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# Multiple Roots of Fruiting Body Formation in Amoebozoa

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**Data deposition:** The genome sequence and gene predictions of *Protostelium aurantium* and *Protostelium mycophagum* were deposited in GenBank under the Accession Numbers MDYQ00000000 and MZNV00000000, respectively. The mitochondrial genome of *P. mycophagum* was deposited under the Accession number KY75056 and that of *P. aurantium* under the Accession number KY75057. The RNAseq reads can be found in Bioproject Accession PRJNA338377. All sequence and annotation data are also available directly from the authors. The *P. aurantium* strain is deposited in the Jena Microbial Resource Collection (JMRC) under accession number SF0012540.

## Abstract

Establishment of multicellularity represents a major transition in eukaryote evolution. A subgroup of Amoebozoa, the dictyosteliids, has evolved a relatively simple aggregative multicellular stage resulting in a fruiting body supported by a stalk. Protosteloid amoeba, which are scattered throughout the amoebozoan tree, differ by producing only one or few single stalked spores. Thus, one obvious difference in the developmental cycle of protosteliids and dictyosteliids seems to be the establishment of multicellularity. To separate spore development from multicellular interactions, we compared the genome and transcriptome of a *Protostelium* species (*Protostelium aurantium* var. *fungivorum*) with those of social and solitary members of the Amoebozoa. During fruiting body formation nearly 4,000 genes, corresponding to specific pathways required for differentiation processes, are upregulated. A comparison with genes involved in the development of dictyosteliids revealed conservation of >500 genes, but most of them are also present in *Acanthamoeba castellanii* for which fruiting bodies have not been documented. Moreover, expression regulation of those genes differs between *P. aurantium* and *Dictyostelium discoideum*. Within Amoebozoa differentiation to fruiting bodies is common, but our current genome analysis suggests that protosteliids and dictyosteliids used different routes to achieve this. Most remarkable is both the large repertoire and diversity between species in genes that mediate environmental sensing and signal processing. This likely reflects an immense adaptability of the single cell stage to varying environmental conditions. We surmise that this signaling repertoire provided sufficient building blocks to accommodate the relatively simple demands for cell–cell communication in the early multicellular forms.

**Key words:** multicellular development, transcriptome, *Protostelium*, Dictyostelia, evolution of development, signaling, Amoebozoa.

## Introduction

Multicellularity was independently acquired in different branches of eukaryotes. This transition from uni-

to multicellular life styles may therefore have been relatively easy achieved compared with, for example, the rare event of organelle capture (Marin et al. 2005; Nowack et al.

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2011). It is unlikely that in each case a new way to this transition to multicellularity was invented. Rather, building blocks already present in unicellular eukaryotes (and their last common ancestor; LCA) were likely recruited to this new task. During evolution, new layers of complexity (i.e., increase of the numbers of cell types) masked these building blocks (Hedges et al. 2004) so that we have difficulties to define them. Genomes of the unicellular ancestors of Metazoa seem to have been already complex. Exaptations of genes were later used for the evolution of multicellular development (Grau-Bove et al. 2017). Interestingly, *Capsaspora owczarzaki*, which can transit to an aggregative state, employs genes for this transition, the orthologs of which establish tissue architectures in Metazoa (Sebe-Pedros et al. 2013; Suga et al. 2013). Thus, aggregative multicellularity seems to have the same roots as true multicellularity at least in this evolutionary branch. Moreover, regulatory changes rather than gene innovations seemed to have played a major role in the establishment of multicellularity in Metazoa (Sebe-Pedros et al. 2016).

There seems to be no significant genomic difference between the unicellular algae *Chlamydomonas* and the multicellular *Volvox* species (Umen and Olson 2012). However, extracellular matrix proteins seem to be important for the establishment of multicellularity in this clade (Prochnik et al. 2010).

Each eukaryote branch or even closely related clades seem to follow its own route to multicellular development and have established their own toolkit for this purpose (Niklas and Newman 2013).

Multicellular organisms generally go through a unicellular spore or fertilized egg stage at least once, which then divides repeatedly to generate the multicellular form. In animals, plants, and fungi, the dividing cells remain attached to each other, representing clonal, multicellular organisms. However, in many other multicellular organisms, dispersed cells can come together when food runs out to construct a fruiting body with aerially borne spores. This type of aggregative multicellularity was invented independently by *Acrasis* spp. in the eukaryote division Discoba (Brown, Silberman, et al. 2012; He et al. 2014), *Fonticula alba* in Holomycota (Brown et al. 2009), *Guttulinopsis* spp. in Rhizaria (Brown, Kolisko, et al. 2012), *Sorodiplophrys stercorea* in Stramenophiles (Tice, Silberman, et al. 2016), *Sorogena stioanovitchae* in Alveolata (Lasek-Nesselquist and Katz 2001), and *Copromyxa protea* and *Dictyostelium* spp. in Amoebozoa (Baldauf et al. 2000; Brown et al. 2011).

Among these organisms, the genetic model system *Dictyostelium discoideum* and its close relatives display the most complex form of aggregative multicellularity with a freely migrating “slug” stage and four to five specialized cell types. Its genome and those of species representative of the five major clades of dictyosteliids have been sequenced (Heidel et al. 2011; Glöckner 2015; Glöckner et al. 2016) and within Amoebozoa, sequenced genomes are also available for the strictly unicellular *Acanthamoeba castellanii* (Clarke et al.

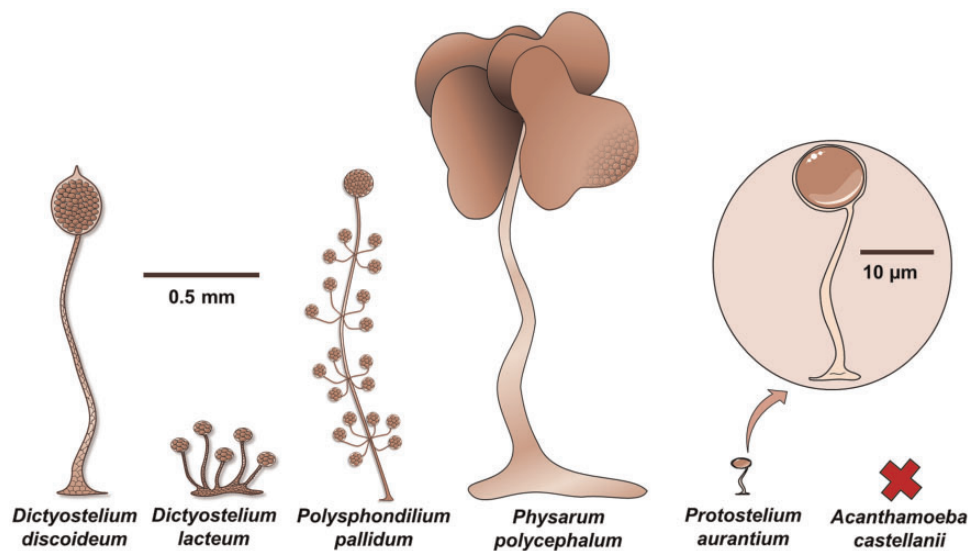
2013) and the syncytial slime mold *Physarum polycephalum* (Schaap et al. 2016). The latter organism alternates between unicellular amoeboid or amoeboflagellate forms, which can either individually encyst or, after mating, transform into a multinuclear syncytium. This syncytium has the choice of forming a dehydrated but viable sclerotium, or fruiting bodies (fig. 1) with multiple spores and acellular stalks (Schaap et al. 2016).

We previously investigated the extent to which genes that are essential for multicellular development of *D. discoideum* are conserved throughout Dictyostelia and unicellular Amoebozoa. This study showed that genes involved in intracellular signal processing were mostly common to both multicellular and unicellular Amoebozoa, but that genes encoding membrane or secreted proteins were more unique to Dictyostelia. This suggested that novel adhesion proteins, signal, and signal sensors are prime requisites for evolving multicellularity. How these were acquired remains often elusive, but of the five known nonpeptide signals which induce cell differentiation in *D. discoideum*, three genes encoding their biosynthetic enzymes were likely to be acquired by Dictyostelia through lateral gene transfer (Glöckner et al. 2016). A parallel retracing genes with conserved upregulation in multicellular development highlighted many conserved genes with likely roles in multicellular development that have not yet been functionally characterized (Schilde et al. 2016).

Multicellular development of Dictyostelia and other organisms depends not only on cell-type specialization but also on coordination of cell movement and/or cell division to generate form. This complicates identification of the mechanisms that uniquely regulate cell differentiation. Comparison of Dictyostelia with protosteloid amoebae, that lack this complication, may uniquely identify the genes involved in differentiation processes.

Fruiting bodies of diverse morphologies are widespread among the Amoebozoa, and the protosteloid type of fruiting body with a stalked spore, or very few spores, but always derived from one cell is the most common (fig. 1). A recent study presented a well resolved tree of Amoebozoa with life cycle characters mapped onto that tree (Kang et al. 2017). Species displaying protosteloid fruiting bodies were found to be scattered throughout the Amoebozoa phylogeny, but also occur within the Eumycetozoa which include the Dictyostelia and *Ph. polycephalum*. Recently, single cell fruiting bodies were also found to occur among the more distant *Acanthamoebidae* (Tice, Shadwick, et al. 2016). It is hypothesized that fruiting body formation was already present in the last common ancestor of all Amoebozoa (Kang et al. 2017), but an independent origin of fruiting body formation in different lineages is also possible (Cavalier-Smith et al. 2016).

We here compare the genome content of *Protostelium aurantium* var. *fungivorum* and *Protostelium mycophagum* with those of Dictyostelia, *Ph. polycephalum*, and *Acanthamoeba castellanii*. We further analyze the



**Fig. 1.**—Amoebozoa fruiting bodies. Schematic view of the different “fruiting bodies” in the Amoebozoa. Depicted are only species where a genome sequence is available including the here presented *Protostelium* spp. genomes. The fruiting bodies are drawn to scale, the inset shows an enlargement of the tiny *Protostelium* fruiting body. No fruiting body is known for *Acanthamoeba castellanii*, indicated by a cross. However, other *Acanthamoebae* do form sporocarps, as was recently discovered for *A. pyriformis* (Tice, Shadwick, et al. 2016).

transcriptome changes that occur in *Protostelium aurantium* var. *fungivorum* (further addressed as *P. aurantium*) during its transition from a feeding amoeba into a mature fruiting body. Our study suggests that fruiting body formation seemed to have evolved analogously from the similar toolbox of genes. Furthermore, despite their simple life style, *Protostelium* spp. already display a large variety of cell signaling genes.

## Materials and Methods

### Species Isolation and Maintenance

Protosteloid amoebae were maintained as spore suspensions in 86% [v/v] glycerol at  $-80^{\circ}\text{C}$ . The type strain of *Protostelium mycophaga* Olive and Stoianovitch was obtained from the Culture Collection of Algae and Protozoa (CCAP, Argyll, United Kingdom) and from the American Type Culture Collection (ATCC, Manassas, VA). *Protostelium aurantium* var. *fungivorum* was isolated as described for protosteliids by Spiegel et al. (2006) with slight modifications. Briefly, beech leaves were cut to  $1\text{ cm}^2$  and centrally placed on the surface of buffered wMY agar ( $0.002\text{ g l}^{-1}$  malt extract,  $0.002\text{ g l}^{-1}$  yeast extract,  $2\text{ mM}$  potassium phosphate buffer pH 6.4 [PB], and  $15\text{ g l}^{-1}$  agar) in petri dishes. The basidiomycetous yeast *Rhodotorula mucilaginosa* DSM70404 was obtained from the Jena Microbial Resource collection and was streaked in a distance of 5 mm to the leaf pieces to promote the outgrowth of fungivorous protosteloid amoeba. Plates were sealed with parafilm and incubated at  $22^{\circ}\text{C}$  for 10–14 days with daily microscopic inspections for trophozoite growth or the formation of fruiting bodies. Isolation was achieved by four consecutive transfers of single amoeba or

sporocarps (fruiting body of a single cell) to new petri dishes. Liquid cultures of *Protostelium* sp. were carried out in petri-dishes filled with 2 mM PB buffer (pH 6.4) and *R. mucilaginosa* as the only food source. Cell numbers for both organisms were determined either using a hemocytometer or an automatic cell counter (Casy TT Cell Counter, OLS Bio, Bremen, Germany).

### Induction of the Developmental Cycle and Preparation of Samples for RNA Isolation

Protosteloid amoebae were cultivated in liquid cultures with successive additions of *R. mucilaginosa* as a food source. For *P. aurantium*, the number of trophozoites reached confluency at 48- to 60-h postinoculation. At this time point, the growth medium was aspirated and residual yeast cells were removed by two consecutive washes with PB. Adherent amoebae were scraped from the surface, resuspended in PB and transferred to a wMY-agar surface in  $10\text{ }\mu\text{l}$  droplets. To obtain sufficient material for RNA isolation, cells from 40  $10\text{-}\mu\text{l}$  droplets were pooled for each time point in one of the three biological replicates. Vegetative cells were harvested from plates directly after plating (timepoint 0 h). Starved cells were harvested after 1.5 h and nearly completed fruiting bodies were harvested from plates incubated for 8 h.

### Nucleic Acids Isolation and Sequencing

Chromosomal DNA was isolated from liquid cultures of protosteloid amoebae. Adherent cells were washed in phosphate buffer and harvested by centrifugation at  $200\times g$  which was sufficient to deplete nearly all residual yeast cells. Trophozoites

were lysed in 50 mM Tris–HCl (pH 7.5) with 2% [w/v] of SDS and 0.5 mM Na<sub>2</sub>EDTA, followed by a 1-h incubation at 60°C with RNAase A (Sigma–Aldrich, Taufkirchen, Germany) and Proteinase K (Sigma–Aldrich) at final concentrations of 100 and 200 µg ml<sup>-1</sup>, respectively. Further purification steps like extractions with phenol-chloroform and precipitation with isopropanol were carried out as described (Sambrook and Russell 2001).

For the isolation of total RNA, cells were rapidly harvested from the agar surface, resuspended in 2 ml of phosphate buffer, and centrifuged for 1 min at 6,000×g. Pellets were shock-frozen in liquid N<sub>2</sub> and stored –80°C. RNA was prepared by phenol extraction with subsequent additions of 500 µl of TRIsure (Bioline, Luckenwalde, Germany) and 200 µl of chloroform. Phase separation was achieved by centrifugation followed by an additional extraction with 500 µl of chloroform. Precipitation was as for DNA. RNA quantity and impurities were determined spectrophotometrically. Integrity of the isolated RNA was checked by agarose gel electrophoresis

### DNA Sequencing, Assembly, and Mapping

Library preparation was done using Illumina's TruSeq DNA PCR free library preparation kit following the manufacturer's description for 550-bp insert libraries. Library quality check and quantification was done as described earlier. Sequencing was done in one lane of a HiSeq 2500 running in paired-end/2× 101 cycle/rapid mode. The reads derived from genomic DNA were assembled using abyss-pe with a kmer size of 45–65 with step increments of 2. The assembly sizes for the *P. aurantium* genome (excluding contaminants) was in a range from 39 (kmer 45) to 68 Mb (kmer 65). The kmer 45 assembly was chosen for further improvement after examination of contig lengths, coverage, and contribution of bacterial and fungal sequences to the assembly for both species. Specifically, the kmer45 assembly was the most contiguous of all assemblies and therefore represented the genomes best. The GC content of the protosteliid genomes was <50.1% while contaminating DNA from other sources (bacteria and the food fungus) was found to have a higher GC content. Thus, the GC content was a first indicator to readily discriminate between the *Protostelium* spp. genomes and contaminating sequences. Furthermore, all contigs were compared with fungal and bacterial genomes to exclude contaminating sequences from further analysis. Here, we were using BLAST analysis to identify fungal or bacterial sequences (BLASTn in case of bacteria and BLASTx for fungi). If a contig had a higher similarity to known bacterial or fungal sequences than to any Amoebozoa genome, it was excluded from the analysis. Further gap closure was done with PAGIT (<http://www.sanger.ac.uk/science/tools/pagit>; last accessed January 31, 2018) using default values. For assembly of the mitochondrial genome, we extended the contig using SSPACE (Boetzer et al. 2011). The resulting overlap between contig ends was

then removed to yield a complete, closed circle mitochondrial genome.

### RNA Sequencing

Each of the three time points of the RNAseq experiment was covered by three biological replicates. The quality and amount of RNA was initially checked using the Agilent Bioanalyzer 2100 in combination with a RNA 6000 nano kit (both Agilent technologies). The RIN (RNA integrity number) varies from 8.4 to 10 with an average of 9.44. Total RNA of 1 µg was used for library preparation using Illumina's TruSeq RNA sample preparation kit v2 following the manufacturer's description which included individual labeling by sequence barcodes. The libraries were again quantified and quality checked using the Bioanalyzer 2100 and DNA 7500 kit. For sequencing, libraries were pooled, individually labeled, and loaded in two lanes of a HiSeq 2500 running in single-end/51 cycle/high output mode. Data extraction was done in FastQ format using the tool bcl2FastQ v1.8.4 (provided by Illumina). Mapping of these reads to the genome was done with tophat2. The counts per gene were used to calculate differential expression using DEseq2 (Love et al. 2014) with the Bioconductor package of R.

### Gene Predictions

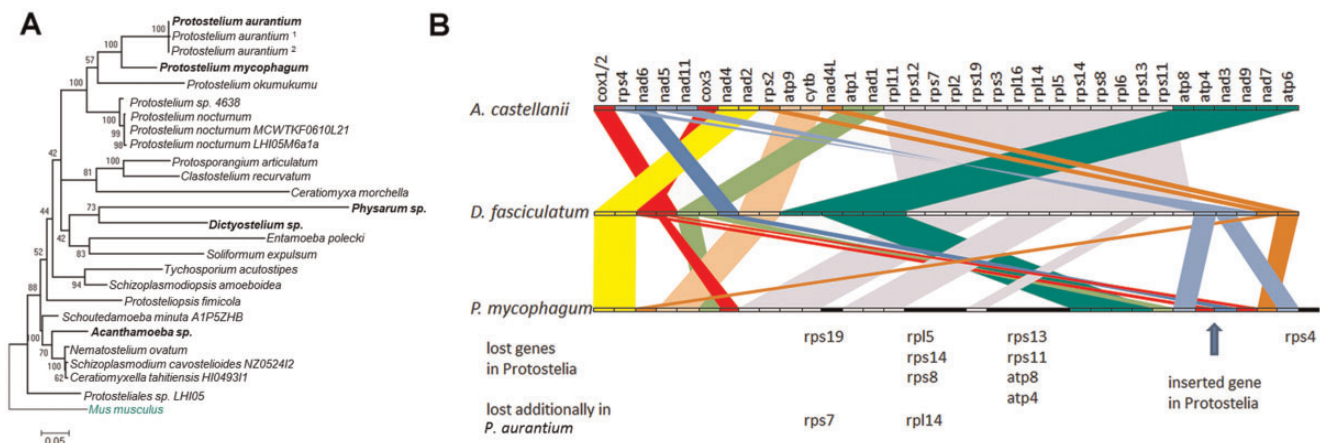
For the annotation of the *P. mycophagum* genome, we used GeneMarkES in self-training mode, which uses no *a priori* knowledge on gene structures. For the annotation of the *P. aurantium* genome, we made use of the RNAseq data, which were mapped to the genome as described earlier. The mapped reads were used to define 52,801 splice sites. These data were used for the GeneMarkET program to enable a better gene prediction.

### ncRNAs

For prediction of nonprotein-coding RNAs (ncRNAs), we used GORAP ([www.rna.uni-jena.de/en/software](http://www.rna.uni-jena.de/en/software); last accessed January 31, 2018), which searches for 2,474 known ncRNA families from Rfam version 12.1 (Nawrocki et al. 2015). GORAP uses improved homology search strategies based on sequence and secondary structures and was run with default settings. All predictions passed Rfam thresholds. All results and files in fasta, gff, and stockholm format are available at <http://www.rna.uni-jena.de/supplements/protostelium>; last accessed January 31, 2018.

### Orthology and Protein Annotation

We defined the orthology between the two *Protostelium* genomes and to the dictyosteliid genomes using OrthoMCL (Li et al. 2003). The protein functions were evaluated by aligning the protein sequences to the protein sequence database obtained from NCBI (version from 5/2016) using BLAST. Domains were searched using InterproScan



**FIG. 2.**—Phylogeny and mitochondrial organization. (A) Phylogeny of selected protosteloid amoeba together with selected other amoebozoa based on 18S rRNA sequences. The species of which the genome sequences were analyzed are given in bold letters. Superscript numbers refer to recently renamed isolates of *Protostelium aurantium* with accession numbers <sup>1</sup>FJ766461.1 and <sup>2</sup>FJ766463.1. The alignment was cleaned of ambiguous positions using GBLOCKS (Talavera et al. 2007) reducing the observable distance between species and clades. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories [+G, parameter = 0.6214]). The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. Evolutionary analyses were conducted in MEGA6 (Tamura et al. 2013). (B) Mitochondrial syntenic organization between selected Amoebozoa. Genes are drawn as boxes of equal length. Syntenic regions are connected by colored bands.

(Jones et al. 2014). The biochemical pathways were analyzed using the KAAS server (Moriya et al. 2007). All programs were run with default values.

### Phylogenetic Analysis of Adenylate and Guanylate Cyclases

The output of an Interproscan of the translated *Protostelium aurantium* transcripts was scanned for proteins harboring Interpro domain IPR001054 for class III adenylate and guanylate cyclases. The *P. aurantium* proteins harboring this domain were aligned with the five *Dictyostelium* cyclases and a few signature *Ph. polycephalum* and *A. castellanii* cyclases using Clustal Omega with five combined iterations (Sievers and Higgins 2014). After deletion of segments that did not align unambiguously, alignments were subjected to Bayesian inference for phylogeny reconstruction using a mixed amino acid model in MrBayes software (Ronquist and Huelsenbeck 2003). Analyses were run until convergence, and trees were rooted at midpoint (Hess and De Moares Russo 2007). For the earliest diverging branches of each major clade, closest homologs were identified by BLASTp of all NCBI nonredundant sequences. These protein sequences were aligned and included in Bayesian inference of the final tree.

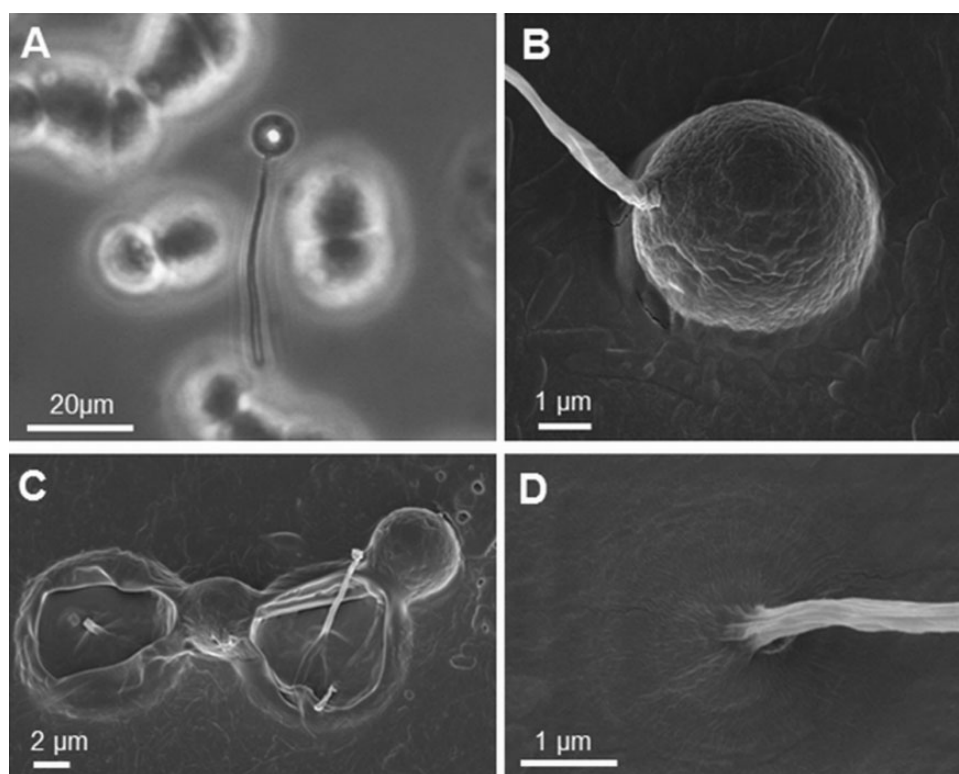
## Results and Discussion

### The Species

*Protostelium mycophagum*, the type strain (Olive and Stoianovitch 1969) was maintained in culture together with the fungus *Rhodotorula mucilaginosa*. As fruiting body formation in this *P. mycophagum* strain occurred at lower

frequencies, and growth was comparably slow, we decided to isolate a wild strain by means of food choice selection, that is, incubating environmental samples together with *Rhodotorula* cultures (Materials and Methods). This way we isolated a protosteloid amoeba in sensu Spiegel (Spiegel et al. 1994, 2006; Shadwick et al. 2009) from a dead aerial beech leaf in the Jena forest. The amoeba revealed striking similarities to the type strain of *Protostelium mycophagum* as described previously (Olive and Stoianovitch 1960). Phylogenetic analysis based on its SSU rRNA (fig. 2A) as well as diagnostic morphological traits identified the isolate as a strain of *Protostelium aurantium* as recently described by Shadwick et al. (2017). Due to its rapid phagocytic feeding on the basidiomycetous yeast *Rhodotorula mucilaginosa* on solid agar surfaces (supplementary data set S1, fig. S1A, Supplementary Material online), we coined this new isolate *Protostelium aurantium* var. *fungivorum*. A detailed taxonomic description of this isolate is available within the supplementary material of this article (supplementary data set S1, Description of *Protostelium aurantium*, Supplementary Material online).

Most characteristic for its identification as a protosteloid amoeba was the abundant formation of globose fruiting bodies carrying a single spore (fig. 3A–C). The sporocarp (fruiting body of a single cell) culminated on an acellular stalk fixed on the substratum (fig. 3D). As described for *P. mycophagum*, asynchronous fruiting body formation occurred on solid surfaces in the absence of the fungal food source. The majority of cells initiated their full developmental cycle 2–4 h following the onset of starvation, passing through the formation of immobile discs, cellular condensation, the formation of the sporocarp, and stalk elevation, with each stage being



**Fig. 3.**—Unicellular fruiting bodies of *Protostelium aurantium*. Light micrograph of a stalked fruiting body (A) and scanning electron micrographs of a tilted, but intact fruiting body (B), which will eventually open and liberate single spores (C), and the rigid base of the stalk (D).

completed in 10–15 min (supplementary data set S1, fig. S2, Supplementary Material online).

### Phylogeny of the Species and Their Genomes

For phylogenetic reconstruction, we sampled 18S RNA sequences of Amoebozoa representatives from the NCBI database, aligned them, and performed a maximum likelihood analysis (fig. 2A). We rooted our tree with the 18S gene from *Mus musculus*. The tree readily identified our environmental isolate as a member of the *Protostelium aurantium* clade and revealed its distant relationship to Dictyostelia and *Physarum*. Despite their phylogenetic distance within the Amoebozoa, all three species share the ability to construct a fruiting body. As gene expression during fruiting body formation of dictyosteliids has been particularly well studied and *P. mycophagum* did not sporulate readily, we focused mainly on the *P. aurantium* genome and expression data and its comparison to Dictyostelia.

Based on our sequencing approach, we assembled the complete mitochondrial genomes and obtained draft nuclear genomes for both *Protostelium* species. The mitochondrial genomes are comparable with those in dictyosteliids with respect to coding capacity and size. The order of genes differs between the two species only for one segment of genes and a few tRNA locus rearrangements (fig. 2B). Furthermore,

*P. aurantium* lacks the genes *rps7* and *rpl14*, located in the rearranged segment. Comparisons to other Amoebozoa mitochondrial genomes show that these additionally lost genes lie in the same region where other ribosomal subunit genes were lost in *Protostelium* spp. (fig. 2B). Thus, loss of such mitochondrial genes seems to be common in the *Protostelium* spp. analyzed.

Based on the gene prediction, both nuclear haploid genomes are comparable in size and also encode a similar number of genes (table 1). For *P. aurantium*, this number was further supported by RNAseq data (supplementary data set S1, fig. S4, Supplementary Material online) using GeneMarkET (Borodovsky and Lomsadze 2011). A distinguishing hallmark of both genomes is the high number of introns per gene (supplementary data set S1, fig. S5, Supplementary Material online). The median intron size is very small with 40–50 bases, but larger than the smallest intron sizes reported so far (Slabodnick et al. 2017). Overall, the *Protostelium* spp. genomes analyzed here harbor circa 30% more genes than Dictyostelia.

In line with the extensive use of introns, we found not only all small nuclear RNAs (snRNAs) of the major spliceosome but also the minor spliceosome to be present in both species. In total, a similar number of noncoding RNAs were annotated for *P. mycophagum* (590) and *P. aurantium* (525) (supplementary data set S1, table S1, Supplementary Material online).

**Table 1**Genomic Features of the Two *Protostelium* Species

	<i>Protostelium mycophagum</i>	<i>Protostelium aurantium</i>
Genome size in Mb (haploid)	39.7	38.3
Scaffolds	2,578	822
Mean GC content	42.2	47.9
Palindrome size (kb)	13.4	16.3
Mitochondrion (kb)	48.6	44.5
Protein coding genes	15,691	17,172
Mean size CDS/AA	1,386/462	1,512/504
tRNAs	426	405
Introns	126,081	145,637
Mean intron length	83.2	63.4
Median intron length	50	46
Introns/gene	8	8.5
Intron space (Mb)	10.5	9.2
CDS space (Mb)	21.8	26

The amount of mobile elements in *P. aurantium* is estimated at ~0.13% of the genome. Although several reverse transcriptase domains of both long terminal repeat (LTR) and non-LTR retrotransposons could be determined, the corresponding elements are highly fragmented and mostly present in single copy. Only four LTR retrotransposons could be analyzed in some detail, and they belong to the Skipper and DGLT-A families of Ty3/gypsy type of LTR retrotransposons known from the dictyosteliid clade (Spaller et al. 2016).

### Gene Families and Orthology Relationships

Both *Protostelium* spp. share among them 6,278 1:1 orthologs and 1,708 gene families with more than one member in at least one species. *Protostelium mycophagum* has 6,264 genes not shared with *P. aurantium*, whereas *P. aurantium* has 5,942 such orphan genes, meaning that they were also not found in other species. These numbers are somewhat comparable to the 4,156 orphan genes reported for *A. castellanii* (Clarke et al. 2013). Interestingly, several of the largest gene families in both species encode proteins with Leucine-rich repeats, which are often involved in protein–protein interactions. Other large gene families comprise Ankyrin repeat containing proteins, nucleotidyl cyclases, or kinases (supplementary data set S1, table S2, Supplementary Material online).

### Gene Regulation during Fruiting Body Formation

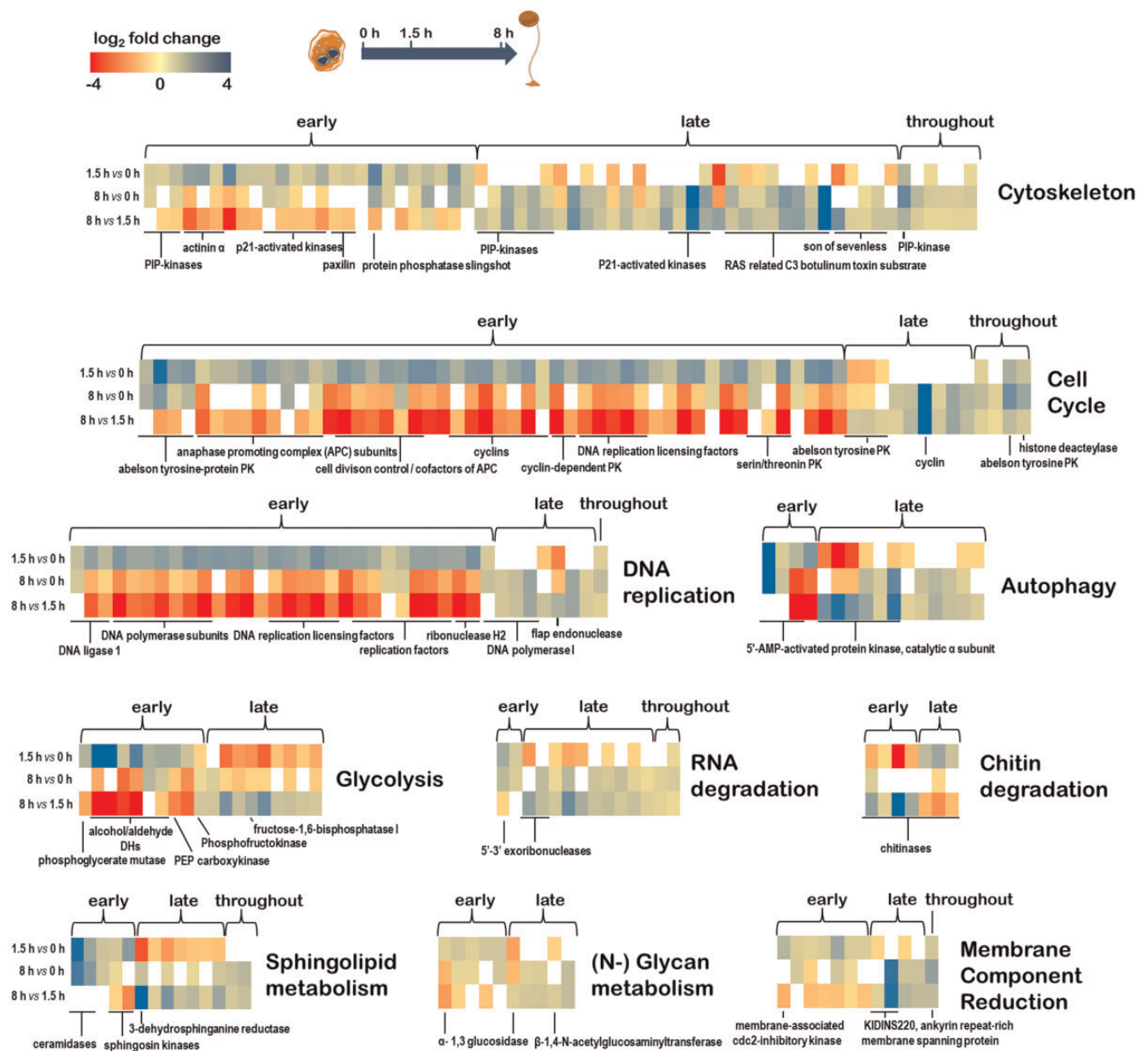
Upon starvation on a solid substratum, *P. aurantium* cells individually differentiate into a spore supported by an acellular stalk. As *P. aurantium* displayed a higher frequency and abundance of these structures relative to *P. mycophagum*, we chose *P. aurantium* to analyze the transcriptional changes during this differentiation process by RNAseq over three time

points (vegetative cells at 0 h, starvation and early differentiation at 1.5 h, and the formation of prespores and mature fruiting bodies at 8 h). Although sporocarp formation was asynchronous among the population, these time points were discernible via the predominant morphology (supplementary data set S1, figs. S2 and S6A, Supplementary Material online). We first assessed which genes are potentially differentially expressed between any of those time points. The biological replicates showed little variation (supplementary data set S1, fig. S6B and C, Supplementary Material online). The DESeq2 analysis (Love et al. 2014) implemented in the BioConductor R package revealed that there are 7,787 genes significantly downregulated and 3,980 genes significantly upregulated. An example of an upregulated gene is given in supplementary data set S1, figure S7, Supplementary Material online. A significance threshold of 0.001 reduces these numbers to 6,579 and 2,685, respectively. We next analyzed which potential functions are encoded in this smaller gene set. This analysis revealed extensive reprogramming of several functions during the developmental cycle (fig. 4). A gene ontology (GO) term enrichment analysis of all upregulated genes using Goterfinder (Boyle et al. 2004) showed that in the GO category “molecular function” ATP and GTP binding proteins are strongly overrepresented (supplementary data set S1, fig. S8, Supplementary Material online). However, no GO terms in the category “biological process” were significantly enriched.

We next divided the genes into three categories (downregulation, early upregulation, and late upregulation). This only roughly approximates the real transcript dynamics since we had only three distinct developmental stages in our time course. The KEGG pathway analysis (Moriya et al. 2007) revealed that up- and downregulated genes can be part of the same pathway and concerted regulation may affect only part of a pathway (fig. 5 and supplementary table S1, Supplementary Material online). Processes related to RNA generation, fatty acid synthesis, RNA and protein transport, and oxidative phosphorylation were uniformly downregulated. This downregulation is a likely hallmark of the fact that the cells are experiencing starvation. Genes that were upregulated early tended to be involved in replication and repair, dNTP synthesis, and fatty acid degradation. These early events likely reflect the fact that the cell tries to recruit energy from fatty acids to complete the developmental cycle. In a later stage, *P. aurantium* devoted resources to the production of membrane components like glycerophospho- and sphingolipids, as well as chitin as a cell wall polysaccharide, indicative of the final stages of spore formation (fig. 5).

We also found that signal transduction cascades that induce cytoskeleton reorganization, survival, adhesion, and migration were upregulated early and this upregulation increased further in differentiation. The second messenger cAMP seems to play a pivotal role here since its production

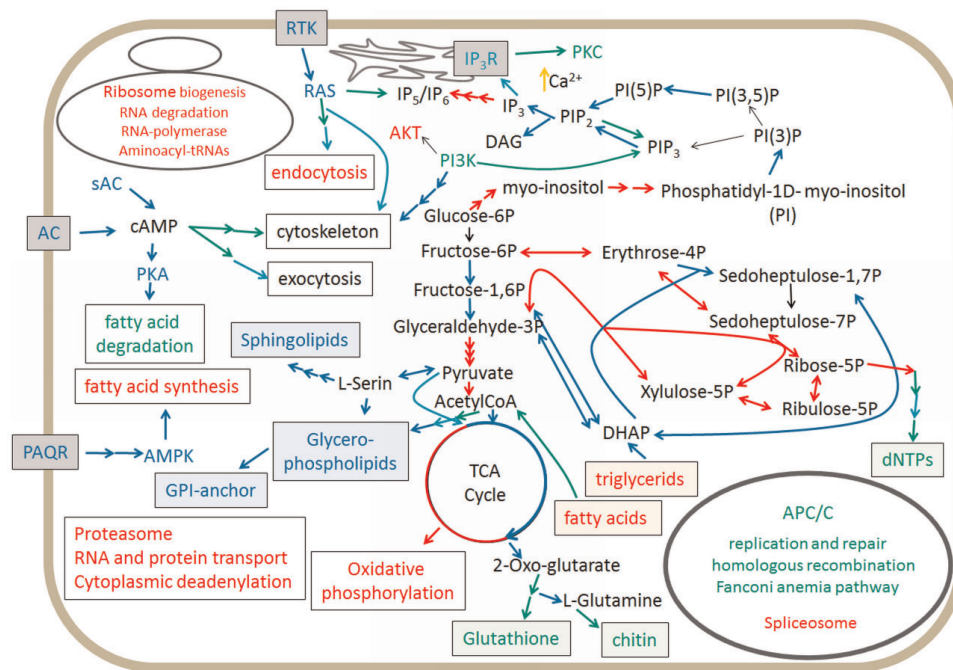




**FIG. 4.**—Transcriptional reprogramming during the developmental cycle of *Protostelium aurantium*. Heat maps indicate higher (blue) and lower (red) expression of genes from major developmental and metabolic pathways when comparing early (0 min vs. 1.5 h, top line) or late stages (0 vs. 8 h, middle line, or 1.5 h vs. 8 h, bottom line) of development. Genes from each cluster are grouped in three temporal categories representing an upregulated expression either early, late, or throughout the developmental cycle. Heat maps illustrate the  $\log_2$  expression ratio based on the mean RPKM values from three biological replicates for each time point. A more detailed table including gene accession numbers, numeric expression ratios, KEGG orthologies, and predicted protein functions are available as [supplementary table S1, Supplementary Material](#) online.

started early in development ([supplementary data set S1, fig. S9, Supplementary Material](#) online). Which of the numerous cyclase encoded in the genome of *P. aurantium* is responsible for this production or whether several cyclases are acting in a concerted way to achieve this, is currently unclear. However, one of the cyclases (PROFUN\_08491) is particularly upregulated during development ([fig. 6](#)) and therefore might be responsible for the early cAMP level rise. PKA activity, which is

involved in common encystation processes in Amoebozoa (Kawabe et al. 2015) is upregulated, presumably via cAMP. Throughout the differentiation process protein modification pathways (N-glycan biosynthesis) were enhanced. Interestingly, N-Glycan biosynthesis plays also a role in *D. discoideum* differentiation (Li et al. 2015). In *P. aurantium*, this pathway may be involved in production of components of the cell wall and the acellular stalk.



**Fig. 5.**—Developmentally regulated pathways in *Protostelium aurantium* based on RNAseq analysis. Red indicates functions represented by genes that are downregulated upon starvation, green and blue highlight functions that are upregulated in early and late development, respectively. Presumed Calcium upregulation is depicted with a yellow arrow. Functions associated with the nucleus are listed in a circle in the right lower corner, and those with the ribosome in the upper left corner. The cytoskeleton and exocytosis are influenced by the depicted pathways, but the impact of this is not clear.

### Comparison between Social Amoebae and *P. aurantium* Developmental Genes

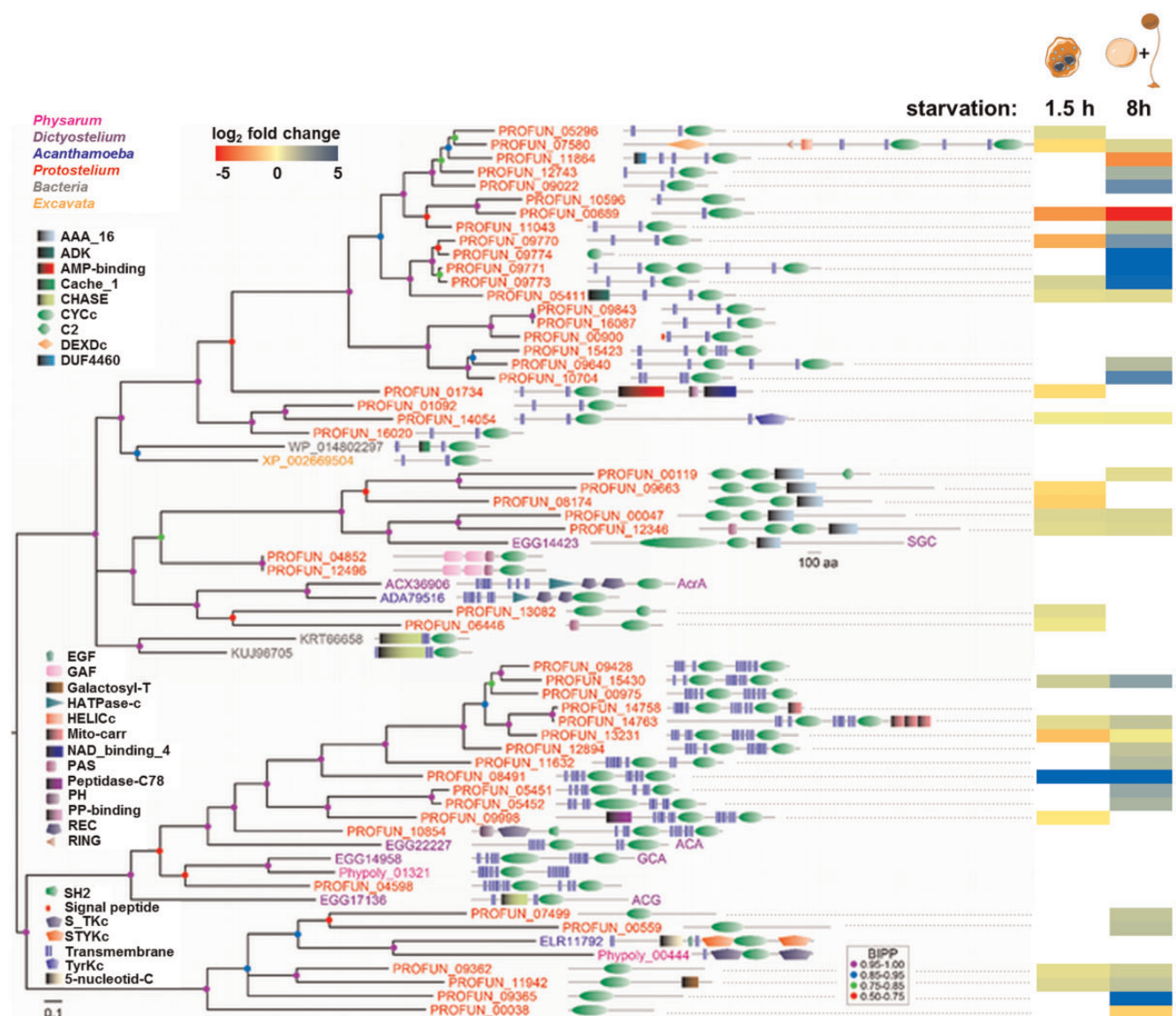
Social amoebae have ~60% orthologs between them as a comparative study showed (Heidel et al. 2011). Since the evolutionary distance between dictyosteliids and the protosteliids analyzed here is even higher than within the dictyosteliids it is not surprising that only 37% of the genes in *P. aurantium* show similarities to genes in *D. discoideum*, as detected by BLAST with a bit score threshold of 200. In two previous studies, we have defined two sets of dictyosteliid genes which are involved in the developmental cycle of social amoebae (Glöckner et al. 2016; Schilde et al. 2016). The set of developmentally essential (DevEs) genes contains 374 genes that upon knock-out cause a developmental defect (Glöckner et al. 2016). The set of developmentally upregulated (DevUp) genes contains 794 genes that are consistently developmentally upregulated across the *Dictyostelium* taxon groups (Schildt et al. 2016). In total, these two sets comprise 1,168 genes (table 2 and supplementary data set S1, fig. S11, Supplementary Material online). A higher percentage of DevEs than DevUp genes has identifiable counterparts in the *P. aurantium* genome (76% vs. 48%). However, of the in total 669 *D. discoideum* proteins with homologs in *P. aurantium*, 172 have the same homolog, indicating gene family expansions in *D. discoideum*. Strikingly, the directionality of developmental regulation of *P. aurantium* counterparts is often opposite (table 2). For example, *statA* with a major role in

*D. discoideum* chemotaxis and stalk formation (Kawata 2011) is upregulated in *D. discoideum*, whereas its *P. aurantium* ortholog (PROFUN\_03920) is slightly downregulated.

Overall, not more than 75 genes were consistently identified as important for development in *D. discoideum* by being upregulated and essential during the developmental process (Glöckner et al. 2016; Schilde et al. 2016). However, for most of these, orthologous genes were found in the genomes of *P. aurantium* (57), *A. castellanii* (47), and *Ph. polycephalum* (57) (table 2). It is therefore conceivable that these genes could belong to the basic toolkit for the evolution of all differentiated forms of development among the Amoebozoa.

The set of in total 233 *D. discoideum* DevUp and DevEs genes with no hits in either the *P. aurantium*, *Ph. polycephalum*, and *A. castellanii* genomes is enriched in several GO terms, for example, “extracellular matrix organization” and “multicellular organismal process” (supplementary data set S1, fig. S10, Supplementary Material online). We next searched for genes in the 1,168 gene set, which are shared exclusively between *D. discoideum* and *P. aurantium* and are not present in *A. castellanii* and *Ph. polycephalum*. This search revealed only 13 genes with a range of putative functions (supplementary data set S1, table S3, Supplementary Material online).

Three of the four genes of *Dictyostelia* (*chlA*, *dgcA*, *dokA*, *iptA*), which are likely derived from horizontal gene transfer from bacteria (Glöckner et al. 2016) and which either sense



**Fig. 6.**—Phylogeny of adenylate and guanylate cyclases. The phylogenetic tree was calculated with Bayesian inference (see Materials and Methods). The protein identifiers are color coded to reflect species names as indicated in the figure, and are annotated with the functional domain architecture of the proteins as determined by SMART (Schultz et al. 1998). PFAM domains are represented as colored to black graded rectangles. Posterior probabilities (BIPP) of tree nodes are indicated by colored dots. The heatmap indicates relative expression levels at 1.5 h (starvation) and 8 h (fruiting body formation) compared with the vegetative state. The cartoon illustrates the predominant cell types at these time points (see [supplementary fig. S7, Supplementary Material](#) online).

(dokA) or synthesize developmental signals, are not present in the *P. aurantium* genome. However, two histidine kinases (PROFUN\_01260 and PROFUN\_15316) have the same domain structure as dokA ([supplementary data set S1, fig. S13, Supplementary Material](#) online) and therefore might encode similar functions even if they are not orthologs.

We also examined, whether transcription factors (TFs) important for the development in Dictyostelia are conserved in *Protostelium* spp. (table 3). Of a total of twelve TFs, orthologous proteins in *P. aurantium* were found for only seven and none of them were highly expressed during development.

Only two TFs which were previously identified in Dictyostelia were slightly conserved only in *P. aurantium* (crtf and stkA) but not found in *Ph. polycephalum* or *A. castellanii*. This indicates a common root for developmental regulation for only few if any TFs and a likely loss in some lineages. Further genome sequences in the Amoebozoa are needed to trace the origin of these TFs to the LCA.

Overall, and somewhat contrary to expectations based on morphological similarities, the conservation of developmental genes between Dictyostelia and *P. aurantium* is low and possible not more than with other unicellular amoebozoan

**Table 2**

Gene Regulation Differences between Social Amoebae, *Protostelium aurantium*, and Other Amoebozoa

Gene Set	<i>Dictyostelium</i> sp.	<i>Protostelium aurantium</i>			Developmental Expression in <i>P. aurantium</i>			<i>Physarum polycephalum</i>	<i>Acanthamoeba castellanii</i>
		Development	Genes with Similarity (score >200)	% of <i>Dictyostelium</i> sp. Genes	Multiple Hits	Up	Down	Neutral	Genes with Similarity (score >200)
Upregulated during development	783	377	48	96	128	199	50	370	310
Essential for development	309	234	76	62	55	138	41	233	193
Upregulated and essential	75	57	78	13	21	21	15	57	47
All	1,167	668	57	171	204	358	106	660	550

NOTE.—About 1,167 genes of *D. discoideum* were identified previously as being either developmentally upregulated (Schilde et al. 2016), developmentally essential (Glöckner et al. 2016), or both (commonly identified in both studies). These genes were searched in the genomes of *P. aurantium*, *Ph. polycephalum*, and *A. castellanii*. To reduce spurious hits a BLAST similarity bit score of 200 was applied when comparing the encoded amino acid sequences. The column “Multiple hits” counts the occasions where different *D. discoideum* genes yielded the same *P. aurantium* hit.

**Table 3**

Developmental TFs from *Dictyostelium discoideum* and Presence of Orthologous Proteins in *Protostelium aurantium*

Gene ID	TF	Best Hit in <i>P. aurantium</i>	Log <sub>2</sub> Change in Expression at 1.5 h, 8.5 h	Comment
DDB_G0278077	crtf	PROFUN_06017	−0.42, 0.45	Expression of aggregation genes; <i>Dictyostelia</i> specific
DDB_G0281387	srfA	PROFUN_09032	−1.13, n.d.	Spore differentiation; MADS box; only box similar
DDB_G0277589	gtaC	PROFUN_10218	0.41, n.d.	pstB cell sorting and basal disc formation; GATA zinc finger; also in <i>Acanthamoeba</i>
DDB_G0278971	dimA	n.d.	−	Development; bZIP transcription factor
DDB_G0291372	dimB	n.d.	−	Development; bZIP transcription factor
DDB_G0279529	bzpf	PROFUN_06600	n.d., 0.58	Spore viability; bZIP transcription factor; common TF
DDB_G0281381	dstA	PROFUN_03920	n.d., −0.29	Culmination, <i>cuda</i> expression; STAT transcription factor; also in <i>A. castellanii</i>
		PROFUN_12834	0.64, n.d.	
		PROFUN_01113	n.d., n.d.	
DDB_G0277147	stkA	PROFUN_04849	n.d., n.d.	Spore formation; GATA zinc finger; <i>dictyosteliid</i> specific
		PROFUN_12502	n.d., n.d.	motif C terminal of zinc finger conserved
DDB_G0284465	cuda	PROFUN_14255	n.d., −0.82	Many prespore genes and for culmination; also in
		PROFUN_08418	n.d., n.d.	<i>A. castellanii</i>
DDB_G0281563	mybC	n.d.	−	Culmination; homology to myb domains only
DDB_G0275445	mybB	n.d.	−	ACA expression; homology to myb domains only
DDB_G0281969	mybE	n.d.	−	Basal disc formation; homology to myb domains only

NOTE.—n.d., orthologous protein or gene expression was not detected.

genomes. Evolution of fruiting body formation could thus have well been independent in the two species. However, analysis of further, phylogenetically more distant, protosteloid amoeba will help to resolve this question.

### Spore and Stalk Genes

The analysis described earlier comprised all known developmentally essential and conserved developmentally upregulated genes, but many of these genes are typically involved in multicellular morphogenesis which has never been observed for *Protostelium* sp. In the following analysis, we

focused entirely on the morphogenetic process that protosteliids and dictyosteliids do have in common, the differentiation of spore(s), supported by a stalk. Although the stalk differs from that of most *Dictyostelia* by not containing any cells, there is also a *Dictyostelium* clade, the acytosteliids, which lack cells in their stalk, but nevertheless have the same stalk-specifying genes as the species with stalk cells (Urushihara et al. 2015).

About 92 genes with roles in spore and stalk formation have been identified in *Dictyostelium discoideum* (Glöckner et al. 2016) and we investigated whether orthologs of these genes were also present in *P. aurantium*, *Ph. polycephalum*,

**Table 4**Conserved *Dictyostelium discoideum* Spore and Stalk Genes in Three Amoebozoa

Required for	<i>Dictyostelium discoideum</i>	<i>Protostelium aurantium</i>	<i>Physarum polycephalum</i>	<i>Acanthamoeba castellanii</i>
Sporulation	72	26	40	32
Stalk formation	20	2	6	2

and *A. castellanii*. Genes in the three species were validated as *D. discoideum* orthologs when 1) the *D. discoideum* and species homologs were each other's best bidirectional BLASTp hit (BBH), 2) they had the same or very similar functional domain architectures, and 3) the species genes clustered with the dictyosteliid orthologs in phylogenetic trees that also included other close homologs (see [supplementary data set S2, Supplementary Material](#) online). All *D. discoideum* spore and stalk genes and their best *P. aurantium*, *Ph. polycephalum*, and *A. castellanii* hits are listed in [supplementary data set S1, table S4, Supplementary Material](#) online, with likely orthologs indicated in bold text. A summary of the analysis is listed in [table 4](#).

Of the 92 *Dictyostelium* genes, 28 had orthologs in *P. aurantium*, but a much larger number (46) had orthologs in *Ph. polycephalum*. Even the nonsporulating *A. castellanii* has slightly more (34) spore and stalk genes in common with *Dictyostelium*. Largely the same genes are shared between all four species, which suggests that they have a common role independent of sporulation. Their role in normal spore and stalk formation in *Dictyostelium* may either result from a dictyosteliid specific recruitment into such a role, or reflect a pleiotropic effect of their loss on spore or stalk formation.

### Cell Signaling

Without undue emphasis on its development into fruiting bodies, we also considered the cell signaling potential of *P. aurantium* in its own right. Similar to most protists, *P. aurantium* amoebae need to sense a range of external stimuli in order to find prey, evade predators, and adapt to environmental change. To assess its sensory potential, we investigated the presence of well-known categories of cell signaling genes. A more detailed description of such genes is provided in [supplementary data set S1, pages 20–30, Supplementary Material](#) online.

### G-Protein Coupled Receptors

Transmembrane receptors that interact with heterotrimeric G-proteins are the most common sensors for external signals in eukaryotes. They are subdivided into six families with family 4 only being found in fungi. *Protostelium aurantium* has only 17 G-protein coupled receptors (GPCRs), mostly belonging to the family 1 rhodopsin-like receptors, and completely lack the family 3 metabotropic glutamate-like receptors ([supplementary data set S1, fig. S12, Supplementary Material](#) online).

This compares rather poorly with *Dictyostelium*, *A. castellanii*, and *Ph. polycephalum*, which have 55, 35, and 146 GPCRs, respectively ([table 5](#)). *Protostelium aurantium* does have 9 heterotrimeric G-proteins, more similar to *Dictyostelium*, which has 12 ([supplementary data set S1, fig. S12, Supplementary Material](#) online).

### Sensor Histidine Kinases/Phosphatases

Histidine kinases/phosphatases with an attached sensor domain are used for detecting and processing a broad range of chemical and physical stimuli in both pro- and eukaryotes. The stimulus activates either phosphorylation or dephosphorylation of a conserved histidine residue in the histidine kinase domain, which respectively triggers forward or reverse histidine–aspartate–histidine relay of the phosphate to/from an aspartate in a receiver domain. This then results in activation or inactivation of an attached effector, often an enzyme or TF (Zschiedrich et al. 2016). *Protostelium aurantium* appears to make up for its low number of GPCRs by having an extremely large number of 71 sensor histidine kinases/phosphatases (SHKPs) ([supplementary data set S1, fig. S13, Supplementary Material](#) online). This is over four times more than *D. discoideum* and also exceeding *Ph. polycephalum* and *A. castellanii* ([table 5](#)). The *P. aurantium* SHKPs contain a variety of sensor domains, among which is also a phytochrome domain. This domain was also found in two SHKPs of *Ph. polycephalum*, where a phytochrome is known to mediate light-induced sporulation (Starostzik and Marwan 1995).

### Cyclic Nucleotide Signaling

Many external stimuli exert their effect by modifying intracellular levels of the cyclic nucleotides (cNMPs) cAMP and cGMP. By binding to the conserved cyclic nucleotide binding domains of protein kinases, ion channels, and other effector proteins, both molecules control a broad range of cellular responses. cAMP is particularly important in the life cycle of *D. discoideum* where it acts both as an extracellular chemoattractant and as an intracellular messenger to regulate aggregation, morphogenesis, spore and stalk cell differentiation, and spore dormancy. In this organism, cAMP and cGMP are synthesized by three adenylate- and two guanylate cyclases, respectively, intracellularly detected by five cNMP binding proteins and hydrolyzed by seven cNMP phosphodiesterases (Saran et al. 2002; Bader et al. 2007). Query of the *P. aurantium* proteome

**Table 5**

Cell Signaling Proteins in Amoebozoa Genomes

Category	<i>Protostelium aurantium</i>	<i>Dictyostelium discoideum</i>	<i>Physarum polycephalum</i>	<i>Acanthamoeba castellanii</i>
G-protein coupled receptors	17	55	146	35
Heterotrimeric G-proteins				
Alpha	9	12	26	6
Beta	1	1	1	n.d.
Gamma	1	1	1	n.d.
Histidine kinases/phosphatases	71	16	51	48
Cyclic nucleotide signaling				
Adenylate/guanylate cyclases	52	5	64	67
cNMP binding domains	27	5	28	7
cNMP phosphodiesterases	16	7	11	10
Protein kinases				
All (S/T, S/T/Y, Y)	827	295	447	377
Sensor tyrosine kinases (Y)	167	0	4	21
SH2 domain proteins	85	15	18	48

revealed a much larger repertoire of cyclases, binding proteins, and phosphodiesterases.

We detected 52 nucleotidyl cyclases in the *P. aurantium* genome (fig. 5), which is somewhat less than the 64 and 67 cyclases detected in the *Ph. polycephalum* and *A. castellanii* genomes, respectively (table 5). However, 66 of the *A. castellanii* cyclases resulted from extensive duplication of a single gene that harbors a cyclase domain flanked by two protein kinase domains (identifier ELR11792 in fig. 5), whereas the remaining cyclase is a homolog of *D. discoideum* AcrA. The kinase-flanked cyclases also represent about half of the *Ph. polycephalum* cyclases, but are not present in *P. aurantium*. Instead, *P. aurantium* has many mammalian-type cyclases with two cyclase domains and two sets of six transmembrane domains. These cyclases are usually regulated by heterotrimeric G-proteins. *Dictyostelium discoideum* ACA and GCA also belong to this category. There are five homologs of the soluble *D. discoideum* guanylate cyclase SGC with two cyclase and one AAA-ATPase domain, which is implicated in chemotaxis (Saran et al. 2002). Although there are no obvious AcrA or ACG representatives, there is a large clade of 23 cyclases with mostly two transmembrane domains and a single cyclase domain. These enzymes have closest homologs in Excavates and Prokaryotes with a similar domain configuration. The two transmembrane domains may, as is the case for ACG (Saran et al. 2002), provide the cyclases with an external sensor domain.

With a total of 27 cyclic nucleotide binding proteins (cNBPs) (supplementary data set S1, fig. S14, Supplementary Material online), *P. aurantium* by far surpasses the 5 and 7 cNBPs of *Dictyostelium* and *A. castellanii*, respectively (table 5). Only *Ph. polycephalum* has one more cNBP, but has less variety in additional functional domains, which are the likely targets for regulation by either cAMP or cGMP (Schaap et al. 2016). The number of cyclic nucleotide phosphodiesterases (PDEs) is also

higher in *P. aurantium* than in the other Amoebozoa (supplementary data set S1, fig. S15, Supplementary Material online).

#### Serine/Threonine and Tyrosine Protein Kinases

Protein kinases that phosphorylate other proteins on either serine/threonine (S/T) or tyrosine residues (Y) or both (S/T/Y) represent the major group of intracellular signal processing intermediates in eukaryotes with ~518 members in humans (Hanks 2003). *Protostelium aurantium* has no less than 827 proteins with S/T, Y, or S/T/Y kinase domains, vastly surpassing the other publically available amoebozoan genomes, as well as humans (table 5). Eukaryotes have in general many S/T or S/T/Y kinases, but the Y specific kinases were previously considered to be present only in animals (Lim and Pawson 2010). Here, particularly the receptor tyrosine kinases play crucial developmental roles as sensors for secreted and exposed peptides that act as growth factors, controlling cell division, or differentiation inducing signals that control cell-type specialization (McDonnell et al. 2015). More recent sequencing of protozoan genomes revealed that tyrosine kinases with and without intrinsic receptor domains are more widespread (Manning et al. 2008; Suga et al. 2012; Clarke et al. 2013; Schaap et al. 2016). Around 167 tyrosine kinases were detected in *P. aurantium* (supplementary data set S1, fig. S16, Supplementary Material online). Most are likely receptor tyrosine kinases with either protein–protein interaction or polysaccharide binding domains in their extracellular regions, suggesting roles in cell–cell recognition or adhesion. The phosphorylated tyrosines in target proteins typically act as binding sites for SH2 domains, causing proteins with SH2 domains to directly interact with the target protein. We detected 85 proteins with SH2 domains in the *P. aurantium* genome, also a 2- to 5-fold increase in numbers compared with the other Amoebozoa (table 5).

## Conclusions

We sequenced the genomes of *Protostelium aurantium* and *Protostelium mycophagum* as the first representatives of the morphologically similar but genetically diverse protosteliids, which are characterized by forming a fruiting structure from a single amoeba consisting of a single spore or few spores on an acellular stalk.

With a size of 38 Mb and 17,000 genes, the *P. aurantium* genome is similar to that of *A. castellanii* (42 Mb, 15,000 genes), markedly smaller than the *Ph. polycephalum* genome (189 Mb, 29,000 genes) and larger than the *Dictyostelium* genomes, which range from 34 Mb, 13,000 genes (*D. discoideum*) to 23 Mb and 10,000 genes (*D. lacteum*). Although *Ph. polycephalum*'s large genome may reflect that this organism displays many alternative morphologies and life cycle stages (amoeba, flagellate, cyst, sporulating syncytium, or sclerotic syncytium), it is quite remarkable that the dictyosteliids with their complex multicellular life cycle and alternative abilities to form either haploid or zygotic cysts have the smallest genomes of currently sequenced amoebozoia. Apparently, the multicellular life style did not require numerically more genes. This is in agreement with analyses of the evolution of multicellularity in other eukaryote lineages (Niklas and Newman 2013; Nguyen et al. 2017). It is even conceivable that social amoebae were freed from an unknown selection pressure due to the evolution of their multicellularity. Thus, they might have been able to jettison surplus genes associated with signaling cascades and reception.

Because the single-celled *P. aurantium* fruiting body might be considered as a prototype for the multicellular *Dictyostelium* fruiting body, we had expected that a significant number of *Dictyostelium* developmental genes originated in the LCA of both lineages as was shown for developmental genes in Metazoa (Sebe-Pedros et al. 2013). However, contrary to this expectation, the conservation of developmentally essential *Dictyostelium* genes in *Protostelium* spp. is limited and actually not higher than for the evolutionary more distant *A. castellanii*. Our data and the recent finding of a sporulating *Acanthamoeba* species (Tice, Shadwick, et al. 2016), makes it conceivable that the entire genus *Acanthamoeba* could comprise at least the genomic capacity for protosteloid fruiting. It is well possible that the lack of apparent conservation in *Dictyostelium discoideum* could be due to the evolutionary distance between these taxa and that Dictyostelia have invented their own toolbox for multicellularity.

Using RNAseq data from discernable time points during development, we were able to dissect early and late events in fruiting body formation. The same processes might be at work in many other Amoebozoia, even if not true orthologous but analogous proteins are being utilized.

All free-living amoeba sequenced thus far do show a large repertoire of sensor histidine kinases, which in *Dictyostelium*

regulate the activity of the intracellular cAMP phosphodiesterase RegA and thereby the activity of PKA (Loomis 2014). All amoebae genomes also contain several to many adenylate cyclases for cAMP production to activate PKA (Clarke et al. 2013; Schaap et al. 2016). In *Dictyostelium*, cAMP acting on PKA critically regulates the transition from growth to development, the encapsulation of spore and stalk cells and the dormancy of spores (Loomis 2014). In both Dictyostelia, which have retained the unicellular survival strategy of encystation, and in *A. castellanii*, cAMP acting on PKA also mediate stress-induced encystation, with RegA antagonizing this process and favoring the trophozoite stage (Du et al. 2014; Kawabe et al. 2015). Since similar to *Dictyostelium* (Brenner 1978), cAMP levels were also found to increase during *P. aurantium* development into fruiting bodies (supplementary data set S1, fig. S9, Supplementary Material online), it is likely that this is one of the core processes that is conserved between Amoebozoa, to mediate their transition into a walled dormant stage, when experiencing stress.

Another conclusion of our analyses is that despite its simple life cycle compared with *Dictyostelium* and *Ph. polycephalum*, and having a five times smaller genome than *Ph. polycephalum*, *P. aurantium* has an extremely large repertoire of proteins for the detection and processing of external stimuli. This implies that *P. aurantium* has many more interactions with other organisms and/or with members of its own species than is currently being realized. It is also remarkable that the multicellular *Dictyostelium* with its complex life cycle has both a smaller number and less variety in its signal detection and processing proteins than the unicellular Amoebozoa. Could this mean that the Dictyostelia evolved multicellularity, because they were less adaptable than other Amoebozoa? Comparisons between genomes of related uni- and multicellular organisms in other eukaryote divisions might reveal whether this is a general trend.

## Supplementary Material

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

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## Author Contributions

The study was conceptualized by F.H., T.W., and G.G. The formal analysis of data was done by T.S., M.M., G.G., P.S., G.F., F.H., M.G., S.N., M.W., I.F., and K.R. All project parts were supervised by F.H. and G.G. The original draft preparation was done by G.G. and the final manuscript review and editing by T.W., F.H., G.G., and P.S.

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