocytosis exists, albeit based on different molecular grounds than eukaryotic phagocytosis⁸⁸. 232 233 In the case of archaea, although Nanoarchaeota can be associated to the inter-membrane space in the archaeon Ignicoccus hospitalis⁸⁹, true endosymbionts remain to be observed. Consequently, regardless 234

the mechanism, these collective observations suggest that prokaryotic endosymbioses, at least within
 bacteria, are feasible.

237

238 Membranes and endomembranes

239 The eukaryotic plasma membrane and endomembrane system, including the endoplasmic reticulum (ER), 240 the nuclear membrane, the Golgi apparatus and other vesicular components (vacuoles, lysosomes, etc.) are interconnected (are either continuous or can fuse and merge). They share a similar composition, with 241 typical bacterial-like phospholipids^{2,90}. The phospholipid bilayer in eukaryotes is particularly flexible and 242 243 can undergo deformation, bending, fusion and fission. This is achieved thanks to a highly developed cytoskeleton⁹¹, coating components (e.g. clathrin/AP1-5, COPI, TSET, COPII, retromer, ESCRT complexes), 244 ARF/ARF-like GTPases and their regulators, and fusion machinery (involving SNARE complex, multisubunit 245 tethering complexes, Rab GTPases and regulatory factors)⁹²⁻⁹⁴. Both, the cytoskeleton and membrane 246 remodeling and fusion complexes were already present in LECA, which was capable of phagocytosis, 247 secretion and trafficking, and have a chimeric origin^{20,93,95-97}. In addition to innovations^{21,95}, some 248 cytoskeletal and membrane-remodeling proteins are archaeal-like (e.g. actin, profilin, ESCRT proteins^{6,98}, 249 perhaps some GTPases⁹⁹, although some of these might be bacterial^{100,101}) but a significant number of 250

endomembrane system-related proteins could also be of bacterial, though not alphaproteobacterial,
 origin⁷⁵. The biosynthesis of sterols is notably of bacterial origin¹⁰².

Most eukaryogenetic models propose a two-partner symbiosis in which the archaeal host 253 254 incorporated the alphaproteobacterial ancestor of mitochondria. This implies a shift of the host membrane from the more rigid archaeal, glycerol-1-phosphate (G1P)-based ether-linked isoprenoid 255 phospholipids to the more flexible and permeable bacterial glycerol-3-phosphate (G3P)-based, usually 256 ester-linked, fatty acid phospholipids^{2,90} (**Box 2**). However, such a transition with, in particular, a G1P-to-257 G3P-based phospholipid shift, has never been observed in nature (some thermophilic bacteria use ether-258 links, long-term-known exceptions⁹⁰). Recently, an engineered *Escherichia coli* strain was forced to 259 express archaeal phospholipids, making up to 30% of the total membrane phospholipids¹⁰³. The 260 engineered heterochiral-membrane strain was viable and, interestingly, the expressed archaeal lipids 261 recruited G1P, suggesting that stereospecificity is somehow linked to the phospholipid composition, in a 262 263 peculiar form of membrane heredity. However, if more than 30% archaeal lipids incorporated to the 264 membrane, severe growth impairment was observed and the shape of cells became aberrant; they produced numerous vesicles and underwent asymmetric cell division¹⁰³. One could therefore ask how the 265 expression of archaeal phospholipids affects E. coli fitness and whether such engineered strain would be 266 267 able to survive competition with normal bacteria in natural environments. Archaeal and bacterial 268 phospholipids impose very different local physicochemical conditions that constrain integral membrane proteins¹⁰⁴. As a consequence, a membrane lipid composition shift implies an extensive adaptation of the 269 whole membrane-associated proteome¹⁰⁵. In this context, neither the stability of heterochiral 270 liposomes¹⁰⁶ nor the (partial) expression of archaeal phospholipids in engineered *E. coli*¹⁰³ can be taken as 271 272 evidence for an archaeal-to-bacterial membrane transition. While the bacterial nature of eukaryotic 273 phospholipids represents a serious difficulty for models invoking an archaeal host, it is naturally explained 274 by the bacterial nature of the host in the Syntrophy hypothesis (Box 2, Fig. 1).

275 In the HS-Syntrophy model, the endomembrane system results from the invagination of the 276 deltaproteobacterium inner membrane and the internalization of the periplasm. The outer bacterial 277 plasma membrane would be retained as the eukaryotic plasma membrane. Many bacteria harbor endomembrane compartments linked to specialized biochemical functions¹⁰⁷. These include the well-278 known cyanobacterial thylakoids, but also compartments in magnetotactic bacteria¹⁰⁸, anammox 279 bacteria¹⁰⁹ and Poribacteria¹¹⁰. Some Planctomycetes develop a thoroughly studied nuclear-like 280 compartment¹¹¹ and a similar structure has been recently described in the candidate phylum 281 Atribacteria¹¹². This implies that the internalization of membranes is relatively common across bacterial 282 phyla. Although Deltaproteobacteria with endomembranes have not been described, their diversity is far 283 from fully explored and they have membrane-remodeling potential. For instance, developed 284 285 cytoskeletons (a prerequisite for extensive membrane remodeling) exist in the predatory *Bdellovibrio*¹¹³ but also in myxobacteria, which are able to generate protruding membrane tubes that interconnect 286 cells¹¹⁴. In the HS-Syntrophy model, similarly to the former HM-Syntrophy¹⁹, the initial driving force for 287 endomembrane evolution is the establishment of a secretory system that connected the endosymbiotic 288 289 archaeon with the periplasm (Fig. 1c-d). As in contemporary heterotrophic deltaproteobacteria, the 290 periplasm was the digestive space of the host deltaproteobacterium in which complex organics uptaken 291 from the environment were hydrolyzed to simpler organics. Some of these simpler organics (e.g. amino 292 acids, short hydrocarbons) were used by the archaeon for its organoheterotrophic metabolism, which

yielded hydrogen used in turn by the SRB host. By being an endosymbiont, the archaeon maximized its 293 294 uptake surface for small organics from the bacterial cytoplasm. In turn, the deltaproteobacterial host 295 maintained an optimal uptake surface for complex organics from the environment while having a ready 296 internal source of hydrogen (or electrons) for sulfate reduction. As the symbiosis evolved, many genes 297 were transferred from the deltaproteobacterium to the archaeon, which progressively centralized genes 298 and gene expression for the whole consortium. This included, notably, many hydrolytic enzymes required 299 for the periplasmic degradation of complex organics. These enzymes were transported from the archaeal 300 compartment towards the original bacterial periplasm via an incipient endomembrane system that, 301 eventually, fully surrounded the archaeon and constituted the future nuclear membrane (see below). This 302 implied the evolution of a transport system only through the archaeal membrane since, on the bacterial 303 side, transporters for the export of newly synthesized hydrolytic enzymes to the periplasm and the 304 environment already existed. Bacterial transporters might have initially been inserted also in the archaeal 305 membranes (following gene transfer to the archaeal genome) but later replaced by channels communicating with bacteria-derived pore-like structures and allowing the export of increasingly bigger 306 307 and varied substrates (see below). The transfer of hydrolytic enzymes to the digestive periplasmic space 308 via the endomembrane system was essential to prevent the hydrolysis of cytoplasmic components (Fig. 309 1d). At the same time, in this way, the digestive space largely increased. Hence, the initial digestive and 310 trafficking-related endomembrane system was the precursor of the nuclear membrane and the ER but also of the different eukaryotic vesicles related to digestive processes (lysosomes, peroxisomes, digestive 311 312 vacuoles). Upon the endosymbiosis of the mitochondrial ancestor and the loss of the archaeal and SRB 313 metabolism, the organics were directly oxidized via aerobic respiration by the alphaproteobacterium and 314 the endomembrane system was retained for the trafficking of proteins synthesized in the proto-nucleus 315 (Fig. 1e) and, with time, in association with the ER itself (Fig. 1f). At the same time, the ancient periplasm was completely internalized and the former digestive periphery transferred to independent vesicular 316 317 compartments (Fig. 1f). The secretory Golgi apparatus as well as other endocytotic and exocytotic 318 systems developed in parallel. The association of archaeal membrane-bending systems (e.g. ESCRT) with 319 the host bacterial membranes facilitated the process of endomembrane formation.

320

321 The origin of the nucleus

Most eukaryogenetic models fail to advance convincing selective forces to explain why the nucleus 322 evolved². We propose, like in the HM-Syntrophy¹⁹, a two-step process entailing two sequential selective 323 forces. First, a proto-nucleus evolved as a different metabolic compartment. This chimeric compartment 324 325 was composed of the endosymbiotic archaeon and the surrounding proto-nuclear membrane of 326 deltaproteobacterial origin (Fig. 1c-d). The first selective force for the evolution of the nuclear membrane 327 was the need to export bacterial enzymes already synthesized by the archaeon (after their genes were 328 transferred to the archaeal genome) to the periplasmic space. Its first role was therefore secretory 329 (export towards the trafficking endomembrane system). Other proteins of archaeal origin also started to be exported, contributing to the evolution of several chimeric eukaryotic systems. Once the archaeal 330 331 genome started to host essential genes from its symbiotic partners, and these genes were lost from the 332 donor genomes, the archaeon started to centralize protein synthesis for the whole consortium. This 333 entailed the development of a transport mechanism from the archaeal cytoplasm to the bacterial

334 endomembrane system, which was at the origin of the nuclear pore. This implied the formation of 335 coordinated apertures through the archaeal membrane and the proto-nuclear membrane, although these apertures might have formed only on the bacterial membrane (future nuclear pores) with archaeal 336 337 membrane transporters facilitating export prior to archaeal membrane loss. The potential to establish communicating pores exists in both, archaea and deltaproteobacteria. Archaea are able to establish 338 intercellular cytoplasmic bridges and fuse¹¹⁵. Myxobacterial deltaproteobacteria are also able to fuse their 339 membranes^{116,117} and develop contact-dependent abilities, including coordinated gliding via junctional 340 pore complexes¹¹⁸⁻¹²¹. Progressively, ribosomes concentrated around these incipient communicating 341 342 pores, eventually migrating to the host's cytoplasm along the endomembrane system (future ER), where 343 protein synthesis started to take place. This led to a progressive decoupling of translation, which became 344 associated to the ER, and transcription, which took place in the archaeal cytoplasm (future nucleoplasm).

345 As the consortium evolved between FECA and LECA, after the mitochondrial ancestor was fixed in the 346 consortium and the archaeal and SRB metabolisms were lost in favor of the more efficient mitochondrial respiration, the archaeal membrane became useless and was completely lost (membrane loss is not 347 infrequent in the framework of endosymbiosis¹²²). However, during this evolutionary process, extensive 348 genome evolution took place¹⁹. This involved (endo)symbiotic gene transfer (EGT) to the archaeal 349 350 genome, likely accompanied by other HGT, which was largely facilitated by active processes fostering 351 genome evolution, such as gene and genome fragment duplication and reshuffling. As transcription and translation decoupled, introns invaded the future eukaryotic genome. They likely derived from the 352 original self-splicing introns of the alphaproteobacterial endosymbiont¹²³, possibly complemented by 353 other mechanisms^{124,125}. Once introns invaded the genome, the proto-nuclear membrane was selectively 354 retained (exapted) to maintain the transcription-translation uncoupling. Therefore, preventing the 355 deleterious massive synthesis of aberrant proteins was the second selective force acting during nuclear 356 evolution¹⁹. Intron invasion has been proposed as exclusive selective force for the origin of the nucleus¹²⁶. 357 358 However, in our view, transcription-translation uncoupling, and therefore a nuclear membrane, must pre-359 exist in order for introns to spread and not the opposite². Not only the insertion of one or a few introns in 360 essential genes would be immediately deleterious, but the evolution of a continuous nuclear membrane requires intermediate steps during which transcription and translation are still coupled that intron 361 invasion as selective force cannot explain. 362

The initial chimeric proto-nuclear pore evolved into the modern nuclear pore as traffic check-point and hub of gene regulation¹²⁷. We view the nucleolus and the ribosomal particle assembly process as remnants of the archaeal origin of the nuclear compartment¹⁹. The assembly of eukaryotic ribosomes is a complex and energy-costly process that takes place in the nucleolus. Ribosomal proteins are synthesized in the cytoplasm and transported to the nucleus. After assembly with rRNA in the nucleolus, ribosomal particles are transported back to the cytoplasm, where they associate to the ER for function¹²⁸. The set of proteins involved (processome) is essentially of archaeal origin¹²⁹.

370

371 The make-up of a composite genome

During eukaryogenesis, various mechanisms shaped the evolving eukaryotic genome. These included
 HGT, EGT, gene duplication, loss and new gene creation, accompanied by the invasion of introns and
 mobile selfish elements. The directionality of gene transfer to the archaeal genome and its establishment
 as future nuclear genome might have been simply dictated by chance; as the consequence of an essential

gene transfer from one genome to the other genome followed by loss in the donor¹⁹. The retention of the 376 377 archaeal endosymbiont genome as future nuclear genome is often criticized on the ground that in extant 378 cases of endosymbioses, endosymbionts tend to reduce their genomes in favor of the host's. However, 379 known extant endosymbioses occur within eukaryotes, which are already composite cells harboring 380 mitochondria and eventually chloroplasts for which many essential genes already reside in the nuclear 381 genome. Therefore, the eukaryotic nuclear genome is essential and must centralize genes coming from any new incoming endosymbiont. The situation was radically different at the origin of eukaryotes when 382 383 organelle reliance on the nuclear genome was not yet established and symbiotic partners were mutually 384 dependent. In the Syntrophy hypothesis, the archaeal genome became the future nuclear genome. 385 Bacterial components were thus included in an archaeal genomic background, leading to the long-term 386 recognized mixed heritage of eukaryotic genomes, with 'informational' genes (related to DNA replication, transcription and translation) being archaeal-like, and 'operational' genes (involved in energy and carbon 387 metabolism) bacterial-like¹³⁰. While true in general terms, a closer look to the bacterial-like genes in 388 eukaryotes poses some questions. 389

390 Most symbiogenetic models invoke only two partners, an archaeal host and the alphaproteobacterial 391 ancestor of mitochondria (Box 1). Consequently, two predictions follow: i) host (archaeal-like) genes must 392 dominate over the endosymbiont (alphaproteobacterial-like) genes and ii) most bacterial-like eukaryotic genes must be of alphaproteobacterial origin. However, neither of them holds. Bacterial-like genes are 393 more abundant than archaeal-like genes in eukaryotic genomes¹³¹ and genes with alphaproteobacterial 394 ancestry only represent a minority of bacterial-like genes in modern eukaryotes^{75,132} and LECA¹³³. To 395 396 explain this 'silent' non-alphaproteobacterial bacterial majority in eukaryotic genomes, the progressive 397 erosion of ancient phylogenetic signal making it difficult to pinpoint the precise origin of those genes and massive HGT from diverse bacterial donors to the archaeal and/or the alphaproteobacterial symbiotic 398 partners have been invoked^{131,134}. High bacteria-to-archaea HGT levels have been observed in several 399 phyla¹³⁵, including the Asgard archaea^{9,10}. However, the patterns observed in eukaryotic genomes could 400 be only explained if genes transferred to the archaeal and/or alphaproteobacterial ancestors of 401 eukaryotes had been subsequently lost in all their sister lineages, which is unlikely¹³³. In addition, 402 eukaryotic alphaproteobacterial-like genes have significantly shorter branches than other bacterial-like 403 genes in phylogenetic trees including prokaryotic homologues⁷⁵. This suggests a late mitochondrial arrival 404 in a host with an already chimeric genome⁷⁵. Moreover, if alphaproteobacterial-like genes mostly relate 405 to mitochondrial functions, bacterial genes of non-alphaproteobacterial ancestry seem to be involved in 406 other essential eukaryotic traits such as the endomembrane system, reinforcing the idea that they 407 evolved prior to the mitochondrion⁷⁵. Non-alphaproteobacterial genes appear to derive from various 408 bacterial phyla (with Deltaproteobacteria and Actinobacteria among the most frequent donors), 409 410 suggesting successive ancient waves of HGT from these phyla and/or the implication of several bacterial 411 symbionts during eukaryogenesis (Box 2e). Symbiogenetic models involving more than two partners are 412 often dismissed applying a simplistic parsimony argument. However, parsimony is not evolutionary 413 evidence and, in most complex ecosystems, multiple symbioses are widespread³. The first cultured 414 Asgard archaeon can indeed grow in symbiosis with one sulfate-reducing deltaproteobacterium and one methanogenic archaeon¹⁸. If we transpose a similar symbiosis at the onset of eukaryogenesis, a significant 415 number of deltaproteobacterial-like genes in eukaryotes might be explained by HGT during the long-term 416 coexistence with a symbiont that later disappeared without integrating the consortium¹⁸. Additional 417

bacterial ecto- or endo-symbionts might have also transferred genes, leading to the mosaic origin of
 eukaryotic bacterial-like genes as proposed in the 'pre-mitochondrial symbiosis' model¹³³ and on the line
 of the 'shopping-bag model' proposed for the evolution of plastid genomes¹³⁶.

However, the presence of many non-alphaproteobacterial bacterial-like genes in eukaryotes is 421 422 compatible with the HS-syntrophy model. Accordingly, deltaproteobacterial genes would have been 423 acquired by EGT. Interestingly, deltaproteobacterial-like genes seem to be the most abundant nonalphaproteobacterial category, and also older⁷⁵. Inferring the precise phylogenetic origin of genes of 424 different ages in eukaryotic genomes is far from trivial due to mutational saturation and the erosion of 425 phylogenetic signal in increasingly older genes¹³⁷. Furthermore, each potential additional symbiont could 426 427 contribute a number of genes acquired by HGT from different donors in such a way that the apparent 428 number of eukaryogenetic symbiotic partners would appear inflated. Nonetheless, strong phylogenetic 429 signal supports the deltaproteobacterial origin of many eukaryotic genes involved in diverse functions and structures. In addition to early identified deltaproteobacterial-like eukaryotic genes²⁶, the list also 430 includes antimicrobial defensins¹³⁸, Ser/Thr/Tyr protein kinases¹³⁹, PPP protein phosphatases¹⁴⁰, high 431 mobility group A proteins¹⁴¹, isoprenoid biosynthesis enzymes¹⁴², cyclitol synthases¹⁴³ and some 432 kinetochore proteins⁹⁷. Mitochondria also recruited some deltaproteobacterial proteins, potentially 433 reflecting an alpha-delta-proteobacterial symbiosis, such as thiolases¹⁴⁴, fatty acid beta-oxidation 434 enzymes¹⁴⁵, and possibly, some proteins involved in anaerobic metabolism. Stemming from the versatility 435 436 of many alphaproteobacteria, we view the mitochondrial ancestor as a facultative aerobe able to carry out anaerobic respiration with various electron acceptors but also substrate-level phosphorylation²⁷. 437 438 Several genes involved in these reactions seem ancestral in eukaryotes, branching close to deltaproteobacteria and other anaerobic bacteria¹⁴⁶. Although they are usually interpreted as 439 independent HGT acquisitions from various donors¹⁴⁶, these observations can alternatively support a 440 deltaproteobacterial anaerobic respiration toolkit in ancestral eukaryotic mitochondria, subsequently lost 441 442 to different degrees in aerobic lineages.

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444 Future prospects

Any model on eukaryogenesis must account for the evolution of key eukaryotic traits (e.g. genome 445 complexity, nature and origin of eukaryotic membranes/endomembranes and the nucleus), in a way that 446 447 i) is mechanistically plausible and ii) explains the observed patterns and the causes (selective forces) for the evolution of those traits in iii) a realistic ecological context. In this framework, our HS-Syntrophy 448 449 model takes into account constraints imposed by the discovery of Asgard archaea and their ancestral 450 metabolic potential to put forward one of the most comprehensive eukaryogenetic models. This model 451 presents some difficulties, notably in the centralization of the genome and protein synthesis by the 452 archaeon with subsequent export to the deltaproteobacterial host. However, it is ecologically relevant, 453 fits well with the observed chimerism of eukaryotic genomes and has the advantage, over archaeal hostbased models (Box 1), of readily explaining the bacterial-like nature of eukaryotic membranes^{2,90}. The HS-454 Syntrophy hypothesis makes several predictions that differentiate it from other hypotheses (Table 1). 455 456 Some of these are shared with the HM-Syntrophy model and include the presence of EGT-derived 457 deltaproteobacterial genes in eukaryotes that should be mostly involved in membrane, cell-signaling and 458 cytoplasmic functions. Others, such as the involvement of a potentially photosynthetic S-oxidizing 459 alphaproteobacterium, specifically characterize the HS-Syntrophy. These predictions suggest that such specific Alpha- and Deltaproteobacterial lineages, phylogenetically closer to eukaryotes than other 460 bacterial lineages, might exist. The nature of the alphaproteobacterial ancestor of mitochondria is indeed 461 still cryptic³¹. If such lineages were discovered through environmental studies in microbial mats or 462 sediments, the HS-Syntrophy would gain support. These Syntrophy models are realistically based on well-463 464 known metabolic interactions in microbial mats or sediments, but metabolic variants involving a similar 465 tripartite symbiosis and eukaryogenetic process might be also envisaged within a more general Syntrophy 466 model; its distinctive features being the nuclear origin from an archaeal endosymbiont in a bacterial 467 cytoplasm and the independent acquisition of the mitochondrial ancestor.

- 468 Beyond the Syntrophy hypothesis, progress in several areas is needed to answer open questions and differentiate major model types (Table 1). In archaeal host-based symbiotic models^{16,18,147,148}, three major 469 issues need attention. First, the archaeal-to-bacterial membrane transition remains a major drawback. 470 471 Cases of membrane transitions implying complete phospholipid-type replacement and concomitant 472 membrane proteome adaptation are not observed in nature. If bacteria and, more particularly archaea, 473 can be engineered to i) accomplish the full replacement of membrane phospholipids for the opposite 474 type and ii) be competitive in real environmental conditions (permissive fitness cost), the historical 475 feasibility of such transition would be supported. So far, the evidence is lacking. In some models, the 476 emission of cell protrusions and a progressive engulfment of an alphaproteobacterium is preferred over immediate phagocytosis^{18,147}. That long process would favor phospholipid exchange and replacement of 477 the rigid archaeal phospholipids for more flexible bacterial ones¹⁴⁷. However, it is unclear how such a 478 479 "slow phagocytosis" process would occur across prokaryotic generations, which are needed for evolution 480 to take place (phospholipid replacement and proteome adaptation) prior to true engulfment. Second, 481 two-partner models need to propose convincing detailed evolutionary mechanisms and selective forces for the origin of the eukaryotic nucleus, which are so far lacking. Finally, eukaryogenetic models need to 482 483 explain the 'silent bacterial majority' in eukaryotic genomes. In two-partner models, HGT from different bacteria to any of the partners might be the most logical explanation, but this does not necessarily 484 explain why groups of functionally-related genes seem to come from a few bacterial groups^{75,133}, even if 485 486 some HGT and phylogenetic reconstruction noise are likely involved in these observations. In-depth 487 phylogenomic analyses including a broad taxon sampling and, if identified, the closest bacterial and 488 archaeal relatives of eukaryotes will be needed. Collectively, the information gathered from 489 environmental, experimental cell biology and phylogenomic studies should help to discriminate existing 490 models, refine them or create new ones. More than ever, solving the eukaryogenesis riddle seems at 491 hand.
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494 References

495 1 Adl, S. M. et al. The revised classification of eukaryotes. J Eukaryot Microbiol 59, 429-493 (2012).

496 2 Lopez-Garcia, P. & Moreira, D. Open questions on the origin of eukaryotes. *Trends Ecol Evol* 30, 697497 708 (2015).

498 3 Lopez-Garcia, P., Eme, L. & Moreira, D. Symbiosis in eukaryotic evolution. *J Theor Biol* 434, 20-33
499 (2017).

501 (2007). 502 5 de Duve, C. The origin of eukaryotes: a reappraisal. Nat Rev Genet 8, 395-403 (2007). 503 6 Eme, L., Spang, A., Lombard, J., Stairs, C. W. & Ettema, T. J. G. Archaea and the origin of eukaryotes. 504 Nat Rev Microbiol 15, 711-723 (2017). 505 7 Embley, T. M. & Hirt, R. P. Early branching eukaryotes? Curr. Opin. Genet. Dev. 8, 624-629 (1998). 506 Cox, C. J., Foster, P. G., Hirt, R. P., Harris, S. R. & Embley, T. M. The archaebacterial origin of 8 507 eukaryotes. Proc Natl Acad Sci U S A 105, 20356-20361 (2008). 508 Spang, A. et al. Complex archaea that bridge the gap between prokaryotes and eukaryotes. Nature 9 509 **521**, 173–179 (2015). 510 10 Zaremba-Niedzwiedzka, K. et al. Asgard archaea illuminate the origin of eukaryotic cellular 511 complexity. Nature 541, 353-358 (2017). 512 11 Williams, T. A., Cox, C. J., Foster, P. G., Szöllősi, G. J. & Embley, T. M. Phylogenomics provides robust 513 support for a two-domains tree of life. Nat Ecol Evol, 138–147 (2019). 514 12 McInerney, J. O., O'Connell, M. J. & Pisani, D. The hybrid nature of the Eukaryota and a consilient 515 view of life on Earth. Nat Rev Microbiol 12, 449-455 (2014). 516 13 Koonin, E. V. Archaeal ancestors of eukaryotes: not so elusive any more. *BMC Biol* **13**, 84 (2015). 517 14 Williams, T. A. & Embley, T. M. Changing ideas about eukaryotic origins. *Philos Trans R Soc Lond B* 518 Biol Sci 370, 20140318 (2015). 519 15 Libby, E., Hebert-Dufresne, L., Hosseini, S. R. & Wagner, A. Syntrophy emerges spontaneously in 520 complex metabolic systems. PLoS Comput Biol 15, e1007169 (2019). 521 16 Spang, A. *et al.* Proposal of the reverse flow model for the origin of the eukaryotic cell based on 522 comparative analyses of Asgard archaeal metabolism. Nat Microbiol (2019). 523 17 Lopez-Garcia, P. & Moreira, D. Eukaryogenesis, a syntrophy affair. Nat Microbiol 4, 1068-1070 524 (2019). 525 18 Imachi, H. et al. Isolation of an archaeon at the prokaryote-eukaryote interface. bioRxiv, 726976 526 (2019). 527 19 López-García, P. & Moreira, D. Selective forces for the origin of the eukaryotic nucleus. *Bioessays* 28, 528 525-533 (2006). 529 20 Koonin, E. V. & Yutin, N. The dispersed archaeal eukaryome and the complex archaeal ancestor of 530 eukaryotes. Cold Spring Harb Perspect Biol 6, a016188 (2014). 531 21 Koonin, E. V. Origin of eukaryotes from within archaea, archaeal eukaryome and bursts of gene gain: 532 eukaryogenesis just made easier? Philos Trans R Soc Lond B Biol Sci 370, 20140333 (2015). 533 22 Sagan, L. On the origin of mitosing cells. *J Theor Biol* **14**, 255-274 (1967). 534 23 Margulis, L. Origin of eukaryotic cells. (Yale University Press, 1970). 535 24 Margulis, L., Dolan, M. F. & Guerrero, R. The chimeric eukaryote: origin of the nucleus from the 536 karyomastigont in amitochondriate protists. Proc Natl Acad Sci U S A 97, 6954-6959 (2000). 537 25 Martin, W. & Muller, M. The hydrogen hypothesis for the first eukaryote. *Nature* **392**, 37-41 (1998). 538 26 Moreira, D. & López-García, P. Symbiosis between methanogenic archaea and delta-Proteobacteria 539 as the origin of eukaryotes: The syntrophic hypothesis. J. Mol. Evol. 47, 517-530 (1998). 540 27 López-García, P. & Moreira, D. Metabolic symbiosis at the origin of eukaryotes. Trends Biochem Sci 541 **24**, 88-93 (1999). 542 28 Javaux, E. J. Challenges in evidencing the earliest traces of life. *Nature* **572**, 451-460 (2019). 543 29 Eme, L., Sharpe, S. C., Brown, M. W. & Roger, A. J. On the age of eukaryotes: evaluating evidence 544 from fossils and molecular clocks. Cold Spring Harb Perspect Biol 6, a016139 (2014). 545 30 Betts, H. C. et al. Integrated genomic and fossil evidence illuminates life's early evolution and 546 eukaryote origin. Nat Ecol Evol 2, 1556-1562 (2018).

Poole, A. M. & Penny, D. Evaluating hypotheses for the origin of eukaryotes. *Bioessays* 29, 74-84

500

- 547 31 Martijn, J., Vosseberg, J., Guy, L., Offre, P. & Ettema, T. J. G. Deep mitochondrial origin outside the
 548 sampled alphaproteobacteria. *Nature* 557, 101-105 (2018).
- S49 32 Roger, A. J., Munoz-Gomez, S. A. & Kamikawa, R. The Origin and Diversification of Mitochondria. *Curr*S50 *Biol* 27, R1177-r1192 (2017).
- Luo, G. *et al.* Rapid oxygenation of Earth's atmosphere 2.33 billion years ago. *Sci Adv* 2, e1600134
 (2016).
- Knoll, A. H., Bergmann, K. D. & Strauss, J. V. Life: the first two billion years. *Philos Trans R Soc Lond B Biol Sci* 371 (2016).
- 555 35 El Albani, A. *et al.* Organism motility in an oxygenated shallow-marine environment 2.1 billion years
 556 ago. *Proc Natl Acad Sci U S A* 116, 3431-3436 (2019).
- 557 36 Canfield, D. E., Habicht, K. S. & Thamdrup, B. The Archean sulfur cycle and the early history of
 558 atmospheric oxygen. *Science* 288, 658-661 (2000).
- 37 Halevy, I., Johnston, D. T. & Schrag, D. P. Explaining the structure of the Archean mass-independent
 sulfur isotope record. *Science* 329, 204-207 (2010).
- Shen, Y., Knoll, A. H. & Walter, M. R. Evidence for low sulphate and anoxia in a mid-Proterozoic
 marine basin. *Nature* 423, 632-635. (2003).
- 563 39 Poulton, S. W., Fralick, P. W. & Canfield, D. E. The transition to a sulphidic ocean approximately 1.84
 564 billion years ago. *Nature* 431, 173-177 (2004).
- 565 40 Stolper, D. A. & Keller, C. B. A record of deep-ocean dissolved O2 from the oxidation state of iron in
 566 submarine basalts. *Nature* 553, 323 (2018).
- 567 41 Seitz, K. W. *et al.* Asgard archaea capable of anaerobic hydrocarbon cycling. *Nat Commun* 10, 1822568 1822 (2019).
- Saghaï, A. *et al.* Unveiling microbial interactions in stratified mat communities from a warm saline
 shallow pond. *Environ Microbiol* 19, 2405-2421 (2017).
- 571 43 Bulzu, P. A. *et al.* Casting light on Asgardarchaeota metabolism in a sunlit microoxic niche. *Nat Microbiol* (2019).
- 44 Hamilton, T. L., Bryant, D. A. & Macalady, J. L. The role of biology in planetary evolution:
 cyanobacterial primary production in low-oxygen Proterozoic oceans. *Environ Microbiol* 18, 325-340
 (2016).
- 45 Lenton, T. M. & Daines, S. J. Matworld the biogeochemical effects of early life on land. *New Phytol*577 215, 531-537 (2017).
- 578 46 Bolhuis, H., Cretoiu, M. S. & Stal, L. J. Molecular ecology of microbial mats. *FEMS Microbiol Ecol* 90, 335-350 (2014).
- Faerl, H. W., Pinckney, J. L. & Steppe, T. F. Cyanobacterial-bacterial mat consortia: examining the
 functional unit of microbial survival and growth in extreme environments. *Environ Microbiol* 2, 11-26
 (2000).
- 48 Martiny, J. B., Jones, S. E., Lennon, J. T. & Martiny, A. C. Microbiomes in light of traits: A phylogenetic
 584 perspective. *Science* 350, aac9323 (2015).
- 585 49 Gutierrez-Preciado, A. *et al.* Functional shifts in microbial mats recapitulate early Earth metabolic
 586 transitions. *Nat Ecol Evol* 2, 1700-1708 (2018).
- 50 Harris, J. K. *et al.* Phylogenetic stratigraphy in the Guerrero Negro hypersaline microbial mat. *ISME J*588 7, 50-60 (2013).
- 51 Wong, H. L., Smith, D. L., Visscher, P. T. & Burns, B. P. Niche differentiation of bacterial communities
 at a millimeter scale in Shark Bay microbial mats. *Sci Rep* 5, 15607 (2015).
- 52 Dombrowski, N., Seitz, K. W., Teske, A. P. & Baker, B. J. Genomic insights into potential
 interdependencies in microbial hydrocarbon and nutrient cycling in hydrothermal sediments. *Microbiome* 5, 106 (2017).

- 53 Lovley, D. R. Syntrophy Goes Electric: Direct Interspecies Electron Transfer. *Annu Rev Microbiol* 71, 643-664 (2017).
- 54 Fenchel, T. & Finlay, B. J. *Ecology and evolution in anoxic worlds*. (Oxford University Press, 1995).
- 55 Lovley, D. R. Happy together: microbial communities that hook up to swap electrons. *ISME J* (2016).
- 56 Mall, A. *et al.* Reversibility of citrate synthase allows autotrophic growth of a thermophilic bacterium.
 599 *Science* 359, 563-567 (2018).
- 600 57 Krukenberg, V. *et al. Candidatus* Desulfofervidus auxilii, a hydrogenotrophic sulfate-reducing
 601 bacterium involved in the thermophilic anaerobic oxidation of methane. *Environ Microbiol* 18, 3073602 3091 (2016).
- 603 58 Oremland, R. S. & Stolz, J. F. The ecology of arsenic. *Science* **300**, 939-944. (2003).
- 604 59 Muyzer, G. & Stams, A. J. The ecology and biotechnology of sulphate-reducing bacteria. *Nat Rev* 605 *Microbiol* 6, 441-454 (2008).
- 60 Knittel, K. & Boetius, A. Anaerobic oxidation of methane: progress with an unknown process. *Annu Rev Microbiol* 63, 311-334 (2009).
- 61 Hillesland, K. L. *et al.* Erosion of functional independence early in the evolution of a microbial
 609 mutualism. *Proc Natl Acad Sci U S A* 111, 14822-14827 (2014).
- 62 Hillesland, K. L. & Stahl, D. A. Rapid evolution of stability and productivity at the origin of a microbial
 611 mutualism. *Proc Natl Acad Sci U S A* **107**, 2124-2129 (2010).
- 63 Monteil, C. L. *et al.* Ectosymbiotic bacteria at the origin of magnetoreception in a marine protist. *Nat* 613 *Microbiol* 4, 1088-1095 (2019).
- 64 Wang, Y., Wegener, G., Hou, J., Wang, F. & Xiao, X. Expanding anaerobic alkane metabolism in the
 615 domain of Archaea. *Nat Microbiol* 4, 595-602 (2019).
- 65 Shi, T. & Falkowski, P. G. Genome evolution in cyanobacteria: The stable core and the variable shell.
 617 *Proc Natl Acad Sci U S A* 105, 2510-2515 (2008).
- 66 Shih, P. M. *et al.* Improving the coverage of the cyanobacterial phylum using diversity-driven genome
 69 sequencing. *Proc Natl Acad Sci U S A* **110**, 1053-1058 (2013).
- 67 Martins, M. C. *et al.* How superoxide reductases and flavodiiron proteins combat oxidative stress in
 621 anaerobes. *Free Radic Biol Med* 140, 36-60 (2019).
- 68 Slesak, I., Kula, M., Slesak, H., Miszalski, Z. & Strzalka, K. How to define obligatory anaerobiosis? An
 evolutionary view on the antioxidant response system and the early stages of the evolution of life on
 Earth. *Free Radic Biol Med* 140, 61-73 (2019).
- 69 Fischer, W. W., Hemp, J. & Valentine, J. S. How did life survive Earth's great oxygenation? *Curr Opin*626 *Chem Biol* **31**, 166-178 (2016).
- 627 70 Neubeck, A. & Freund, F. Sulfur Chemistry May Have Paved the Way for Evolution of Antioxidants.
 628 Astrobiology (2019).
- 629 71 Berghuis, B. A. *et al.* Hydrogenotrophic methanogenesis in archaeal phylum Verstraetearchaeota
 630 reveals the shared ancestry of all methanogens. *Proc Natl Acad Sci U S A* 116, 5037-5044 (2019).
- 631 72 Borrel, G. *et al.* Wide diversity of methane and short-chain alkane metabolisms in uncultured
 632 archaea. *Nat Microbiol* 4, 603-613 (2019).
- 633 73 Evans, P. N. *et al.* An evolving view of methane metabolism in the Archaea. *Nature Reviews*634 *Microbiology* 17, :219-232 (2019).
- 635 74 McKay, L. J. *et al.* Co-occurring genomic capacity for anaerobic methane and dissimilatory sulfur
 636 metabolisms discovered in the Korarchaeota. *Nat Microbiol* 4, 614-622 (2019).
- 637 75 Pittis, A. A. & Gabaldon, T. Late acquisition of mitochondria by a host with chimaeric prokaryotic
 638 ancestry. *Nature* 531, 101-104 (2016).
- 639 76 Canfield, D. E. & Des Marais, D. J. Aerobic sulfate reduction in microbial mats. *Science* 251, 1471640 1473 (1991).

- 641 77 Visscher, P. T. *et al.* Formation of lithified micritic laminae in modern marine stromatolites
 642 (Bahamas): The role of sulfur cycling. *Am Mineral* 83, 1482-1493 (1998).
- 643 78 Munoz-Gomez, S. A., Wideman, J. G., Roger, A. J. & Slamovits, C. H. The Origin of Mitochondrial
 644 Cristae from Alphaproteobacteria. *Molecular biology and evolution* 34, 943-956 (2017).
- 645 79 Cavalier-Smith, T. Predation and eukaryote cell origins: a coevolutionary perspective. *Int J Biochem*646 *Cell Biol* 41, 307-322 (2009).
- 647 80 Martin, W. F., Garg, S. & Zimorski, V. Endosymbiotic theories for eukaryote origin. *Philos Trans R Soc*648 *Lond B Biol Sci* 370, 20140330 (2015).
- 649 81 Martijn, J. & Ettema, T. J. From archaeon to eukaryote: the evolutionary dark ages of the eukaryotic
 650 cell. *Biochem Soc Trans* 41, 451-457 (2013).
- 651 82 von Dohlen, C. D., Kohler, S., Alsop, S. T. & McManus, W. R. Mealybug beta-proteobacterial
 652 endosymbionts contain gamma-proteobacterial symbionts. *Nature* 412, 433-436 (2001).
- 83 Sassera, D. *et al.* '*Candidatus* Midichloria mitochondrii', an endosymbiont of the tick Ixodes ricinus
 with a unique intramitochondrial lifestyle. *Int J Syst Evol Microbiol* 56, 2535-2540 (2006).
- 655 84 Wujek, D. E. Intracellular bacteria in the blue-green alga *Pleurocapsa minor Trans Am Microscop Soc*656 98, 143-145 (1979).
- Larkin, J. M., Henk, M. C. & Burton, S. D. Occurrence of a *Thiothrix* sp. attached to mayfly larvae and
 presence of parasitic bacteria in the Thiothrix sp. *Appl Environ Microbiol* 56, 357-361 (1990).
- 659 86 Larkin, J. M. & Henk, M. C. Filamentous sulfide-oxidizing bacteria at hydrocarbon seeps of the gulf of
 660 Mexico. *Microsc Res Tech* 33, 23-31 (1996).
- 661 87 Yamaguchi, M. *et al.* Prokaryote or eukaryote? A unique microorganism from the deep sea.
 662 *Microscopy* 61, 423-431 (2012).
- 663 88 Shiratori, T., Suzuki, S., Kakizawa, Y. & Ishida, K.-I. Phagocytosis-like cell engulfment by a
 664 planctomycete bacterium. *Nat Commun* 10, 5529-5529 (2019).
- 665 89 Heimerl, T. *et al.* A Complex Endomembrane System in the Archaeon Ignicoccus hospitalis Tapped by
 666 Nanoarchaeum equitans. *Frontiers in microbiology* 8 (2017).
- 667 90 Lombard, J., López-García, P. & Moreira, D. The early evolution of lipid membranes and the three
 668 domains of life. *Nat Rev Microbiol* 10, 507-515 (2012).
- 91 Jekely, G. Origin and evolution of the self-organizing cytoskeleton in the network of eukaryotic
 organelles. *Cold Spring Harb Perspect Biol* 6, a016030 (2014).
- 671 92 Dacks, J. B. & Field, M. C. Evolutionary origins and specialisation of membrane transport. *Curr Opin*672 *Cell Biol* 53, 70-76 (2018).
- 673 93 Dey, G., Thattai, M. & Baum, B. On the archaeal origins of eukaryotes and the challenges of inferring
 674 phenotype from genotype. *Trends in Cell Biology* 26, 476-485 (2016).
- 675 94 Rout, M. P. & Field, M. C. The Evolution of Organellar Coat Complexes and Organization of the
 676 Eukaryotic Cell. *Annu Rev Biochem* 86, 637-657 (2017).
- 95 Yutin, N., Wolf, M. Y., Wolf, Y. I. & Koonin, E. V. The origins of phagocytosis and eukaryogenesis. *Biol Direct* 4, 9 (2009).
- 679 96 Lombard, J. The multiple evolutionary origins of the eukaryotic N-glycosylation pathway. *Biology*680 *Direct* 11 (2016).
- 681 97 Tromer, E. C., van Hooff, J. J. E., Kops, G. & Snel, B. Mosaic origin of the eukaryotic kinetochore. *Proc*682 *Natl Acad Sci U S A* 116, 12873-12882 (2019).
- 683 98 Akıl, C. & Robinson, R. C. Genomes of Asgard archaea encode profilins that regulate actin. *Nature*684 562, 439-443 (2018).
- 685 99 Klinger, C. M., Spang, A., Dacks, J. B. & Ettema, T. J. Tracing the archaeal origins of eukaryotic
 686 membrane-trafficking system building blocks. *Molecular biology and evolution* 33, 1528-1541 (2016).
- 687 100 Jekely, G. Small GTPases and the evolution of the eukaryotic cell. *Bioessays* 25, 1129-1138. (2003).
- 101 Low, H. H. & Lowe, J. A bacterial dynamin-like protein. *Nature* 444, 766-769 (2006).

- 689 102 Santana-Molina, C., Rivas-Marin, E., Rojas, A. M. & Devos, D. P. Origin and evolution of polycyclic
 690 triterpene synthesis. *Molecular biology and evolution*, pii: msaa054. doi:
- 691 010.1093/molbev/msaa1054. [Epub ahead of print] (2020).
- 692 103 Caforio, A. *et al.* Converting Escherichia coli into an archaebacterium with a hybrid heterochiral
 693 membrane. *Proc Natl Acad Sci U S A* 115, 3704-3709 (2018).
- 694 104 Pogozheva, I. D., Tristram-Nagle, S., Mosberg, H. I. & Lomize, A. L. Structural adaptations of proteins
 695 to different biological membranes. *Biochim Biophys Acta* 1828, 2592-2608 (2013).
- 696 105 Makarova, M. *et al.* Delineating the rules for structural adaptation of membrane-associated proteins
 697 to evolutionary changes in membrane lipidome. *bioRxiv*, 762146 (2019).
- 698 106 Shimada, H. & Yamagishi, A. Stability of heterochiral hybrid membrane made of bacterial sn-G3P
 699 lipids and archaeal sn-G1P lipids. *Biochem* 50, 4114-4120 (2011).
- 107 Diekmann, Y. & Pereira-Leal, J. B. Evolution of intracellular compartmentalization. *Biochem J* 449, 319-331 (2013).
- T02 108 Greene, S. E. & Komeili, A. Biogenesis and subcellular organization of the magnetosome organelles of
 T03 magnetotactic bacteria. *Curr Opin Cell Biol* 24, 490-495 (2012).
- 109 van Niftrik, L. A. *et al.* The anammoxosome: an intracytoplasmic compartment in anammox bacteria.
 FEMS Microbiol Lett 233, 7-13 (2004).
- 110 Jahn, M. T. *et al.* Shedding light on cell compartmentation in the candidate phylum Poribacteria by
 high resolution visualisation and transcriptional profiling. *Sci Rep* 6, 35860 (2016).
- **708** 111 Fuerst, J. A. Intracellular compartmentation in Planctomycetes. *Annu Rev Microbiol*, 299-328 (2005).
- 709 112 Katayama, T. *et al.* Membrane-bounded nucleoid discovered in a cultivated bacterium of the
 710 candidate phylum 'Atribacteria'. *bioRxiv*, 728279 (2019).
- 711 113 Borgnia, M. J., Subramaniam, S. & Milne, J. L. Three-dimensional imaging of the highly bent
 712 architecture of Bdellovibrio bacteriovorus by using cryo-electron tomography. *J Bacteriol* 190, 2588713 2596 (2008).
- 714 114 Remis, J. P. *et al.* Bacterial social networks: structure and composition of Myxococcus xanthus outer
 715 membrane vesicle chains. *Environ Microbiol* 16, 598-610 (2014).
- 716 115 Naor, A., Lapierre, P., Mevarech, M., Papke, R. T. & Gophna, U. Low species barriers in halophilic
 717 archaea and the formation of recombinant hybrids. *Curr Biol* 22, 1444-1448 (2012).
- 718 116 Nudleman, E., Wall, D. & Kaiser, D. Cell-to-cell transfer of bacterial outer membrane lipoproteins.
 719 Science 309, 125-127 (2005).
- 117 Cao, P. & Wall, D. Direct visualization of a molecular handshake that governs kin recognition and
 tissue formation in myxobacteria. *Nat Commun* 10, 3073 (2019).
- 118 Jakobczak, B., Keilberg, D., Wuichet, K. & Sogaard-Andersen, L. Contact- and protein transfer dependent stimulation of assembly of the gliding motility machinery in *Myxococcus xanthus*. *PLoS Genet* 11, e1005341 (2015).
- 119 Wolgemuth, C. W. & Oster, G. The junctional pore complex and the propulsion of bacterial cells. J
 Mol Microbiol Biotechnol 7, 72-77 (2004).
- 120 Nan, B. & Zusman, D. R. Uncovering the mystery of gliding motility in the myxobacteria. *Annu Rev Genet* 45, 21-39 (2011).
- 729 121 Munoz-Dorado, J., Marcos-Torres, F. J., Garcia-Bravo, E., Moraleda-Munoz, A. & Perez, J.
- 730 Myxobacteria: Moving, Killing, Feeding, and Surviving Together. *Frontiers in microbiology* 7, 781 (2016).
- 732 122 Patron, N. J. & Waller, R. F. Transit peptide diversity and divergence: A global analysis of plastid targeting signals. *Bioessays* 29, 1048-1058 (2007).
- Rogozin, I. B., Carmel, L., Csuros, M. & Koonin, E. V. Origin and evolution of spliceosomal introns. *Biol Direct* 7, 11 (2012).

- 736 124 Catania, F., Gao, X. & Scofield, D. G. Endogenous mechanisms for the origins of spliceosomal introns.
 737 *J Hered* (2009).
- 738 125 Vosseberg, J. & Snel, B. Domestication of self-splicing introns during eukaryogenesis: the rise of the
 739 complex spliceosomal machinery. *Biol Direct* 12, 30 (2017).
- 740 126 Martin, W. & Koonin, E. V. Introns and the origin of nucleus-cytosol compartmentalization. *Nature*741 440, 41-45 (2006).
- 742 127 D'Angelo, M. A. Nuclear pore complexes as hubs for gene regulation. *Nucleus* 9, 142-148 (2018).
- 743 128 Peña, C., Hurt, E. & Panse, V. G. Eukaryotic ribosome assembly, transport and quality control. *Nature* 744 *Structural & Amp; Molecular Biology* 24, 689 (2017).
- 745 129 Feng, J. M., Tian, H. F. & Wen, J. F. Origin and evolution of the eukaryotic SSU processome revealed
 746 by a comprehensive genomic analysis and implications for the origin of the nucleolus. *Genome Biol*747 *Evol* 5, 2255-2267 (2013).
- 748 130 Rivera, M. C., Jain, R., Moore, J. E. & Lake, J. A. Genomic evidence for two functionally distinct gene classes. *Proc. Natl. Acad. Sci. USA* 95, 6239-6244 (1998).
- 750 131 Pisani, D., Cotton, J. A. & McInerney, J. O. Supertrees disentangle the chimerical origin of eukaryotic
 751 genomes. *Molecular biology and evolution* 24, 1752-1760 (2007).
- 752 132 Gabaldon, T. & Huynen, M. A. From endosymbiont to host-controlled organelle: the hijacking of
 753 mitochondrial protein synthesis and metabolism. *PLoS Comput Biol* 3, e219 (2007).
- 754 133 Gabaldon, T. Relative timing of mitochondrial endosymbiosis and the "pre-mitochondrial symbioses"
 755 hypothesis. *IUBMB Life* 70, 1188-1196 (2018).
- 134 Ku, C. *et al.* Endosymbiotic gene transfer from prokaryotic pangenomes: Inherited chimerism in eukaryotes. *Proc Natl Acad Sci U S A* 112, 10139-10146 (2015).
- 135 López-García, P., Zivanovic, Y., Deschamps, P. & Moreira, D. Bacterial gene import and mesophilic
 adaptation in archaea. *Nat Rev Microbiol* 13, 447-456 (2015).
- 760 136 Larkum, A. W., Lockhart, P. J. & Howe, C. J. Shopping for plastids. *Trends Plant Sci* 12, 189-195 (2007).
- 761 137 Philippe, H. *et al.* Comparison of molecular and paleontological data in diatoms suggests a major gap
 762 in the fossil record. *J Evol Biol* 7, 247-265 (1994).
- 763 138 Zhu, S. Evidence for myxobacterial origin of eukaryotic defensins. *Immunogenetics* 59, 949-954
 764 (2007).
- 765 139 Perez, J., Castaneda-Garcia, A., Jenke-Kodama, H., Muller, R. & Munoz-Dorado, J. Eukaryotic-like
 766 protein kinases in the prokaryotes and the myxobacterial kinome. *Proc Natl Acad Sci U S A* 105,
 767 15950-15955 (2008).
- 768 140 Kerk, D., Uhrig, R. G. & Moorhead, G. B. Bacterial-like PPP protein phosphatases: novel sequence
 769 alterations in pathogenic eukaryotes and peculiar features of bacterial sequence similarity. *Plant*770 *Signal Behav* 8, e27365 (2013).
- 141 Elias-Arnanz, M., Padmanabhan, S. & Murillo, F. J. The regulatory action of the myxobacterial
 CarD/CarG complex: a bacterial enhanceosome? *FEMS Microbiol Rev* 34, 764-778 (2010).
- 142 Bock, T., Kasten, J., Muller, R. & Blankenfeldt, W. Crystal Structure of the HMG-CoA Synthase MvaS
 from the Gram-Negative Bacterium Myxococcus xanthus. *Chembiochem* 17, 1257-1262 (2016).
- 143 Osborn, A. R. *et al.* Evolution and Distribution of C7-Cyclitol Synthases in Prokaryotes and Eukaryotes.
 ACS Chem Biol 12, 979-988 (2017).
- 144 Pereto, J., Lopez-Garcia, P. & Moreira, D. Phylogenetic analysis of eukaryotic thiolases suggests
 multiple proteobacterial origins. *J Mol Evol* 61, 65-74 (2005).
- 145 Schluter, A., Ruiz-Trillo, I. & Pujol, A. Phylogenomic evidence for a myxococcal contribution to the
 mitochondrial fatty acid beta-oxidation. *PloS one* 6, e21989 (2011).
- 146 Stairs, C. W., Leger, M. M. & Roger, A. J. Diversity and origins of anaerobic metabolism in
 mitochondria and related organelles. *Philos Trans R Soc Lond B Biol Sci* 370, 20140326 (2015).
- 783 147 Baum, D. A. & Baum, B. An inside-out origin for the eukaryotic cell. BMC Biol 12, 76 (2014).

- 784 148 Sousa, F. L., Neukirchen, S., Allen, J. F., Lane, N. & Martin, W. F. Lokiarchaeon is hydrogen dependent.
 785 Nat Microbiol 1, 16034 (2016).
- 786 149 Searcy, D. G. in *The origin and evolution of the cell* (eds H. Hartman & K. Matsuno) 47-78 (World
 787 Scientific, 1992).
- 788 150 Searcy, D. G. Metabolic integration during the evolutionary origin of mitochondria. *Cell Res* 13, 229789 238 (2003).
- fould, S. B., Garg, S. G. & Martin, W. F. Bacterial vesicle secretion and the evolutionary origin of the
 eukaryotic endomembrane system. *Trends Microbiol* 24, 525-534 (2016).
- T92 152 Embley, T. M. & Martin, W. Eukaryotic evolution, changes and challenges. *Nature* 440, 623-630 (2006).
- Field, M. C. & Rout, M. P. Pore timing: the evolutionary origins of the nucleus and nuclear pore
 complex. *F1000Res* 8 (2019).
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802 Author contributions

- 803 P.L.-G. and D.M. conceived and discussed the ideas presented in the manuscript. P.L.-G. wrote the
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- 805

806 Competing interests

807 The author declares no competing financial interests.

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812 Figure Legend

Figure 1 | Environmental context, metabolic interactions and (endo)membrane evolution during 813 814 eukaryogenesis according to the HS-Syntrophy hypothesis. a, eukaryogenesis took place in phototrophic 815 microbial mats where steep redox gradients occur. Syntrophic interactions based on interspecies 816 hydrogen and/or sulfur transfer are widespread depending on local physicochemistry; they notably 817 involve methanogens, Asgard archaea, sulfate-reducing bacteria (SRB) and sulfide-oxidizing bacteria 818 (SOB). b-f, different eukaryogenesis steps. b, initial facultative symbiosis stage involving a hydrogen-819 producing Asgard archaeon, a sulfate-reducing deltaproteobacterium and a sulfide-oxidizing 820 alphaproteobacterium possibly able to carry out oxygenic photosynthesis. c, first integration of the 821 Asgard archaeon as an endosymbiont (future nucleus). d, second integration step involving the 822 endosymbiosis of the alphaproteobacterium. Stages c and d might have been coetaneous (first eukaryotic 823 common ancestor - FECA - stage) or, more likely, decoupled in time (FECA 1 and 2). e, advanced 824 integration stage involving important changes in metabolism (the consortium relies on aerobic 825 respiration, all other previous metabolic interactions between partners being lost) and endomembrane 826 evolution. f, last eukaryotic common ancestor stage (LECA). The position of H_2 or any other substrate by 827 an arrow (over or under) implies transfer in the sense of the arrow; when it is on two arrows of opposed 828 directionality, transfer may occur either way. HGT, horizontal gene transfer; EGT, endosymbiotic gene 829 transfer; aa, amino acids; ER, endoplasmic reticulum; conc., concentration; Ox., oxygenic; Anox., 830 anoxygenic; SRB, sulfate-reducing bacteria; SOB, sulfide-oxidizing bacteria; FECA, first eukaryotic common 831 ancestor; LECA, last eukaryotic common ancestor; hv, photon-derived energy.

833 Box 1 | Symbiogenetic models for the origin of eukaryotes based on metabolic exchange.

A variety of models propose that the eukaryotic cell evolved from a metabolic symbiosis (or syntrophy) 834 835 established between archaeal and bacterial cells in anoxic or microoxic environments. Most of them 836 involve only two partners, one archaeon and the alphaproteobacterial ancestor of mitochondria. 837 However, some models invoke the participation of one additional bacterium, either transiently, as facilitator of the eukaryogenetic symbiosis¹⁸, or as an integral part of it^{19,26}. One of the oldest proposals 838 based on explicit syntrophy was that of D. Searcy, who stated that eukaryotes derived from a sulfur-839 mediated symbiosis between a wall-less, sulfur-respiring Thermoplasma-like archaeon and photo- or 840 chemoautotrophic H₂S-utilizing bacterium^{149,150}. The original Hydrogen hypothesis postulated a hydrogen-841 842 mediated symbiosis between a hydrogenoclastic methanogenic archaeon and a hydrogen-producing alphaproteobacterium, hydrogen being used to reduce the CO₂ also released by the bacterium for 843 methanogenesis²⁵. In a more recent version of the Hydrogen hypothesis, the initial methanogenic host 844 was abandoned in favor of an autotrophic, non-methanogenic, archaeon that would use the Wood-845 Ljungdahl pathway to fix carbon using the hydrogen released by the mitochondrial ancestor¹⁴⁸. Based on 846 the inferred ancestral metabolism of Asgard archaea, which likely were organoheterotrophs with flexible 847 848 potential for hydrogen consumption and production, Spang and co-workers put forward the Reverse Flow 849 model. In this model, the eukaryogenetic syntrophy was based on hydrogen transfer (or electrons, i.e. 850 reducing equivalents, which might be also mediated by formate or acetate) from the anaerobic 851 heterotrophic archaeon to the alphaproteobacterium¹⁶. The recent Entangle-Engulf-Enslave $- E^3 \mod^{18}$ favors a dual symbiosis of an Asgard archaeon that degraded amino acids to short-chain fatty acids and 852 853 hydrogen with a sulfate-reducing bacterium (SRB) and an aerobic organotrophic alphaproteobacterium in 854 microoxic environments that scavenged toxic O_2 . As the consortium progresses towards increasingly oxic zones, the interaction with the alphaproteobacterium becomes stronger until it is engulfed. The SRB 855 symbiosis is transient and eventually lost¹⁸. Finally, the original version of the Syntrophy hypothesis (HM-856 857 Syntrophy) postulated a tripartite integrative symbiosis. First, a syntrophy based on interspecies H₂-858 transfer was established between a fermentative, ancestrally sulfate-reducing, myxobacterium 859 (Deltaproteobacteria) and a methanogenic archaeon using the fermentation-derived hydrogen for methanogenesis. Subsequently, a metabolically versatile alphaproteobacterium able to carry out 860 facultative aerobic respiration but also to oxidize methane (methanotroph) incorporated stably into the 861 consortium^{19,26}. In the revised variant of the Syntrophy model (HS-Syntrophy model), we hypothesize a 862 symbiosis between a hydrogen-releasing Asgard archaeon able to degrade small organics and a complex, 863 864 myxobacterial-like, deltaproteobacterial host scavenging hydrogen (reducing equivalents) for sulfate 865 reduction. The alphaproteobacterial ancestor of mitochondria was a sulfide-oxidizing facultative aerobe, 866 recycling sulfur in the consortium. Possibly, it was also a mixotrophic organism able to carry out 867 anoxygenic photosynthesis using H₂S as electron donor.

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872 Box 2 | Symbiogenetic models for the origin of eukaryotes according to the timing and mode of evolution of 873 key eukaryotic traits.

874 Regardless the metabolic basis of the symbiosis established between the Asgard archaeon and its 875 bacterial partner(s) during eukaryogenesis, models differ in the proposed mechanisms that resulted in the 876 physical integration of two or more cells in one (future eukaryotic cell) and the evolution of typical 877 eukaryotic traits as well as in the relative timing of the involved events (a-e). The Syntrophy hypothesis is 878 the only model where a membrane transition is not needed, since the host (future cytoplasm) is a 879 deltaproteobacterium naturally endowed with bacterial phospholipids (a). The future nucleus has an early 880 origin in this model; it would derive from a distinct metabolic compartment (endosymbiotic archaeon) that is progressively confined by a host-derived secretory membrane (Med., medium). Mitochondria 881 882 appear relatively late. In the Hydrogen hypothesis²⁵, the endosymbiosis of the alphaproteobacterial ancestor of mitochondria within an archaeon by means independent of classical, eukaryotic-like 883 884 phagocytosis (Ek-phagocytosis* in the figure) is the starting event triggering eukaryogenesis (b). The nucleus and the associated membrane system would form de novo from lipid vesicles produced by the 885 alphaproteobacterium^{126,151}. The archaeal membrane phospholipids would have been fully replaced by 886 the bacterial phospholipids by the fusion of those bacterial vesicles with the outer plasma membrane and 887 the progressive displacement of archaeal lipids¹⁵¹. In the currently most widely accepted type of 888 eukaryogenetic models, which include the 'phagocytosing archaeon'⁸¹ hypothesis (**c**) and the 'reverse 889 flow' model (d), the development of a complex cytoskeleton and endomembrane system predate the 890 acquisition of the mitochondrial ancestor by classical phagocytosis^{6,9,16}. Although these two models show 891 similarities, in the 'phagocytosing archaeon' model the nucleus would appear before the mitochondrial 892 acquisition, in contrast with the 'reverse flow' model . At any rate, in these models (c, d) the 893 894 mitochondrial endosymbiosis would be the consequence of an already well engaged eukaryogenetic 895 process. From this perspective, these models represent the transposition of past scenarios based on the existence of a proto-eukaryotic lineage different from archaea and bacteria endowed with all typical 896 eukaryotic traits but mitochondria^{4,5,79,152} to a nucleus-lacking proto-eukaryotic Asgard archaeon that is 897 already seen as the first eukaryotic common ancestor (FECA)^{6,153}. These two models and the hydrogen 898 899 hypothesis clearly differ in the timing of the mitochondrial acquisition and the endomembrane system but, in the three cases, the archaeal membrane phospholipids are replaced relatively late by bacterial-900 type phospholipids (**b-d**). In another set of models, including the inside-out¹⁴⁷ and the E³ hypotheses, the 901 mitochondrial ancestor is acquired relatively late by a slow process of engulfment. This involves archaeal 902 903 membrane extrusions that progressively surround, entangle and end up by enslaving the future 904 mitochondria (e). At least in the inside-out model, the archaeal-to-bacterial membrane phospholipid transition would occur relatively early, facilitated by close cell-cell contact and symbiotic gene transfer. 905 906 According to this model, only bacterial membranes, much more flexible than the archaeal ones, would be able to form an endomembrane system and carry out phagocytosis¹⁴⁷. Finally, in the Serial Endosymbiosis 907 model, several sequential symbioses intervene^{75,133}. Although the details remain undetermined, the 908 mitochondrion would be acquired late (f). 909 910

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915 Table 1 | Key open questions and possible ways of progress to discriminate or refine current

916 symbiogenetic models of eukaryogenesis

	Test - Means of obtaining answers	Models favored or disfavored		
Predictions of the HS-syntrophy hypothesis				
Existence of a versatile SRB deltaproteobacterial lineage, possibly sharing complex traits in common with myxobacteria, closer to eukaryotes	Explore microbial ecosystems relevant for eukaryogenesis (sediments, microbial mats) in search for novel deltaproteobacterial lineages followed by phylogenomic analyses	The detection of deltaproteobacterial lineages sharing a common and stronger phylogenetic signal with eukaryotes as compared to other bacteria would be consistent with the HS-Syntrophy model		
Existence of an alphaproteobacterial lineage of versatile S-oxidizers, perhaps photosynthetic, closer to mitochondria	Explore microbial ecosystems relevant for eukaryogenesis in search for novel alphaproteobacterial lineages followed by phylogenomic analyses	The detection of sulfide-oxidizing, potentially photosynthetic, alphaproteobacterial lineages sharing a common phylogenetic signal with eukaryotes to the exxclusion of other bacteria would be consistent with the HS-Syntrophy model		
A large fraction of bacterial genes in eukaryotes predates the mitochondrial endosymbiosis and derives from Deltaproteobacteria	Improve phylogenetic analyses of bacterial genes across eukaryotes, particularly those present in LECA	The presence, function and relative amount of deltaproteobacterial-like genes in eukaryotes as compared to other bacterial-like genes might support the involvement of a deltaproteobacterial symbiont in eukaryogenesis		
Deltaproteobacterial genes mostly relate to membrane, cell signaling and cytoplasm functions	Improve phylogenetic analyses and functional annotation of deltaproteobacterial-like genes in eukaryotes	The involvement of deltaproteobacterial genes in membrane, cell signaling and cytoplasmic functions would be supportive of the HS- Syntrophy model		
Bacterial genes widespread in eukaryotes (present in LECA) largely derive from EGT, not HGT	Improve phylogenetic analyses of bacterial genes present in LECA and look for potential homologues in Asgard archaea and in the closest alphaproteobacterial ancestors of mitochondria	If those bacterial-like genes in eukaryotes are missing in Asgard archaea or in Alphaproteobacteria, or the potential homologues are more distantly related than genes from other prokaryotic lineages, models invoking bacterial symbioses prior to the mitochondrial symbiosis would be favored		
More general questions / problems				
Do endosymbiotic prokaryotes exist in free-living prokaryotes? Do endosymbiotic archaea exist within bacteria?	Look for potential prokaryotic endosymbionts in anoxic/redox-transition ecosystems such as sediments or microbial mats.	If prokaryotic endosymbionts occur within prokaryotes, models proposing early prokaryotic endosymbionts would be as favored as models for which eukaryotic-like phagocytosis is a prerequisite. Finding archaeal endosymbionts within bacteria would relieve constraints for models proposing the endosymbiosis of one archaeon within a bacterium during eukaryogenesis		
Mitachandria: original matabalism and timi				
What was the metabolism of the alphaproteobacterial mitochondrial ancestor like? Did it have genes from other bacteria?	Explore microbial ecosystems in search for novel alphaproteobacterial lineages closely related to the mitochondrial lineage	If the closest alphaproteobacteria to the mitochondrion are identified, they might provide clues about the metabolic properties of the mitochondrial ancestor, potentially favoring specific eukaryogenesis models. If non-alphaproteobacterial genes in eukaryotes can be mapped back to these alphaproteobacteria, the origin of those genes would be more easily explained by HGT to the mitochondrial ancestor from other bacteria		
Will the inclusion of more bacterial and archaeal genomes potentially more closely related to eukaryotes lead to the discovery of genes displaying similarly long branches in phylogenetic trees as compared to alphaproteobacterial genes?	Enrich the taxonomic sampling of Asgard archaea and bacteria having close homologues in eukaryotic genomes	The discovery of an Asgard and/or bacterial lineage closer to eukaryotes and displaying branches of equivalent length to that of alphaproteobacterial-like genes in eukaryotes in phylogenetic trees might imply a simultaneous or temporally close symbiotic interaction of archaea and/or other bacteria during eukaryogenesis		
Urigin and nature of eukaryotic membranes				
Can bacteria expressing archaeal phospholipids be stably maintained?	Carry out experiments progressively expressing more archaeal phospholipids in bacteria until the complete replacement of bacterial phospholipds (eventually knocking out bacterial phospholipid synthesis genes)	If bacteria bearing membranes where bacterial phospholipids have been fully replaced by archaeal phospholipids can be experimentally produced, a bacterial-to-archaeal membrane transition would have been historically feasible		
If so, what is the fitness cost?	Study fitness of bacteria with archaeal phospholipids in their membrane in long-term experiments with and without competition with wild/other strains and as a function of environmental conditions	If fitness decreases and if bacteria cannot compete under any tested environmental conditions with wild strains and/or other bacteria, a membrane transition would have been historically unlikely		
Does the whole proteome evolve?	Study how the proteome change in eperimental evolution as a function of archaeal phospholipid content in bacterial membranes	If important changes in the proteome are observed, this imposes constraints for models invoking a membrane-transition		
Can we answer in the same way to the three previous questions for archaea expressing bacterial phospholipids?	Engineer archaeal cells with bacterial phospholipids and carry out similar experiments as described above for bacteria expressing archaeal phospholipids	If if archaea bearing membranes with only bacterial phospholipids can be produced, an archaeal-to-bacterial type membrane transition could have been historically feasible. If not, the fitness is too high for archaea to compete with wild-type archaea and/or in natural environments and the proteome is significantly affected, an archaeal-to-bacterial type membrane transition would have been historically unlikely		

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Figure Box 1. Lopez-Garcia & Moreira (R2)



Figure Box 2. Lopez-Garcia & Moreira (R2)



Figure 1. Lopez-Garcia & Moreira