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3	A eukaryote-wide perspective on the diversity and evolution of the ARF GTPase protein
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#### 35 Abstract

The evolution of eukaryotic cellular complexity is interwoven with the extensive 36 37 diversification of many protein families. One key family is the ARF GTPases that act in 38 eukaryote-specific processes, including membrane traffic, tubulin assembly, actin dynamics, 39 and cilia-related functions. Unfortunately, our understanding of the evolution of this family is limited. Sampling an extensive set of available genome and transcriptome sequences, we 40 41 have assembled a dataset of over 2,000 manually curated ARF family genes from 114 42 eukaryotic species, including many deeply diverged protist lineages, and carried out 43 comprehensive molecular phylogenetic analyses. These reconstructed as many as 16 ARF 44 family members present in the last eukaryotic common ancestor (LECA), nearly doubling the 45 previously inferred ancient system complexity. Evidence for the wide occurrence and 46 ancestral origin of Arf6, Arl13 and Arl16 is presented for the first time. Moreover, Arl17, 47 Arl18 and SarB, newly described here, are absent from well-studied model organisms and as 48 a result their function(s) remain unknown. Analyses of our dataset revealed a previously unsuspected diversity of membrane association modes and domain architectures within the 49 50 ARF family. We detail the step-wise expansion of the ARF family in the metazoan lineage, 51 including discovery of several new animal-specific family members. Delving back to its 52 earliest evolution in eukaryotes, the resolved relationship observed between the ARF family 53 paralogs sets boundaries for scenarios of vesicle coat origins during eukaryogenesis. 54 Altogether, our work fundamentally broadens the understanding of the diversity and 55 evolution of a protein family underpinning the structural and functional complexity of the 56 eukaryote cells.

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59 Key words: ARF family, eukaryotic cell, evolution, GTPases, last eukaryotic common60 ancestor, post-translational modifications

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## 62 Significance

ARF Family GTPases are crucial regulations of a diversity of cellular compartments and
processes and as such the extent of this system in eukaryotes reflects both cellular complexity
in modern eukaryotes and its evolution. Strikingly, a comprehensive comparative genomic
analysis of the protein family is lacking, leaving its recent and ancient evolution poorly
resolved. We performed a comprehensive molecular evolutionary analysis, reconstructing a
highly complex ARF family complement in the Last Eukaryotic Common Ancestor,

69 including a number of paralogs never before identified as such, and we find resolved
70 relationships between the paralogs. This work has implications for cellular evolution from
71 eukaryogenesis to cellular complexity in metazoans.

72

#### 73 Introduction

74 Understanding how the eukaryotic cell evolved in all its complexity is one of the greatest 75 open questions in evolutionary biology. Eukaryogenesis involved both the origin of new 76 genes and the diversification of key building blocks (Dacks et al. 2016; Eme et al. 2017). 77 Among the different building blocks, particular groups of proteins radiated early in the evolution of eukaryotes and are represented by a large number of pan-eukaryotic orthologs, 78 79 presumably with conserved functions. One of the largest groups of proteins, acting in an 80 incredibly diverse array of cellular pathways, is the Ras superfamily of GTPases. This 81 superfamily is frequently equated with familiar and extensively studied eukaryotic "small 82 GTPases". However, the more appropriate, i.e. evolutionary, definition conceives it as a 83 major monophyletic subgroup of the vast TRAFAC class of GTPases that also includes 84 prokaryotic representatives, larger proteins combining a Ras-related GTPase domain with 85 other functional domains, and – surprisingly to many in the field – the alpha subunits of 86 heterotrimeric G-proteins (Leipe et al. 2002). Because of its central role in so many 87 fundamental cellular functions, understanding the origin and evolution of this complex 88 superfamily of proteins is necessary for uncovering the processes by which eukaryotes 89 evolved and diversified.

90 The internal classification of the Ras superfamily is unsettled. In many overviews, 91 especially those concentrating on the eukaryotic small GTPases, the content of the 92 superfamily is pigeonholed into five major families (Ras, Rho, Rab, Ran, Arf/Sar; Colicelli 93 2004; Rojas et al. 2012), but this scheme ignores the prokaryotic superfamily members 94 (Wuichet and Søgaard-Andersen 2014), multi-domain proteins (such as the ROCO family; 95 Bosgraaf and Van Haastert 2003), and various other lineages clearly distinct from or not 96 easily classified into the well known families, such as the Gtr/Rag family (Klinger, Spang et 97 al. 2016) or RJL proteins (Elias and Archibald 2009). Understanding the diversity and the evolutionary origin of the Ras superfamily in eukaryotes is a challenging task, given the 98 99 presence of tens to hundreds of Ras superfamily genes in each extant eukaryote genome 100 (Rojas et al. 2012). Disregarding potential (presently unknown) cases of horizontal gene 101 transfer from prokaryotic sources into particular eukaryote lineages, the wealth of Ras 102 superfamily genes in eukaryotes ultimately derives from a set of genes present in the Last

103 Eukaryote Common Ancestor (LECA). Several evolutionary analyses have attempted to 104 reconstruct LECA's complement of particular Ras superfamily subgroups and detail the 105 downstream innovation within eukaryotes. Prominent examples include analyses of the Rab 106 (Diekmann et al. 2011; Elias et al. 2012; Klöpper et al. 2012) and Ras families (van Dam et 107 al. 2011), and some isolated lineages like RJL (Elias and Archibald 2009), Miro (Vlahou et 108 al. 2011), or RABL2 (Eliáš et al. 2016). These investigations demonstrated that a large 109 number of functionally investigated paralogs were present in the LECA, emphasizing the role of loss or streamlining of genomic complement in many eukaryotic lineages. They also 110 111 identified ancient LECA paralogs of unknown function that have been lost in lineages leading 112 to conventional model systems but which are present in diverse eukaryotic lineages of 113 ecological and medical importance. Paralogs with such an evolutionary distribution were 114 recently coined jotnarlogs (More et al. 2020). Finally, these studies also inevitably shed light 115 on the diversification of GTPases in the post-LECA expansion phase. For example, divergent paralogs of unclear evolutionary relationships are found in various taxa (e.g., Pereira-Leal 116 117 2008), most likely resulting from rapid sequence evolution of lineage-specific paralogs linked 118 to their neofunctionalization. Additionally, the inherently small nature of the GTPases makes 119 them particularly susceptible to molecular tinkering, such as accretion of additional domains 120 or gain/loss of motifs mediating specific post-translational modifications (e.g., Záhonová et 121 al. 2018).

122 Not yet addressed in a comparable evolutionary framework is the ARF protein family. This large protein family is comprised of the "true" ADP Ribosylation Factors (i.e., Arfs), as 123 124 well as Arf-like proteins (Arls), Arf-related protein 1 (Arfrp1), and Sar1. Clearly related are 125 the beta subunits of the signal recognition particle receptor (SRβ; Schwartz and Blobel 2003). 126 Sequence analyses have also revealed that an Arf-like ancestor, modified by insertion of a 127 novel  $\alpha$ -helical region into its GTPase domain and high sequence divergence, gave rise to the 128 alpha subunits of heterotrimeric G-proteins (abbreviated Ga; Neuwald 2007; Anantharaman 129 et al. 2011). The distinction between Arf and Arf-like (Arl) proteins was originally made 130 based upon activity in the cholera toxin-catalyzed ADP-ribosylation of the stimulator of 131 adenylyl cyclase,  $G\alpha_s$ , as all tested Arfs retain this functionality while the Arls did not 132 (Tamkun et al. 1991; Clark et al. 1993). However, this activity has proven of very limited 133 utility in studies of cellular functions for ARF family members as greater appreciation of both 134 the size of the family in model organisms as well as the diversity of functions became clear. 135 Thus, little if any weight should be given to whether a gene is named as an Arf, an Arl, an Arfrp1, or a Sar. The ARF family is functionally heterogeneous and comprises proteins 136

137 involved in membrane vesicle formation (Arfs, Sar1), other aspects of vesicle traffic and maintenance of membranous organelle morphology (e.g., Arl1, Arl5, or Arfrp1), microtubule 138 139 dynamics and mitochondrial fusion (Arl2), and cilium biogenesis and function (Arl3, Arl6, 140 Arl13) (Gillingham and Munro 2007; Donaldson and Jackson 2011; Francis et al. 2016). 141 Members of this family are critical to these diverse cellular activities and dysfunction results 142 in numerous human diseases. Family members are generally considered to be single-domain 143 small GTPases. Post-translational modifications (N-terminal myristoylation or acetylation) are also often critical to the protein's localization and function. 144

145 An early phylogenetic study on the ARF family, limited by a lack of taxonomic 146 breadth in available genomic sequences, provided an early estimate of the ancient complexity 147 of the family in LECA and identified putative lineage-specific expansions in metazoans (Li et 148 al. 2004). The analyses showed that LECA contained at least eight ancient groups of orthologs inferred from representatives being present in metazoans and at least one non-149 150 opisthokont (protist or plant) eukaryote. This analysis also demonstrated that some of the 151 metazoan family members lacked close relatives in other eukaryotes, suggesting that lineage-152 specific expansions related to metazoan multicellularity occurred. Perhaps most familiar is 153 expansion yielding the well-known and founding members of the family, Arfs 1-5. These 154 have been shown as deriving from a single ancestral gene (here referred to as Arf1 for 155 simplicity) which duplicated prior to choanoflagellates, yielding Arfs 1-3 (sometimes named 156 Class I Arfs but for convention referred to here as Arf1) and Arfs 4-5 (sometimes named 157 Class II Arfs but for convention referred to here as Arf4), with each of those diversifying into 158 five Arf paralogs around the whole genome duplications in the vertebrate lineage (Manolea et 159 al. 2010). However, since these early studies, several family members from the target species 160 (including humans) have been identified (Kahn et al. 2006) and methods of phylogenetic 161 analyses of protein sequences have advanced, including the development of the ScrollSaw 162 approach facilitating analyses of complex paralog-rich families (Elias et al. 2012). Thus, the 163 time is ripe for obtaining a much better picture of the evolution of ARF family than in the previous studies. 164

To this end, we assembled, extensively curated and phylogenetically analysed a
dataset of ARF family sequences from a taxonomically broad selection of eukaryotic species.
This enabled us to revise the set of ancestral eukaryotic ARF family paralogs, which has now
expanded to between 14 and 16 genes. Two paralogs, described here for the first time, are not
represented in well studied models and point to hitherto unstudied molecular functions
mediated by the ARF family. We observed an unexpected diversity of domain architectures

- 171 challenging the dogma that ARF family proteins are only small and single-domain proteins.
- 172 Our analyses also unveiled a range of predicted post-translational modifications (PTMs),
- 173 including but not limited to well-established N-terminal myristoylation, and other molecular
- adaptations that facilitate membrane association as a central feature of ARF family biology.
- 175 Finally, we identified well supported relationships between the paralogs, which have
- 176 implications for the inferred function of the primordial family members during
- 177 eukaryogenesis.
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#### 179 Results and Discussion

#### 180 A comprehensive dataset and phylogeny of the ARF family

181 We first gathered all ARF family sequences (including SR $\beta$  but excluding the highly

divergent Gα proteins) from a broad diversity of eukaryotes, exploiting both publicly

available and privately curated genomes and transcriptomes. We did not rely solely on

184 predicted protein sequence sets but also checked the genome and transcriptome assemblies to

185 ensure maximal accuracy when it comes to statements about the absence of particular genes

- in different taxa. All sequences were carefully validated, as described under Materials and
- 187 Methods, and when needed, edited (by modifications of the originally predicted gene models

188 or by changes in the assembled nucleotide sequences based on inspection of raw sequencing

- data) to ensure maximal quality and completeness of the data. Our final dataset, provided as
- 190 supplementary dataset 1 (Supplementary Material online), included >2,000 manually curated
- 191 sequences from 114 species (supplementary table 1, Supplementary Material online). The
- 192 number of ARF family genes in individual species ranged from 5 in the yeast
- 193 Schizosaccharomyces pombe to 70 in the rotifer Adineta vaga (this high number apparently

194 reflecting the tetraploid origin of its genome; Flot et al. 2013).

195 The genes were initially annotated based on their similarity to previously 196 characterized or named ARF family genes in model organisms scored by BLAST. While this 197 procedure enabled us to recognize candidate groups of orthologs and to assign most of the 198 genes into these groups, the assignment of many sequences was uncertain or unclear and a 199 more rigorous method for establishing orthologous relationships – phylogenetic analysis of a 200 multiple sequence alignment – was required to corroborate the proposed groups of orthologs 201 and to possibly identify additional ones not readily apparent from sequence-similarity comparisons. Such an analysis of the whole dataset was impractical, if not impossible, for its 202 203 size and the existence of divergent sequences that tend to disrupt the results of phylogenetic 204 inference. We therefore utilized of the ScrollSaw protocol previously developed to deal with

205 a similarly complex family of Rab GTPases (Elias et al. 2012) and applied by others to 206 resolve deep relationships within protein families (e.g., Vosseberg et al. 2021). This protocol 207 enables one to infer a "backbone" phylogeny of a protein family by concentrating on 208 preselected sequences likely representing slowly-evolving members of the main clades of the 209 family conserved across distantly related organismal lineages. Briefly (see Materials and 210 Methods for details), we divided the sampled species into 13 groups corresponding to major 211 eukaryotic lineages, and for each pair of groups we identified all pairs of sequences (the two sequences representing the two different groups) that had mutually minimal genetic distances 212 213 calculated by the maximum likelihood method from a multiple sequence alignment. We then 214 gathered all the sequence pairs of all the comparisons, removed redundancies, and inferred 215 trees from the full resulting dataset (supplementary fig. 1, Supplementary Material online) or 216 after pruning sequences from selected species to further decrease the complexity of the analysis (fig. 1; supplementary fig. 2, Supplementary Material online). This resulted in a 217 taxonomically rich and generally well resolved final phylogeny, which enabled us to infer 218 219 various aspects about the evolutionary and diversity history of the ARF family in eukaryotes.

220

#### 221 LECA possessed an extensive array of ARF family paralogs

222 Dissection of the "ScrollSaw" trees indicated the existence of 13 potentially monophyletic 223 groups (Sar1 and SarB are counted as a single putative clade for the moment, see below). 224 Each group is represented by genes from all or a majority of the major eukaryote lineages, in 225 all cases spanning both putative principal clades of eukaryotes (Opimoda and Diphoda) 226 defined by the most recent hypothesis on the position of the root of the eukaryote phylogeny (Derelle et al. 2015). As such these groups all are candidates for separate ARF family 227 228 paralogs differentiated before the radiation of extant eukaryotes and perhaps present in the 229 LECA, provided that they are monophyletic (i.e. that the root of the ARF family tree lies 230 outside of them). Our trees are inherently unrooted due to the absence of a suitable outgroup, 231 as other GTPases, including the presumably most closely related group,  $SR\beta$ , are too 232 divergent and their inclusion into these analyses limits the resolution of the trees. Hence, to 233 formally rule out the possibility that the root lies in any of the 13 putative clades, we 234 employed the outgroup-independent minimal ancestor deviation (MAD) method (Tria et al. 235 2017), which placed the root onto a branch separating the Arl16 group from all other groups 236 combined (fig. 1). We also note that the rooting outside any of the 13 groups implies a much 237 simpler evolutionary scenario than a root positioned into any of the groups, so hereafter we 238 treat the 13 groups as clades. Most of them have high statistical support (posterior

239 probability, SH-aLRT support, and ultrafast bootstrap values greater than or equal to 0.98, 240 98, and 98, respectively) (fig. 1). An exception is the clade denoted Arf1 and comprising 241 prototypical Arf sequences, but there is little doubt that it constitutes a coherent group of 242 orthologs. The weak signal for its monophyly may stem from a very slow evolution of Arf1 243 sequences (apparent also from very short branches in the tree) having precluded 244 accumulation of paralog-specific sequence features that would enable strong phylogenetic 245 separation from the related, more rapidly evolving (and much more strongly supported) 246 paralogs. Nevertheless, a focused analysis restricted to Arf1, Arf6, Arl1 and Arl5 allowed us 247 to use a protein alignment with more positions and recovered Arf1 as a supported 248 monophyletic clade (supplementary fig. 3, Supplementary Material online).

249 The existence of two separate clades of Arfs originated before the divergence of 250 metazoans, fungi, and plants was hypothesized previously but not convincingly demonstrated 251 (Li et al. 2004). We show that mammalian Arf6 has robustly supported orthologs in various 252 protists spanning the phylogenetic breadth of eukaryotes. The existence of a separate eukaryotic Arf6 clade is further supported by comparison of intron positions in Arf genes 253 254 (supplementary fig. 4, Supplementary Material online). In contrast, as expected, the 255 mammalian Arf1-Arf5 proteins (class I and II Arfs) all cluster into the Arf1 clade. Our 256 analyses further demonstrate that the metazoan Arl16 has orthologs present in diverse protists 257 and thus represents a novel ancient ARF family paralog. Another previously unrecognized 258 ancient paralog, which we propose to call Arl18, was missed because it is not represented in 259 metazoans and has no characterized or named member. It is most closely related to Arl8, yet 260 the separation of Arl8 and Arl18 is apparent not only from the phylogenetic analysis (fig. 1; 261 supplementary fig. 2, Supplementary Material online) but also from their distinct exon-intron 262 structures (supplementary fig. 5, Supplementary Material online).

263 Two additional ancient eukaryotic ARF family paralogs seem to exist, although they 264 were not unambiguously supported by our phylogenetic analyses. The broader clade 265 including Sar1 proteins and their relatives has a somewhat unusual internal structure with a 266 strongly supported subclade, comprised of typical Sar1 proteins found in all taxa investigated, and a more basal paraphyletic group of proteins representing different Sar1-like 267 268 paralogs from phylogenetically diverse protist lineages (fig. 1; supplementary fig. 2, 269 Supplementary Material online). These are not simply divergent Sar1 orthologs, as they 270 always co-occur with a bona fide Sar1 in each species analyzed, and multiple lines of 271 evidence suggest they constitute a separate ancient paralog of their own, which we call SarB 272 (adopting the name proposed before for a respective *Dictyostelium discoideum* 

273 representative; Week et al. 2003). Specifically, some intron positions in SarB genes are 274 exclusive for this group and not shared with Sar1 (supplementary fig. 6, Supplementary 275 Material online) and the functionally important Walker B motif of SarB generally exhibits a 276 conserved tryptophan residue shared by other ARF family members and Ga proteins, as 277 opposed to a phenylalanine residue typical for Sar1 proteins (Vetter 2014; supplementary fig. 278 7, Supplementary Material online). Furthermore, a ML tree with SarB sequences constrained 279 to form a clade could not be rejected by AU test, as opposed to trees imposing topologies that 280 would correspond to the origin of SarB genes by multiple independent duplications of Sar1 281 genes proper (supplementary table 2, Supplementary Material online). Hence, it is most 282 parsimonious to interpret SarB as a bona fide ancient ARF family paralog different from 283 Sar1, with the phylogenetic signal for its monophyly virtually vanished over the eons. Such a 284 situation is not uncommon in phylogenetic analyses of families of short proteins with an 285 inherently limited phylogenetic signal. For instance, a similar behaviour was previously 286 observed with the highly conserved Rab1 GTPase paralog, whose undoubted monophyly was 287 also difficult to recover (Elias et al. 2012).

288 The second additional potential ancient paralog, here proposed to be called Arl17, is 289 present in various protists, certain fungi, and a single metazoan lineage, and its representative 290 contain one to three non-identical copies of a novel conserved domain C-terminal to the 291 GTPase domain (fig. 2; supplementary fig. 8, Supplementary Material online). The novel 292 ~100 residue, C-terminal domain displays no discernible homology to previously described 293 domains (even when tested by the highly sensitive HHpred searches), but occurs also in other (non-Arl17) proteins from some opisthokonts and bacteria, either as a stand-alone protein 294 295 (e.g., EGF92317.1) or in combination with various non-GTPase domains (e.g., 296 XP\_004347279.1). Despite their unique domain architecture, no Arl17 sequences passed the 297 ScrollSaw filter, hence they are absent from the tree presented in fig. 1, and although forming 298 a clade in phylogenetic analysis, statistical support for their monophyly is lacking (fig. 2). 299 Still, the most parsimonious interpretation of our analyses is that Arl17 is an ancient ARF 300 family GTPase that was present already in LECA and had evolved from a duplication of the Arf1 gene, but the tendency of the GTPase domains in Arl17 proteins to be very divergent 301 302 (supplementary fig. 9, Supplementary Material online) has weakened the signal for their

303 monophyly.

Having established the main lineages of the ARF family, we attempted to assign all other genes in our full dataset (i.e. those that were excluded by the ScrollSaw protocol) into them by considering sequence similarity scored by BLAST, comparison to lineage-specific 307 profile HMMs by HMMER, and by targeted phylogenetic analyses. The majority of genes in 308 our dataset could be allocated with confidence to a specific, ancient ARF family paralog, 309 enabling us to evaluate the pattern of retention of the ancient paralogs in modern eukaryotes 310 and to map the presumed gene losses to the eukaryote phylogeny (fig. 3; supplementary table 311 3, Supplementary Material online). Nevertheless, a relatively small number of genes (160 out 312 of > 2,000 sequences) remained unclassified. A majority of these likely correspond to taxon-313 specific duplications of the standard ARF family members that have diverged substantially, obscuring their actual evolutionary origin. Some cases, however, may represent excessively 314 315 divergent, unrecognized direct orthologs of the widespread genes. For example, several 316 unclassified genes showed potential affiliation to Arf6, yet without significant support in 317 phylogenetic analyses. These sequences all share one or more intron positions specific to 318 Arf6 (supplementary fig. 4, Supplementary Material online), supporting their annotation as 319 highly derived Arf6 genes. Future studies with a more comprehensive sampling may help 320 resolve cases such as these.

321

## 322 Complex cellular repertoire inferred from the LECA complement

323 The analyses presented above indicate that the LECA possessed at least 15 ARF family genes; Arf1 and 6, Arl1, 2, 3, 5, 6, 8, 13, 16, 17, and 18, Arfrp1, Sar1, and SarB. In addition, 324 325 it certainly encoded SR<sup>β</sup>, excluded from our ScrollSaw analysis (hence absent from the trees 326 in fig. 1 and supplementary fig. 2, Supplementary Material online) due to its marked 327 divergence from the (core) ARF family and because SR<sup>β</sup> orthologs can be unambiguously recognized by sequence similarity. Eight of these clades (Arf1, Arl1, 2, 3, 5, and 8, Arfrp1, 328 329 and Sar1) were previously recognized as likely ancient (Li et al. 2004) and the existence of 330 orthologs of the metazoan Arl13 in protists was also noted (e.g., Miertzschke et al. 2014), 331 although perhaps never documented by phylogenetic analyses. Our analysis thus indicates 332 that the complement of ARF family paralogs in LECA may have been twice as big as 333 previously identified, and further strengthens the idea that the LECA was a fully-fledged 334 eukaryotic cell making broad use of complex molecular machinery.

The cellular functions of many of the 16 ARF family GTPases in the LECA in principle can be considered from what has been learned about their descendants in modern eukaryotes, although our present knowledge about the function of various GTPases comes from a limited number of phylogenetically biased model eukaryotes (primarily metazoans and the yeast *Saccharomyces cerevisiae*, i.e. the opisthokonts) and it is not always certain to

340 what extent we can generalize from them to eukaryotes as a whole. In addition, each ARF 341 family member studied in any depth in mammalian cells has been found to act in more than 342 one pathway and typically with multiple downstream effectors (Kahn et al. 2009; Sztul et al. 343 2019), often making it difficult to assess which of these are ancient and which were acquired 344 later. Finally, we recognize that any inferences about ancient functional roles relies on an assumption of functional homology across eukaryotes and an assumption of parsimonious 345 346 retention of pleisiomorphic traits. From a large assessment of membrane-trafficking proteins 347 that have been tested in model systems from across the eukaryotic tree, this assumption of 348 functional homology appears to be justified (Klinger et al. 2016), but does warrant being 349 explicitly named. With this caveat in mind, we summarize the key findings about the 350 different paralogs to paint a hypothetical picture of the cellular engagement of the ARF 351 family members in the LECA.

352 Most of the ARF family paralogs clearly play a role in the endomembrane dynamics. 353 As a subunit of the receptor of the signal recognition particle, SR<sup>β</sup> mediates co-translational import of proteins into the ER (Schwartz and Blobel 2003). Sar1 also associates with the ER 354 355 and recruits subunits of the COPII coat complex to promote budding of transport vesicles from the ER (Miller and Barlowe 2010). Four paralogs – Arf1, Arfrp1, Arl1 and Arl5 – are 356 physically and functionally associated with the Golgi/*trans*-Golgi network (TGN). One key 357 358 function of Arf1 (including the metazoan Arf1 to Arf5) is to recruit different types of vesicle 359 coats (COPI, AP-1/clathrin, AP-3) to different parts of the Golgi (Jackson and Bouvet 2014). 360 Arl1 and Arfrp1 (confusingly called Arl3p in the yeast S. cerevisiae) are functionally linked, 361 the latter shown to be critical for Arl1 recruitment to the trans-Golgi in both yeast and 362 mammalian cells (Panic et al. 2003; Setty et al. 2003; Zahn et al. 2006). Arl1 recruits several 363 effectors (e.g. golgins, arfaptins, and Arf-GEFs) to the trans-Golgi network (TGN) and is important for endosome-to-TGN traffic (Yu and Lee 2017). The function of Arl5 is less-well 364 365 understood, but it may partly overlap with that of Arl1, as it also localizes to the *trans*-Golgi 366 (Houghton et al. 2012), and both the fly Arl5 and the yeast Arl1 each interact with the GARP tethering complex (Panic et al. 2003; Rosa-Ferreira et al. 2015). In contrast to the Golgi 367 368 localizing and acting members of the ARF family, Arf6 acts predominantly at the cell surface 369 and endosomes to mediate endosome recycling, cell motility, and membrane extensions, 370 which together influence cell division, lipid/cholesterol metabolism, and changes in actin 371 dynamics (D'Souza-Schorey and Chavrier 2006; Cotton et al. 2007; Funakoshi et al. 2011; 372 Schweitzer et al. 2011). Arl8 has been implicated in controlling lysosomal motility and traffic in metazoan cells (Khater et al. 2015). Its localization to the vacuolar membranes in *A*.

- thaliana (Heazlewood et al. 2007) suggests that functional association of Arl8 with the
- 375 lysosomal/vacuolar compartment is ancestral and conserved.

376 Three paralogs, Arl3, Arl6, and Arl13 have been implicated in flagellar function 377 (Fisher et al. 2020). Arl3 has been proposed to regulate the delivery of N-myristoylated and prenylated proteins to the cilium (Fansa and Wittinghofer 2016; Stephen and Ismail 2016). 378 379 Arl6 (also called BBS3) regulates the function of the BBSome (a protein complex involved in intraflagellar transport; Mourão et al. 2014). Arl13 is involved in ciliary protein import and 380 381 export, purportedly mediated by its activity as a positive regulator (guanine nucleotide 382 exchange factor, GEF) for Arl3 (Gotthardt et al. 2015; Ivanova et al. 2017). Arl2 shares some 383 effectors with Arl3 and is probably involved in traffic of lipidated proteins (Van Valkenburgh 384 et al. 2001; Fansa and Wittinghofer 2016), but it has its own specific agenda, as it regulates 385 the assembly of  $\alpha\beta$ -tubulin dimers (Al-Bassam 2017; Francis et al. 2017a; Francis et al. 2017b) and mitochondrial fusion (Newman et al. 2017). 386

Only a single study addressing the function of Arl16 has been published, reporting 387 388 that the mammalian Arl16 inhibits the function of the RIG-I protein, involved in the defence 389 against RNA viruses (Yang et al. 2011), but more specific functional insights are lacking. 390 Functions for of the newly discovered paralogs SarB, Arl17, and Arl18 are completely 391 unknown, as these paralogs are missing from all common model eukaryotes and thus 392 represent examples of "jotnarlogs", proteins that are present across eukaryotes, but missing in 393 well-studied cell biological models (More et al. 2020). This adds further credence to the 394 proposal that this is a substantial evolutionary cell biological phenomenon and highlights the gap in our understanding of the cell biology of the ARF family in eukaryotes. Nevertheless, 395 396 some clues as to the function of these proteins are provided by the phylogenetic relationship 397 to other paralogs, as relatedness within the ARF family appears to signify some level of 398 functional similarity, despite exceptions. Indeed, the aforementioned functional aspects 399 shared by the pairs Arl2-Arl3 and Arl1-Arl5 are reflected by close relationship of the 400 paralogs in the pairs (fig. 1). Likewise, the related Arf1 and Arf6 paralogs, although different 401 in terms of the intracellular localization and effectors they deploy (Jackson and Bouvet 402 2014), share the same class of GEFs and GTPase activating proteins (GAPs), though to a 403 very incompletely characterized extent (Casanova 2007; Kahn et al. 2008; Sztul et al. 2019). 404 Hence, by analogy we speculate that Arl18 may have similar functional attributes as its 405 closest paralog Arl8 (e.g. it may likewise function in the lysosomal/vacuolar sector of the 406 endomembrane system), and that SarB functions similarly to the canonical Sar1 protein in the

secretory pathway (Sato and Nakano 2007; Melville et al. 2020). The specific relationship of
Arl17 and true Arfs may be less informative concerning the function of the latter, given the
unique domain architecture of Arl17 proteins and the generally divergent nature of their
GTPase domains (compare the branch lengths of Arl17 sequences in the tree in fig. 2).

411

#### 412 Phylogenetic profiles of some ancestral eukaryotic ARF family paralogs illuminate

differential simplification of endomembrane system functions in eukaryote evolution
A detailed scrutiny of the taxonomic distribution of some of the ancestral ARF family
paralogs in extant eukaryotes provides interesting insights into the variation of their roles in
cell functions across eukaryotes. While a hallmark of the ARF family perhaps is that
members are commonly found to be active in multiple, distinct pathways in the same cells
(Francis et al. 2016; Sztul et al. 2019), here we discuss their known or predicted

419 functionalities with respect to their best known activities, recognizing the limitations that420 result.

421 Arfs (specifically the Arf1 paralog), Sar1, and SR $\beta$  are all found in every eukaryote sampled (with one exception in case of SR $\beta$ , most likely due to incompleteness of the data; 422 supplementary table 3, Supplementary Material online), indicating that they belong to the 423 424 functional core of the eukaryotic protein toolkit. Nearly ubiquitous is Arl2, being absent only 425 from Entamoeba histolytica. Inspection of genomes of other Entamoeba species suggest that 426 Arl2 loss is not an artefact and predates the radiation of the genus. Given the role of Arl2 in 427 the assembly of tubulin dimers and in mitochondrial fusion (Francis et al. 2016), its absence in Entamoeba may be related to a unique combination of traits of this taxon including 428 divergent tubulin sequences and a highly reduced microtubular cytoskeleton (Roy and Lohia 429 430 2004; Meza et al. 2006), and a simplified mitochondrion (i.e., a mitosome; Makiuchi and 431 Nozaki 2014).

432 Five of the ancestral paralogs functionally linked to the endomembrane system (based 433 on data from model eukaryotes) show various degrees of patchiness in their occurrence (fig. 434 3A; supplementary table 3, Supplementary Material online). Arl1, Arl5, and Arfrp1, all 435 associated with the Golgi apparatus, have been preserved in all main eukaryote lineages 436 sampled, but have been lost from some more terminal branches. Arl1 is missing from the fission yeast (S. pombe), diplomonads, and some apicomplexans. Arfrp1 is absent from the 437 438 same set of species plus two more (the highly reduced endosymbiotic kinetoplastid 439 Perkinsela sp. CCAP 1560/4 and the tiny green alga Micromonas commoda). The similar

440 patterns of loss of these two GTPases may reflect the fact that they were shown to work in 441 the same functional cascade (see above). How Arl1 functions in the absence of Arfrp1 in 442 *Perkinsela* or *Micromonas* remains an open question but may reflect the multiplicity of 443 pathways each GTPase may influence and the potential differences in their means of 444 localization and activation. Arl5 is missing from many more eukaryotes, including even some metazoans (e.g., the flatworm Schmidtea mediterranea). A minimum of 20 independent 445 446 losses of Arl5 is required to explain its distribution in our dataset (supplementary table 3, 447 Supplementary Material online), suggesting that this GTPase is a less critical component of 448 the basic infrastructure of the eukaryotic cell. In accord, disruption of the Arl5 gene in 449 Drosophila melanogaster does not alter the fly's viability or fertility (Rosa-Ferreira et al. 450 2015). Arl5 is closely related to Arl1 and the two GTPases may share some effectors (see 451 above). It is thus possible that Arl5 loss is facilitated by partial functional redundancy with 452 Arl1.Similar to Arl5, the distribution of Arl8 in extant eukaryotes has been shaped by 453 multiple (at least 14) independent losses, including one in the lineage leading to the main eukaryotic taxon Stramenopiles (fig. 3B; supplementary table 3, Supplementary Material 454 455 online). Comparison of phylogenetic profiles of Arl8 and the related uncharacterized paralog 456 Arl18 reveals that the former paralog has been retained more frequently than the latter, but in 457 a few taxa (e.g., stramenopiles) Arl18 occurs in the absence of Arl8 (fig. 3A; supplementary 458 table 3, Supplementary Material online). It would be interesting to investigate whether a level 459 of functional redundancy might allow Arl18 to have taken over some of the Arl8 functions in 460 these organisms. The presence of both Arl8 and Arl18 in model systems like *Tetrahymena* 461 thermophila and Trypanosoma cruzi (supplementary table 3, Supplementary Material online) provides a chance that functional dissection of these closely related paralogs is possible. 462

463 The patchy distribution of Arf6 is somewhat surprising, at least in part because it 464 contrasts with the near universal distribution of Arf1 paralogs. While Arf6 is perhaps most 465 commonly associated with endocytosis and plasma membrane dynamics (see above) we 466 speculate that perhaps it is its role in pericentriolar localization of specific subsets of recycling endosomes that are required for midbody formation and abscission (Fielding et al. 467 468 2005; Wilson et al. 2005; Turn et al. 2020) that vary amongst species. The nature and 469 composition of centrioles, as well as associated components are known to vary, including 470 losses or differences in Archaeplastida and SAR (Nabais et al. 2020).

The unexpected discovery of the sporadically distributed, yet potentially ancestral
SarB paralog (figs 1 and 3A; supplementary table 3, Supplementary Material online) raises
an interesting possibility of a specific elaboration of the ER function in the LECA lost for

474 some reason(s) by most major eukaryotic groups. Direct functional characterization of SarB 475 in suitable model organisms is necessary before the causes behind the retention/loss pattern 476 of the gene may be understood. However, it is interesting to compare SarB with the recently 477 uncovered complexity of the ancestral set of paralogs of the COPII coat complex, including 478 the Sec24III paralog as patchily distributed as SarB (Schlacht and Dacks 2015). The 479 phylogenetic profiles of SarB and Sec24III do not overlap well (e.g., SarB is missing from 480 Chloroplastida and Sec24III is absent from diatoms), so we are not suggesting a specific functional link between these two proteins. Nevertheless, the existence of both proteins 481 482 implies the existence of an interesting degree of variation in the COPII vesicle formation at 483 the ER in different eukaryotes.

484

#### 485 Arl17 provides a rare example of horizontal transfer of a Ras superfamily gene

486 The newly recognized functionally uncharacterized Arl17 group of ARF family protein is 487 unusual not only because of its unique domain architecture (fig. 2), but also due to its very patchy taxonomic distributions (fig. 3A; supplementary table 3, Supplementary Material 488 489 online). Based on our current sampling, Arl17 is completely missing from several major 490 eukaryotic clades (Malawimonadida, Metamonada, Discoba, Stramenopiles, Haptophyta, and 491 Rhodophyta), whereas its occurrence in the other groups is typically sporadic. Particularly 492 interesting is identification of a group of four closely related Arl17 homologs in the rotifer A. 493 vaga (Fig. 2; supplementary table 3, Supplementary Material online), which is the sole 494 representative of the densely sampled Holozoa clade possessing Arl17 (supplementary table 495 3. Supplementary Material online). Transcriptome data from A. vaga relatives indicate that 496 Arl17 is not restricted to a single rotifer species (data not shown), ruling out contamination in 497 the A. vaga genome data. Hence, the isolated occurrence of Arl17 in a rotifer lineage strongly 498 indicates gain via horizontal gene transfer (HGT) from a protist or fungal lineage, with 499 subsequent gene duplications (at least partly accounted for by tetraploidy of the A. vaga 500 genome, see above). Indeed, analyses of rotifer genomes revealed propensity of these 501 peculiar microscopic animals for gene gain from various sources, and three of the four A. 502 vaga Arl17 paralogs were included in the list of HGT candidates in the A. vaga genome (Flot 503 et al. 2013). To our knowledge, this is the first convincing case of a eukaryote-to-eukaryote 504 HGT in the whole Ras superfamily. Even though phylogenetic analysis of the Arl17 GTPase 505 domain did not shed light on the origin of rotifer's Arl17 (fig. 2), a specific relationship to 506 Arl17 proteins from *Physarum polycephalum* is suggested by a phylogeny inferred for the

507 different copies of the C-terminal novel domains (supplementary fig. 10, Supplementary508 Material online), suggesting that rotifers acquired Arl17 from an amoebozoan.

509

#### 510 Expansion of the ARF family in Holozoa

511 Given the prominent position of metazoan model systems (humans, *Mus musculus*, *D*. 512 melanogaster, Caenorhabditis elegans) in research on the ARF family, we carried out a 513 separate analysis concentrating on the family members in widely sampled representatives of Metazoa and their closest protist relatives, together constituting the taxon called Holozoa. 514 515 Analogously to our eukaryote-scale ScrollSaw analysis described above, we compared 18 516 groups of sequences corresponding to the main holozoan lineages (phyla). This approach 517 narrowed our original holozoan dataset of nearly 550 sequences to ~320 sequences and 518 phylogenetic analysis of this reduced dataset revealed a set of strongly supported clades that 519 provided a basis for defining ARF family paralogs conserved across the main holozoan or metazoan lineages (fig. 4). All ancient eukaryotic paralogs represented in this taxon, except 520 521 Arf1, form supported clades (note that Arl17 failed to pass the ScrollSaw step as it is present 522 only in rotifers). Furthermore, six additional groups could be identified based on this 523 analysis, namely Arf4 (class II Arf), Arl4, 10, 15, 19 and TRIM23. Most of them are named 524 according to previously annotated vertebrate genes (Gillingham and Munro 2007). An 525 exception is a novel group, here named Arl19, which is not a resolved clade, but seems to 526 represent a coherent evolutionary lineage based on additional evidence (see below). Analysis of intron positions in a subgroup of ARF family genes corresponding to Arfs and their closest 527 528 relatives supported the delimitation of the main groups, but also suggested that several 529 sequences initially annotated as Arf1 (based on BLAST searches) may constitute a novel 530 conserved group in unicellular holozoans and several invertebrate lineages (supplementary 531 fig. 11, Supplementary Material online). Specifically, this group is characterized by three 532 unique intron positions, and a focused phylogenetic analysis supported its monophyly and 533 separation from Arf1 and other clades (supplementary fig. 12A, Supplementary Material 534 online). We thus named this novel clade Arl20.

Establishment of novel ARF lineages provided a basis for the assignment of sequences excluded by the ScrollSaw protocol by the same approach as described for the ancient eukaryotic paralogs. Moreover, inspection of the exon-intron structures facilitated assignment of some of the problematic genes. For example, *Takifugu rubripes* harbours several standard Arfs and one additional Arf-like paralog (TruArf4L in supplementary table 1, Supplementary Material online) with an almost equal similarity to the Arf1 and Arf4

groups. Both phylogenetic and HMMER-based analyses were inconclusive concerning the
origin of this gene, but the exon-intron structure of TruArf4L exhibits the pattern typical to
the Arf4 group (supplementary fig. 11, Supplementary Material online), supporting
annotation of this gene as a divergent representative of the Arf4 group. Combining such
different forms of evidence allowed us to annotate the majority of sequences, to establish the
phylogenetic distribution of the main groups, and to map their origins and losses onto the
holozoan phylogeny (fig. 5; supplementary table 3, Supplementary Material online).

Altogether we could recognise seven groups that apparently originated after the split 548 549 of the holozoan lineage from their relatives (Holomycota), that is in the holozoan stem itself 550 (Arf4), at a later step but still before the divergence of Metazoa and their sister group 551 choanoflagellates (Arl15, 19, 20), in the metazoan stem (Arl10), or after the divergence of the 552 deepest metazoan phyla (Arl4, TRIM23). This stepwise build-up of complexity of the ARF 553 family (fig. 5B) contrasts with a somewhat different evolutionary pattern documented for the 554 Rab family, which experienced a wave of expansion concentrated in the metazoan stem 555 lineage (Elias et al. 2012). The novel ARF family members in Holozoa apparently emerged 556 by modification of duplicated copies of specific ancient eukaryotic paralogs, although the 557 exact sources may be difficult to determine. Sequence similarity and phylogenetic analysis 558 (fig. 4) point to the Arl2/3 clade as the most likely cradle of Arl10 and 15, but the position of 559 these two paralogs is unstable in different phylogenies (e.g. supplementary fig. 12B, 560 Supplementary Material online). Evidence is more solid for the origin of Arf4, Arl4, 19, 20 561 and TRIM23, suggesting these are offshoots stemming from Arf1/6-like ancestors (fig. 4; 562 supplementary fig. 12, Supplementary Material online).

563 A common origin of Arf1 and Arf4 groups was already reported (Li et al. 2004; 564 Manolea et al. 2010), but our analysis placed this event before the divergence of 565 ichthyosporeans to the common ancestor of Holozoa (fig. 5B), which probably possessed 566 Arf1, Arf4, and Arf6 as single-copy genes. While Arf4 and Arf6 seem to duplicate only 567 sporadically, Arf1 is often present in more than one copy, suggesting a high propensity for 568 duplication; this tendency is in fact seen for eukaryote lineage in general (supplementary 569 table 3, Supplementary Material online). Phylogenetic analyses usually do not recover Arf1 570 and Arf4 as supported monophyletic clades (e.g., fig. 4), which is probably a result of their 571 high sequence similarity reflected also in partial functional overlap of Arf1 and Arf4 (Jackson 2014; Jackson and Bouvet 2014). However, their separation is obvious from the comparison 572 573 of the exon-intron structures of the respective genes (supplementary fig, 11, Supplementary 574 Material online).

575 Two more holozoan or metazoan GTPase groups are likely evolutionarily derived 576 from the ancestral Arf1 gene, yet have diverged to the point it seems inappropriate to call 577 them "Arfs". One is Arl20, a previously unrecognized group of genes sharing three specific 578 intron positions (supplementary fig. 11, Supplementary Material online). Their relationship to 579 Arf1 cannot be conclusively inferred from our phylogenetic analysis (supplementary fig. 580 12A, Supplementary Material online), but an intron position shared with Arf1 (and Arf4) and 581 outcomes of similarity searches support this hypothesis. TRIM23 (also called ARD1) is an 582 unusual protein including not only the GTPase domain, but also a block of domains 583 characteristic for the TRIM family (RING-type E3 ubiquitin ligase, a tandem of BBbox 584 domains, and the BBC domain forming a coiled-coil) at the N-terminus. The GTPase domain 585 is highly similar to true Arfs (Vichi et al. 2005) and its specific relationship to Arf1 is 586 obvious from the virtually identical exon-intron structure (of the gene part encoding the 587 GTPase domain; supplementary fig. 11, Supplementary Material online).

588 Arl4 and the Arl19 group newly recognized here constitute a sister group to Arf6 in our trees (fig. 4; supplementary fig. 12A, Supplementary Material online). While Arl4 forms 589 590 a highly supported monophyletic group, its placement disrupts the monophyly of Arl19, 591 perhaps due to an insufficient phylogenetic signal that would unite all Arl19 sequences in the 592 analyses. The origin of Arl4 and Arl19 from Arf6 is conceivable and there are also potential 593 functional links between Arl4 and Arf6; e.g., mammalian Arl4 proteins can recruit the Arf6 594 GEFs cytohesins to the plasma membrane (Hofmann et al. 2007) and each GTPase can influence actin dynamics (Cotton et al. 2007; Li et al. 2007; Patel et al. 2011). The exon-595 596 intron structure of Arl4 and Arl19 are not helpful in unveiling their origin. Only a minority of 597 Arl4 genes contain introns, the intron positions are not conserved between Arl4 genes, and do 598 not match the rest of examined Arf genes (supplementary fig. 11, Supplementary Material 599 online). This suggests that Arl4 may have originated through retroposition (Kaessmann et al. 600 2009), that is by integration of a reverse-transcribed mRNA into the genome of an early 601 metazoan, with the few non-conserved introns gained secondarily and independently in 602 different metazoan lineages. The exon-intron structure of Arl19 is rather puzzling, as several 603 genes share an intron with Arf1 (supplementary fig. 11, Supplementary Material online), but 604 the whole clade branches off close to Arf6 (fig. 4).

In addition to the aforementioned novel ARF family members broadly conserved
across Holozoa or Metazoa, various metazoan lineages exhibit still other novelties suggesting
further functional elaboration. Here we focus on vertebrates. First, the vertebrate ARF family
complement has been expanded by duplications of Arf1 and Arf4, yielding the well-known

609 two groups of paralogs (Arf1, 2, 3 versus Arf4 and 5). Together with multiple duplications of 610 Arl4, vertebrates are thus endowed with a battery of lineage-specific paralogs that are 611 generally highly similar in sequence and (presumably) function (supplementary tables 1 and 612 3, Supplementary Material online). Second, vertebrates have experienced duplication of the 613 Arl10 gene inherited from their invertebrate ancestor, giving rise to two in-paralogs that 614 diverged from each other to such an extent that they were not initially recognized as closely 615 related and which is reflected in their different names: Arl9 and Arl10 (supplementary fig. 12B, Supplementary Material online). Finally, vertebrates encode two divergent ARF family 616 617 members of a common origin, called Arl11 and Arl14, that seems to have evolved by 618 duplication and divergence from Arl4 (fig. 5; supplementary fig. 13, Supplementary Material 619 online). The functional significance of these novelties is unclear, owing to limited knowledge 620 of the function of the respective proteins in any vertebrate species including humans. It is, however, important to stress that the vertebrate ARF family complement has been sculpted 621 also by gene loss, as vertebrates (in contrast to their sister group tunicates represented in this 622 623 study by Ciona intestinalis) lack Arl19 and Arl20 (fig. 5).

624

# The emergence of other major eukaryotic clades was accompanied by limited evolutionary novelty in the ARF family

627 Given the identification of multiple novel ARF family paralogs in Holozoa/Metazoa, we also 628 applied the ScrollSaw protocol to other eukaryote groups to uncover possible lineage-specific 629 innovations. Interestingly, while gene duplications specific to terminal organismal lineages 630 are common in the ARF family, only three higher-level taxa – rhodophytes, glaucophytes, and Chloroplastida – seem to have evolved novel family members by gene duplication in 631 632 their stem lineages (fig. 3B; supplementary table 4, Supplementary Material online). The 633 genome of red algal ancestors underwent massive reductive evolution (Yoon et al. 2017), 634 which is reflected also by their highly reduced set of Rab GTPases (Petrželková and Eliáš 635 2014) as well as of ARF family proteins (fig. 3; supplementary table 3, Supplementary Material online). Somewhat opposite to this trend, a novel ARF family member, here denoted 636 637 ArlRhodo, is shared by distantly related rhodophyte taxa and apparently emerged before the 638 radiation of the whole group. Their origin remains elusive, as the phylogenetic analysis 639 placed ArlRhodo as a separate clade of the ARF family with no specific affinities to any of 640 the ancestral clades (fig. 6A). By contrast, the glaucophyte innovation, in fact represented by 641 multiple paralogs in individual glaucophyte species, can clearly be traced as a highly 642 divergent offshoot of Arl13 (supplementary fig. 14, Supplementary Material online).

643 The only previously documented innovation of the ARF family specific for a major 644 eukaryotic group other than metazoans is the plant ArfB (Vernoud et al. 2003). It was 645 proposed to be an Arf6 ortholog (Li et al. 2004), and indeed our phylogenetic analysis places 646 ArfB as sister group to Arf6 (supplementary fig. 15, Supplementary Material online). 647 However, this topology is not statistically supported and can be an artefact resulting from the apparently rapid initial evolution of the ArfB gene reflected by the long stem branch 648 649 subtending the ArfB subtree. Moreover, ArfB genes share one intron position with Arf1, but none with Arf6 (supplementary fig. 4, Supplementary Material online). Hence, we leave the 650 651 origin of the ArfB group as unresolved. This not withstanding, the timing of the ArfB 652 emergence coincides with a duplication of the ARF GEF BIG in the Chloroplastida (Pipaliya 653 et al. 2019). The duplication of the ArfB paralogs in embryophytes also coincides with the 654 duplication of GBF1 proteins in that same lineage. As both of these GEFs act on Arf1-655 derived paralogs in metazoans at least, this lends itself to the hypothesis that ArfB is derived 656 from Arf1. It raises the further speculation that one of the BIG duplicates acts specifically on 657 ArfB in green algae and suggests that the ArfB, BIG, and GBF1 duplicates should all be 658 included in any activity assays aimed at understanding how this network functions in plant 659 cells.

660

## Extensive molecular tinkering in the evolution of membrane attachment mechanisms inthe ARF family

It is currently understood that a large fraction of ARF family members act within
endomembrane traffic pathways through their actions on the surface of source membranes
(Gillingham and Munro 2007). This necessitates specific, and (typically) transient, membrane
attachment, typically relying on specific PTMs, employed by different ARF family members.
Our analyses illuminate the origins of the previously described means of membrane
association, but also finds evidence consistent with diversity in the mechanisms involved in
membrane association (summarized in fig. 7A-F).

N-terminal myristoylation (N-myristoylation) is the most common lipid modification mediating the reversible membrane attachment of ARF family proteins (Kahn et al. 1988; Liu et al. 2009). Two necessary prerequisites for N-myristoylation are the glycine residue at the second position of the protein and specific sequence motif downstream that is recognised by the myristoyl transferase catalysing the addition of the myristate moiety to the N-terminal glycine (Duronio et al. 1991; Resh 1999). Once acted upon by N-myristoyl transferase, the myristate group is attached through an amide bond that is permanent for the life of the

677 protein. Reversibility in membrane association is tightly linked to the activation status of the 678 ARF family protein, as the myristoylated N-terminal α-helix is accommodated in a 679 hydrophobic channel when the protein is inactive (GDP-bound) but becomes solvent exposed 680 in response to activation (GTP-binding), resulting in its propensity to bury the freed myristate 681 in a lipid bilayer (Pasqualato et al. 2002; Seidel et al. 2004; Liu et al. 2009; Liu et al. 2010). 682 Using dedicated bioinformatic tools (see Materials and Methods), we predicted this 683 post-translational modification for the majority of the proteins representing the ancestral eukaryotic paralogs Arf1, Arf6, Arl1, and Arl5 (fig. 7G; supplementary tables 1 and 5, 684 685 Supplementary Material online), in keeping with previous experimental data from yeast and 686 mammalian proteins (Kahn et al. 1988; D'Souza-Schorey and Stahl 1995; Lee et al. 1997; Lin 687 et al. 2002). Virtually all Arf6, Arl1 and Arl5 proteins possess the conserved glycine residue 688 at the second position, and the negligible minority of those not predicted as N-myristoylation 689 targets may be false negatives. From almost 450 Arf1 genes investigated, 40 do not possess 690 the expected glycine residue and cannot be modified by myristoylation in a standard manner. We note that a recent study found N-myristoyltransferase capable of acylating lysine in the 691 692 third position (Dian et al. 2020), though the predicting algorithms employed here did not 693 consider this possibility. Regardless, only three of the 40 Arf1 proteins without a 694 myristoylatable glycine have a lysine residue at the third position. All of them are 695 accompanied by two or more Arf1 genes that are N-myristoylated in the given organism 696 (supplementary table 1, Supplementary Material online), so they apparently represent 697 lineage-specific paralogs with a changed behaviour towards membranes. The newly 698 recognised Arl18 paralog, though not closely related to the previous four paralogs, also is 699 predicted to be ancestrally myristoylated, as all genes contain a glycine residue at the second 700 position and the majority of them are predicted as N-myristoylated (fig. 7G; supplementary 701 tables 1 and 5, Supplementary Material online). Interestingly, the Arl18 sister group Arl8 702 seems to ancestrally lack glycine at the second position (fig. 7G; supplementary table 1, 703 Supplementary Material online) and the only putatively N-myristoylated Arl8 can be found in 704 rhizarians, suggesting secondary acquisition of the myristoylation motif in this lineage. The 705 majority of Arl2 and Arl3 proteins do harbour a glycine residue at the second position, but N-706 terminal myristoylation is predicted only for a few Arl3 proteins (fig. 7G; supplementary 707 tables 1 and 5, Supplementary Material online) and these may be false positives, considering 708 the experimental evidence for the lack of N-myristoylation in representative Arl3 proteins 709 (Sharer et al. 2002; Setty et al. 2004). The Arl6 group is clearly heterogeneous, including 710 members that certainly are not myristoylated as well as members that likely have this

modification. Thus, the evolutionary course leading to the distribution of N-myristoylation in
different ARF family members is not always clear. One possibility is an early origin of this
modification in an ancestor of all the clades with N-myristoylated members, followed by its
multiple secondary losses. However, multiple independent acquisitions is certainly a likely,
and mutually non-exclusive, alternative.

716 S-palmitoylation (i.e., addition of a palmitoyl moiety to one or more cysteine 717 residues) also mediates protein association with membranes, though unlike N-myristoylation there are enzymes capable of reversing this acylation making it a more transient modification 718 719 (Zhou and Cox 2014). We again employed a suite of dedicated algorithms to predict the 720 presence of this modification in ARF family members, as described under Materials and 721 Methods. Arl15 proteins typically harbour several N-terminal cysteine residues, usually 722 predicted as S-palmitoylated (supplementary fig. 16, Supplementary Material online), and 723 approximately half of the Arl13 and Arl16 sequences analysed also contain one or more 724 putative S-palmitovlated cysteine residues in their N-terminal region (fig. 7G; supplementary 725 tables 1 and 5, Supplementary Material online). S-palmitoylation of Arl13 from C. elegans 726 and mammals has been confirmed experimentally and demonstrated as crucial not only for 727 the proper localization of the proteins, but also for stability and function (Cevik et al. 2010; 728 Roy et al. 2017). In a few cases, such as in the red algae-specific paralog ArlRhodo, S-729 palmitoylation seems to accompany N-myristoylation (figs 6B and 7G; supplementary table 730 1, Supplementary Material online), similar to various other proteins, including GTPases (e.g., 731 some Ga proteins; Zhou and Cox 2014).

732 In addition to employing covalently attached saturated fatty acids, proteins also can be 733 permanently (absent proteolytic cleavage) anchored in the membrane via a transmembrane 734 domain. Of the proteins investigated here, this was previously demonstrated for SR $\beta$ , a 735 protein anchored in the ER membrane via its N-terminal transmembrane region (Keenan et 736 al. 2001) that appears to be conserved in all SRB sequences investigated (fig. 7G). An N-737 terminal transmembrane region was independently acquired by the Metazoa-specific Arl10 738 (see above) and several other ARF family members in various eukaryotes (fig. 7G; 739 supplementary table 1, Supplementary Material online). In some cases, we could confirm 740 conservation of such putative N-terminally anchored GTPases in a broader organism clade 741 beyond the species primarily targeted by our analysis, as is the case of divergent putative 742 Arf1 paralogs from *Bigelowiella natans* and other chlorarachniophytes (supplementary fig. 743 17A, Supplementary Material online) and from *Pavlova pinguis* and other haptophytes of the

744 class Pavlovophyceae (supplementary fig. 17B, Supplementary Material online). Another mode of membrane attachment utilized by some ARF family members is accretion of specific 745 746 membrane-binding domains. This is exemplified by unusual proteins from choanoflagellates 747 and trypanosomatids that contain an N-terminal phosphoinositide-binding PH domain 748 (Lemmon 2007) connected to the ARF family GTPase domain by a long linker region (fig. 749 7E; supplementary table 1, Supplementary Material online). Finally, the eustigmatophyte 750 *Vischeria* sp. encodes a unique ARF family protein (VisArlX2 in supplementary table 1, 751 Supplementary Material online) with a long N-terminal extension lacking any detectable 752 conserved protein domain or functional motif and with a C-terminal tail ending with the 753 amino acid sequence CSIM (fig. 7F), which is reminiscent of the so-called CaaX motif (or 754 box) directing prenylation of the cysteine residue in diverse proteins (Fu and Casey 1999). A 755 similar protein, including this motif, is encoded by additional eustigmatophytes (not shown), 756 and two different prediction programs proposed the cysteine residue to be prenylated (see 757 Material and Methods for details). C-terminal prenylation is a common modification ensuring membrane attachment of GTPases belonging to Rab, Ras and Rho families (Zhou and Cox 758 759 2014), but to our knowledge it has not been reported previously for an ARF family protein.

760 The well-studied mammalian members of the ARF family are subject to other post-761 translational modifications (e.g., see Phosphosite Plus; https://www.phosphosite.org/), though 762 these either lack consensus motifs that prevent predicting their existence in other organisms 763 or have no known functional consequences, or both. One exception to this is N-terminal 764 acetylation of Arl8, which has been shown to be important for its association with lysosomal 765 membranes (Hofmann and Munro 2006). Similarly, in S. cerevisiae the Arfrp1 protein 766 (unfortunately named Arl3p only in this organism) is also acetylated and this is required for 767 its association with Golgi membranes (Behnia et al. 2004). Future development of 768 appropriate prediction tools, perhaps combined with dedicated biochemical investigations, 769 will be instrumental in grasping the full breath and evolutionary conservation of PTMs in the 770 ARF family.

In summary, the use of several different means of membrane attachment is consistent with ARF family proteins acting predominantly on a membrane surface, and the diversity of various membrane attachment mechanisms exhibited by this family is surprisingly extensive and reminiscent of what has been described for the distantly related GTPase Rheb (Záhonová et al. 2018). It is perhaps worth noting that eukaryotic organisms can vary widely in their lipid composition and the same is true of different organelles in an organism, making different means of membrane association likely important for this family of cell regulators

that most often act on membrane surfaces and can even modify the lipid composition via

779 direct activation of lipid kinases and lipases.

780

#### 781 Extensive diversity of multi-domain ARF family members

782 The existence of the PH domain-containing ARF family proteins or the aforementioned 783 multi-domain TRIM23 protein (Vichi et al. 2005) counter the paradigm of ARF family 784 members being limited to single (GTPase) domain proteins with only short N- and C-terminal 785 extensions. In fact, our analyses challenge this dogma further. Although they represent a 786 minority (75 out of >2,000 sequences in our dataset), multi-domain ARF family members 787 represent a much greater number of different protein architectures involving combinations of 788 the GTPase domain of the ARF family with other functional domains than thought previously 789 (see column S in Supplementary table 1, Supplementary Material online).

790 The novel, presumably ancestral eukaryotic, Arl17 group characterized by combining 791 an Arf-related domain with varying numbers of tandemly arrayed copies of a novel uncharacterized domain (fig. 2) was introduced above. Additional domain architectures are 792 793 found in proteins that generally seem to be lineage-specific innovations restricted to 794 particular taxa; some examples are provided in fig. 8. Similar to TRIM23, some include 795 domains linked to ubiquitination, namely the BTB domain or the F-box domain (see 796 Genschik et al. 2013), indicating recurrent recruitment of ARF family members into 797 ubiquitin-dependent regulatory circuits. Ciliates exhibit a unique protein with an ARF family 798 GTPase domain fused to a segment homologous to radial spoke protein 3 (RSP3), a 799 component of radial spokes in the axoneme (see Wirschell et al. 2008). This predicts ciliary localization of this protein, and indeed, it is among the proteins detected in the ciliary 800 801 proteome of *T. thermophila* (Smith et al. 2005). *Entamoeba histolytica* possesses a protein 802 with a divergent C-terminal ARF family domain preceded by the VPS9 domain. The latter 803 domain is known to act as a GEF of the endosomal Rab GTPase Rab5 (Ishida et al. 2016), so 804 this protein may be part of a pathway with multiple sequentially acting GTPases similar to 805 regulatory GTPase cascades known from mammalian or yeast cells (Jones et al. 1999; Mizuno-Yamasaki et al. 2012). Another unique domain combination occurs in one of the Arf 806 807 paralogs in the haptophyte *Emiliania huxleyi*, which is fused to the C-terminus of a block including a domain of the 2OG-Fe(II) oxygenase superfamily. It is possible that the GTPase 808 809 domain regulates the enzyme activity of the N-terminal part of the protein. The ARF family 810 domain can combine also with other Ras superfamily GTPase domains, as demonstrated by a 811 protein from Malawimonas californiana with an N-terminal Rab domain and a C-terminal 812 Arf domain linked by a region containing detectable BTB and BACK domains (fig. 8A). Tinkering with protein domains in ARF family proteins can be encountered in a 813 814 different evolutionary context than the emergence of lineage-specific paralogs. In the case of 815 Arl13, domains were acquired or lost without gene duplication, resulting in differences in domain architectures between orthologous Arl13 genes. Previously characterized orthologs 816 817 from mammals and Chlamydomonas reinhardtii exhibit a poorly conserved C-terminal extension that includes a region forming a coiled-coil followed by a proline-rich region (Hori 818 819 et al. 2008; Miertzschke et al. 2014; fig. 8B). Inspection of the large collection of Arl13 820 sequences amassed for this study revealed that this arrangement is distributed broadly across 821 the eukaryote phylogeny and likely ancestral. However, some species (represented by eleven 822 Arl13 genes out of 70 included in our dataset) depart in various way from this structure, e.g. 823 by lacking the proline-rich region or the coiled-coil. Recently, Zhang et al. (2018) identified a 824 non-canonical Arl13 gene from Trypanosoma brucei containing the DD RI PKA domain (Dimerization/Docking domain of the Regulatory subunit of protein kinase A (PKA)) that is 825 826 essential for targeting of T. brucei Arl13 to the cilium. Our analysis revealed that the same 827 protein architecture is present also in *Euglena gracilis*, suggesting it is a synapomorphic 828 character for the whole Euglenozoa phylum (fig. 8B). Meanwhile, a subset of Stramenopiles 829 (oomycetes and ochrophytes) independently acquired DD\_RI\_PKA domain as two tandemly 830 arrayed copies (fig. 8B). DD\_RI\_PKA mediates interaction of PKA with A-kinase-anchoring 831 proteins (AKAPs), which regulate PKA localization in the cell (Sarma et al. 2010). Given the 832 ciliary function of Arl13 (see above), we speculate that the DD\_RI\_PKA domain in some Arl13 proteins interacts with a cilium-localized AKAP, such as the aforementioned RSP3 833 834 protein (Gaillard et al. 2001; Jivan et al. 2009). In contrast, mammalian Arl13b contains the 835 simpler VxP motif in the large C-terminal domain that is required for ciliary localization 836 (Higginbotham et al. 2012; Cevik et al. 2013; Gigante et al. 2020). DD RI PKA domains in 837 Phytophthora sojae Arl13 are followed by the TUDOR domain, known for the ability to bind to the methylated lysine and/or arginine residues (Botuyan and Mer 2016). The TUDOR 838 839 domain was independently accreted also to the C-terminus of the Arl13 from 840 Aurantiochytrium limacinum (fig. 8B). Another notable variant is encountered in Arl13 from 841 *B. natans* (fig. 8B) and other chlorarachniophytes (supplementary fig. 18, Supplementary Material online), which exhibit a novel form of the C-terminal extension including the  $Ca^{2+}$ -842 binding EF-hand motif. Interestingly, the N-terminus of chlorarachniophyte Arl13 proteins 843 844 appears to be related to calcineurin B, a Ca<sup>2+</sup>-binding regulatory subunit of the protein

phosphatase calcineurin (Guerini 1997). It thus seems likely that Arl13 function is regulated

- by  $Ca^{2+}$  in chlorarachniophytes. Exceptional is an *E. gracilis* gene (co-occurring in this
- species with a typical Arl13 gene) that we named Arl13Triple, as it is composed of a tandem
- triplication of a divergent Arl13-reated GTPase domain (fig. 8B). The varying domain
- architecture of Arl13 in different eukaryotes points to a substantial degree of functional
- 850 divergence of this key ciliary component.
- 851

## 852 Insights into the early radiation of the ARF family

853 In the analysis of protein family evolution, resolution between the paralogs is a tremendously 854 informative result as it allows the inference of cellular evolution of the associated organellar 855 compartments. However, such resolution has been difficult to obtain for many families. The 856 ScrollSaw methodology was a step forward in obtaining resolution for datasets with many 857 paralogs and short sequence length; e.g., Rabs and TBC proteins (Rab GAPs; Elias et al. 2012; Gabernet-Castello et al. 2013). Here, our application of the ScrollSaw methodology 858 also yielded a partially resolved backbone topology (fig. 1). We observed the robust 859 860 sisterhood of Arl8 and Arl18 and of these both to Arl16. We also observed the sisterhood of 861 Arl2 and Arl3 plus the moderately supported node uniting Arf1 with Arf6. Most notably, 862 there was a strongly resolved node grouping together Arf1, Arf6, Arl1, Arl2, Arl3 and Arl5 863 and separating them from the remainder of the paralogs.

864 This resolution provides the basis for several key inferences about the ancestral role 865 of the ARF family progenitor and some implications about the role of these proteins during 866 eukaryogenesis. Taking only the most broadly conserved biochemical and cellular features of the various ARF family members, and assuming basic functional homology in orthologs to 867 868 their roles in LECA (Klinger et al. 2016), what is likely ancestral is a GTPase that changes 869 conformation to relocate from the cytosol to a membrane and which binds other proteins as 870 effector(s). Given the widespread role of ARF family members, this may mean a role in 871 membrane-traffic. However, with at least one resolved node separating the best-known 872 family members Arf1 and Sar1, a simple scenario of a single primordial GTPase that nucleates a primordial vesicle coat-forming complex is ruled out. This suggests that the 873 proto-coatomer hypothesis (Devos et al. 2004) may well need to be modified to take a more 874 complicated scenario, including possible convergence, parallel evolution, and even merging 875 876 of architectures into account (Dacks and Robinson 2017; Field and Rout 2019).

877

#### 878 Conclusions

879 Our comprehensive analysis of an extensive, well-curated dataset of ARF family proteins has 880 provided evolutionary insights and raised questions to be addressed by future molecular cell 881 biological exploration. The identification of 16 ancient ARF family paralogs both extends the 882 inferred complexity of LECA and sets a framework of what components can be expected to 883 be acting when delving into cellular function in diverse eukaryotes. By contrast the 884 identification of expanded complements, including novel paralogs, e.g. the metazoan Arl19 885 and Arl20, provide specific new candidates for investigation in some of the best explored and 886 heavily utilized cell biological model systems. The diversity of domain architecture 887 challenges the paradigm of this family strictly as small GTPases and begs probing of new 888 protein-protein interactions. Altogether, our work thus establishes a solid basis for future 889 more detailed investigations into the biology of ARF family proteins at a eukaryote-wide

- scale.
- 891

## 892 Materials and Methods

## 893 Building and curation of the ARF family dataset

894 ARF family GTPases were searched in genome and/or transcriptome assemblies from 114 895 eukaryotic species selected such as to cover as many main eukaryotic lineages as possible 896 (sequence identifiers, source databases, and further comments are provided in supplementary 897 table 1, Supplementary Material online). The selection of taxa reflected the availability of 898 relevant data as of 2018, when the sampling was frozen to obtain a final sequence dataset for all the subsequent analyses. As a result, several main eukaryote lineages, for which genome 899 900 or transcriptome data became available more recently (e.g. the CRuMs supergroup, 901 Telonemia, Rhodelphidia etc.), are not represented in our dataset. ARF family sequences 902 were identified using BLAST and its variants (Altschul et al. 1997). Each organism-specific 903 dataset was queried with reference members of the family and significant hits were evaluated 904 by reverse BLAST searches against an in-house extensively curated taxonomically-rich 905 database of GTPases. Query sequences being more similar to previously annotated members 906 of the ARF family (including SR $\beta$ ) were kept for further analysis. Existing protein sequence 907 predictions were carefully evaluated and in a many cases revised by modifying the predicted 908 exon-intron structure of the underlying gene model based on information from transcriptomic 909 data or comparison to homologous sequences. To identify genes potentially missing from 910 existing genome annotations, tblastn searches of nucleotide sequence data were carried out 911 and gene models were created anew for previously missed genes. If possible, truncated

912 sequences were completed using EST/TSA data or by iteratively recruiting raw

913 genomic/transcriptomic sequencing reads. Revised, newly predicted or extended sequences

are provided in supplementary table 1, Supplementary Material online. Putative pseudogenes

- 915 (except for the human Arf2 pseudogene sequence, which can be reconstructed) as well as
- extremely divergent sequences with disrupted ARF family motif(s) were not included into thedataset and are not listed in supplementary table 1, Supplementary Material online.

918 Each gene was initially annotated by considering results of blastp searches against our 919 comprehensive database of Ras superfamily proteins (iteratively updated by adding 920 sequences newly annotated in the course of the study). In most cases, the blastp output 921 enabled unambiguous assignment of the query sequence into one of the previously delineated 922 ortholog groups or into novel orthogroups that emerged during the study. Sequences most 923 similar to true Arfs, yet difficult to assign into the Arf1 or Arf6 groups or being visibly 924 divergent were provisionally annotated as "ArfX". Still more divergent ARF family members 925 that did not show an apparently consistent affinity to a particular ARF family orthogroup when examined by BLAST searches were provisionally annotated as "ArlX". The annotation 926 927 of some of the ArfX and ArlX sequences was subsequently revised after the employment of 928 the ScrollSaw protocol described below.

929 Sequence data from the glaucophyte *Gloeochaete wittrockiana* were included despite 930 the fact that we noticed contamination of both available transcriptome assemblies 931 (MMETSP0308 and MMETSP1089; https://www.imicrobe.us/#/projects/104) by sequences 932 from an amoebozoan. The putative contaminants were identified by careful examination of 933 individual sequences and excluded from the dataset. Another potential contaminant (the 934 contig PCB a545736;2 K: 25), showing a high similarity to Arl4 genes from primates, was 935 noticed in the transcriptome assembly from the breviate Pygsuia biforma and removed from 936 analyses.

937

## 938 The ScrollSaw protocol and phylogenetic analyses

A master multiple alignment was built for the identified ARF family members, excluding short incomplete sequences and also all SR $\beta$  sequences, as this group is noticeably different from the core of the ARF family and many of its members tend to be rather divergent in their sequence. Altogether, the master alignment included 1931 sequences. It was built iteratively, starting with separate alignments for each group of sequences initially assigned to the same (potential) orthologous group using the on-line program MAFFT (version 7), with default parameters (Katoh and Standley 2013). All alignments were checked by eye and further edited manually using BioEdit (Hall 1999). The set of separate alignments was merged into
one large alignment using the on-line Merge function of MAFFT. Divergent (ArlX)
sequences were added to the alignment at the end. The final alignment was then manually
trimmed to remove poorly conserved and unreliably aligned positions. After removing
redundancies, the alignment comprised 1891 non-identical sequences and 148 aligned
positions (all falling within the GTPase domain shared across the family).

952 The alignment was subjected to an analysis essentially following the previously published ScrollSaw protocol (Elias et al. 2012). The sequences were divided according to 953 954 the source species into 13 taxonomic groups covering the known diversity of eukaryotes: 955 Holozoa, Holomycota, Apusomonadida, Breviatea, Amoebozoa, Malawimonadida, 956 Planomonadida, Discoba, Metamonada, Archaeplastida, Cryptista, Haptista, and SAR. The 957 sizes of the groups differ substantially, since many evolutionarily important lineages were 958 represented only by a small number of species with genomic or transcriptomic data available 959 at the time when we initiated the study (in the case of Breviatea and Planomonadida by only a 960 single species). The master alignment was then subsampled by keeping only sequences from 961 each possible pair of the taxa listed above, corresponding to 78 combinations. For each of the 962 78 alignments, genetic distances between the sequences were inferred using the maximum 963 likelihood (ML) method (with the WAG+ $\Gamma$ +I substitution model) implemented in Tree-964 Puzzle 5.3 (Schmidt et al. 2002). Each resulting distance matrix was analysed using a custom 965 Python script to identify the so-called minimal-distance pairs. A minimal-distance pair 966 consists of two sequences from the two different taxonomic group compared that have 967 mutually minimal distances when distances to sequences from the other taxon are considered. 968 Minimal-distance pairs from all 78 pairwise taxon comparisons were gathered and 969 redundancies were removed, resulting in a set of 568 sequences. To further reduce the 970 complexity of the dataset we then removed all sequences that formed only one minimal-971 distance pair in all 78 pairwise taxon comparisons combined. This step yielded the final full 972 ScrollSaw dataset comprising 354 sequences. A reduced ScrollSaw variant was prepared by 973 removing sequences from the majority of metamonads exhibiting generally divergent genes, 974 including Monocercomonoides exilis, Giardia intestinalis, Spironucleus spp. and 975 Trichomonas vaginalis.

976 The two variants of the final ScrollSaw dataset were used for inferring the ML
977 phylogenetic trees using the program IQ-TREE (Nguyen et al. 2015). The substitution model
978 (LG+I+G4) was selected by the program itself based on specific optimality criteria. Branch
979 support was assessed by the SH-aLRT test (Guindon et al. 2010) and the ultrafast bootstrap

980 approximation (Minh et al. 2013). The branch support of the reduced dataset was further 981 examined by MrBayes 3.2 (Ronquist et al. 2011) using the CIPRES Science Gateway (Miller 982 et al. 2010) with the following settings: prset aamodelpr=fixed(WAG); lset rates=gamma 983 Ngammacat=4 mcmc ngen=1000000 printfreq=10000 samplefreq=1000 nchains=4 984 burnin=80. A number of additional alignments for specific dedicated analyses, derived by 985 subsampling the master alignment or aligning the selected sequences de novo (using MAFFT 986 with subsequent manual editing as described above) were used for ML phylogenetic 987 inference using the same or similar approach. The alignments were in most cases trimmed 988 according to the mask applied to the master alignment. Smaller phylogenetic analysis with 989 only a subset of paralogous groups were trimmed either manually or by stand-alone version 990 of trimAl (version 1.2rev57; option automated1; Capella-Gutierrez et al. 2009) in order to 991 retrieve more positions for the ML phylogenetic analysis. The stand-alone version of IQ-992 TREE or the IQ-TREE web server (http://iqtree.cibiv.univie.ac.at/; Trifinopoulos et al. 2016) 993 were used for the analyses. The substitution models were selected by the model selection 994 program implemented in the IQ-TREE. Branch support was assessed by the SH-aLRT test 995 and the ultrafast bootstrap approximation.

996 Tree topology testing was employed to test the hypothesis that SarB sequences form a 997 monophyletic group sister to the Sar1 group. ML trees were inferred from the reduced 998 SrollSaw alignment with a different topological constraints (specified in supplementary table 999 2, Supplementary Material online) using IQ-TREE and the same procedure as used for 1000 computing the unconstrained tree (shown in fig. 1). The unconstrained and constrained trees, 1001 together with a sample of 1,000 trees obtained as ultrafast bootstrap replicates in the 1002 unconstrained ML search on the alignment, were then compared in IQ-TREE (-au option) 1003 with the substitution models and its parameters optimized from the original alignment (-m 1004 TEST) and using 10,000 RELL replicates. The p-values of the alternative topologies obtained 1005 with the Kishino-Hasegawa (KH), Shimodaira-Hasegawa (SH), and approximately unbiased 1006 (AU) tests were considered.

1007

#### 1008 Annotation of sequences

1009 The full and reduced ScrollSaw datasets were used as a basis for annotation of the rest of the

1010 sequences. The identity of individual sequences or their groups was tested by adding them to

1011 the reduced ScrollSaw dataset and inferring a ML tree with IQ-TREE. The scrutinized

1012 sequences were assigned to a particular ancestral eukaryotic paralog and annotated

1013 accordingly if they clustered together with reference representatives of the given paralog

1014 group and the relationship was supported by SH-aLRT and ultrafast bootstrap values of  $\geq 80$ 1015 and 95, respectively. Not all genes could be annotated by this approach, hence the HMMER 1016 package (stand alone version 3.0; hmmer.org) was employed as an alternative. The aligned 1017 full ScrollSaw dataset was divided into 14 separate alignments, each representing one 1018 ancestral paralog (Arf1, 6, Arl1, 2, 3, 5, 6, 8, 13, 16, 18, Arfrp1, Sar1, and SarB). A profile 1019 HMM was constructed for each alignment using hmmbuild and a database of profile HMMs 1020 was created using hmmpress. The unannotated sequences were then used as queries in hmmscan searches against the database and the "best 1 domain" score difference between the 1021 1022 first and the second best hits was determined. If this difference was equal to or higher than 1023 20, the sequence was annotated according to the best hit. Sequences annotated based on the 1024 phylogenetic analyses or hmmscan searches are marked by asterisk (\*) in the column 1025 "Conclusively annotated" in the supplementary table 1, Supplementary Material online. 1026 Proteins representing SR $\beta$  and Arl17, which were not represented by reference sequences in 1027 the ScrollSaw dataset, were unequivocally identified owing to the distinct characters of these 1028 sequence groups, which makes them easy to recognise by BLAST-based similarity searches 1029  $(SR\beta)$  or by considering the presence of the novel conserved C-terminal domain (Arl17; see 1030 the main text). All SRβ and Arl17 proteins are therefore also considered as conclusively 1031 annotated. A single truncated sequence (Arl17b gene from Chromera velia) lacked the C-1032 terminal extension with the characteristic C-terminal domain, but was assigned to the Arl17 1033 group based on its close sequence similarity to undisputed Arl17 sequences. A combination 1034 of BLAST searches, ML phylogenetic analyses and comparison of exon-intron structure was 1035 used to obtain the most likely annotation of the sequences that could not be conclusively 1036 annotated by the aforementioned approaches. Several sequences were annotated as Arf1/6, as 1037 they showed affinity to the Arf1/6 clade, but it was impossible to decide whether they 1038 originated from ancestral Arf1 or Arf6 paralogs. Only 160 out of more than 2000 ARF family 1039 sequences analysed could not be annotated with any confidence, so they remained unassigned 1040 to any ancestral paralog (supplementary tables 1 and 3, Supplementary Material online). 1041

## 1042 Taxon-specific ScrollSaw analyses and annotation of lineage-specific paralogs

1043 To detect lineage-specific paralogs, we applied the ScrollSaw protocol separately to sets of

1044 sequences from the following main eukaryote taxa: Chloroplastida, Rhodophyta,

- 1045 Glaucophyta, Cryptista, Haptista, SAR, Discoba, Metamonada, Amoebozoa, Holomycota and
- 1046 Holozoa. Lineages represented by only one or two species (Apusomonadida, Breviatea,
- 1047 Planomonadida, and Malawimonadida) were not included. The ScrollSaw protocol and

1048 phylogenetic analyses of the resulted datasets were performed generally as described above 1049 for the whole dataset. For each main eukaryote taxon analysed, species representing it were 1050 assigned to predefined monophyletic subgroups specified in supplementary table 4, 1051 Supplementary Material online. Sequences from these species were extracted from the 1052 trimmed master alignment of the ARF family protein (except for sequences from Holozoa, 1053 which were aligned *de novo* using MAFFT and then trimmed according to the mask used for 1054 the whole dataset), the ScrollSaw protocol was applied to identify minimal-distance pairs, and ML phylogenetic trees were calculated on the filtered sequences. In contrast to the pan-1055 1056 eukaryotic ScrollSaw analysis, sequences that formed only one minimal-distance pair were 1057 not omitted (except for the analysis of the Holozoa dataset, where the criterion of the 1058 sequence belonging to at least two minimal-distance pairs was kept). The ML trees were 1059 inspected to identify robustly supported clades that would define conserved paralogs 1060 ancestral for the focal eukaryotic taxon but different from the previously defined ancestral eukaryote paralogs. In the case of Holozoa, paralogs specific for individual subgroups were 1061 considered, too. Further representatives of these paralogs (i.e., specific orthologs of the 1062 1063 constituent sequences identified in the ScrollSaw trees) were then identified among the 1064 sequences that did not pass the ScrollSaw step by a combination of BLAST searches, 1065 phylogenetic analyses and (in case of Holozoa) HMMER-based comparisons. Candidates for ancestral taxon-specific paralogs were detected only in Chloroplastida, Rhodophyta, and 1066 1067 Glaucophyta, as described in detail in the main text.

1068

## 1069 Prediction of transmembrane regions and post-translation modifications

1070 The presence of transmembrane (TM) regions in ARF family proteins was examined using

1071 the online TMHMM Server v. 2.0 (http://www.cbs.dtu.dk/services/TMHMM/). In the case of

1072 sequences with suspicious TM absence or presence (i.e., when the result was untypical for the

1073 respective ARF family subgroup), the on-line tool TMpred

1074 (<u>https://www.ch.embnet.org/software/TMPRED\_form.html</u>) was additionally employed. The

- 1075 predictions are listed in supplementary table 1, Supplementary Material online. N-terminal
- 1076 myristoylation of sequences with the glycine residue at the second position was evaluated
- 1077 using the on-line ExPASy Myristoylator tool (http://web.expasy.org/myristoylator/; Bologna
- 1078 et al. 2004), NMT The MYR Predictor
- 1079 (http://mendel.imp.ac.at/myristate/SUPLpredictor.htm; Maurer-Stroh et al. 2002), and the
- 1080 stand-alone version of GPS-Lipid (v1.0, http://lipid.biocuckoo.org/index.php; Xie et al.
- 1081 2016). Only those proteins predicted as N-terminally myristoylated by at least two tools were

1082 considered as significant candidates. Setting of all tools was default except for NMT - The 1083 MYR Predictor where only N-terminal glycine residues were considered, and fungal 1084 sequences were predicted with the "Fungi specific" option. In case of GPS-Lipid, the 1085 threshold was set to "low". Possible S-palmitoylation was predicted using SeqPalm 1086 (http://lishuyan.lzu.edu.cn/seqpalm/; Li et al. 2015), the stand-alone version of CKSAAP-1087 Palm programme (http://doc.aporc.org/wiki/CKSAAP-Palm; Wang et al. 2009), PalmPred 1088 (http://proteininformatics.org/mkumar/palmpred/index.html; Kumari et al. 2014), stand-alone version of GPS-Lipid, and WAP-Palm (http://bioinfo.ncu.edu.cn/WAP-Palm.aspx; Shi et al. 1089 1090 2013). Only those sites predicted as S-palmitoylated by at least three tools were considered as 1091 significant candidates. Setting of all tools was default except for GPS-Lipid with the 1092 threshold set to "high". Complete results from all tools are showed in supplementary table 5, 1093 Supplementary Material online, consensual results are included in supplementary table 1, 1094 Supplementary Material online. Possible prenylation was assessed only for VisArlX2 from the alga *Vischeria* sp., as it is the only protein from our dataset with a typical C-terminal 1095 1096 prenylation motif. The online programs iPreny-PseAAC (http://app.aporc.org/iPreny-1097 PseAAC/index.html; Xu et al. 2017) and GPS-Lipid were used with default settings; both 1098 tools predicted VisArlX2 as a prenylated protein.

1099

#### 1100 Other sequence analyses

1101 Intron positions were investigated in four groups of ARF family genes (Sar1/SarB; Arl8/Arl18; Arfs and the GTPase domain of Arl17; Arfs and selected Arf-like in Holozoa) as 1102 a means to illuminate the origin and relationships of these genes. The positions of introns 1103 1104 (including their phases) were mapped onto a multiple alignment of respective protein 1105 sequences using a custom Java script. The multiple sequence alignments were constructed de 1106 *novo* using MAFFT, inspected visually and adjusted manually whenever necessary 1107 (Sar1/SarB, Arl8/Arl18, Arf, and Arf-like in holozoans). For the analysis of Arf and Arl17 genes, the respective protein sequences were extracted from the master alignment. Sequences 1108 1109 with no introns in the coding sequence or represented only by transcriptomic data were 1110 omitted. A manually curated dataset of gene exon-intron structures was used as the input for 1111 the intron positions mapping. For presentation purposes, regions corresponding to 1112 unconserved N- and C- termini of the sequences were trimmed and long sequence-specific 1113 insertions were collapsed. To highlight the pattern of protein sequence conservation, 1114 CHROMA (ver. 1.0 Goodstadt and Ponting 2001) was used for processing some of the 1115 multiple sequence alignments presented. Sequence logos of the Walker B motif were

obtained using the on-line tool WebLogo 3 (http://weblogo.threeplusone.com/create.cgi;
Crooks et al. 2004) from the multiple sequence alignment of the respective sequences after
removing sequence-specific insertions present in a few sequences.

1119 Conserved protein domains and other structural features in ARF family proteins were 1120 identified using searches of Pfam (http://pfam.xfam.org/; Finn et al. 2016), the Conserved 1121 Domains database (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi; Marchler-Bauer 1122 et al. 2017), and the SMART database (http://smart.embl-heidelberg.de/; Letunic and Bork 2018). Domain predictions provided by the three tools were compared and spurious results 1123 1124 (low-significance with only a single tool) were ignored. The identity of unusual N-terminal 1125 extensions present in some Arl13 proteins were evaluated using HHpred 1126 (https://toolkit.tuebingen.mpg.de/; Söding et al. 2005). Multiple sequence alignments of the 1127 different forms on the N-terminal extensions conserved within different taxa were used as queries in the HHpred searches. In case of the N-terminal extension conserved in Arl13 1128 proteins from Euglenozoa, the sampling was expanded beyond the focal set of taxa (including 1129 1130 only three euglenozoans) by adding to the alignment several additional euglenozoan Arl13 sequences to improve the representativeness of the alignment. Similarly, additional 1131 1132 chlorarachniophyte Arl13 sequences were identified and aligned with the sole representative 1133 in the focal dataset (that from *B. natans*), and additional stramenopile (oomycete and 1134 ochrophyte) Arl13 sequences with the same conserved N-terminal extension as the 1135 stramenopile sequences in the focal set were included to increase the sensitivity of the analysis. Some TRIM23 sequences were predicted by the standard tools to contain only one 1136 1137 BBOX domain rather than the two common in most members of this group, but inspection of 1138 a multiple sequence alignment revealed high similarity of all sequences in the respective 1139 region, suggesting that all TRIM23 sequences likely conform to the same domain architecture 1140 with two BBOX domains.

1141

## 1142 Supplementary Material

1143 Supplementary data are available at Genome Biology and Evolution online.

1144

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- 1157

## 1158 Data availability

- 1159 ARF family gene sequences extracted from unpublished genome assemblies of *Gefionella*
- 1160 okellyi, Planomonas micra, and Paratrimastix pyriformis, and from our unpublished
- transcriptome assembly of Vicheria sp. CAUP Q 202, were deposited at GenBank with
- 1163 protein sequences is available in supplementary dataset 1, Supplementary Material online.
- 1164

## **1165 Competing interests**

- 1166 No competing interests declared.
- 1167

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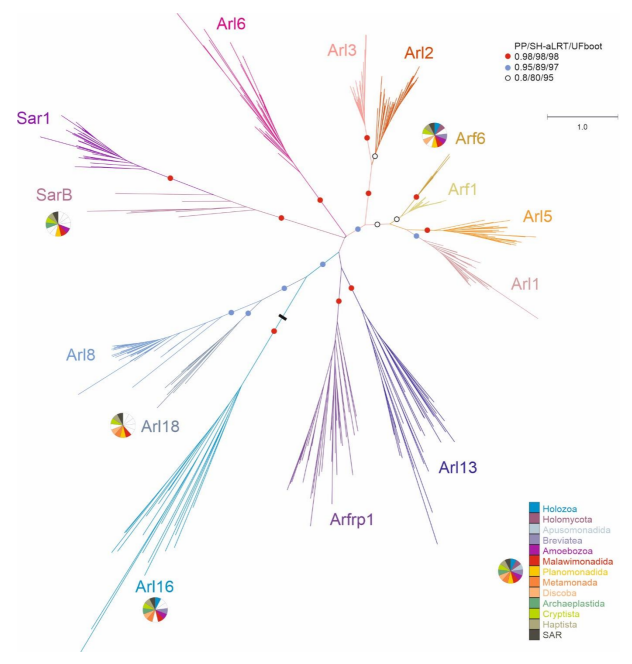
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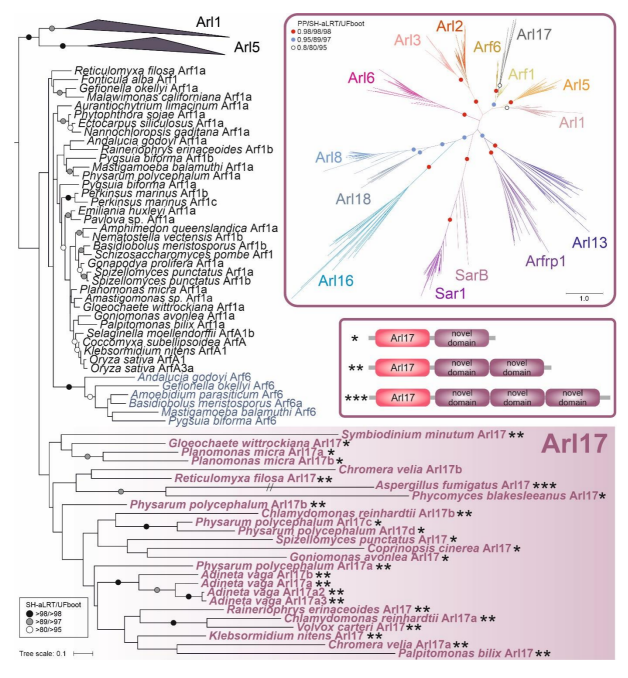
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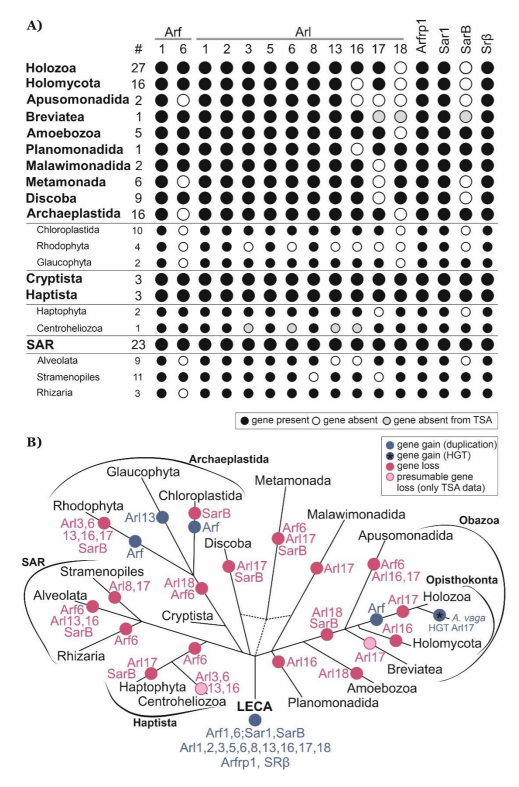
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Fig. 1. Maximum likelihood phylogenetic tree of the ARF family based on a reduced ScrollSaw 1580 1581 dataset. The tree was inferred using IQ-TREE with LG+I+G4 model (the model selected by the program itself) based on a multiple alignment of 348 protein sequences. Brach support was evaluated 1582 with MrBayes (posterior probability, PP) and with IQ-TREE using the SH-aLRT test and the ultrafast 1583 (UF) bootstrap algorithm (both 10,000 replicates), as described under Materials and Methods. Dots at 1584 branches represent bootstrap values as indicated in the graphical legend (top right), the black bar 1585 indicates the position of the root of the tree as determined with the MAD method. The bar on the top 1586 corresponds to the estimated number of substitutions per site. The pie charts indicate the occurrence of 1587 Arf6, Arl16, Arl18 and SarB in main eukaryotic lineages (indicated by different colours explained in the 1588 graphical legend in the lower right). The remaining paralogs have ubiquitous distribution (i.e., are 1589 present in all main lineages analysed). A full version of the tree is provided in supplementary fig. 2, 1590 Supplementary Material online. 1591



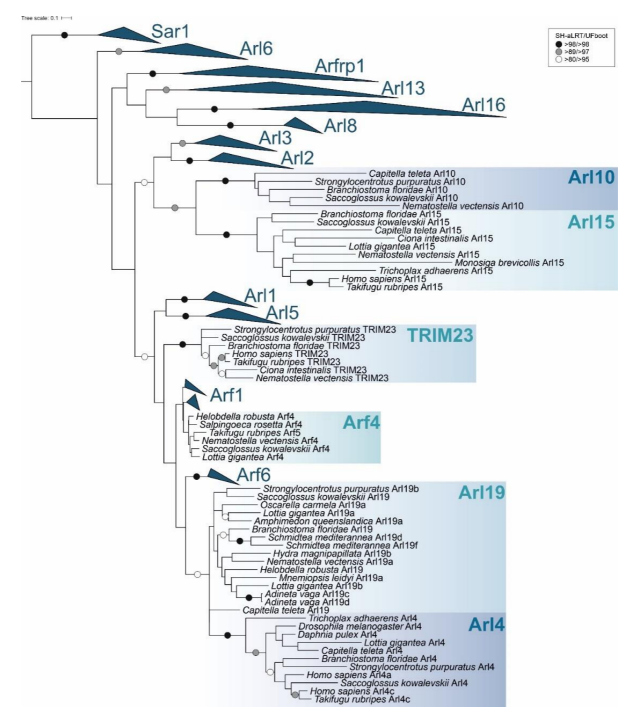
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1594 Fig. 2. Phylogenetic analysis and domain architecture of Arl17. The tree shown is a result of a ML analysis of all Arl17 sequences and a subset of the reduced "scrollsawed" dataset restricted to Arf1. 1595 Arf6, Arl1, and Arl5 sequences (the latter two collapsed as triangles), altogether 127 protein sequences. 1596 1597 The alignment was trimmed manually. The tree was inferred using IQ-TREE with LG+I+G4 model (the model selected by the program itself) with the ultrafast bootstrap algorithm and the SH-aLRT test (both 1598 1599 10,000 replicates). Dots at branches represent bootstrap values as indicated in the graphical legend (top right). The upper inset shows the ML tree inferred from a full reduced "scrollsawed" dataset 1600 combined with a subset of Arl17 sequences (picking one representative per each major eukaryote 1601 1602 group), altogether 356 protein sequences. The tree was inferred using the same approach as the tree shown in fig. 1. The inset beneath provides a schematic representation of three different variants of the 1603 1604 Arl17 domain architecture (correspondence to specific proteins in the tree is indicated by the asterisks). The exact architecture of the Ch. velia Arl17b protein could not be determine due to incompleteness of 1605 1606 the genome assembly.



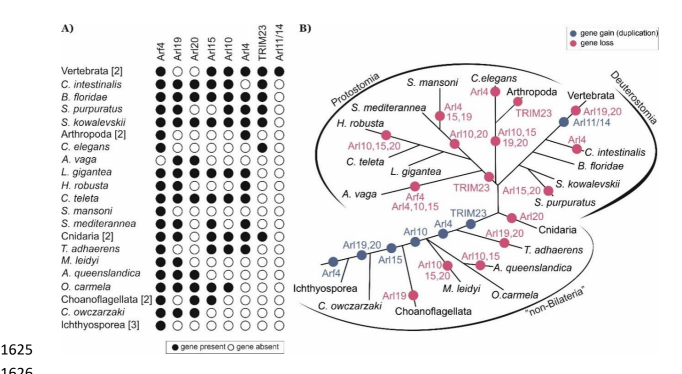


1609 Fig. 3. Retention of ancient paralogs of the ARF family in main lineages of eukaryotes. (A) Black circle: the paralog is present in at least one member of the lineage. White circle: the gene is absent 1610 from the lineage (evidenced by genome sequence data). Grey circle: the gene was not found in the 1611 transcriptome data available (lineages with transcriptome assemblies only). The hashtag (#) indicates 1612 1613 the number of species included in the analysis. (B) Gene gains (blue circles) and losses (pink circles) mapped onto the eukaryote phylogeny. Only duplications specific to whole lineages listed in the picture 1614 are considered. The acquisition of Arl17 via HGT in rotifers (here represented by A. vaga) is indicated 1615 with a blue circle with an asterisk within. 1616



1617 1618

- 1619 Fig. 4. Maximum likelihood phylogenetic tree of the ARF family based on a ScrollSaw dataset in
- 1620 Holozoa. The tree was inferred using IQ-TREE with LG+I+G4 model (the model selected by the
- 1621 program itself) from a multiple alignment of 323 protein sequences with the ultrafast bootstrap algorithm
- and the SH-aLRT test (both 10000 replicates), as described under Materials and Methods. Dots at
- 1623 branches represent bootstrap values as indicated in the legend shown in the bottom left. Eukaryotic
- ancestral paralogs are collapsed as triangles.



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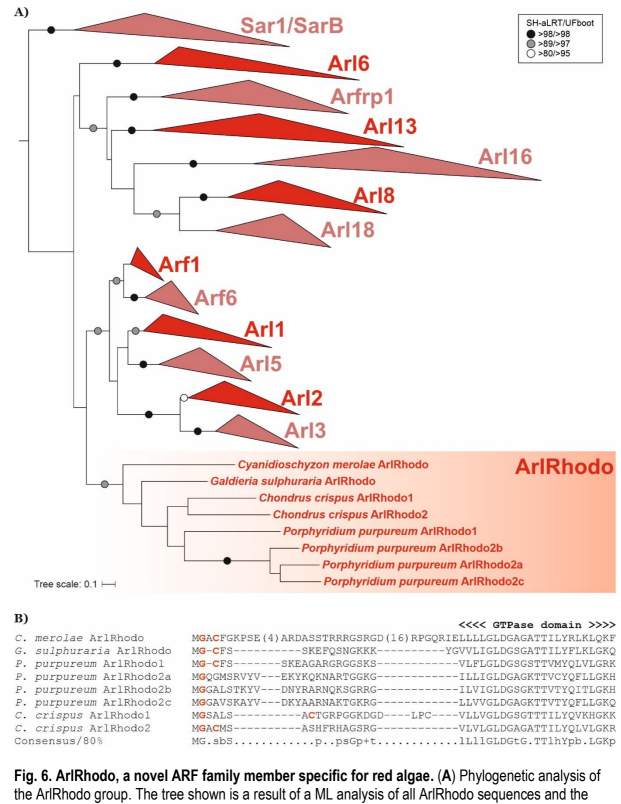
Fig. 5. Retention of lineage-specific paralogs of the ARF family in main lineages of Holozoa. (A) 1627

Black circle: the paralog is present in at least one member of the lineage; white circle: the gene is 1628

absent from the lineage (evidenced by genome sequence data). Species with identical distribution are 1629

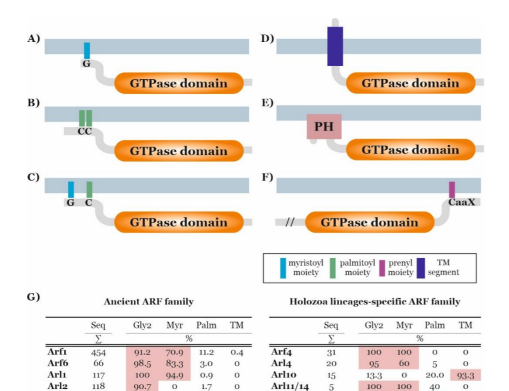
collapsed into higher taxa with the number of species indicated in the square brackets. (B) Gene gains 1630

(blue circles) and losses (pink circles) mapped onto the holozoan phylogeny. 1631



**Fig. 6. ArlRhodo, a novel ARF family member specific for red algae.** (**A**) Phylogenetic analysis of the ArlRhodo group. The tree shown is a result of a ML analysis of all ArlRhodo sequences and the reduced "scrollsawed" dataset (altogether 356 sequences). The tree was inferred using IQ-TREE with LG+I+G4 model (the model selected by the program itself) with the ultrafast bootstrap algorithm and the SH-aLRT test (both 10,000 replicates). Dots at branches represent bootstrap values as indicated in the graphical legend (top right). (**B**) N-terminal region of ArlRhodo proteins with the characteristic configuration of glycine and cysteine residues (highlighted in red) predicted to be N-myristoylated and S-palmitoylated, respectively.

1632 1633



1.8

0

1.1

0

0

2

0

0

0

0.6

0

100

6.4

14.5

12.5

0.9

55.7

54

15.4

21.4

9.8

1.9

0

20.3

Arl15

Arl19

Arl20

TRIM23

14

44

19

0

Seq

16

8

Chloroplastida ArfB

Glaucophyta Arl13L 6

Rhodophyta ArlRhodo

92.9

18

5

33.3

Palm

0

62.5

0

0

0

0

0

TM

0

12.5

0

7.1

77.3

36.8

0

Myr

100

0

%

14.3

100

89.5

0

Archaeplastida lineages-specific

**ARF** family

Glv2

100

100

16.7

Arl3

Arl5

Arl6

Arl8

Arl13

Arl<sub>16</sub>

Arl<sub>17</sub>

Arl<sub>18</sub>

Arfrp

Sarı

SarB

SRB

110

83

88

100

70

50

26

28

112

161

15

118

89.1

100

77

21.1

24.3

6

26.9

100

1.8

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0

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8.2

78

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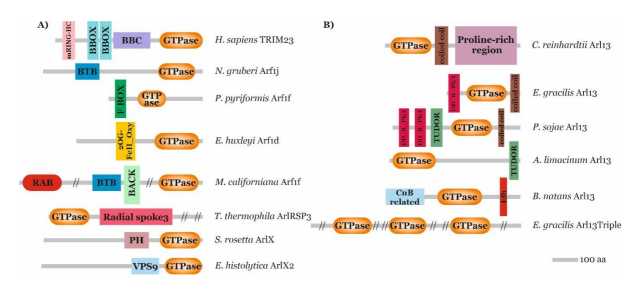
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1643

Fig. 7. Membrane attachment mechanisms of ARF family proteins. Examples of different broadly 1644 conserved mechanisms of membrane attachments of ARF family members are depicted. (A) N-1645 1646 terminally myristoylated glycine residues, common for Arfs and several Arf-like proteins. (B) One or two S-palmitoylated cysteine residues near the N-terminus, typical for Arl16 and also common in Arl13. (C) 1647 N-terminally myristoylated glycine residue coupled with S-palmitoylated cysteine residue near the N-1648 1649 terminus, typical for ArlRhodo. (D) N-terminal transmembrane region, typical for SRβ and Arl10. (E) N-1650 terminally accreted PH domain, present in divergent Arf-like proteins in kinetoplastids and choanoflagellates. (F) Prenvlation motif (CaaX) at the C-terminus of certain eustigmatophyte-specific 1651 ARF family members (characterized also by a long N-terminal extension, in the figure marked with "//"). 1652 supplementary table 1. Supplementary Material online lists all identified ARF family proteins predicted 1653 to be N-myristoylated or S-palmitoylated, or to contain a transmembrane region or PH domain. (G) 1654 Summary of the results of prediction of N-myristoylation, S-palmitoylation and presence of the 1655 transmembrane (TM) region in particular subgroups of the ARF family. For each subgroup (group of 1656 orthologs), the number of sequences (Seq) and the percentages of sequences with glycine residues at 1657 1658 the second position (Glv2), sequences predicted as N-myristoylated (Myr), sequences predicted as Spalmitoylated on at least one cysteine residue (Palm), and sequences with predicted transmembrane 1659 1660 region(s) (TM) are given. Values above 50% are highlighted in pink. For complete data see supplementary tables 1 and 5, Supplementary Material online. These predictions were done as 1661 described under Materials and Methods. 1662



1663 1664

1665 Fig. 8. Multi-domain architectures of ARF family proteins. (A) Examples of lineage-specific ARF

1666 family proteins with extra domains accreted to the GTPase domain. Sequence IDs of the proteins listed

are provided in supplementary table 1, Supplementary Material online. (B) Variation in the domain

architecture of Arl13 proteins across the eukaryote diversity. The Arl13 from *Chlamydomonas* 

1669 *reinhardtii* represents the most common and presumably ancestral state.