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The zoospores of the thraustochytrid *Aurantiochytrium limacinum*: transcriptional reprogramming and lipid metabolism associated to their specific functions.

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Running title: Transcriptional footprint of *A. limacinum* zoospores
Abstract

* Aurantiochytrium limacinum* (Thraustochytriaceae, class Labyrinthulomycetes) is a marine Stramenopile and a pioneering mangrove decomposer. Its life cycle involves a non-motile stage and zoospore production. We observed that the composition of the medium, the presence of amino acids in particular, affects the release of zoospores. Two opposite conditions were defined, one with a cell population mainly composed of zoospores and another one with almost only non-motile cells. *In silico* allelic frequency analysis and flow cytometry suggest that zoospores and non-motile cells share the same ploidy level and are diploid. Through a RNA-seq approach, the transcriptional reprogramming accompanying the formation of zoospores was investigated, with a particular focus on their lipid metabolism. Based on a differential expression analysis, zoospores are characterized by high motility, very active signal transduction, an arrest of the cell division, a low amino acid metabolism and low glycolysis. Focusing on lipid metabolism, genes involved in lipase activities and peroxisomal β-oxidation are up-regulated. qRT-PCR of selected lipid genes and lipid analyses during the life span of zoospores confirmed our observations. These results highlight the importance of the lipid dynamics in zoospores and show the metabolic processes required to use these energy-dense molecules as fuel for zoospore survival during their quest of new territories.

Key words: Fatty acids; GPCR pathway; Lipid metabolism; PUFA synthase; RNA-sequencing; Signal transduction; Thraustochytrids; Zoospores; ω3-docosahexaenoic acid (DHA).
Introduction

Mangrove forest environments sustain a composite ecosystem (Isa et al., 2017), with a food web primed by decomposers (Demopoulos et al., 2007). Thraustochytrids are osmo-saprotrophic protists belonging to the phylum Heterokonta, class Labyrinthulomycetes (Cavalier-Smith et al., 1995) and are among the first organisms that colonize and decompose a fallen mangrove leaf (Raghukumar et al., 1994). The life cycle of thraustochytrids is characterized by vegetative reproduction and a motile stage, the zoospores (Honda et al., 1998, Iwata et al., 2017, Morabito et al., 2019). The biflagellated zoospores released from sporangia are essential to colonize new environments and propagate the population. Thraustochytrids are nutritionally valuable preys for microzoo- and zoo-plankton, especially because of their high content in very long chain polyunsaturated fatty acids (VLC-PUFAs) (Kimura and Naganuma, 2001 and references therein, Dellero et al., 2018a). Furthermore, the coastal zone they inhabit is the interface between the land-fixed and the ocean captured carbon, the so-called ‘Blue Carbon’ (Nelleman et al., 2009). Therefore, thraustochytrids may play a pivotal role in the ‘mycoloop’, a recently described energy pathway based on chytrid zoospores (Kagami et al., 2014).

Through the mycoloop refractory or inaccessible carbon can be transferred to the zooplankton (Kagami et al., 2007).

Although several organisms living in aquatic environments, such as algae, fungi, oomycetes and bacteria are able to produce zoospores (Agrawal, 2012), the literature concerning their metabolism and physiology is quite scarce. Most of the available studies concerning eukaryotic zoospores were carried out on oomycetes because several oomycetes, such as Phytophthora infestans, are plant pathogens and zoospores initiate the infectious interaction with the hosts (Hardham, 2007). This feature has fostered research to decipher the physiology of infective zoospores, with the aim to prevent infection, improve cultivation and reduce crop lost. The release of zoospores from sporangia may depend on several parameters such as light, nutrients, temperature shifts or intracellular pH variations (Hyde et al., 1991, Suzaki et al., 1996, Agrawal, 2012). Once produced, zoospores sense and reach new territories via phototaxis, electrotaxis or chemotaxis (Agrawal, 2012, Swafford and...
In the last decade, several studies on oomycetes focused on the identification of infection-related factors using transcriptomics approaches (Judelson et al., 2008; Chen et al., 2013, 2014; Sun et al., 2017). A panel of predicted genes, including transcription factors, protein kinases, putative signaling proteins, ion channel, or calcium binding proteins were suggested to be involved in the formation, survival and physiology of zoospores (Tani et al., 2004; Ah-Fong et al., 2017). In contrast with oomycetes, little information is available about the zoospore physiology and metabolism of non-infective thraustochytrids. Recently, thraustochytrids have attracted industrial interest for their ability to synthesize high levels of VLC-PUFAs, in particular docosahexaenoic acid (DHA, 22:6), an essential compound for human health (Simopoulo, 2008). Indeed, thraustochytrids have the unique feature to synthesize FA via two enzymatic systems; the classical fatty acid synthase (FAS) involved in the synthesis of relatively short saturated FA chains (16 to 18 carbons) and the PUFA synthase involved in the synthesis of longer unsaturated chains, such as DHA (Morabito et al., 2019).

The biotechnological interest towards thraustochytrids has directed efforts to unravel their physiology and metabolism, with the main aim to enhance lipid production (Raghukumar, 2008; Taoka et al., 2009; Aasen et al., 2016). This has prompted genomic and transcriptomic studies (Ma et al., 2015; Liu et al., 2016; Bi Z.-Q. et al., 2018; Iwasaka et al., 2018; Liang et al., 2018; Seddiki et al., 2018) which provide today new databases for metabolic engineering purposes and for ecophysiological understanding as well.

In a recent work, we investigated the ecophysiology of *Aurantiochytrium limacinum* zoospores and the triggers for their formation, settlement and maturation. Zoospores are released upon a transfer to a fresh nutrient-poor medium. They can swim for several days, relying on their energy storage for cell survival and motility (Dellero et al., 2018a). These observations raise questions about the nutritional factors that trigger a shift from a sedentary to a motile state, and about the associated transcriptional reprogramming and metabolic changes. Here we further describe the role of the external medium in the release of zoospores and depict the metabolic and transcriptional signatures...
associated with the vegetative-to-zoospore transition, with an emphasis on lipid metabolism. Our results reveal that such a transition implies profound transcriptional and metabolic rearrangements.
Results

Zoospore release

As previously shown (Dellero et al., 2018a), in a rich medium (R) containing glucose and yeast extract (YE) less than 3% zoospores were observed at best (D1), whereas in a poor medium (P), in which the glucose and YE concentrations were reduced to 1/40 (Supplementary Table 1), zoospores represented the main cell type (Table 1). A different medium, named R2, was produced in order to rule out the possibility that one or several components of the YE might repress zoosporulation. In the R2 medium YE was replaced by nitrate and phosphate solutions at concentrations comparable to those found in YE (Supplementary Table 1). A six-day-old R-grown culture, i.e. which did not contain any zoospores, was used as inoculum. During the first 24 hours in R2, zoospores were massively produced and represented 92% of the population (Table 1). A high zoospores/non-motile cells ratio was maintained until day 4 (D4). Thereafter, however, zoospore fraction decreased to nil and non-motile cells dominated the population. It can be concluded that one of the YE components may impair the release of zoospores.

In order to discriminate whether amino acids (the main source of nitrogen in R) hamper zoosporulation, nitrate was removed from R2 and replaced by a cocktail of amino acids in concentrations that mimic YE (R3 medium, Supplementary Table 1). The zoospore/non-motile cells ratio in R3 was comparable to that recorded in R, i.e. very few zoospores were observed. Inversely, when amino acids were added in a P culture containing almost exclusively zoospores, the non-motile cell occurrence increased at D4 (i.e. 48 hours after the addition of amino acids). At D6, a 3.4% zoospore/non-motile cells ratio was recorded. Altogether, these results highlight a potential role of amino acids in controlling the initiation/maturation of the motile stage of A. limacinum.

Ploidy level of the zoospores and non-motile cells

A variant calling analysis was performed by mapping the transcriptomes of non-motile cells and zoospores produced in the present work against the Aurantiochytrium limacinum ATCC® MYA1381™ reference genome. 38110 (Supplementary file 1) and 53181 (Supplementary file 2) variants were
identified in non-motile cells and zoospores, respectively. Single Nucleotide Polymorphisms (SNPs) represented about 94% of the variants in both groups. A vast majority of the variants (64% and 63% of total variants for non-motile and zoospores, respectively; Supplementary Files 1 and 2) were identified as alternative genotypes, i.e. all the reads in the transcriptome of non-motile cells and zoospores presented the same sequence, different from that in the reference genome. Such sites can be classified as intraspecific polymorphisms between the Aurantiochytrium limacinum ATCC® MYA1381™ used as reference and A. limacinum CCAP 4062/1 strain. In order to estimate the ploidy level of zoospores and non-motile cells, allele frequency distributions were calculated (Fig. 1A). The peak of ‘homozygous variants’ (allele frequency = 100) on the right-hand side of the plots (Fig. 1A) actually identifies the intraspecific polymorphic sites described above. Noteworthy, 13829 (36%) and 19854 (37%) variants were classified as heterozygous (0/1 or 1/0 genotype, Supplementary Files 1 and 2) in non-motile cells and zoospores, respectively. The allele density (y-axis in Fig. 1A) is an indication of the number of reads bearing one or the other allele. A Mann Whitney Test was performed to compare the distribution of the allele frequencies of non-motile cells and zoospores and the difference resulted not significant (p = 0.35). The portion of the allele frequency curve of the heterozygous variants only (Fig. 1A) is centered on 50 in both samples, therefore, it can be stated that zoospores and non-motile cells share the same ploidy level and that they are both diploid. In order to validate the latter result, flow cytometry was used to measure the quantity of DNA in zoospores vs. non-motile cells.

In flow cytometry, light scatter analysis of a fast growing population in R showed a large panel of cell sizes and internal complexity (granulometry) (Fig. 1B). The fluorescence pattern of propidium iodide (PI) of R-grown cells displayed a main peak at an arbitrary value of 280 and a second one representing cells containing about twice as much DNA as the cells forming the first peak (Fig. 1C). The right-hand side of the figure indicates the presence of cells containing even more DNA. Assuming that A. limacinum is diploid (as suggested by allele frequency analysis), the first peak would represent diploid mononucleated cells and the second either mononucleated cells in G2 phase or binucleated
cells. Because of the presence of multinucleated cells, the interpretation of the following peaks is cumbersome. Each peak could represent either a cell population with n nuclei in G2 phase or cells with 2n nuclei in G1 phase. On the contrary, zoospore size and granulometry was more uniform (Fig. 1D). PI fluorescence (Fig. 1E) showed a main and relatively sharp peak and a second small peak representing cells with twice as much DNA. By comparing Figures 1C and 1E, it is clear that zoospores did not contain half the DNA than non-motile cells. These results strongly suggest that zoospores are diploid, corroborating the allele frequency analysis (Fig. 1A).

**Genome wide expression and transcriptional footprints in zoospores**

A differential expression analysis based on RNA-seq data showed that in zoospores 4280 and 3330 genes were up- and down-regulated (cut-off arbitrarily set at log2,FC = 1), respectively (Supplementary Table 2). 7610 genes represent about half of the whole genome. RNA-seq differential gene expression analysis was validated by qRT-PCR of 22 selected genes (Supplementary Table 3) on RNA extracted from a completely independent experiment carried out exactly as described for RNA-seq. The Pearson’s correlation coefficient ($r = 0.93$) showed a strong positive linear correlation ($p$-value $< 10^{-5}$) between qRT-PCR and RNA-seq analyses. The strong linear correlation was coupled with a strong monotonic relation between the two sets of results by Spearman’s Rho ($r_s = 0.88$, $p$-value = 0.05).

From a Gene Ontology Enrichment Analysis (GOEA) (Supplementary Fig. 1), most of the significantly enriched up-regulated GO terms were ‘signal transduction’ (GO:0007165) and several descendants like ‘G-protein coupled receptor signaling pathway’ (GO:0007186), or ‘signal transducer activity’ (GO:0097677) which groups mainly transcription factors. This is indicative of an increased ability of the zoospores to sense and react to their external environment. Zoospores are biflagellated cells and some of the highly enriched GO terms among the up-regulated genes are, not surprisingly, ‘microtubule associated complex’ (GO:0005875), ‘dynein complex’ (GO:0030286), ‘motor activity’ (GO:0003774). Among the down-regulated genes, ‘translation’ (GO:0006412), ‘aminoacyl-tRNA ligase
activity’ (GO:0004812), ‘structural constituent of ribosome’ (GO:0003735) were the most enriched GO terms, suggesting an arrest of the cell division and a lower turnover of transcripts.

**Signal transduction**

Genes involved in signal transduction and signaling cascades were strongly expressed in zoospores (Fig. 2A). They included GPCRs (G protein-coupled receptor), with 30 highly up-regulated transcripts encoding for different GPCRs. Interestingly, seven out of these transcripts showed almost no expression in the non-motile cells (fgenesh1_pg.9_421, estExt_fgenesh1_pg.C_6_t10488, gm1.13383_g, fgenesh1_pg.8_348, fgenesh1_pg.36_3, fgenesh1_pg.3_318, gm1.10294_g, gm1.4442_g), with levels below 20 Transcripts Per Kilobase Million (TPM) compared to values ranging from 696 to 3702 in the zoospores (Supplementary Table 2). In addition, many transcripts encoding the signaling transduction route were up-regulated, e.g. 14 Ras GTPases as well as several proteins linked to the cAMP signaling pathway (present in the G protein item in Fig. 2A; Supplementary Table 2).

**Cytoskeleton and molecular motors**

Thirty transcripts encoding for dynein (present in the intraflagellar transport and flagellar movement items in Fig. 2A), 24 for myosin and more than 35 for kinesin were up-regulated in zoospores (Fig. 2A and Supplementary Table 2). Genes involved in the tubulin-dependent cytoskeleton as well were largely up-regulated, as expected for flagellated cells. These findings, together with the up-regulation of transcripts encoding for proteins involved in the regulation of cellular motors (putative CAP-GLY domain containing linker protein, CLIP (estExt_fgenesh1_kg.C_140042)) and actin cytoskeleton (formins), reveal a highly dynamic cytoplasm. Actin-Myosin transportation are important for the perception of external stimuli in higher plants as well as in microalgae (Duan and Tominaga, 2018).

**DNA replication and transcription**

DNA replication was reduced in zoospores (Fig. 2B), with eight down-regulated transcripts encoding for DNA polymerase, five transcripts encoding for DNA topoisomerase and 11 transcripts encoding for histones (Supplementary Table 2). Genes encoding for RNA polymerases, helicases or involved in...
chromatin remodeling were also significantly down-regulated (Fig. 2C), suggesting that DNA transcription was lowered. Transcripts involved in the translation process were down-regulated as well. These results suggest that zoospores do not divide, as already suggested by the flow-cytometry data.

**Nitrogen and amino acid metabolism**

Genes involved in nitrogen and amino acid metabolisms were highly differentially expressed (Fig. 2D). About 20 genes involved in nitrate and ammonium transport and nitrogen assimilation (such as nitrate (fgenesh1_pm.38_1) and nitrite reductases (e_gw1.38.18.1) or glutamate (estExt_Genewise1.C_17_t10014) and glutamine synthetases (fgenesh1_pm.1_555, fgenesh1_kg.1_553_isotig07286, estExt_Genewise1.C_17_t10014e_gw1.5.701.1)) were strongly up-regulated (Supplementary Table 2), possibly denoting the nitrogen deficient medium in which zoospores were produced. In contrast, most of the genes related to the metabolism and turnover of amino acids (amino acid biosynthesis and degradation, urea cycle) were down-regulated. This presumably indicates a reduced need of protein synthesis because of the arrest of the cell division. Genes identified as ‘amino acid transport’ were predominantly down-regulated, but a few of them were significantly up-regulated with $\log_2$FC ranging from 4.4 (fgenesh1_pg.18_302) to 1.6 (e_gw1.22.185.1) (Supplementary Table 2). Their function and localization were not further investigated here.

**Carbohydrate metabolism**

Genes involved in key steps of the glycolysis (Fig. 2E) were down-regulated, such as triosephosphate isomerase (fgenesh1_kg.10_22_isotig07893), 3-phosphoglycerate kinase enolase (estExt_fgenesh1_kg.C_30041), and pyruvate kinase (estExt_fgenesh1_kg.C_90258) (Supplementary Table 2). The same pattern was observed in several genes coding for the Krebs cycle enzymes, including components of the pyruvate dehydrogenase complex, succinate dehydrogenase (fgenesh1_pm.8_94), citrate synthase (fgenesh1_pg.13_25), isocitrate dehydrogenase (estExt_fgenesh1_kg.C_210136), the T protein of the glycine decarboxylase complex, as well as
oxidases such as cytochrome c oxidase (fgenesh1_pm.10_34) and alternative oxidase (gw1.27.100.1).

Taken as a whole, these results suggest that carbohydrate catabolism and respiratory activities were
down-regulated in zoospores compared with non-motile cells. This assumption was confirmed by the
direct measurement of the respiratory activities (Table 2). In R-grown cells, oxygen consumption at
D1 considerably augmented, in correspondence to the resuming of cell divisions. Conversely,
zoospore respiration slowly declined with time (Table 2).

Fatty acid synthesis and degradation

FAS is involved in the synthesis of 16:0 which can be thereafter elongated and desaturated to
produce VLC-PUFAs; PUFA synthase directly produces VLC-PUFAs. Only a few genes involved in the
biosynthesis of FAs were differentially expressed in zoospores, at least at D1. The interpretation of
the results to decipher how the biosynthetic pathway to VLC-PUFAs is regulated in zoospores is
challenging (Fig. 2F). Two desaturases (estExt_fgenesh1_kg.C_1_t10086; gw1.29.176.1) were up-
regulated whereas e_gw1.11.243.1 was down-regulated, and the only elongases (e_gw1.5.731.1;
estExt_fgenesh1_kg.C_60194) found in A. limacinum were down-regulated (Supplementary Table 4).
Desaturases and elongases are likely involved in the FAS pathway (for a review, see Morabito et al.,
2019), but FAS-1 (e_gw1.21.366.1) was not differentially expressed. Likewise, subunits A and B of the
PUFA synthase (PUFA-A fgenesh1_pg.14_.#_251, PUFA-B estExt_fgenesh1_kg.C_140136) were not
differentially expressed, and subunit C (PUFA-C estExt_fgenesh1_kg.C_190026) was slightly down-
regulated (Supplementary Table 4).

FA degradation takes place in both mitochondria and peroxisomes via β-oxidation. The hydrolysis of
FA esterified to glycerolipids is mediated by lipases. About 30 genes were identified as lipase or
lipase-like, out of which 17 were up-regulated and 11 down-regulated. Their expression levels
showed that lipases were globally upregulated, suggesting that glycerolipid degradation was
potentially increased as a whole (Fig. 2F and Supplementary Table 4). Most of the up-regulated
lipases belong to class 3 lipases, i.e. involved in the hydrolysis of ester bonds of TAGs or DAGs.

However, phospholipases A (Pla1 e_gw1.1.605.1, Pla2 e_gw1.24.47.1), B (Plb e_gw1.6.53.1) and D
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(Pld2 estExt_Genewise1Plus.C_1_t20024, Pld3 gm1.46_g), involved in the degradation of phospholipids, were either non differentially expressed or down-regulated. This suggests that only storage lipids were part of the catabolic processes, whereas membrane-bound glycerolipids are not. FAs released from triacylglycerids (TAGs) are activated as acyl-CoA by acyl-CoA ligases before entering β-oxidation. An overview of the transcripts involved in FA degradation (Fig. 2F) suggests a global down-regulation. However, a closer inspection indicates that the peroxisomal pathway was up-regulated. Indeed, out of the three genes encoding for Long chain Acyl-CoA ligase, Acsl2 (fgenesh1_kg.4_#_23_#_isotig03564) was the only presenting a peroxisomal localization and the only up-regulated one. In addition, most of the transcripts involved in the peroxisomal β-oxidation (acyl-CoA ABC transporter; acyl-CoA oxidase Acox1, Acox3; bifunctional enzyme Ehhadh1, Ehhadh3; ketoacyl-CoA thiolase Acaa1, Acaa2, Acaa4) were globally up-regulated (Supplementary Table 4). The mitochondrial β-oxidation did not show a clear pattern: out of the five genes encoding the acyl-CoA dehydrogenase (Acad), three were down-regulated, one was up-regulated and one did not significantly vary (Supplementary Table 4). Altogether, these results suggest that TAGs are more actively degraded in the zoospores, liberating FAs that are conveyed preferentially to the peroxisome to be oxidized.

Lipid dynamics in zoospores and non-motile cells

The expression levels of five genes involved in lipid metabolism (Tagl3 e_gw1.7.228.1, Acox1 fgenesh1_kg.9_#_301_#_isotig03404, Acad1 estExt_fgenesh1_kg.C_330018, FAS-1, PUFA-A) were followed for six days in zoospores and non-motile cells (Fig. 3). A ΔΔCt analysis was performed by comparing the expression level of each transcript at day 1 (D1), day 2 (D2), day 4 (D4), and day 6 (D6) with the expression level of the same transcript at D1 in R medium.

The TAG lipase 3 (Tagl3) was strongly up-regulated at D1 in zoospores and represented at this point the most expressed TAG lipase (Supplementary Table 4). Its expression level did not vary sensibly thereafter (Fig. 3A), suggesting that TAGs were constantly degraded throughout the whole zoospore life. Tagl3 expression in non-motile cells increased progressively to reach a maximal value.
at D4 and D6 (Fig. 3B), i.e. when TAGs accumulate (see below). Highly likely, TAG synthesis and
degradation may occur simultaneously in non-motile cells.

**β-oxidation** is initiated in peroxisomes and mitochondria through the activities of Acox and Acad,
respectively. The peroxisomal Acox1 (Dellero et al., 2018b) and mitochondrial Acad1 (Dellero et al.,
2018b) transcripts were the most expressed Acox and Acad in both cell types (Supplementary Table
4). Acox1 was slightly up-regulated in zoospores at D1 and increased thenceforward. Similarly, in
non-motile cells, Acox1 expression increased with time, but at a much lower extent than in
zoospores. In contrast, the Acad1 expression was down-regulated in zoospores, especially at D6,
whereas it did not significantly change in non-motile cells. These results emphasize again the role of
peroxisomes vs. mitochondria for the oxidation of FAs in zoospores.

In thraustochytrids, the synthesis of FAs is driven by two enzymatic systems, FAS and PUFA synthase
(see Introduction). In zoospores, FAS-1 and PUFA-A genes showed no significant regulation at D1 and
D2 compared to non-motile cells (Fig. 3). In the following days, however, their expression decreased
in zoospores, especially PUFA-A. In contrast, FAS-1 and PUFA-A expression increased over time in
non-motile cells. In both cell types, PUFA-A expression varied significantly more than FAS-1, possibly
indicating the higher relevance of the PUFA synthase synthetic pathway in the production of PUFAs.

**Fatty acid analyses**

To correlate gene expression with metabolic changes, the FA content was analyzed during a six-day
growth in both P and R media (Fig. 4). The major FA species (16:0, 22:5, 22:6) in zoospores increased
at D1 (the transfer in P medium occurred on D0) (Fig. 4A), then declined thereafter. In R-grown cells
all FA species declined the first day (Fig. 4B) then constantly raised thenceforth to get almost to the
initial value at D6. This illustrates the strong relationship between nitrogen availability and lipid
metabolism described earlier (Dellero et al., 2018a). In R, cells first consumed their TAGs, then, upon
nitrogen limitation, TAGs were synthesized again. In P, nitrogen was limiting right at the beginning of
the experiment and TAGs were immediately synthesized until glucose was exhausted, then were
consumed to sustain the energy demand. The distribution (%) of main FAs did not vary much (Figs.
Although slight differences could be still observed between the two cell populations. Indeed, zoospores showed lower 15:0 and higher 16:0 contents. Odd-numbered FAs (mainly 15:0) are produced from branched-chain amino acids because their degradation leads to propionyl-CoA (Crown et al., 2016) that can replace acetyl-CoA for the synthesis of FAs. In R, the presence of YE, rich in amino acids, corroborates the hypothesis that 15:0 synthesis is driven by the propionyl-CoA derived from branched-chain amino acids. The lack of 15:0 in zoospores might be due to the reduced YE concentration in P medium. Another intriguing difference is the accumulation in zoospores of 18:0 at D1 and 18:1 at D6. The accumulation along the zoospore life span of 18:1 could be due to a residual desaturation activity of 18:0, although no Δ9-desaturase was found in the genome of A. limacinum (Dellero et al., 2018b).

The increase of lipids in P-grown cells at D1 raises the question whether the sporangium or the zoospore is responsible for the recorded lipid accumulation. The release of zoospores and lipid accumulation during the first 24 hours were compared. The release of zoospores started about 8 hours after the transfer in P, and rapidly raised to almost 100% of the population at 24 hours (Fig. 4E). The FA content in P-grown cells did not vary during the first 12 hours (Fig. 4F), then it nearly doubled at 24 hours. Thus, FAs accumulated after the release of zoospores was initiated, suggesting that they could originate from de novo synthesis.

Discussion

Zoospores were observed mostly in culture media deprived of amino acids. A nitrogen deficiency cannot explain alone the production of zoospores because replacement of amino acids by NO₃ triggered the release of zoospores during the first days of culture and high biomass accumulation. Amino acids could play a specific role in the life cycle of thraustochytrids. Indeed, thraustochytrids feed on decaying mangrove leaves (Raghukumar et al., 1995), and protein degradation in decaying leaves likely releases amino acids in the close vicinity of the leaf itself. Because nitrogen availability is one of the prime factor limiting cell growth (Alongi, 2009; Rao et al., 1994), the presence or absence of nitrogen-rich organic compounds might be one of the switches controlling the release of
zoospores from zoosporangia. Oomycete zoospores reach and settle in new territories via chemotaxis, following attractants such as amino acids (Hardham, 2007; Swafford and Oakley, 2017).

Similarly, it was demonstrated that zoospores of different thraustochytrid species show a positive chemotactic response to glutamic acid and pectin (Fan et al., 2002). This suggests a specific ability of Aurantiochytrium zoospores to detect the presence of amino acids in the external environment (Dellero et al., 2018a). Indeed, eleven genes involved in amino acid transport were down-regulated and four were up-regulated with fgenesh1_pg.18_302 showing a very low expression level in non-motile cells. In zoospores its expression strongly increased with a log$_2$FC of 4.4.

The ploidy of A. limacinum zoospores and non-motile cells was investigated via two different approaches, namely the distribution of SNPs through the whole exome and flow cytometry. Both methods suggest a diploid status. Flow cytometry experiments showed that the main peak of fluorescence was much sharper in zoospores (Fig. 1E) than in non-motile cells (Fig. 1C), suggesting that zoospores were in G1 phase and that division/DNA replication was blocked. The latter hypothesis is in agreement with transcriptomic data. Diploidy of A. limacinum zoospores corroborates the results reported for A. acetophilum for which sexual reproduction has been described (Ganuza et al., 2019). In A. acetophilum biflagellated cells produced by specialized sporangia (Type II sporangia) can act as gametes. The physico-chemical as well as physiological conditions triggering sexual reproduction were not elucidated and the ploidy level of such ‘putative gametes’ was not tested (Ganuza et al., 2019).

Differential gene expression analyses indicate a profound reprogramming of in zoospores. The transcriptomic data were summarized into a simplified scheme representing some aspects of the metabolism (Fig. 5). Zoospores are potentially characterized by their motility, highly active signal transduction, a low DNA replication activity, a high nitrogen transport activity but a low amino acid metabolism, a low carbohydrate catabolism, high lipase activities and a high β-oxidation of FAs in peroxisomes.
G proteins were markedly up-regulated. They are often associated with membrane-spanning receptors (Weis and Kobilka, 2018), and it is possible that the increase of expression level connotes the pioneering activity of zoospores. Indeed, zoospores are likely released to explore new territories, implying a high sensitivity to probe their environment in order to find new substrates or leaves. G proteins also regulate, among others, transcription, cellular differentiation, secretion and motility (Neves et al., 2002). As expected for flagellated cells, most of the genes involved in the cytoskeleton dynamics were strongly up-regulated. The signaling pathway of GPCRs is spread through the whole living kingdom (de Mendoza et al., 2014), e.g. it has been demonstrated that diatoms sense the presence of predators via the GPCR signal transduction pathway (Amato et al., 2018) and that in the rhizarian protist *Plasmodiophora brassicae* GPCR signal transduction pathways underwent a robust expansion and plays a pivotal role in the germination of resting spores (Bi K. et al., 2019).

Furthermore, in oomycetes a very peculiar GPCR system evolved (Meijer and Govers, 2006). Conversely, genes involved in DNA replication (such as polymerase, topoisomerase, replication factor etc.) were markedly down-regulated, suggesting that zoospores were blocked in the G1 phase. The arrest of the cell division was presumably associated with a decrease of several metabolic activities. Noteworthy, the amino acid metabolism was largely down-regulated, indicating a potential decline of protein synthesis and turnover. This is true for carbohydrate catabolism as well, since many genes involved in the glycolysis and the Krebs cycle showed a decreased expression in zoospores. Carbohydrate catabolism is strongly connected to the energy metabolism. Thus, results presented here indicate that zoospores have a lower energy demand than dividing cells, despite their swimming activity, an assumption supported by the direct recording of their respiratory activities.

The differential expression of genes involved in lipid metabolism did not show a clear tendency at D1, except for lipase activities, which was up-regulated. In zoospores, the time point D1 corresponded to a metabolic turn, where the FA metabolism shifted from synthesis to catabolism. This may explain the difficulty to correlate gene expressions and FA analyses. Nevertheless, following the expressions of key genes for either FA synthesis or degradation during the whole growth experiment, gene...
expressions and FA analyses correlated well. The gene expression of PUFA-A always varied more
strongly than FAS-1 although 22:6 and 16:0, their respective products, did not. A very fast turnover of
either the transcripts or the proteins may be reason of such observation. Indeed, the level of
transcripts of the PUFA-A (fgenesh1_pg.14_251) in non-motile cells at D1 (about 56000 TPM) was
one of the highest compared to other genes involved in lipid synthesis. A 32-fold increase in
expression (log_{2}FC = 5) at D6 in non-motile cells was observed, strongly suggesting that a very high
number of transcripts is required to sustain a high enzymatic activity. In zoospores, FA catabolism
was associated with an increased expression of the the peroxisomal β-oxidation genes, whilst the
mitochondrial β-oxidation did not follow the same pattern. In many (but not all) organisms,
peroxisomes oxidize very long FA chains down to a certain chain length (Reddy and Hashimoto,
2001). The products of the peroxisomal β-oxidation are then shuttled to mitochondria for a complete
oxidation to CO_{2} and H_{2}O (Wanders et al., 2016). In thraustochytrids, 16:0 β-oxidation would be
initiated in peroxisomes because the 16:0 and 22:6 content (in percentage, Fig. 4C) in zoospores
dealed at comparable pace.

Potential differences between oomycete and thraustochytrid zoospores
One of the critical points in the transition from the sedentary to the motile state is the synthesis of
flagella. In oomycetes, zoosporogenesis is often induced by a cold shock, and flagella are synthesized
from basal bodies at the apex of the nucleus (Hardham, 2007). Zoospores are released upon
formation, and the whole process takes about one hour (Hyde et al., 1991). A. limacinum zoospores
appear eight hours after transfer in P medium (Fig. 4E). Assuming that zoosporogenesis is induced
soon after the transfer, zoospore formation and release appear slower in thraustochytrids than in
oomycetes, suggesting that different processes could be involved.

Oomycete zoospores can swim for hours or even days (Hardham, 2007; Kagda et al., 2018), like A.
limacinum’s. Thus, in both organisms, zoospore survival may depend on the endogenous energy
stores. Oomycetes supposedly store energy in mycolaminarin or lipids. Mycolaminarin is a (1,3,1,6)-
β-glucan polysaccharide that may represent up to 13% of the dry weight in oomycetes (Du and
The lipid content in the oomycete *Phytophthora infestans* is rather reduced, representing at best 5% of the dry weight, with TAGs accounting for half of the total (Griffiths *et al.*, 2003). This is a striking difference with *A. limacinum* where TAGs may represent up to 30% of the dry weight (Dellero *et al.*, 2018b), suggesting that the energetic metabolisms in these two organisms rely on different pathways. Interestingly, in *P. infestans*, a taurocyamine kinase (phosphagen kinase of the creatine kinase family) was found to be highly expressed and targeted towards flagellar axonemes during zoosporogenesis, possibly to fulfill the ATP demand associated with the flagellar movement (Kagda *et al.*, 2018). A BLAST search on *A. limacinum* genome identified three genes of the creatine kinase family homologous to the *P. infestans* gene (estExt_fgenesh1_kg.C_140197, fgenesh1_kg.3_231_isotig05435, fgenesh1_kg.12_175_isotig10864) and up-regulated in zoospores, suggesting that a similar function may be hypothesized in *A. limacinum*. Recently, a differential expression analysis of the of *P. infestans* zoospores vs. mycelia, the vegetative dividing stage (Ah-Fong *et al.*, 2017) revealed that, like in *A. limacinum*, signal transduction and flagellar proteins were up-regulated, while many metabolic pathways, including glycolysis, TCA cycle and amino acid metabolism were down-regulated. In contrast with results presented here, oomycete zoospores displayed a down-regulation of the β-oxidation and higher levels of transcripts involved in DNA replication. Although the pathways that generate energy are probably different, it is surprising that DNA replication could increase in non-dividing cells. It was postulated that the developmental program of *P. infestans* zoospores anticipates the need to resume replication after encystment and maturation (Ah-Fong *et al.*, 2017). This was obviously not the case for *A. limacinum* zoospores. Taken as a whole, the present work shows for the first time that *A. limacinum* zoospores are mainly diploid and that zoosporulation is a complex event depending on external factors, among which the lack of amino acids could play an important role. Here we highlight how the transcriptional reprogramming that occurs in *A. limacinum* zoospores, contributes to the metabolic changes required to migrate, probe their environment and survive; three essential skills to find new territories.
and propagate the population. The transcriptional signature of the zoospore reveals the main role
played by the G-protein signal transduction pathway in perceiving the external environment,
although the actual mediators are still unknown. Lipid metabolism plays an essential role in providing
the energy required to swimming for several days and the exploration of new areas.
Experimental procedures

Strain and media
CCAP 4062/1 strain was collected in Mayotte island (Indian Ocean, 12°48'51.8''S, 45°14'21.7''E) and routinely cultivated at 20 °C in 250 ml Pyrex® Erlen-Meyer flasks filled with 50 mL of R medium (Supplementary Table 1) (Dellero et al., 2018b) with 100 rpm orbital shaking. For all experiments, six day-old axenic cultures grown on R were transferred to fresh culture media at an initial cell concentration of 5×10⁵ cells·mL⁻¹.

The R and P media used in the experiments presented here were prepared as described in Dellero et al. (2018a, b); the R2 and R3 recipes are reported in Supplementary Table 1, along with the recipes of R and P. All the culture media were prepared using autoclaved MilliQ water.

Cell enumeration was performed with a Malassez hemocytometer (ca. 100 to 200 cells counted per sample) under a Zeiss AxioScopeA1 (Carl Zeiss SAS, Oberkochen, Germany) microscope. For zoospore enumeration, a sample of 2 mL was fixed with a drop of 2.5 % glutaraldehyde to immobilize zoospores. All the experiments were run in triplicate.

RNA extraction and sequencing
For RNA-seq experiments, a six-day-old culture was inoculated in triplicate in 50 mL of either R or P media, at an initial cell concentration of 5×10⁵ cells·mL⁻¹. After 24 hours 1.5×10⁷ cells were harvested by centrifugation, snap frozen in liquid nitrogen and then stored at -80 °C until use. RNA was extracted using the TRI Reagent (Sigma Aldrich) as described by Amato et al. (2017). RNA extracted from biological triplicates of R- and P-grown cultures were sent out for Illumina sequencing. Libraries were produced and processed following the manufacturer’s instructions and sequenced on single-end 75 bp mode on NextSeq500 (Illumina, San Diego, CA). The CASAVA 1.8.2 version of the Illumina pipeline was used to process raw data for both format conversion and de-multiplexing.

Bioinformatics analyses of RNA-seq data
Raw reads were processed with FASTQC (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and BBDuk (https://jgi.doe.gov/data-
and-tools/bbtools/) in order to check quality and remove low quality bases and adapters. A minimum
quality of 25 and a minimum read length of 35 are required. High quality reads were then mapped
against the *Aurantiochytrium limacinum* ATCC® MYA1381™ reference genome
(https://genome.jgi.doe.gov/Aurli1/Aurli1.home.html) with STAR (Dobin et al., 2012). FeatureCounts
(Liao et al., 2013) was used to perform read summarization at gene level, only reads with quality
higher than 30 were used. In addition, strand-specific and paired-end mode are included. Statistical
analyses and plots were generated with R software. Lowly expressed genes were filtered out with the
HTSFilter package (Rau et al., 2013), then a differential expression analysis was performed using
edgeR (Robinson et al., 2009). Genes with an FDR less than or equal to 0.05 were considered
significantly differentially expressed. The lists of up- and down-regulated genes were used to perform
Gene Ontology Enrichment Analysis (Du et al., 2010) using in-house scripts. The assembled and
annotated transcriptome was submitted to NCBI SRA with the accession number PRJNA590015.

**Estimation of the heterozygosity level**

The mapping files (BAM) obtained by aligning the trimmed RNA-seq reads from the six samples were
processed with the Opossum pipeline (Oikkonena and Lise, 2017) in order to remove reads with low
quality mapping (MAPQ < 40) and duplicates and to split reads mapping across introns. The three
replicates of the R-grown cultures were pooled and the same was done with the three replicates of
the P-grown cultures in order to obtain two final BAMs. The final mapping files were then analyzed
with Platypus (Rimmer et al., 2014) in order to perform variant calling. Only variants with a depth of
at least 10 reads, minimum base quality of 30, a minimum posterior probability of 30 and classified as
PASS were considered. Allele frequencies were calculated as the ratio between the NV and NR fields
of the VCF file.

**Validation of RNA-seq data and differential gene expression analyses**

qRT-PCR validation of RNA-seq data was carried out on a completely independent experiment as
described above for RNA-seq. RNA samples were reverse transcribed using the SuperScript IV VILO
Mastermix with ezDNAse kit (ThermoFisher) according to the manufacturer’s instructions. Reactions
were run in a final volume of 10 µL containing 10 ng of cDNA, 5 µL of Power SYBR® Green PCR Master Mix (Applied Biosystems, ThermoFisher) and 600 nM of each primer. Reactions were performed in a CFX ConnectTM Real-Time System (BioRad®) with the following program: initial denaturation step at 95 °C for 10 minutes; 40 denaturation-amplification-elongation cycles (95 °C, 10 s; 55 °C, 10 s; 72 °C, 30 s), followed by melting curve assessment (65 °C to 95 °C, with a 0.5 °C increment). All primer sequences are available in Supplementary Table 3. Transcript levels were normalized against the geometric mean of three reference genes, with similar expression in both conditions: Pacifastin (estExt_fgenesh1_kg.C_160075), Cystein desulfurase NFS1 (estExt_fgenesh1_kg.C_30063), Protein involved in Snf1 protein kinase complex assembly (gw1.10.847.1) (Supplementary Table 5). For the gene differential expression experiment along a six day-growth, a six-day-old culture grown on R was inoculated in 50 mL of either R or P fresh culture media at an initial cell concentration of 5×10⁵ cells·mL⁻¹. 1.5 mL were gathered by centrifugation at the following time points day 1 (D1), day 2 (D2), day 4 (D4), and day 6 (D6). RNA was extracted and reverse transcribed as described above. qRT-PCR was performed as depicted above. Five genes were analyzed; Tagl3, Acox1, Acad1, FAS-1, PUFA-A. The list of primer sequences is reported in Supplementary Table 5. Gene abbreviations follow Dellero et al. (2018b).

The differential expression analyses and statistics were performed using the Pair Wise Fixed Reallocation Randomisation Test method developed in the Relative Expression Software Tool REST® (Pfaffl et al., 2002). Biological triplicates and technical triplicates of each reaction were performed. Flow cytometry, respiratory activities, lipid extraction and fatty acids analyses

To reduce variability, a six-day old R-grown culture was inoculated at 5×10⁵ cells·mL⁻¹ in 50 mL of P medium to induce zoospore formation. After 24 hours, 200 µL of the culture were spread onto a 1 % agar-R plate. One of the single colonies was picked and cultivated in R medium for six days. Cells were inoculated in triplicate into 50 mL of either fresh R or fresh P media at a concentration of 5×10⁵ cells·mL⁻¹. R and P cultures were harvested at time points 15 hours and 24 hours, corresponding to the production peaks of mononucleated non-motile cells and zoospores, respectively (Dellero et al.,
Three million cells were collected from each sample by centrifugation (5 minutes at 3500 xg for R cultivated cells and 7000 xg for P cultivated cells). Pellets were washed with 1 mL of PBS-EDTA 2 mM, and then fixed with 500 µL of 70 % ethanol for at least 30 minutes at room temperature. After fixation, cells were washed twice with 1 mL of PBS-EDTA 2 mM and resuspended in 500 µL of PBS-EDTA 2 mM. Five microlitres of propidium iodide (PI) and 5 µL of RNase I were added and cells were incubated 30 minutes in the dark. All the stained samples were analyzed with a BD FacsCalibur (Benton Dickinson) flow cytometer (excitation 488 nm, 585/42 emission filter).

Respiration was measured using a Clark-type oxygen electrode (Hansatech, Oxygraph) at 20°C. Electrode was first calibrated using media saturated with either air or argon for the 100 % and the 0 %, respectively. Oxygen consumption was directly monitored in 1 mL of culture medium.

Lipids were extracted according to Folch et al. (1957). FAs were converted into methyl esters (FAME), then analyzed by gas chromatography (GC-MS/FID) on a BPX70 (SGE) column as previously described (Dellero et al., 2018a, b), using 21:0 as internal standard.
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Figure legends

Figure 1: The ploidy of *Aurantiochytrium limacinum* cells. (A) Density plot of allele frequencies calculated in non-motile cells and zoospores. Flow cytometry of non-motile cells (B and C) and zoospores (D and E). Light scatter analysis of cell populations showing cell size and granulometry (B and D). Propidium iodide fluorescence (C and E).

Figure 2: Differentially expressed genes (cut-off Log$_2$FC = |1|) for selected items. (A) signal transduction and cell motility; (B) DNA replication; (C) DNA transcription; (D) nitrogen and amino acid metabolisms; (E) carbohydrate metabolism; (F) lipid metabolism. GPCR: G protein-coupled receptors; RGS: regulator of G protein signaling.

Figure 3: Relative expression of key genes involved in FA synthesis and FA degradation in zoospores (A) and non-motile cells (B). *Tagl3* (TAG lipase 3) was chosen as a representative of TAG degradation. *FAS-1* and *PUFA-A* were chosen as representatives of FA synthesis. *Acad1* (acyl-CoA dehydrogenase, β-oxidation in mitochondria) and *Acox1* (acyl-CoA oxidase, β-oxidation in peroxisomes) were chosen for FA degradation. Dotted lines indicate significance threshold. Error bars indicate standard deviation of three independent biological repeats.

Figure 4: FA content in P-grown (red) and R-grown (green) cells. A and B) FA content, expressed as nmoles FA per mg dry weight. C and D) FA distribution, expressed as % of total FAs. E) Number of zoospores during the first 24 hours expressed as % of the total cell number. F) FA content evolution during the first 24 hours. Error bars indicate standard deviation of three independent biological repeats.

Figure 5: A schematic representation of *Aurantiochytrium limacinum* zoospore differential expression analysis based on RNA-seq. Metabolic pathways are illustrated with different colors: pink for DNA replication; purple for FA synthesis; turquoise for elongation and desaturation of FAs; yellow for lipase activities; light orange for signal transduction and G protein pathways; brown for mitochondrial β-oxidation; grey for peroxisomal β-oxidation; bright orange for glycolysis and the Krebs cycle; green for nitrogen and ammonium uptake; blue for amino acid metabolism; and dark
blue for flagellar activity and the cytoskeleton dynamics. The colors of full arrows indicate the log$_2$FC value (color scale on the right bottom of the figure) for each gene involved in lipid metabolism. The empty arrows indicate the trend of the metabolic pathway. Abbreviations: SFA, saturated FAs; PUFA, polyunsaturated FAs; TAG, triacylglycerol; DAG, diacylglycerol; MAG, monoacylglycerol. Gene abbreviations (in alphabetic order): abcd, peroxisomal ABC transporter; ACC, acetyl-CoA carboxylase; Acaa, 3-ketoacyl-CoA thiolase; Acad, acyl-CoA dehydrogenase; Acox, acyl-CoA oxidase; Acsl, long chain acyl-CoA ligase; CACT, carnitine acylcarnitine transferase; CPT, carnitine palmitoyl transferase; Δelo, fatty acid Δ elongase; ΔFAD, Δ fatty acid desaturase; Dagl, diacylglycerol lipase; Ech, enoyl-CoA hydratase; Ehhadh, bifunctional enzyme; Fas, fatty acid synthase; Hadh, 3-hydroxyacyl-CoA dehydrogenase; HADH, trifunctional enzyme; Magl, monoacylglycerol lipase; Ω3FAD, Ω3 fatty acid desaturase; Tagl, triacylglycerol lipase.
Table 1: Effect of the medium composition on the production of zoospores. The initial cell concentration was set at $5 \times 10^5$ cells mL$^{-1}$ in all conditions. P* = after 48 hours growth, amino acids were added to the culture at the same concentration as R3 (Supplementary Table 1). The number of zoospores is expressed as percentage of total cells at day 1 (D1), day 2 (D2), day 4 (D4) and day 6 (D6). Experiments were run in triplicate.

| Media | C source | N and P sources | Zoospores at D1 | Zoospores at D2 | Zoospores at D4 | Zoospores at D6 | Total number of cells at D6 (x10$^7$ cells mL$^{-1}$) |
|-------|----------|----------------|-----------------|----------------|----------------|----------------|-----------------------------------------------|
| R     | 6% Glucose | 2% YE | <3% | <1.5% | <1.5% | 0 | 7.2 |
| P     | 0.15% Glucose | 0.05% YE | 85% | 98% | 96% | 88% | 0.45 |
| R2    | 6% Glucose | NaNO$_3$ | 92% | 83% | 80% | 25% | 10 |
| R3    | 6% Glucose | NaH$_2$PO$_4$ | <4% | <2% | 0 | 0 | 8.9 |
| P*    | 0.15% Glucose | AA | 90% | 98% | 4.8% | 3.4% | 0.36 |
Table 2: Respiratory activity in R- and P-grown cells along a six-day growth. Oxygen consumption rates are expressed as nmoles O$_2$ min$^{-1}$ per million cells ± SD. Experiments were run in triplicate.

| Days | R-grown | P-grown |
|------|---------|---------|
| 0    | 1.71 ± 0.16 | 1.71 ± 0.16 |
| 1    | 6.16 ± 1.27  | 1.76 ± 0.22  |
| 2    | 2.75 ± 0.62  | 0.53 ± 0.1   |
| 4    | 3.06 ± 0.52  | 0.44 ± 0.05  |
| 6    | 2.96 ± 0.24  | 0.16 ± 0.05  |
The ploidy of *Aurantiochytrium limacinum* cells. (A) Density plot of allele frequencies calculated in non-motile cells and zoospores. Flow cytometry of non-motile cells (B and C) and zoospores (D and E). Light scatter analysis of cell populations showing cell size and granulometry (B and D). Propidium iodide fluorescence (C and E).
Differentially expressed genes (cut-off Log2FC = |1|) for selected items. (A) signal transduction and cell motility; (B) DNA replication; (C) DNA transcription; (D) nitrogen and amino acid metabolisms; (E) carbohydrate metabolism; (F) lipid metabolism. GPCR: G protein-coupled receptors; RGS: regulator of G protein signaling.
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