The *Dictyostelium* Model for Mitochondrial Disease

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Abstract

Mitochondrial diseases are a diverse family of genetic disorders caused by mutations affecting mitochondrial proteins encoded in either the nuclear or the mitochondrial genome. By impairing mitochondrial oxidative phosphorylation, they compromise cellular energy production and the downstream consequences in humans are a bewilderingly complex array of signs and symptoms that can affect any of the major organ systems in unpredictable combinations. This complexity and unpredictability has limited our understanding of the cytopathological consequences of mitochondrial dysfunction. By contrast, in Dictyostelium the mitochondrial disease phenotypes are consistent, measurable “readouts” of dysregulated intracellular signalling pathways. When the underlying genetic defects would produce coordinate, generalized deficiencies in multiple mitochondrial respiratory complexes, the disease phenotypes are mediated by chronic activation of an energy-sensing protein kinase, AMPK (AMP-activated protein kinase). This chronic AMPK hyperactivity maintains mitochondrial mass and cellular ATP concentrations at normal levels, but chronically impedes growth, cell cycle progression, multicellular development, photosensory and thermosensory signal transduction. It also causes the cells to support greater proliferation of the intracellular bacterial pathogen, Legionella pneumophila. Notably however, phagocytic and macropinocytic nutrient uptake are impervious both to AMPK signalling and to these types of mitochondrial dysfunction. Surprisingly, a Complex I-specific deficiency (midA knockout) not only causes the foregoing AMPK-mediated defects, but also produces a dramatic deficit in endocytic nutrient uptake accompanied by an additional secondary defect in growth. More restricted and specific phenotypic outcomes are produced by knocking out genes for nuclear-encoded mitochondrial proteins that are not required for respiration. The Dictyostelium model for mitochondrial disease has thus revealed consistent patterns of sublethal dysregulation of intracellular signalling pathways that are produced by different types of underlying mitochondrial dysfunction.
1. Introduction

Mitochondrial diseases are complex degenerative disorders caused by mutations affecting nuclear genes for mitochondrial proteins or mitochondrial genes that encode subunits of the oxidative complexes or translational machinery [1]. In humans, the pathological outcomes include blindness, deafness, epilepsy, heart disease, stroke-like episodes, ataxia, muscle weakness, exercise intolerance, diabetes and kidney disease [2,3]. Mitochondrial dysfunction also plays pathological roles in neurological disorders such as Parkinson’s disease and Alzheimer’s disease [3-6].

Although much is known about the mtDNA mutations associated with human mitochondrial diseases, the relationship between genotype and phenotype is complicated and poorly understood. To gain further insight into this relationship, D. discoideum has been employed as a model organism to study mitochondrial biogenesis and disease [7-11]. The mitochondrial genome of Dictyostelium has been fully sequenced [12], while mitochondrial transcription and RNA processing in this organism have been thoroughly examined [13-15]. Mitochondrial disease has been created in Dictyostelium by a variety of methods and the pathological consequences studied. Since Dictyostelium has motile unicellular and multicellular stages with multiple cell types, many phenotypes can be examined including phototaxis, thermostaxis, macropinocytosis, phagocytosis, cell cycle progression and growth, amoeboid motility and chemotaxis, morphogenesis and intracellular growth of pathogens such as Legionella pneumophila. These varied phenotypes represent measurable, reproducible “readouts” of the intracellular signaling pathways that regulate them. By assaying them in mitochondrially diseased Dictyostelium lines, we can gain a better understanding of genotype-phenotype relationships in mitochondrial disease, without the overlaid complexities associated with mammalian systems.

To understand the cytopathological pathways involved in mitochondrial diseases it is important to identify and study proteins that play roles in these pathways. One such protein is AMP-activated protein kinase (AMPK), a ubiquitous, highly conserved protein kinase that maintains cellular energy homeostasis in healthy and diseased cells [9,16]. Bokko et al. (2007) [9] and Francione et al. (2009) [11] have shown that signalling by this protein is responsible for diverse cytopathologies seen in Dictyostelium mitochondrial disease. However, Carilla-Latorre et al. (2010) [17] reported recently that, in Complex I-specific disease, additional aberrant signalling pathways become involved so that additional AMPK-independent phenotypes are also observed. This review will highlight the contributions made by the Dictyostelium model to our understanding of mitochondrial biology and disease. The genetic manipulation strategies that have been
brought to bear and the associated phenotypes in *D. discoideum* are discussed, the role of AMPK is featured and our emerging understanding of Complex I-specific disease is described.

2. Mitochondrial Biology

Mitochondria are ubiquitous in eukaryotic cells where they are central to the maintenance of cell function and viability. These organelles, numbering from one to thousands depending on the cell type, are responsible for generating most of the cell’s ATP which constitutes the direct or indirect energy source for most cellular functions. ATP is generated by oxidative phosphorylation (OXPHOS) at the inner mitochondrial membrane. The proteins involved in OXPHOS are encoded by genes of both the mitochondrial and nuclear genomes (see Supplementary Table 1). Mutations in these or other genes that affect OXPHOS lead to an array of mitochondrial diseases [18].

2.1. The mitochondrial genome of humans and *D. discoideum*

Each *D. discoideum* cell contains about 200 copies of the mitochondrial genome which, like that of humans, is a circular molecule. The mitochondrial genome of humans is 16,569 bp [19] whereas in *D. discoideum* it is larger at 55,564 bp [12]. The average coding capacity of the mitochondrial genome in eukaryotes is 40-50 genes. The mitochondrial gene products are mainly required for five processes: respiration and/or OXPHOS, translation and in some organisms also transcription, RNA maturation and protein import [1,20]. In mammals, the majority of protein subunits (> 80) of the respiratory chain are encoded in the nucleus whereas only 13 subunits are encoded in the mitochondrial genome [18,20]. Production of these mitochondrially-encoded proteins requires the import of a variety of nuclear encoded proteins such as the single subunit mitochondrial RNA polymerase, amino acyl tRNA synthetases and the mitochondrial transcription factor A. Molecules required for mitochondrial protein synthesis which are encoded by the human and *D. discoideum* mitochondrial genomes are shown in Supplementary Table 2.

The mitochondrial genes and ORFs in *Dictyostelium* are all transcribed in the same orientation [12] (Figure 1). Some overlapping of genes is evident in the tightly packed genome, which also exhibits intergenic spacing from several nucleotides to greater than 2 kb. The genome also encodes several ORFs that do not exist in most other organisms and presumably do not play essential universal roles.

All of the genes in the *Dictyostelium* mitochondrial genome are transcribed to one of eight large polycistronic mRNA transcripts which are further processed to form a variety of smaller monocistronic, dicistronic or tricistronic mature RNA molecules [14]. Recently these eight transcripts were shown to be derived from processing of a single primary RNA transcript [15,21]. The transcription initiation site in the
Dictyostelium mitochondrial genome is located in a noncoding region upstream of rnl which encodes the large ribosomal subunit RNA (Figure 1) [15].

2.2. The mitochondrial protein import machinery

The majority of the ca. 1500 resident mitochondrial proteins are encoded by nuclear DNA, translated by cytoplasmic ribosomes and imported by the mitochondrial protein import machinery [20,22]. These include not only the other metabolic enzymes and “housekeeping” proteins, but also the remaining 70 subunits of the OXPHOS complexes (that assemble with those encoded by mtDNA) as well as factors required for expression and replication of the mitochondrial genome [23]. Defective protein import may lead to mitochondrial respiratory dysfunction, reduced ATP production and mitochondrial disease (e.g. Human Deafness Dystonia Syndrome [24]). In turn impairment of respiration can inhibit mitochondrial protein import, since import is energized both directly by the mitochondrial membrane potential (Δψₘ) and by ATP hydrolysis involving proteins such as the mitochondrial heat-shock protein (Hsp) 70 (mtHsp70) and mitochondrial import stimulation factor (MSF) [25,26]. The mitochondrial protein import machinery also assists in the correct sorting and translocation of proteins to the correct mitochondrial compartments - the matrix, the inner membrane, the intermembrane space and the outer membrane [22].

Although our knowledge of protein import into mitochondria is based mostly on research in S. cerevisiae and N. crassa, similar processes are likely to occur in other species. Tom40, Tom70 and Tom22 were found to be common elements of the mitochondrial protein machinery amongst eukaryotic genome sequences and orthologues are also encoded in the Dictyostelium genome [27].

Interestingly Ahmed et al. (2006) [28] reported that import of some mitochondrial proteins in Dictyostelium occurs cotranslationally in vivo. Cotranslational import of mitochondrial proteins has also been observed in yeast [29,30]. Current evidence suggest that it occurs as a result of transport and binding of the mRNA to the mitochondrial surface in a manner that depends on secondary stem-loop structures in the mRNA, often in untranslated regions of the transcript [29].

3. Human mitochondrial diseases

Mutations in mitochondrial or nuclear genes that encode components of the translational machinery or the OXPHOS complexes have been associated with human mitochondrial diseases. Due to the presence of mitochondria in all cells of the human body, every tissue can be adversely affected by mutations in the mitochondrial genome. However, there is tissue-specific variation in the numbers of mitochondria, the proportion of mutant mitochondrial genomes, the energy demands of the cell, the isoforms of nuclear-encoded mitochondrial proteins that are expressed and the extent of cellular and mitochondrial
proliferation. Depending on which tissues are affected most in a given individual, this leads to a bewildering complexity in mitochondrial diseases which exhibit a very broad range of possible symptoms (Supplementary Tables 3 and 4). Individuals with the same mitochondrial disease mutation can exhibit huge variation in clinical symptoms while different mutations can result in similar clinical manifestations. Thus phenotypes associated with human mitochondrial diseases are unpredictable from specific genetic defects.

In mitochondrial diseases, pathogenic mutations in mitochondrial genes usually affect only a subset of the genomes (a state referred to as heteroplasmy) and the extent of the phenotypic outcomes is sometimes related to how large this subset is (the mutant load). This can contribute to a phenomenon whereby a particular pathological outcome is only observed when the mutant load exceeds a particular threshold. A good example of such a threshold is shown by NARP (Neuropathy, Ataxia and Retinitis Pigmentosa) and LS (Leigh Syndrome) which can result from the same mutation. NARP is caused by 70-90% of mutated mtDNA whereas LS develops if the mutation load is above 90–95% [31,32]. In some cases, mtDNA mutations must be above a particular threshold for significant ATP depletion and/or symptoms of the disease to occur [33]. Biochemical thresholds that influence the extent to which ATP levels are reduced as a consequence of gene mutations, have been described in detail by Rossignol et al. (2003) [34].

4. D. discoideum as a model organism for mitochondrial disease

The genetics, biochemistry and signal transduction processes of D. discoideum have been studied extensively and are well understood. With fully sequenced mitochondrial [12] and nuclear [35] genomes, the organism is one of a small number of nonmammalian model organisms recognized by the National Institute of Health (NIH) in the U.S.A. for their importance in biomedical research (http://www.nih.gov/science/models/). Many Dictyostelium genes are orthologues of disease-related genes [35] and this has facilitated the establishment of Dictyostelium models for investigating a variety of human diseases [10,39,40], including mitochondrial disease [21]. In those cases where the relevant experiments have been done, heterologous expression and study of Dictyostelium genes in other organisms, or vice versa, has confirmed directly that the functions of orthologous proteins have been conserved during evolution [10].

4.1. Life cycle and phenotypes

What makes Dictyostelium so valuable a model organism is that it combines genetic, biochemical and cell biological tractability with a unique life cycle providing a great variety of readily assayed, reproducible disease phenotypes. The organism grows as isolated amoeboid cells that divide mitotically and obtain nutrients either by phagocytic consumption of bacteria or by macropinocytosis of liquid medium. Starvation induces a developmental programme in which the amoebae differentiate and consequently
become responsive to an attractant, cAMP, which they now synthesize and secrete. The resulting aggregation process leads to formation of a motile, multicellular organism (the “slug”) containing multiple cell types organized in a well-defined spatial pattern. The migratory slug exhibits highly sensitive phototactic and thermotactic responses under control of its specialized anterior cells. After a variable period of migration it forms a fruiting body by a combination of morphogenetic movements and further cell differentiation. This life cycle, with its motile unicellular and multicellular stages, offers diverse phenotypic “readouts” of the signaling pathways that regulate them. These include cell growth and division [38]; amoeboid motility and chemotaxis [39]; phagocytosis and macropinocytosis [40]; phototaxis and thermotaxis [41,42]; cell and tissue differentiation and pattern formation [43]; autophagic cell death [44]; multicellular tissue movement and morphogenesis [45,46]. Dictyostelium has also been used to study the interactions between microbial pathogens, such as Legionella pneumophilia, and their hosts [47]. The molecular mechanisms of Legionella pathogenesis in D. discoideum are similar to those in human cells [48,49]. Despite this phenotypic richness, many of the complexities associated with mammalian systems are eliminated in the Dictyostelium model, since all stages of the life cycle can be readily studied in clonally derived, genetically identical cell lines. The study of mitochondrially diseased Dictyostelium strains has accordingly revealed consistent, reproducible outcomes that contrast with the unpredictability of mitochondrial disease in humans.

As mentioned previously human mitochondrial diseases are characterized by pathological or biochemical thresholds. Mitochondrial disease in Dictyostelium has exhibited similar thresholds in that some phenotypes appear more sensitive to mitochondrial dysfunction than others [7,8,50]. Although genetic manipulations that would impair respiration resulted in similar phenotypic outcomes, the severity of the phenotypes varied amongst the mitochondrially diseased strains so that particular defects such as slow growth were not observed in every strain [7-9,50]. Much of this variation in the severity of the phenotypic aberrations in mitochondrially diseased Dictyostelium was shown to be caused by differences in the severity of the underlying genetic defect. As described in the next section, such differences are readily created and quantitatively measured in Dictyostelium.

4.2. Genetic manipulation to produce mitochondrial disease in Dictyostelium

In order to study mitochondrial disease in Dictyostelium various methods have been employed to create sublethal mitochondrial dysfunction. These include RNAi (interfering RNA) [51] or antisense inhibition of expression of specific mitochondrial proteins [8,9], heteroplasmic disruption of mitochondrial genes [7,11,52] and knockout of nuclear genes encoding nonessential mitochondrial proteins [53-56]. Mitochondrial defects in Dictyostelium have also been generated pharmacologically through the use of ethidium bromide to deplete the cells of mitochondrial DNA [57].
4.2.1. Heteroplasmic targeted disruption

The first *Dictyostelium* mitochondrial gene targeted for heteroplasmic disruption was *rnl*, the mitochondrial large ribosomal subunit RNA gene [7]. The original disruption of this gene was a result of a nontargeted insertion into a gene important for phototaxis. Subsequent targeted disruptions confirmed the effect on phototaxis and initiated the study of mitochondrial disease in *Dictyostelium* [7]. Another *Dictyostelium* mitochondrial gene *rps4*, encoding a ribosomal protein, was targeted for heteroplasmic disruption by Inazu et al (1999) [52]. Apart from *rnl* and *rps4*, 8 other mitochondrial genes have been disrupted and regardless of which gene was targeted, the expression of the entire mitochondrial genome was reduced [50]. The strength of signals in Southern blots of the mitochondrial DNA indicates that only a subset of the mitochondrial genomes is affected in these strains i.e. the disruptions are heteroplasmic [7,50] allowing *Dictyostelium* to be used as a model for human heteroplasmic mitochondrial disease.

4.2.2. Antisense and RNAi inhibition of nuclear genes encoding essential mitochondrial proteins

Mitochondrial disease can also be caused by mutations affecting nuclear genes that encode mitochondrial proteins. An example is *hspA*, which encodes chaperonin 60 (Cpn60), a protein located in the mitochondrial matrix. Cpn60 is required for correctly folding proteins newly imported into the mitochondrial matrix. A deficiency of Cpn60 is linked to a mitochondrial disease in humans, the symptoms of which include severe neurological and developmental defects and depletion of many respiratory enzymes [58-60]. In *D. discoideum* a Cpn60 insufficiency was created by antisense inhibition of expression of the protein [8,9]. Transformation of *Dictyostelium* with an antisense-inhibition construct produces transformants in each of which a different number of copies of the plasmid are integrated into the nuclear genome. The copy number is stable, easily determined using several techniques [61,62] and correlates with the reduction of expression of the target gene and the phenotypic outcomes [8,9]. RNAi constructs have been used similarly to reduce the expression of the catalytic subunit of mitochondrial succinate dehydrogenase (respiratory Complex II) (Lay & Fisher, unpublished results) and Dd-TRAP1 (*D. discoideum* Tumor necrosis factor receptor-associated protein), a Hsp90 homologue which translocates to the mitochondria in early differentiation [51]. By analogy with Hsp90 in other organisms, it may play a role in chaperoning some proteins *en route* to the mitochondria.

4.2.3. Targeted disruption of nuclear genes encoding nonessential mitochondrial proteins

While the heteroplasmic disruption of essential mitochondrial genes is possible, it is lethal to disrupt a nuclear gene in *Dictyostelium*’s haploid genome if it encodes an essential mitochondrial protein. Such a gene must supply its encoded protein to every mitochondrion in the cell. However some mitochondrial proteins are not essential for viability. Genes encoding these proteins can be disrupted through
homologous recombination and the associated phenotypes examined. Such genes include $cluA$ [54], $torA$ (Tortoise) [53], $fszA$ and $fszB$ [55], $midA$ [17,63] and $aoxA$ [64].

4.3. *Dictyostelium* mitochondrial disease pathology

Several broad categories of mitochondrial dysfunction have been studied in the *Dictyostelium* model (Table 1). The first is generalized oxidative phosphorylation (OXPHOS) defects that are expected to impair multiple respiratory complex deficiencies. Such defects have been studied by sublethal genetic or pharmacological manipulations that reduce but do not eliminate the oxidative phosphorylation (OXPHOS) capacity of the mitochondria. The second category of mitochondrial defects studied in *Dictyostelium* are those that affect mitochondrial functions other than OXPHOS. Finally, in very recent work, a class of mutant has been found that produces a specific reduction in the activity of only one of the respiratory complexes (Complex I). Each of these categories of mitochondrial dysfunction has contributed to our growing understanding of the cytopathological pathways underlying mitochondrial disease.

4.3.1. Generalised OXPHOS defects

The most obvious biochemical consequence of a generalized deficiency in oxidative phosphorylation is a reduced capacity to synthesize ATP. In humans the phenotypic outcomes of mitochondrial disease were thought accordingly to result from a depletion of ATP and the different energy requirements of cellular functions. However, in *Dictyostelium* it has been shown that mitochondrial disease phenotypes arising from generalized OXPHOS defects (Table 1) do not result simply from a depletion of ATP but from disturbances in intracellular signalling networks. In mitochondrial disease the energy generating capacity of the mitochondria is compromised which results in activation of intracellular energy stress signals. These signals are relayed into various signalling pathways to produce phenotypic affects, some of which are more sensitive to mitochondrial dysfunction than others.

The first mitochondrial disease phenotype to be discovered in *Dictyostelium* was impaired phototaxis and thermotaxis in the multicellular slug stage of the life cycle [7]. *Dictyostelium* slugs display extremely sensitive, accurate orientation towards light and in temperature gradients [47,48]. Phototaxis and thermotaxis pathways converge early so that almost all of the proteins involved are required for both processes. The photo/thermosensory transduction pathways in *Dictyostelium* are not completely understood, but many participating molecules have been identified which could be downstream targets of energy stress signalling. These include heterotrimeric G proteins [47,48]; the second messengers cAMP, cGMP, IP$_3$ and Ca$^{2+}$ [47,48]; signalling proteins such as RasD [65], GefE and Gefl [66]; protein kinases such as PKB and ErkB [67]; and cytoskeletal proteins such as GRP125 [68], villidin [69], CAP [70], filamin [71] and FIP [72]. A number of the proteins involved form a photosensory signalling complex that is assembled on
the scaffolding protein filamin [67,73]. The phototaxis and thermotaxis signalling pathways were impaired by mitochondrial disease in all cases tested, regardless of how the mitochondrial dysfunction was caused. Thus heteroplasmic disruption of any of 10 different mitochondrial genes and down regulation of chaperonin 60 all resulted in phototaxis and thermotaxis defects [7,8,11,50,52]. Likewise phototaxis was impaired by depletion of mitochondrial DNA using ethidium bromide treatment [57].

The second phenotype to be affected by mitochondrial disease is growth. *Dictyostelium* cells can grow either on bacterial lawns (obtaining nutrients by phagocytosis) or axenically in liquid medium (obtaining nutrients by macropinocytosis). Mitochondrially diseased *Dictyostelium* cells showed impaired growth in liquid and on bacterial lawns [7-9]. However these growth defects were not a result of altered phagocytosis or macropinocytosis [9], nor did the cells show alterations to their size, implying a coordinate inhibition of both cell growth and cell cycle progression (Ahmed and Fisher, unpublished data).

The proteins controlling the cell cycle that could be dysregulated in mitochondrial disease include seven cyclins, six cyclin-dependent kinases identified in *Dictyostelium* and homologues of inhibitors of cell cycle progression such as the retinoblastoma protein Rb, a homologue of which is encoded by rblA [35]. The major signalling pathway controlling cell growth in metazoans involves the protein kinase TOR in a multiprotein complex called TORC1. Inhibition of this pathway involves activation of TSC2 which in turn inactivates Rheb and thereby inactivates TOR [38,74]. These proteins are conserved in *Dictyostelium* [35] but their functional roles in signalling pathways involved in growth control or mitochondrial disease have not been determined. In mammals AMPK inhibits TORC1 directly through the RapTOR subunit [75] and indirectly by activating TSC2 [74]. Because mitochondrial dysfunction compromises ATP generation, AMPK should be chronically activated in mitochondrially diseased cells. This indeed appears to be the case for mitochondrially diseased *Dictyostelium* cells [9] as well as Parkinson’s [76], Alzheimer’s [77] and Huntington’s [78] disease neurons. As described in greater detail in later sections, chronic AMPK signalling was found to be responsible for the slow growth of mitochondrially diseased *Dictyostelium* cells [9].

The third phenotype which is consistently affected by mitochondrial dysfunction is the differentiation of cells into stalk and spore cells, with mitochondrially diseased cells showing an increased number of cells directed into the stalk differentiation pathway and mislocalisation of these cells in the multicellular slug. At culmination this results in fruiting bodies with thick short stalks. This effect was seen in mitochondrially diseased cells created either by genetic manipulation [8,9] or by treatment with ethidium bromide [57]. In addition, almost all cells were directed into the stalk differentiation pathway by inhibitors of cyanide-resistant respiration that also impaired the mitochondrial membrane potential, loss of which is known to stimulate autophagy [79]. Stalk cell differentiation in *Dictyostelium* involves a programmed cell death accompanied by the accumulation of autophagic vacuoles. Mitochondrial dysfunction in *Dictyostelium* thus
appears to direct cells into an autophagic cell death pathway that may be analogous to the autophagic cell death implicated in neurodegeneration in humans [44].

The last phenotype to be associated with mitochondrial disease in Dictyostelium is altered transition from growth to development and subsequent chemotactic aggregation. Inazu et al (1999) [52] showed that knock down of the mitochondrial rps4 gene resulted in greatly impaired cell aggregation and significantly lower levels of car1, an early developmental gene. Depletion of mitochondrial DNA by ethidium bromide treatment impaired aggregation significantly and reduced the expression of early developmental genes including carA [57]. Aggregation was also impaired in chaperonin 60 or succinate dehydrogenase antisense-inhibited cells resulting in fewer, smaller aggregates [8,9,Lay & Fisher, unpublished].

4.3.2. Defects not known to affect OXPHOS

Mutations affecting the mitochondria have been created which do not damage the energy generating capability of the mitochondria (Table 1). One example is knockout of the mitochondrial fission proteins FszA and FszB [55], whose closest homologues are FtsZ proteins that mediate binary fission in the α-proteobacteria, the presumed ancestors of mitochondria. In some eukaryotic lineages such as fungi, animals and plants, no mitochondrial FtsZ proteins have been identified and mitochondrial division is primarily undertaken by FIS1 and DNM1L (dynamin-like protein also known as DRP1) [80]. Mitochondrial fission in Dictyostelium involves both dynamin-like proteins [81] and FtsZ proteins [55], indicating that the latter were retained in the amoebozoan lineage but discarded in some others. Disruption of FszA or FszB impaired mitochondrial division, producing more elongated mitochondria, while disruption of FszB but not FszA also caused growth defects [55]. However, none of the other characteristic mitochondrial disease phenotypes described in the preceding section were observed.

FszB was found localised to submitochondrial bodies of unknown function [55] which are also enriched in another mitochondrial protein TorA (tortoise) [53]. TorA null amoebae move slowly and fail to suppress inappropriate lateral pseudopod formation during chemotaxis, defects which may explain why they grow slowly on bacterial lawns. The specificity of the motility and chemotaxis defects suggests that TorA null cells still produce wild type levels of ATP [53]. There is no human homologue for TorA and it is unknown whether TorA mutant cells display any of the phenotypes typically associated with mitochondrial respiratory dysfunction in Dictyostelium.

A nuclear gene required for normal subcellular localization of mitochondria is cluA. First discovered and functionally characterized in Dictyostelium, cluA has homologues in all eukaryotes whose genomes have been sequenced. CluA null mutants exhibit pronounced perinuclear clustering of the mitochondria in mutant cells [54]. CluA is also involved in cytokinesis since mutant cells were multinucleate and slightly
impaired in growth [54]. The involvement in cytokinesis is likely to be an indirect result of the contractile ring being physically occluded by large aggregates of mitochondria that block completion of cleavage [82].

Another mitochondrial protein that has been studied genetically is TRAP1, a homologue of mammalian TRAP1 (Tumour Necrosis Factor Receptor-Associated Protein 1). TRAP1 belongs to the Hsp90 molecular chaperone family and localises to the mitochondria as well as to extramitochondrial sites such as the nucleus, secretory granules and cell membranes [83-85]. The protein is localised to the cell cortex of cells growing at low cell densities and translocates to the mitochondria as the cell density of growing cells increases [56]. RNAi inhibition of TRAP1 expression resulted in slow vegetative growth and delayed aggregation, while overexpression resulted in premature aggregation.

The alternative oxidase encoded by aoxA in Dictyostelium is a mitochondrial protein with homologues in animals, plants, fungi and bacteria [86]. It can accept electrons from ubiquinone in place of Complex III and pass them directly to molecular oxygen, thus providing an alternative pathway for electron flow in which protons are pumped across the mitochondrial inner membrane only at Complex I. This process of cyanide-resistant respiration appears to serve primarily as an electron sink that prevents leakage of electrons in single electron transfers to oxygen that would form superoxide anions and hence other reactive oxygen species (ROS). Kimura et al. (2010) [64] knocked out aoxA in Dictyostelium and found no obvious phenotypic changes in growth or development under normal conditions. However development in the mutant was completely blocked by millimolar concentrations of KCN that merely delayed development in the wild type.

4.3.3. The role of AMPK

It was first proposed by Wilczynska et al. (1997) [7] that a cytopathological effect of mitochondrial disease is impairment of signal transduction pathways with some pathways being more sensitive than others. Recently a molecular link between mitochondrial disease and cellular signalling was identified in Dictyostelium in the form of AMP-activated protein kinase (AMPK) [9]. AMPK is an essential sensor and homeostatic regulator of cellular energy status [87]. The serine/threonine protein kinase is a heterotrimer composed of a catalytic α subunit, a regulatory γ subunit and a β subunit which acts as a scaffold for the other subunits. In mammalian cells there are three genes encoding isoforms of the γ subunit and two for each of the α and β subunits, whereas in Dictyostelium only one of each subunit is encoded in the genome.

Activation of AMPK occurs when AMP binds to the two Bateman domains on the γ subunit relieving its inhibition of the catalytic α subunit. The α subunit is now able to be phosphorylated at a threonine residue by an upstream kinase and is also resistant to dephosphorylation by phosphatases. Because of competition between AMP and ATP for binding to the Bateman domains, AMPK activation is very sensitively regulated
by AMP/ATP ratios with high ATP levels producing the inactive heterotrimer. Several upstream AMPK kinases have been identified to date - LKB1 [88,89], CaMKKα and β [90,91] and TAK1 [92]. Of these, LKB1 is reported to be the major activator [89].

Once activated, AMPK homeostatically rectifies low cellular energy levels by activating pathways that produce ATP while inhibiting others that consume it. These effects are exerted by inhibiting key enzymes in biosynthetic pathways (e.g. acyl CoA carboxylase in fatty acid synthesis and 3-hydroxy-3-methylglutaryl-CoA reductase in cholesterol biosynthesis), activating protein translocation (e.g. the GLUT4 glucose transporter to the plasma membrane, increasing glucose uptake) [93], and altering gene expression (e.g. repression of glucose responsive genes and protein synthesis [94,95], induction of mitochondrial biogenesis [96,97]). These actions enable AMPK to regulate the AMP/ATP ratio and maintain healthy energy levels within the cell.

Because of its exquisite sensitivity to AMP/ATP ratios, AMPK exerts its effects prior to a serious depletion in energy. Bokko et al. (2007) [9] provided molecular genetic evidence that AMPK is chronically activated in mitochondrial disease. Mitochondrial disease was created in Dictostelium by antisense inhibition of chaperonin 60 to produce the defective phenotypes discussed earlier, including defective photo/thermotaxis, growth and morphogenesis. These defective phenotypes were phenocopied by overexpressing a constitutively activated form of AMPK (AMPKαT). Conversely, when AMPKα expression was knocked down in mitochondrially diseased cells (chaperonin 60 antisense inhibition), the defective phenotypes were suppressed. AMPKαT-expressing strains also exhibited increased mitochondrial mass and higher ATP levels, as expected since AMPK stimulates biogenesis and ATP production in mammalian cells. Mitochondrially diseased cells did not show changes in mitochondrial mass or steady state ATP levels presumably due to a balance between reduced biogenesis and ATP production caused by antisense inhibition of chaperonin 60 and increases due to chronic activation of AMPK. However this chronic AMPK hyperactivity in mitochondrially diseased cells leads to permanent dysregulation of the downstream energy-consuming pathways under AMPK’s control (Figure 2).

4.3.4. Dictostelium and Complex I dysfunction

NADH:ubiquinone oxidoreductase or Complex I (CI) is the first and the largest complex of the respiratory chain and couples the oxidation of NADH to reduction of ubiquinone and the transport of protons. It is formed by 45 protein subunits in mammalian cells that assemble together into a structure of approximately 1 Mda [98]. CI has important implications in human pathology due to its role in the generation of reactive oxygen species and the increasing number of diseases whose cause is directly or indirectly associated with CI function. About 40% of inherited disorders of the OXPHOS system involve isolated or combined
deficiencies in Cl. As described previously these deficiencies present diverse combinations of clinical manifestations, including fatal childhood disorders such as Leigh syndrome. Complex I dysfunction has also been linked to neurodegenerative disorders such as Parkinson’s and Alzheimer’s diseases [99-101]. Despite its importance in human pathology many aspects of Cl function, including its regulation, assembly and structure are poorly understood.

Cl is present in prokaryotes in its minimal version formed by the so-called core subunits. This bacterial complex is able to perform the essential redox reactions required for its function. However eukaryotic Cl contains additional subunits (known as accessory subunits) that might have been acquired during evolution to enrich Cl with new functions and regulatory properties. Interestingly Cl has been lost in certain eukaryote lineages such as the one leading to the yeast S. cerevisiae. However, other fungi such as Yarrowia lipolytica and Neurospora crassa do contain Cl and have been used as simple models for its study [102,103].

Like Yarrowia and Neurospora, Dictyostelium possesses Cl and homologous genes encoding Complex I subunits can be recognized in the mitochondrial and nuclear genomes. Also like them, Dictyostelium has proteins similar to all core subunits and a subset of accessory proteins (Supplementary Table 5). Interestingly the Dictyostelium mitochondrial genome codes for three Cl subunits that are nuclear encoded in other organisms [104]. This is consistent with divergence of major eukaryotic lineages before the process of transferring mitochondrial genes to the nuclear genome had been completed.

Despite its complexity only few assembly factors have been described to be required for the correct assembly and stability of this huge multiprotein complex. Six have been described to be involved in human Cl deficiency: NDUFAF1, NDUFAF2, NDUFAF3 and NDUFAF4 [105-108], C8orf38 [109] and C20orf7 [110,111]. Others are required for Cl assembly but have not yet been implicated in human disease such as Ecsit [112], AIF [113], Indl [114,115] and MidA/C2orf56 [17]. Interestingly, two (C20orf7 and MidA) are putative methyltransferases suggesting that methylation might play a relevant role in Cl function.

MidA is a conserved protein that was described for the first time in Dictyostelium as a mitochondrial protein required for cellular bioenergetics since null mutants have reduced ATP levels and are compromised in growth and development [63]. Bioinformatics modeling and site directed mutagenesis suggested that MidA contains a methyltransferase domain required for its function. The protein’s ability to complement the mutant phenotype is abolished by mutations in a critical glycine in the catalytic core that is predicted to interact with the methyl donor S-adenosyl methionine (SAM) [17]. Dictyostelium cells lacking MidA show a specific defect in Complex I activity and this role has also been confirmed in mammalian cells since the knock-down of human MidA in HEK293T cells results in an isolated defect in the activity and assembly of Complex I. Both Dictyostelium and human MidA interact with the Cl core subunit Ndufs2, but the consequences of this interaction and whether or not MidA is involved in Ndufs2 methylation remain
open questions [17]. Only two protein methylation modifications have been detected in Complex I subunits. One of them is in the bovine NDUFB3 (B12) subunit that is methylated at conserved His residues but has no homologue in Dictyostelium. Interestingly, the other one is the human NDUFS2 subunit that harbors a methylated arginine, R323 [116].

MidA knockout in Dictyostelium causes a complex phenotype that shows similarities but also differences with that of other mitochondrially diseased strains (Table 1 and Figure 2). Defects in phototaxis and thermotaxis, a hallmark of Dictyostelium mitochondrial defects, are present in the MidA mutant and can be rescued by AMPK antisense inhibition, similarly to the other described mitochondrial mutants [9,17]. However, midA cells are also defective in phagocytosis and macropinocytosis, a phenotype that is not rescued by AMPK antisense inhibition and is not even present in the other described mitochondrial mutants. These results suggest that isolated defects in CI might cause abnormal phagocytosis and pinocytosis. Alternatively, MidA might have additional functions beyond CI that could affect these cellular activities. The study of other CI-specific mutants in Dictyostelium is necessary to distinguish between these possibilities and to clarify the complex cytopathology associated with CI disease.

5. Conclusion

The Dictyostelium model for mitochondrial disease is proving to be a rich field for investigation. It is simple and tractable enough to yield consistent, reproducible phenotypes in clonally derived mutants, yet has a sufficiently complex life cycle with unicellular and multicellular motile forms to provide a rich lode of phenotypic readouts of varied intracellular signalling pathways. It has taught us that sublethal mitochondrial dysfunction at the cellular level produces phenotypic outcomes from disturbed intracellular signalling networks not merely ATP depletion, cumulative oxidative damage and cell death. The implication is that whole organism pathology may arise earlier and be more nuanced than is often supposed - an outcome of sublethal cellular dysfunction not simply the result of cell death. It is to be hoped that with this insight and greater understanding of the nature of the cellular disturbances produced by mitochondrial disease, new treatment strategies will present themselves for these currently untreatable disorders.

Acknowledgements

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### Tables.

**Table 1. Phenotypes associated with mitochondrial dysfunction in *Dictyostelium***.

<table>
<thead>
<tr>
<th>Method of generating mitochondrial dysfunction</th>
<th>Phenotype</th>
<th>Growth on bacteria</th>
<th>Growth in broth</th>
<th>Phagocytosis</th>
<th>Pinocytosis</th>
<th>Phototaxis</th>
<th>Thermotaxis</th>
<th>Morphogenesis</th>
<th>Aggregation</th>
<th>Chemotaxis</th>
<th>Legionella susceptibility</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pharmacological, expected to affect respiration</strong></td>
<td>Ethidium bromide inhibition of mtDNA replication</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>- (Stalky)</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td>57</td>
</tr>
<tr>
<td><strong>Genetic, expected to affect respiration</strong></td>
<td>Heteroplasmic mitochondrial gene disruption (<em>rnl, nad5, cob, nad2, atp6, atp1, cox3, ORF1740, ORF796</em>)</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>- (Stalky)</td>
<td>-</td>
<td>-</td>
<td></td>
<td>7,11,50</td>
</tr>
<tr>
<td></td>
<td>Heteroplasmic <em>rps4</em> disruption</td>
<td>+/-</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chaperonin 60 antisense inhibition</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>- (Stalky)</td>
<td>-</td>
<td>-</td>
<td></td>
<td>8,9,11</td>
</tr>
<tr>
<td><strong>Genetic, respiratory complex-specific defect in respiration</strong></td>
<td>MidA knockout producing specific Complex I deficiency</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+/- (Stalky)</td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td>63,17</td>
</tr>
</tbody>
</table>
### Genetic, not known to affect respiration

<table>
<thead>
<tr>
<th>Genetic Factor</th>
<th>Nuclear fszA, fszB disruption</th>
<th>Nuclear cluA disruption</th>
<th>Nuclear torA disruption</th>
<th>Nuclear Dd-TRAP1 RNAi inhibition</th>
<th>Nuclear aoxA disruption</th>
<th>55, Fisher (unpublished)</th>
</tr>
</thead>
<tbody>
<tr>
<td>+ (fszA); - (fszB)</td>
<td>+ (fszA); - (fszB)</td>
<td>+ (Defective cytokinesis)</td>
<td>- (Defective cytokinesis)</td>
<td>-</td>
<td>+</td>
<td>54</td>
</tr>
<tr>
<td>- (fszA); - (fszB)</td>
<td></td>
<td></td>
<td></td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>53</td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td>-</td>
<td>56</td>
<td></td>
</tr>
</tbody>
</table>

+ : Wild type phenotype. - : Aberrant phenotype. +/- : Mildly aberrant phenotype. Shaded cells, Phenotype not reported.
References:


83. Cechetto JD, Gupta RS. Immunelectron microscopy provides evidence that tumor necrosis factor receptor-associated protein 1 (TRAP1) is a mitochondrial protein which also localises at specific extramitochondrial sites. Exp Cell Res 2000;260:30-9.


Figure legends.

Figure 1. Organisation, transcription and transcript processing of the *Dictyostelium* mitochondrial genome.

The *Dictyostelium* mitochondrial genes as well as introns and noncoding regions are shown. The genome is transcribed from a single start site as indicated by “Transcription start” [15]. The primary transcript may be rapidly and cotranscriptionally processed into smaller mature transcripts. Only the secondary transcripts A (3.1 kb), B (4.6 kb), C (5.6 kb), D (9.5 kb), E (6.0 kb), F (6.5 kb), G (3.7 kb), H (8.7 kb) and their smaller derivatives (tertiary transcripts) have been detected in northern hybridisation studies (Barth *et al.*, 2001). From Fig. 1 of Barth *et al.* (2007) [21].

Figure 2. Cytopathological pathways in *Dictyostelium* mitochondrial disease.

Two pathways are shown. 1. Generalized mitochondrial respiratory dysfunction affecting multiple oxidative phosphorylation complexes compromises mitochondrial ATP generation, leading to chronic AMPK activation. The resulting dysregulation of intracellular signalling produces multiple cytopathological outcomes. Not shown is the homeostatic feedback by which AMPK stimulates mitochondrial biogenesis and ATP production. In mitochondrially diseased cells chronic AMPK activation can thereby maintain ATP at normal levels while at the same time causing chronic downstream cytopathologies. 2. In addition to AMPK-dependent pathways, a MidA-dependent dysfunction (possibly through Complex I-specific deficiency indicated by the question mark) impairs endocytic pathways (phagocytosis and macropinocytosis) in an AMPK-independent manner. Different, more limited cytopathologies may be caused by mutations that affect other aspects of mitochondrial biology without impairing ATP production.