The organellar genome and metabolic potential of the hydrogen-producing mitochondrion of *Nyctotherus ovalis*.

Research Article

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Running head: Hydrogen-producing mitochondria of Nyctotherus.
Abstract

It is generally accepted that hydrogenosomes (hydrogen-producing organelles) evolved from a mitochondrial ancestor. However, until recently, only indirect evidence for this hypothesis was available. Here we present the almost complete genome of the hydrogen-producing mitochondrion of the anaerobic ciliate *Nyctotherus ovalis* and show that, except for the notable absence of genes encoding electron-transport chain components of Complexes III, IV and V, it has a gene content similar to the mitochondrial genomes of aerobic ciliates. Analysis of the genome of the hydrogen-producing mitochondrion, in combination with that of more than 9,000 gDNA and cDNA sequences, allows a preliminary reconstruction of the organellar metabolism. The sequence data indicate that *N. ovalis* possesses hydrogen-producing mitochondria that have a truncated, two step (Complex I and II) electron-transport chain that uses fumarate as electron acceptor. In addition, components of an extensive protein network for the metabolism of amino-acids, defense against oxidative stress, mitochondrial protein synthesis, mitochondrial protein import and processing, and transport of metabolites across the mitochondrial membrane were identified. Genes for MPV17 and ACN9, two hypothetical proteins linked to mitochondrial disease in humans, were also found. The inferred metabolism is remarkably similar to the organellar metabolism of the phylogenetically distant anaerobic Stramenopile *Blastocystis*. Notably, the *Blastocystis* organelle and that of the related flagellate *Proteromonas lacertae* also lacks genes encoding components of Complexes III, IV and V. Thus, our data show that the hydrogenosomes of *N. ovalis* are highly specialized, hydrogen-producing mitochondria.
Introduction

Mitochondria are essential organelles in aerobic eukaryotes. They play a central role in ATP production as well as in many other cellular processes, such as apoptosis, cellular proliferation, heme synthesis, steroid synthesis and Fe/S cluster assembly (Miller 1995; Lill and Kispal 2000; Scheffler 2001; Duchen 2004). Aerobic mitochondria perform oxidative phosphorylation, i.e. they use oxygen as terminal electron acceptor, oxidizing NADH and FADH₂ while producing ATP with the aid of an ATP synthase that exploits the proton gradient generated by the electron-transport chain. Eukaryotes living under anaerobic circumstances have evolved a spectrum of organelles that are related to mitochondria and have names such as anaerobic mitochondria (both temporarily and permanently anaerobic), hydrogenosomes, mitosomes, mitochondrial remnants, modified mitochondria or mitochondria-like organelles, reflecting their metabolic peculiarities (Tielens et al. 2002; Barbera et al. 2007; Tachezy and Dolezal 2007; Tachezy and Smid 2007; Tielens and van Hellemond 2007; Hjort et al. 2010).

Mitochondria that produce ATP via oxidative phosphorylation have a mitochondrial genome, as the protein complexes of proton-pumping electron-transport chains contain essential subunits whose genes were never translocated to the nuclear genome during evolution, but always remained part of the mitochondrial genome. The canonical (textbook) mitochondrion uses oxygen as final electron acceptor, but there are also anaerobically-functioning mitochondria that use compounds other than oxygen as final electron acceptor. Most organisms with anaerobic mitochondria use an endogenously produced electron acceptor, such as fumarate (Tielens et al. 2002). Some anaerobic functioning mitochondria possess a hydrogenase besides a proton-pumping electron-transport chain and, therefore, they can use protons as final electron acceptors resulting in the production of hydrogen. These organelles are the so-called hydrogen-
producing mitochondria and examples are found in *Nyctotherus ovalis* and *Blastocystis* sp.. Hydrogen production in the latter organelles, however, has not been demonstrated so far. (Lantsman et al. 2008; Stechmann et al. 2008) These organelles and those of *Proteromonas*, for which currently nothing is known about hydrogen production, are considered to be bona-fide mitochondria as they use a proton-pumping electron-transport chain and hence possess a mitochondrial genome encoding essential parts of it (Perez-Brocal, Shahar-Golan, and Clark 2010).

Hydrogenosomes are mitochondrion-related-organelles that, by definition, produce ATP and hydrogen using protons as electron acceptors. In contrast to hydrogen-producing mitochondria, hydrogenosomes have no mitochondrial genome, do not possess a proton-pumping electron-transport chain and produce ATP exclusively via substrate-level phosphorylation. Hydrogenosomes are found in various unrelated anaerobic eukaryotes such as anaerobic flagellates, chytridiomycete fungi and several anaerobic ciliates (Hackstein et al. 2001; Embley et al. 2003; Embley and Martin 2006; Hackstein et al. 2008a; Hackstein et al. 2008b; de Graaf et al. 2009a).

Mitosomes produce neither hydrogen nor ATP, but in general have retained components of a Fe/S cluster synthesizing machinery, which is now believed to be the reason why mitochondria, hydrogenosomes and mitosomes are essential to eukaryotic life (Henze and Martin 2003; Yarlett 2004; Tachezy and Dolezal 2007; Goldberg et al. 2008). Even for *Entamoeba* (Aguilera, Barry, and Tovar 2008) that has been regarded as the potential exception to this pattern, evidence is now mounting that its mitosomes also assemble Fe/S clusters (Maralikova et al. 2010), although Mi-ichi et al. (Mi-ichi et al. 2009) could not confirm this finding using a proteomics approach.
All eukaryotic organisms studied so far in sufficient molecular and ultrastructural detail appear to possess either mitochondria, hydrogenosomes or mitosomes. Furthermore, no species has been found that contains both mitochondria and mitosomes or mitochondria and hydrogenosomes, supporting the hypothesis that these organelles have a common origin. Despite this common origin, mitochondria, hydrogenosomes and mitosomes are very diverse: not only in the size of their proteomes, but also by virtue of their protein composition and function (Gabaldón and Huynen 2004). However, classical aerobic mitochondria share two characteristics: they have retained a genome, allowing unambiguous documentation of their descent from an α-proteobacterium by phylogenetic analyses (Lang, Gray, and Burger 1999), and they contain an electron-transport chain with Complexes I through IV as well as an ATPase (Complex V). In the evolution of mitochondria, hydrogenosomes and mitosomes from an α-proteobacterium, a large number of genes and proteins have been lost, gained from various sources, or retargeted to other organelles, thus shaping the huge diversity of current mitochondria and their homologs in various species (Gabaldón and Huynen 2004). Within the evolutionary “gap” between species with genome-containing mitochondria and species with mitosomes or hydrogenosomes (both lacking an organellar genome), the species N. ovalis provides an interesting link. N. ovalis is an anaerobic ciliate that lives in the hindgut of various cockroach species. It has numerous hydrogen producing organelles that are intimately associated with endosymbiotic methane-producing archaea that use the hydrogen produced by the organelles. Despite the hydrogenosomal metabolism of this organelle, we have found that it actually has a genome (Akhmanova et al. 1998a), that, similar to mitochondrial genomes (Gray 2005), encodes elements of a electron-transport chain (Boxma et al. 2005). Phylogenetic analyses of the organellar genome of N. ovalis showed that it has evolved from the genome of an aerobic mitochondrion of a ciliate ancestor.
(Boxma et al. 2005), providing direct evidence that hydrogenosomes can evolve from mitochondria. This blurs the distinction between mitochondria and hydrogenosomes. A comparable situation is found in the mitochondrion-like organelles of the phylogenetically only distantly related Stramenopile Blastocystis that is clearly distinct from the ciliates, which belong to the Alveolata. Blastocystis also retains a mitochondrial genome and parts of an electron-transport chain (Perez-Brocal and Clark 2008; Stechmann et al. 2008; Wawrzyniak et al. 2008). Since this organelle also hosts a [FeFe] hydrogenase, it can be regarded as a hydrogenosome with genome, although thus far no hydrogenase activity has been demonstrated in this species (Lantsman et al. 2008; Stechmann et al. 2008).

Also for other (genome-less) types of hydrogen producing organelles evidence is accumulating that they have evolved from mitochondria (Embley and Martin 2006). In the absence of an organellar genome, the evidence for homology is based on phylogenetic analyses of proteomes. Based on phylogenetic analyses of Complex I subunits, the hydrogenosomes of the parabasalid Trichomonas are inferred to share a common ancestor with mitochondria (Hrdý et al. 2004). Also, the hydrogenosomes of anaerobic chytrids seem to have evolved from the mitochondria of their aerobic ancestors (Akhmanova et al. 1998b). It is therefore likely that hydrogenosomes arose repeatedly by evolutionary tinkering as an adaptation to the particular requirements of hosts, which thrive in rather different anaerobic environments.

Here we describe the isolation and sequence analysis of the major part (41,666bp) of the organellar genome of N. ovalis, and compare it with the mitochondrial genomes of the ciliates Paramecium aurelia, Tetrahymena spp. and Euplotes minuta as well as with the
mitochondrial/hydrogenosomal genome of the Stramenopiles Blastocystis sp. and Proteromonas lacertae. We show that the organellar genome of N. ovalis is a typical ciliate mitochondrial genome except for the obvious absence of genes encoding components of Complexes III, IV, and V. The lack of evidence for the presence of these genes in N. ovalis coincides with the definitive absence of these genes from the organellar genomes of Blastocystis sp. and P. lacertae (Perez-Brocal and Clark 2008; Stechmann et al. 2008; Wawrzyniak et al. 2008; Perez-Brocal, Shahar-Golan, and Clark 2010). By comparing the hydrogenosomal proteins and metabolic pathways of N. ovalis with the various hydrogenosomes, mitosomes, mitochondria-like organelles and mitochondria, we show that the N. ovalis organelle is as complex as some mitochondria.

Materials and Methods

Cell isolation

Nyctotherus ovalis cells were isolated from hindgut of the cockroach Blaberus sp. var. Amsterdam. Total DNA was isolated by dissolving living cells in 8M guanidinium chloride and separation on a hydroxyapatite column using standard procedures (de Graaf et al. 2009b). The organellar DNA was isolated from the total DNA by pulsed field gel electrophoresis. This organellar DNA was used for a partial Sau 3A I digest and cloning in the Bam H1 restriction site from the vector pUC18c. From this library 500 clones were sequenced in both directions and additional 500 in one direction. Less than 3% of these clones contained a mitochondrial sequence. Finally, larger pieces of the organellar genome were reconstructed and the gaps were filled by long range PCR with specific primers resulting in a contig of 41,666bp representing the major part of the organellar genome (accession number: GU057832). Due to the very limited amount of DNA available, we did not attempt to sequence the chromosome ends.
Identification of \textit{N. ovalis} sequences likely encoding hydrogenosomal proteins

\textit{N. ovalis} sequences translated in the 6 frames were compared to the mitochondrial proteomes of human, yeast, rat and \textit{Tetrahymena}. The mitochondrial proteomes were retrieved from the mitoproteome (http://www.mitoproteome.org), the SGD database (Cherry et al. 1998), the supplementary material from Sickmann et al. (Sickmann et al. 2003), from Forner et al. (Forner et al. 2006) and from Smith et al. (Smith et al. 2007). The accession numbers of the \textit{N. ovalis} sequences are displayed in Supplementary Table 1. \textit{N. ovalis} sequences with a reciprocal best hit with one of the mitochondrial proteins (Smith Waterman, E < 0.01) were selected for further phylogenetic analysis to confirm orthology with a mitochondrial protein. Each nucleotide sequence selected was compared by the SWX algorithm to the 165 complete proteomes and the first 100 hits were kept and used together with the \textit{N. ovalis} sequence to produce a multiple alignment with Muscle v3.7 (default parameters; (Edgar 2004)). Positions of the alignments that did not contain gaps were automatically selected and a tree was subsequently derived using PhyML (Guindon and Gascuel 2003) using the JTT model and an estimated number of invariable sites with four substitution rate categories. 100 bootstraps were performed.

Phylogenetic position

To determine the phylogenetic position of the \textit{N. ovalis} hydrogen-producing mitochondrion, we composed a concatenated alignment phylogeny of the Complex I proteins that tend to be encoded on the mitochondrial genome (nad1, nad2, nad3, nad4, nad4L, nad5 and nad6), using the gamma-proteobacterium \textit{Escherichia coli} and the \textit{α}-proteobacterium \textit{Rickettsia prowazekii} as outgroups.
The sequences in each protein family were first aligned with Muscle (Edgar 2004). We then concatenated the alignment blocks, where gaps were inserted in the rare cases that a protein was missing from a certain species (e.g. nad6 from N. ovalis). Given the length of the alignment, we opted to compute a bootstrapped Maximum Likelihood (ML) phylogeny. For this we first selected the best-fit model of amino acid replacement, using the Akaike Information Criterion (AIC) as goodness of fit measure, as implemented in ProtTest (version 2.2) (Abascal, Zardoya, and Posada 2005). Accordingly, a ML phylogeny, 100 times bootstrapped, was computed with PhyML (version 3.0.1) (Guindon and Gascuel 2003) using the previously selected LG model of amino acids substitution, with a discrete gamma distribution approximated by 4 rate-categories (+4G), estimated proportion of invariable sites (+I) and observed amino acid frequencies (+F).

**Identification of sequences derived from HGT**

In order to identify Horizontal Gene Transfers present in the N. ovalis genome, we built phylogenies for all the proteins that had best hits with non-eukaryotic proteins in a Blast search. We carefully selected the homologous dataset from the 250 best hits in the non-redundant data base (NRDB), supplemented with 10 phylogenetically well distributed, best Blast hits from eukaryotes only, aligned the sequences using ClustalW (version 2.0.10) (Thompson, Higgins, and Gibson 1994) and subsequently pruned the alignment for redundancy. Each alignment was then submitted to ProtTest (version 2.4) (Abascal, Zardoya, and Posada 2005) in order to statistically infer the best-fit model of amino acid substitution, according to the Akaike Information Criterion (AIC). Finally, a Maximum Likelihood phylogeny was computed with PhyML (version 3.0.1) (Guindon and Gascuel 2003) employing the previously chosen model and bootstrapped 100 times.
Subcellular localization

Mitop2 (Andreoli et al. 2004) was used when the N-terminal part of the *N. ovalis* sequence was present to examine the presence of a putative Mitochondrial Import Signal.

*N. ovalis* cDNA and gDNA

*N. ovalis* cDNA and gDNA libraries were constructed as described earlier (Ricard et al. 2008). The macronuclear gDNA clones were obtained after amplification of total ciliate DNA with telomere primers. In general, only gDNA clones with at least one telomere were used in order to exclude contamination by bacterial sequences.

Results and Discussion

I. Gene content and structure of the organellar genome of *Nyctotherus ovalis*

In 2005, Boxma et al. showed that a 14Kb fragment of the organellar genome of the anaerobic hydrogen-producing ciliate *N. ovalis* that had been isolated from the cockroach *Blaberus* sp. var. Amsterdam was of mitochondrial origin (Boxma et al. 2005). Here we describe an analysis of the nearly complete organellar genome. Pulsed field gel electrophoresis followed by Southern blot hybridization with two different $^{32}$P-labeled probes, *nad7* and 12S (*rns*), indicated that the size of the organellar genome of *N. ovalis* exceeds 48 Kb (Supplementary Fig. 1).

Mitochondrial genes present in the organellar library were identified by Blast analysis (Altschul et al. 1997) and, in the case of non-overlap, subjected to long-range PCR to fill the gaps between the fragments. Using this approach, the major part of this genome (41,666bp) has been sequenced and reconstructed as a single contig. This part of the organellar genome contains an almost complete set of genes found in the mitochondrial genomes of other ciliates (Table 1, Fig. 1).
1). In addition, we found a stretch larger than 11 Kb that possesses 7 open reading frames (orfs) with no significant sequence similarity to any known genes.

The organellar genome contains 9 genes encoding elements of mitochondrial Complex I: \textit{nad1}, \textit{nad2}, \textit{nad3}, \textit{nad4}, \textit{nad4L}, \textit{nad5}, \textit{nad7}, \textit{nad9} and \textit{nad10}. This set of genes is also found in other ciliates (Table 1). The \textit{nad1} gene is split in the mitochondrial genomes of all ciliates studied so far into a larger (\textit{nad1a}) and a smaller (\textit{nad1b}) part (de Graaf et al. 2009b). In \textit{N. ovalis} we could identify the larger \textit{nad1a} gene piece while the \textit{nad1b} gene is likely encoded by \textit{orf224} (Fig. 2).

The other Complex I genes have a similar size as in the \textit{Tetrahymena} species and \textit{Paramecium aurelia}. This is in contrast to the situation in \textit{Euplotes minuta} and \textit{Euplotes crassus} where many Complex I genes have extended 5’ends (de Graaf et al. 2009b). The \textit{nad4} gene of \textit{N. ovalis} is a bit shorter than in other ciliates, and it is very likely that \textit{orf110} represents an additional part of the \textit{nad4} gene (Fig. 1).

Phylogenetic analysis of seven concatenated Complex I genes reveals that the Complex I of \textit{N. ovalis} is closely related to that of \textit{E. crassus} and \textit{E. minuta} and that it clusters together with the Complex I genes of \textit{Tetrahymena} and \textit{Paramecium} (Fig. 2), while it is -as expected- only distantly related to Complex I of \textit{Blastocystis} and \textit{Proteromonas}.

Organellar genes encoding components of Complexes III, IV or V could not be identified. These genes are consistently present in the mitochondrial genomes of the aerobic ciliates \textit{P. aurelia}, \textit{Tetrahymena} spp. and \textit{E. minuta} that have a size around 48 kb (Table 1; (de Graaf et al. 2009b)). Moreover, as a rule, all mitochondrial genomes sequenced to date host genes belonging to Complexes III, IV and V (Burger, Gray, and Lang 2003). The loss of genes encoding Complexes
III, IV and V from the mitochondrial genome has been regarded as indicative of a loss from the species, rather than a transfer to the nuclear genome in the mitochondrion-like organelles of Blastocystis (Perez-Brocal and Clark 2008; Stechmann et al. 2008; Wawrzyniak et al. 2008). The absence of Complex III and IV genes in the organellar genome of N. ovalis is consistent with the observation that activity of Complexes III and IV could not be observed in inhibitor studies (Boxma et al. 2005).

A total of 15 unique orfs were identified, 7 of which form a cluster with a total length greater than 11 Kb with no significant sequence similarity to any known genes. No open reading frames with sequence similarity to orfs of Tetrahymena spp, P. aurelia and E. minuta were identified.

We detected three transfer RNA genes in the organellar genome (trnF, trnY and trnW), similar to the mitochondrial genome of P. aurelia in which four tRNA genes were found (Pritchard et al. 1990). It is possible that additional tRNA genes are present in the missing part of the organellar genome. A large complex repeat region is present in the organellar genome between orf479 and ssu/rns genes. This region contains 12 repeats of 34 nucleotides followed by a (not perfectly) duplicated stretch of around 400 bp (Supplementary Fig. 2). It is unlikely that this repeat region plays the same role in initiation of transcription as suggested for the mitochondrial genome of the E. minuta and E. crassus (de Graaf et al. 2009b), because all genes found on the 41,666 bp part of the organellar genome of N. ovalis are oriented in the same direction, both before and after the repeat. In E. minuta and E. crassus, the repeat separates the open reading frames into two blocks, each having a single direction of transcription, from the repeat towards the ends of the chromosome.
We identified seven ribosomal proteins in the organelar genome of *N. ovalis*. This seems to be an average number for ciliates, since nine ribosomal proteins were found in *Tetrahymena* spp., seven in *P. aurelia* and six in *E. minuta* (de Graaf et al. 2009b). Three of these ribosomal proteins were found in all four ciliate species with sequenced mitochondrial genomes: Rps12, Rpl2 and Rpl14. In contrast, the *rps8* gene is unique for *N. ovalis* (Table 1). Notably, as in *E. minuta*, all organelar ribosomal protein genes of *N. ovalis* have an N-terminal extension that is similar to a mitochondrial targeting signal (not shown, cf. (Ueda et al. 2008; de Graaf et al. 2009b)). The overall A-T content of the organelar genome of *N. ovalis* (58.5%) is identical to the A-T content of the mitochondrial genome of *P. aurelia*. The organelar genes are not tightly packed and intergenic spacers represent 11.7% of the genome. In contrast, the genomes of the other ciliates contain no more than about 4% of intergenic spacers. The A-T content of these spacers is almost identical to the overall A-T content: 57.8%. Spacer lengths vary from 0 to 2,159 base pairs with an average size of 320 bp. In 6 cases genes have an overlap, ranging from 3-61 bp. The 11 Kb fragment that harbors the cluster of 7 orfs with no homology to other known genes contains large open reading frames with small intergenic spacers ranging from -3 to 9 bp. These 7 genes do not appear to be the remains of genes encoding components of Complex III, Complex IV and Complex V. Neither Blast searches against mitochondrial encoded proteins, nor profile-based searches against domain databases (PFAM, SMART) produced hits, even at an insignificant E value cutoff of 10. Since none of the proteins has a pI > 10, it is also unlikely that they are ribosomal proteins.
Thus, the sequenced fragment of the organellar genome of *N. ovalis* exhibits all characteristics of a ciliate mitochondrial genome, except for the absence of genes encoding components of the mitochondrial Complexes III, IV, and V.

**II. Nuclear encoded organellar proteins**

As in other mitochondria, most of the organellar proteins in *N. ovalis* are encoded by the nuclear genome, synthesized in the cytoplasm and imported into the organelle. The signal to direct these proteins into the hydrogen-producing mitochondrion resembles mitochondrial targeting signals (Boxma et al. 2005) and can be an indication that a certain protein is targeted to the organelle. Furthermore, because mitochondrial proteins tend to preserve their cellular location in evolution (Calvo et al. 2006; Szklarczyk and Huynen 2009), the orthology of nuclear encoded mitochondrial proteins from species like *Homo sapiens*, *Saccharomyces cerevisiae* and *T. thermophila* with *N. ovalis* proteins can be used to predict hydrogenosomal proteins. Here we describe in more detail nuclear encoded proteins that are probably targeted to the hydrogen-producing mitochondrion of *N. ovalis*.

Since our initial analysis (Boxma et al. 2005), we have obtained ~4,500 additional *N. ovalis* gDNA and cDNA sequences (Ricard et al. 2008). To identify new proteins involved in the hydrogenosomal metabolism we first performed a Bidirectional Best Hit analysis. 1,914 non-redundant cDNAs and 2,841 non-redundant gDNAs from *N. ovalis* were compared to the mitochondrial proteomes of human, yeast and rat. This allowed us to identify orthologs of mitochondrial proteins, for which 161 phylogenetic trees were constructed with PhyML and
subsequently analyzed by hand to determine orthology relations (not shown). This phylogenetic analysis allowed us to predict a number of new hydrogenosomal proteins as well as to provide stronger evidence for some that had been predicted previously (Boxma et al. 2005). From the phylogenetic trees, we observed that most of the *N. ovalis* sequences with close relatives among the mitochondrial proteins also cluster with their counterparts from the aerobic ciliates *Tetrahymena thermophila* and *Paramecium tetraurelia*. This indicates their recent descent from proteins functioning in aerobic mitochondria (data not shown). Furthermore, to compare the metabolic complexity of the *N. ovalis* hydrogen-producing mitochondrion with that of mitochondria, genome-less hydrogenosomes and mitosomes, we analyzed the distribution of *N. ovalis* proteins with sequence similarity to organellar proteins in other species. These were the mitochondria-possessing species *Homo*, *Saccharomyces*, *Reclinomonas*, *Plasmodium*, *Tetrahymena* and *Paramecium*, the species possessing a mitosome *Cryptosporidium*, *Encephalitozoon*, *Giardia* and *Entamoeba*, and the species with a hydrogenosome *Piromyces*, *Trichomonas*, *Psalteriomonas* and *Blastocystis* (Supplementary Table 1). Here we describe some of the proteins and their roles. Figure 3 shows how they might function together, shaping the organellar metabolism:

- Pyruvate dehydrogenase (PDH)

The PDH complex converts pyruvate and CoA into acetyl-CoA while reducing NAD$^+$ to NADH. In addition to the three subunits which have already been described (E1α, E1β and E2) we found a fourth subunit, a dihydrolipoyl dehydrogenase, also known as subunit E3, thereby completing the complex. The presence of a complete PDH complex contrasts strongly with the pyruvate catabolizing enzymes in other hydrogenosomes. Although the genome analysis of *Blastocystis*
revealed a PDH and a pyruvate:ferredoxin oxidoreductase (PFO) (Stechmann et al. 2008), Lantsman et al. measured pyruvate:NADP+ oxidoreductase (PNO) activity (Lantsman et al. 2008). Thus, the situation in Blastocystis is unclear. In Trichomonas pyruvate is decarboxylated by PFO (Hrdý, Tachezy, and Müller 2007). In contrast, in anaerobic chytridiomycete fungi (Piromyces sp. E2 and Neocallimastix sp. L2), pyruvate is catabolized by pyruvate-formate lyase (PFL) and not by PFO (Akhmanova et al. 1999). Furthermore, a large cDNA library of Piromyces constructed by the DOE Joint Genome Institute does not contain a single PFO but many PFL sequences (data not shown). The acetyl-CoA generated in N. ovalis by PDH is further metabolized by an ASCT (acetate:succinate CoA-transferase) belonging to the subfamily 1A with sequence similarity to the ASCT from Trypanosoma brucei (Tielens et al. 2010). Finally, ATP is generated by the action of succinyl-CoA synthetase (SCS), an enzyme that is also part of the TCA cycle, but this cycle is not operative in N. ovalis (Fig. 3). Also in Blastocystis and Trichomonas acetyl-CoA is metabolized to acetate by the cyclic action of ASCT and SCS. However, Blastocystis contains genes for ASCTs of the subfamilies 1B and 1C and Trichomonas contains a gene for an ASCT of the subfamily 1C that does not exhibit sequence similarity to the subfamily 1A ASCT of N. ovalis (Supplementary Table 1; (Stechmann et al. 2008; Tielens et al. 2010)).

- Tricarboxylic Acid Cycle (TCA cycle):

Based on experiments with 14C labeled glucose (Boxma et al. 2005), the TCA cycle in N. ovalis is not complete, using only the malate-fumarate-succinate part in a reductive direction (Fig. 3). Thus far, however, we have only found succinyl CoA synthetase α and β subunits (the same
enzyme as mentioned above) as well as the a and b subunit of succinate dehydrogenase / fumarate reductase. These enzymes are also present in the mitochondrion-like organelle of *Blastocystis* in which succinyl CoA synthetase, succinate dehydrogenase, fumarate hydratase and malate dehydrogenase have been identified (Supplementary Table 1) (Stechmann et al. 2008; Wawrzyniak et al. 2008). These enzymes are universal in species with mitochondria. Note that among the gDNA sequences of *N. ovalis* we did encounter a malate dehydrogenase that is orthologous to cytosolic malate dehydrogenases from other species and was therefore considered to be also cytoplasmic in *N. ovalis* as well.

- Electron-transport chain

Complex I (NADH-quinone oxidoreductase), consists of 14 subunits in eubacteria and of 33-45 subunits in mitochondria (Friedrich and Böttcher 2004; Gabaldon, Rainey, and Huynen 2005). We found 12 subunits of Complex I in *N. ovalis* (Nad1, Nad2, Nad3, Nad4, Nad4L, Nad5, Nad7, Nad9, Nad10, 24kDa*, 51kDa* and 75kDa*), among them 3 that are encoded by the nuclear genome (marked by an asterisk); the others are encoded by the hydrogenosomal genome (Fig. 2). All the *N. ovalis* Complex I proteins are part of the 14 proteins that compose the core bacterial Complex I. Also in *Blastocystis* 10 Complex I genes have been identified that are located on the organelle genome, and an additional 6 that are nuclear-encoded (Perez-Brocal and Clark 2008; Stechmann et al. 2008; Wawrzyniak et al. 2008). In *Proteromonas*, 10 Complex I proteins are encoded by the organellar genome (Perez-Brocal, Shahar-Golan, and Clark 2010).

Of complex II we detected SDHa and SDHb. In aerobic mitochondria, the four subunits of Complex II catalyze the oxidation of succinate to fumarate and the reduction of ubiquinone to ubiquinol. In *N. ovalis*, SDHa and SDHb have been proposed to function as fumarate reductase,
reversing the reaction and allowing the oxidation of quinols (rhodoquinol) (Boxma et al. 2005). In *Blastocystis*, all 4 components of Complex II have been identified (Stechmann et al. 2008).

Previous biochemical analyses have failed to detect the presence of Complex III/IV activity in the *N. ovalis* hydrogen-producing mitochondrion (Boxma et al. 2005) and, accordingly, the analysis of its genome did not provide any evidence for the presence of Complex III/IV genes (Fig. 1). Surprisingly, however, we did identify a nuclear-encoded homolog of cytochrome c1, an electron transporter related to Complexes III and IV. We sequenced the complete gene-sized-piece as well as the complete cDNA, confirming that the cytochrome c protein is transcribed. However, the cDNA appears to contain a stop codon, albeit one that is very rarely used (TGA). Therefore, with our current knowledge we cannot exclude (Ricard et al. 2008) that the cDNA is translated notwithstanding the presence of a potential stop codon. Accordingly, an alignment of the cytochrome c sequence with homologs from various species reveals that the sequence is highly conserved, but that the cysteines that normally hold the heme in cytochrome c1 are not present in the *N. ovalis* sequence, probably rendering the protein non-functional in electron transfer (Supplementary Fig. 3). Therefore, if the gene is translated, it should exhibit an alternative function. This is the first report of a cytochrome c1 that has lost the two crucial cysteines, as well as the histidine directly following the second cysteine. Only some *Trypanosoma, Leishmania* and *Euglena* species miss the first cysteine, while the second cysteine and the histidine are universally conserved among species (Priest and Hajduk 1992).

We did not find genes encoding Complex V proteins in *N. ovalis* that could use the proton motive force generated by Complex I (Boxma et al. 2005). An organelle-encoded component of
Complex V could not be identified in *Blastocystis* and *Proteromonas* either (Perez-Brocal and Clark 2008; Stechmann et al. 2008; Wawrzyniak et al. 2008; Perez-Brocal, Shahar-Golan, and Clark 2010).

Another interesting protein involved in electron transfer is the $\alpha$ subunit of the Electron Transfer Flavoprotein (ETF). ETF is an electron acceptor for various mitochondrial dehydrogenases, for example dehydrogenases that oxidize amino acids. It transfers electrons to the electron-transport chain via an ETF-ubiquinone oxidoreductase. The presence of ETF underscores the importance of the *N. ovalis* electron-transport chain in its mitochondrial catabolism.

With the notable exception of *Blastocystis* and *Proteromonas*, complete, functional Complexes I and II appear to be absent from hydrogenosomal species such as *Trichomonas*, which has only retained two subunits of Complex I (the 24kD and the 51kD subunits) (Hrdý et al. 2004; Carlton et al. 2007; Stechmann et al. 2008). The electron-transport chain is also absent from all mitosomal species. All these species lack a mitochondrial genome, whereas *N. ovalis*, *Blastocystis* and *Proteromonas* possess elements of the electron-transport chain as well as a mitochondrial genome. This supports the conclusion that all mitochondrial genomes carry genes encoding proteins of the electron-transport chain (Burger, Gray, and Lang 2003), and likewise, that all organisms with an electron-transport chain have a mitochondrial genome.

- Propionate metabolism
The predicted mitochondrial proteome contains a propionyl CoA carboxylase (EC: 6.4.1.3), that catalyses the carboxylation reaction of propionyl CoA to D-methylmalonyl CoA, and a methylmalonyl CoA mutase (also found in Blastocystis, (Stechmann et al. 2008)) that converts methylmalonyl CoA to succinyl CoA. Both propionyl CoA carboxylase and methylmalonyl CoA mutase are absent from Paramecium and Tetrahymena (Supplementary Table 1), indicating that the hydrogen-producing mitochondrion of N. ovalis does not just possess a subset of proteins present in aerobic mitochondria.

- Fatty acid metabolism

In the fatty acid activation pathway we detected a long-chain-fatty-acid-CoA ligase, which catalyzes the reversible reaction:

\[
\text{ATP} + \text{a long-chain carboxylic acid} + \text{CoA} \rightleftharpoons \text{AMP} + \text{diphosphate} + \text{acyl-CoA}.
\]

This enzyme is present in all the 11 organisms in supplementary Table 1 for which complete genomes are available, regardless of whether these organisms possess mitochondria, hydrogenosomes, or mitosomes. In contrast, the glycerol kinase is only present in the 7 organisms that possess mitochondria or hydrogenosomes. Organisms with mitosomes lack glycerol kinase, with the exception of Giardia lamblia (Supplementary Table 1).

- Amino Acid metabolism

Besides the glycine cleavage system (GCS, see below) we found several other enzymes involved in the metabolism of amino acids and specifically in the degradation of valine, methionine and isoleucine. These were a branched-chain aminotransferase (EC: 2.6.1.42), the 2-oxoisovalerate
dehydrogenase α and β (EC: 1.2.4.4) subunits and a branched-chain α-keto acid dihydrolipoyl acyltransferase (EC: 2.3.1.-). The organelle of Blastocystis also contains enzymes involved in valine, leucine and isoleucine metabolism (Stechmann et al. 2008). In addition we identified a glutamate dehydrogenase and a cystathionine β-synthase. In general, amino-acid metabolism appears much reduced in mitosomes and genome-less hydrogenosomes relative to mitochondria (Supplementary Table 1).

- Glycine cleavage system (GCS)

The dihydrolipoyl dehydrogenase of PDH is also involved in the glycine cleavage system that catalyzes the reversible reaction:

\[
glycine + \text{tetrahydrofolate} + \text{NAD}^+ \Leftrightarrow 5,10\text{-methylene-tetrahydrofolate} + \text{NH}_3 + \text{CO}_2 + \text{NADH} + \text{H}^+
\]

It provides the 5,10-methylene-tetrahydrofolate that is required for nucleotide synthesis. Of the glycine cleavage system we detected: the H-protein, a lipoic acid containing protein that carries the aminomethyl moiety of glycine that is bound to the sulfhydryl group by an S-C bond; the T-protein, which degrades reversibly the aminomethyl moiety attached to the H-protein to methylene-tetrahydrofolate and ammonia, using tetrahydrofolate in the process; and the L-protein, a dihydrolipoamide dehydrogenase that catalyzes the oxidation of the dihydrolipoyl group of the H-protein. We did not detect the substrate determining pyridoxal phosphate-containing protein (P-protein) of this complex, casting some doubt on whether the system actually uses Glycine. Nevertheless, we did find other components linked to this pathway: a
cystathionine β-synthase (CβS), a serine hydroxymethyl transferase (SHMT) that converts serine to glycine, and a 5,10-methenyltetrahydrofolate synthetase. Interestingly, two proteins of the glycine cleavage system have also been identified in the hydrogenosome of *Trichomonas* (L- and H-proteins) (Carlton et al. 2007), while the L, T and H proteins also appear to be present in the genome of the minimal mitochondria-containing *Plasmodium falciparum* (Supplementary Table 1). The mitochondrion-like organelle of *Blastocystis* contains all four components of the glycine cleavage system, including the P-protein (Stechmann et al. 2008). The glycine cleavage system is completely absent from mitosome-harboring species such as *Encephalitozoon cuniculi, Entamoeba histolytica, Giardia lamblia* and *Cryptosporidium parvum*.

- **Fe/S cluster synthesis**

The only characteristic known so far that is shared by all mitochondria, mitosomes and hydrogenosomes is the synthesis of Fe/S clusters (Lill and Mühlenhoff 2005; Tachezy and Dolezal 2007). Unfortunately, we found only a mitochondrial type I [2Fe2S] ferredoxin, which might provide reducing equivalents for Fe/S cluster synthesis as well as for other processes. Furthermore we found the mitochondrial Hsp70, which has a role in Fe/S assembly and in other processes such as protein import and folding. Although we have no reason to doubt their presence, we did not detect genes specific for Fe/S cluster assembly like Isu1/2, Nfs1, Isd11, Isa1/2 or frataxin, and therefore have no specific evidence for canonical mitochondrial Fe/S cluster synthesis in *N. ovalis*.

- **Mitochondrial protein synthesis and turnover**
We found eight mitochondrial ribosomal proteins in *N. ovalis*: Rps 8, Rps 12, Rps 14; Rpl 2, Rpl 6, Rpl 14, Rpl 11, Rpl 15 and Rpl 20; the latter three are nuclear encoded. In agreement with Smits and coworkers (Smits et al. 2007), we did not find these mitochondrial ribosomal proteins in species lacking a mitochondrial genome such as *G. lamblia, E. cuniculi, E. histolytica, C. parvum* or *T. vaginalis*. Notably, 16 ribosomal proteins are encoded in the genome of the mitochondrion-like organelle of *Blastocystis* (Perez-Brocal and Clark 2008; Stechmann et al. 2008; Wawrzyniak et al. 2008). In *N. ovalis*, we found, besides the ribosomal proteins, FtsJ (MRM2), a well-conserved heat shock protein that is responsible for methylating 23S (*rnl*) rRNA.

- Mitochondrial carriers

We detected three mitochondrial carrier proteins: one orthologous to the 2-oxoglutarate/malate carrier that imports malate into the organelle and exports oxoglutarate out of the mitochondrial matrix, one orthologous to an ADP/ATP carrier, and one carrier that does not specifically cluster with any carrier of known specificity (PET8). The oxoglutarate/malate carrier (Coll et al. 2003) also transports other dicarboxylates and tricarboxylates (such as oxaloacetate, malate, malonate, succinate, citrate, isocitrate, *cis*-aconitate, and *trans*-aconitate). This carrier may also play an important role in several metabolic functions requiring organic acid flux to or from the mitochondria, such as nitrogen assimilation, transport of reducing equivalents into the mitochondria, and fatty acid elongation (Picault et al. 2002). *N. ovalis* may use this carrier to import malate into the hydrogen-producing mitochondrion, which is transformed into fumarate and used by Complex II enzymes. In the sequenced genomes in supplementary Table 1 the oxoglutarate/malate carrier is present in several mitochondria-bearing species but absent from yeast, species with mitosomes and species with hydrogenosomes without an organellar genome.
(Supplementary Table 1). The oxoglutarate/malate carrier and a putative ADP/ATP carrier are also found in *Blastocystis* (Stechmann et al. 2008).

Additional mitochondrial proteins that are lacking in *Trichomonas* and all species with mitosomes are MPV17 and ACN9, which have homologues in human and yeast mitochondria (not shown in Fig. 3). They are, however, present in *Blastocystis*. Consistent with its presence in *N. ovalis*, MPV17 has been associated with mitochondrial genome stability (Spinazzola et al. 2006), while ACN9 has been associated with respiration (Steinmetz et al. 2002).

- **Protein import and processing**

  We detected the mitochondrial import machinery components TOM 34, MAS 5, MPP, FtsH, peptidyl-prolyl cis-trans isomerase and protein disulfide isomerase. Hsps 10, 60, 70 and 90 were also identified.

- **Mitochondrial division**

  One gene, a dynamin-related protein, has been found that might be engaged in mitochondrial division since it is an ortholog of the plant dynamin ADL2 that is known to be involved in mitochondrial division (Arimura et al. 2004).

- **Reactive oxygen species (ROS) defense**

  To our surprise, we found three systems used to cope with the detrimental effects of oxygen radicals. First of all, we found thioredoxins that act as antioxidants by facilitating the reduction of cysteines in other, oxidized proteins by thiol-disulfide exchange. Second, of the glutathione
system we found a glutathione reductase (EC 1.8.1.7) that reduces glutathione disulfide (GSSG) to the sulfhydryl form GSH. Finally, we found a peroxiredoxin.

The presence of enzymes supposed to react against oxygen radicals in an anaerobic environment might seem strange, but ROS can be produced endogenously. Complex I is a major source of ROS, although in mitochondria Complex III seems to be the principal site of $O_2^{-}$ (superoxide) formation (Chen et al. 2003).

III. Horizontal Gene Transfer (HGT)

To detect Horizontal Gene Transfer, we followed the same procedure as previously described (Ricard et al. 2006). After clustering the sequences to remove redundancy, we determined the Best Hit (BH) of 1,914 cDNAs and 2,841 gDNAs against a set of 165 complete proteomes. We identified 31 cDNAs and 25 gDNAs with a bacterium as Best Hit, and 5 cDNAs and 6 gDNAs with an archaeon as Best Hit. We analyzed the 67 *N. ovalis* sequences that have a bacterium or an archaeon as best hit further, by reconstructing the evolution of the proteins in a phylogenetic tree generated with PhyML. When requiring that the *N. ovalis* sequence clustered with a high bootstrap value with only non-eukaryotic sequences, 4 genes could be concluded to likely have been acquired through Horizontal Gene Transfer (Table 2; Supplementary Figures 4a-4d). These genes are an acetylCoA synthase (AMP-forming), a β-lactamase, a glycosylhydrolase, and an ornithine carbamoyltransferase. None of the proteins encoded by these genes has however an N-terminal targeting signal. In contrast, for the [FeFe] hydrogenase evidence for HGT and organellar location have been documented before (Boxma et al. 2007). This [FeFe] hydrogenase is remarkable since it is fused with the 24kD and 51kD subunits of a bacterial Complex I. This allows the use of NADH as electron donor. That we found only one gene obtained by HGT (the
hydrogenase) that unequivocally encodes an enzyme located in the hydrogen-producing mitochondrion, suggests that the role of HGT in the transformation to hydrogen-producing mitochondrion is limited.

**Conclusions**

With the bioinformatic analysis of the organellar genome and of the gDNAs and cDNAs we are beginning to get a comprehensive view of the metabolism of the hydrogen-producing mitochondrion of *N. ovalis*. First of all, the organelle genome is a typical ciliate mitochondrial genome, with the exception of the notable lack of components of Complexes III, IV, and V of the electron-transport chain. The latter feature is shared with the genome of the unrelated mitochondrion-like organelles of *Blastocystis* and *Proteromonas*. This phenomenon is a striking example of convergent evolution. Consequently, given the situation with *Nyctotherus*, *Blastocystis* and *Proteromonas*, we can no longer use the production of ATP via oxidative phosphorylation as a defining feature of mitochondria, rather what sets them apart from mitosomes and hydrogenosomes is the proton pumping electron-transport chain of their mitochondrion-like organelles.

Moreover, the organelle of *N. ovalis* appears to host, besides an incomplete TCA cycle, other “classical” mitochondrial pathways such as amino-acid metabolism and fatty acid metabolism. A number of these proteins are typical for mitochondria and are not present in species with mitosomes or hydrogenosomes without a genome. These include, of course, proteins of the electron-transport chain and of the mitochondrial ribosome, but also enzymes such as succinyl-CoA synthetase (however present in *Trichomonas*), pyruvate dehydrogenase, malate/a
ketoglutarate antiporter and 5,10-methylenetetrahydrofolate synthetase (see also (Ginger et
al. 2010)). What emerges is a metabolism that is in some aspects (Complex III, Complex IV,
Complex V) reduced relative to the metabolism of aerobic ciliate mitochondria and that is
strikingly similar to the metabolism of the mitochondrion-like organelle of Blastocystis.

However, N. ovalis also possesses some extra proteins relative to aerobic ciliates. A few of these
appear to have been acquired by Horizontal Gene Transfer. The case of the [FeFe] hydrogenase
is well documented (Boxma et al. 2007) but our analysis has also unearthed a few others, like an
acetyl-CoA synthetase (AMP-forming). Nevertheless, their role in the adaptation to the
anaerobic ecological niche of N. ovalis is not obvious. Not all N. ovalis proteins that we found
and that are absent from the aerobic ciliates have been gained by HGT from bacteria. We also
observe a pathway for propionate metabolism that contains a propionyl CoA carboxylase and a
methylmalonyl CoA mutase. This pathway has not yet been observed in ciliates, and besides its
presence in metazoa, has a patchy distribution among the eukaryotes, suggesting multiple loss
events rather than Horizontal Gene Transfer. Another example is a hybrid cluster protein
(prismane) that is absent from the sequenced aerobic ciliates and that among the eukaryotes is
mainly present in (facultatively) anaerobic species. However, the N. ovalis sequence does not
specifically cluster with bacterial sequences (data not shown).

By definition, the organelle of N. ovalis is a hydrogenosome, since it generates hydrogen with
the aid of an [FeFe] hydrogenase (Lindmark and Müller 1973). We have shown here that this
organelle is a mitochondrion that produces hydrogen, representing an intermediate state between
mitochondria and genome-less hydrogenosomes. This allows us to outline a hypothetical scheme
for the evolution of hydrogenosomes. A possible scenario for eukaryotes with a textbook
mitochondrion genome and metabolism to evolve into an organism with hydrogenosomes could be as follows (where the order of events could be different):

1. Evolution of a reverse action of the TCA cycle (fumarate respiration) as shown in a number of anaerobic mitochondria (Tielens et al. 2002; Tielens and van Hellemond 2007).

2. Recruitment of a hydrogenase like that shown in *Naegleria gruberi* (Fritz-Laylin et al. 2010) and *N. ovalis*.

3. Loss of Complexes III, IV and V of the electron-transport chain as shown in *N. ovalis*, *Blastocystis* and *Proteromonas*.

4. Loss of the remaining part of the organellar genome as shown, for example, in *Trichomonas*, *Psalteriomonas* and other organisms with hydrogenosomes.

It is clear that hydrogenosomes, hydrogen-producing mitochondria and mitochondrion-like organelles evolved several times from different aerobic progenitors. The eukaryotic cell is clearly characterized by the presence of a mitochondrion that has the capacity to adapt to anaerobic environments by reductive evolution to yield hydrogenosomes (and mitosomes).
Table 1: Various genes encoded by the organellar genomes of *Nyctotherus ovalis* (Nov), *Blastocystis* sp. (Blas), *Proteromonas lacertae* (Pro), *Euplotes minuta* (Emi) and *Plasmodium falciparum* (Pfa).

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■: present in the organellar genome  
□: absent in the organellar genome
Table 2: *N. ovalis* genes that likely have been acquired by Horizontal Gene Transfer from Bacteria/Archaea. The proteins associated with the genes were selected based on having a best hit with Bacteria or Archaea, followed by a phylogenetic analysis using the 250 best hits in the complete NRDB supplemented with the 10 best hits in Eukaryotes and requiring a highly supported clustering of the *N. ovalis* gene with non-eukaryotic genes, see methods for details. With each HGT candidate are indicated the genus name of the species with the best hit, the NCBI-identifier of the best hit, the source of the genetic material for the *N. ovalis* gene (cDNA or gDNA) and the extra evidence that the sequence is indeed derived from *N. ovalis* (the presence of telomeres specific for *N. ovalis* genes or of a poly A tail for a cDNA). Y indicates the presence of an ortholog of the mitochondrial protein in the corresponding species, N its absence. A blank cell is inserted if no answer is possible due to the incomplete genome. Species with mitochondria: (Has) *Homo sapiens*, (Sce) *Saccharomyces cerevisiae*, (Tth) *Tetrahymena thermophila*, (Pte) *Paramecium tetraurelia*, (Ram) *Reclinomonas americana* and (Pfa) *Plasmodium falciparum*, species with hydrogenosomes: (Tva) *Trichomonas vaginalis*, (Bla) *Blastocystis*, (Pla) *Psalteriomonas lanterna*, (Psp) *Piromyces* sp. and species with a mitosomes: (Cpa) *Cryptosporidium parvum*, (Ehi) *Entamoeba histolytica*, (Gla) *Giardia lamblia* and (Ecu) *Encephalitozoon cuniculi*. 

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<th>Ehi</th>
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Legends to the figures

Figure 1: Organellar gene maps of Nyctotherus ovalis and Euplotes minuta.

Figure 2: Maximum likelihood phylogeny of the N. ovalis hydrogen-producing mitochondrion, based on a concatenated alignment of seven mitochondrial Complex I encoded proteins (Nad1, Nad2, Nad3, Nad4, Nad4L, Nad5 and Nad6). Only bootstrap values of 50% and higher are indicated.

Figure 3: Tentative reconstruction of the metabolism of the hydrogen-producing mitochondria of N. ovalis. The metabolism is based on proteins that are orthologous to mitochondrial proteins (Methods) and proteins derived from HGT that are likely to have an organellar location based on their metabolic function. It includes the results of metabolic experiments described in Boxma et al. (2005). In red: metabolism linked to NADH (green) oxidation and electron transfer, as well as
solute carriers. In yellow are the proteins that seem to have been acquired by HGT. In blue: glycine metabolism. In orange: intermediate metabolism. Dotted arrows stand for metabolic steps we assume to be present based on biological experiments, broken arrows for the inferred metabolism for which we did not find the gene yet.

AAC: ADP/ATP carrier; ACS: acetyl CoA synthetase, AMP-forming (EC: 6.2.1.1); ALT: alanine amino transferase; ASCT: acetate:succinate CoA transferase; Cyt c1: cytochrome c1; CβS: cystathione β synthase; EfTu: elongation factor Tu; ETF: electron transfer flavoprotein; Fe-hyd: FeFe hydrogenase; FRD/SDH: fumarate reductase/ succinate dehydrogenase; GDH: glutamate dehydrogenase; GLO1: glyoxalase I; MCF Pet 8: mitochondrial carrier family; MDH: malate dehydrogenase; ME: malic enzyme; Met tRNA ft: methionyl-tRNA formyltransferase; MMM: methyl malonyl CoA mutase; MOC: malate:oxoglutarate carrier; NADH-DH: NADH :quinone oxidoreductase; PCC: propionyl CoA carboxylase; PDH: pyruvate dehydrogenase; RQ: rhodoquinone; SCS: succinyl CoA synthetase; SHMT: serine hydroxymethyl transferase;

* includes several enzymes involved in branched-chain amino acid metabolism

Supplementary material:

Supplementary Table 1: Proteins and RNAs likely involved in *N. ovalis* hydrogenosomal metabolism and their presence or absence in 14 other species.
**Supplementary Figure 1:** Pulsed field gel electrophoresis of total DNA of *N. ovalis* and Southern blot hybridizations with $^{32}$P labeled Nad7 and 12S (*rns*) probes.

**Supplementary Figure 2:** Dot plot from the repeat region of the organelar genome of *N. ovalis* (position: 22600-24600). The plot has been prepared with the aid of [http://www.vivo.colostate.edu/molkit/dnadot/](http://www.vivo.colostate.edu/molkit/dnadot/).

**Supplementary Figure 3:** Alignment of cytochrome c1. The conserved cysteines and histidine are indicated by black arrows. Alignment obtained with Clustal (Thompson 1994)

**Supplementary Figures 4a-d:** Evidence for a Bacterial or Archaeal origin of *N. ovalis* acetyl CoA synthetase (a), $\beta$ lactamase (b), glycosylhydrolase (c) and ornithine carbamoyltransferase (d). Maximum Likelihood phylogenies were computed using a LG+G+I model (LG matrix of amino acid replacement with a Gamma distribution approximated by 4 discrete rate categories and with the empirical proportion of Invariant sites).

For Figure 4b-4d the genbank identifiers are supplied with the tree. For Figure 4a the genbank identifiers, with the species initials are: Ap:284161372, Av:226946300, Bb:156084806, Bf:154310341, Ch:78044614, Cm:149194553, Cs:257060413, Ec:218702735, Es:299470777, Gv:37519728, Kk:256821575, Mb:145608328, Ms:117923908, Ms:126667192, Nc:164424614, No:60417330, Ol:145343719, Pf:229592164, Pi:301123135, Pt:229592164, Rx:108803698, Se:134098939, Se:56750212, Ss:16331700, St:51892019, Tt:269925997, Tt:55981217, Xa:154245426, Yl:50555297

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Literature cited


Figure 1

Euplotes minuta
mitochondrial genome
≈ 42Kb

N. ovalis (p.a.) partial
hydrogenosomal genome
41666 bp.
 Concatenated NAD1, NAD2, NAD3, NAD4, NAD4L, NAD5 and NAD6

Figure 2
Figure 3