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# **Regulation of ATPase dimer formation and optogenetic control of metabolism**

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# **Regulation of ATPase dimer formation and optogenetic control of metabolism**

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El Doctor José Antonio Enríquez Domínguez, líder del grupo de investigación “Genética Funcional del Sistema de Fosforilación oxidativa” del Centro Nacional de Investigaciones Cardiovasculares (CNIC),

CERTIFICA

Que la Tesis Doctoral “Regulation of ATPase dimer formation and Optogenetic control of metabolism” ha sido realizada en la Fundación Centro Nacional de Investigaciones Cardiovasculares Carlos III, bajo su tutela, y que reúne las condiciones para optar al grado de Doctor.

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Fdo. José Antonio Enríquez



# Abstract



## ABSTRACT

The oxidative phosphorylation system comprises three fundamental processes: electron transport, proton pumping and ATP synthesis. Electron transport is required for both anabolic and catabolic processes, which require the delivery of electrons to the mitochondrial electron chain. This flux of electrons drives proton pumping and generates an electrochemical gradient across the mitochondrial inner membrane. This gradient is used by ATP synthase (ATPase, complex V) to generate ATP. Electron transport and proton pumping can be detached in many non-mammalian eukaryotes because they have non-proton-pumping oxidases. Mitochondrial ATPase is structured in two domains, the  $F_0$  domain, hydrophobic and embedded in the mitochondrial inner membrane, and the  $F_1$  domain, hydrophilic, which protrudes through the mitochondrial matrix and is responsible for the ATP synthesis from ADP and Pi. In recent years, new ATPase subunits have been discovered: DAPIT and 6.8kDa protein (MP68). Very little is known about them, but DAPIT has been related with glucose metabolism and diabetes. They have been recently reported to form an ATPase subdomain called the intermembrane space domain (IMD), formed by only these two proteins. In this thesis, the functional role of DAPIT and MP68 was evaluated by generating knockout mouse models. An opposite role for each one was demonstrated: while DAPIT promotes ATPase dimerisation, MP68 prevents it. In the absence of DAPIT, an ATPase subcomplex is dissociated, which indicates that DAPIT promotes ATPase structure maintenance.

Next we focused on developing cellular tools where mitochondrial ATP synthesis capacity could be studied separately from electron transport. For this purpose, we designed optogenetic tools to generate an electrochemical gradient in mitochondria. Optogenetic tools were initially used in the neural field to modify and control the channel flux in neurons to alter their activity. Here we transformed light energy into an energy, that cells can use to grow. By expressing deltarhodopsin in mammalian cells, we were able to rescue energy deficiencies using light as a source of energy.



# Resumen



## RESUMEN

El sistema de fosforilación oxidativa concentra tres funciones principales: el transporte de electrones, el bombeo de protones y la síntesis de ATP. El transporte de electrones es requerido para procesos anabólicos y catabólicos en los que participan las reacciones de oxidación–reducción que se llevan a cabo en la mitocondria. El bombeo de protones está impulsado por el flujo de electrones, y genera un gradiente electroquímico en la membrana interna mitocondrial. Este gradiente es utilizado por la ATP sintasa (ATPasa o complejo V) para la síntesis de ATP. Los procesos de transporte de electrones y el bombeo de protones solo pueden desacoplarse en organismos eucariotas no mamíferos, ya que presentan oxidasas que no bombean protones. La ATPasa mitocondrial está formada por dos dominios, el dominio  $F_0$ , hidrofóbico, embebido en la membrana interna; y el dominio  $F_1$ , hidrofílico, localizado hacia la matriz mitocondrial, y que contiene la estructura responsable de la síntesis de ATP a partir de ADP y fosfato inorgánico. En los últimos años, se han descubierto nuevas subunidades, DAPIT y la proteína 6.8kDa (MP68). La función de ambas es poco conocida, pero DAPIT ha sido relacionada con el metabolismo de la glucosa y diabetes. Recientemente, se ha descrito un nuevo subdominio de la ATPasa llamado “dominio del espacio intermembrana” (IMD) que está formado únicamente por ambas proteínas. En esta tesis, se ha evaluado la función de DAPIT y MP68 generando ratones con falta de función para cada una de ellas, demostrando su función opuesta entre ellas. Mientras que DAPIT promueve la formación de dímeros de ATPasa, MP68 promueve su disociación. Además, en ausencia de DAPIT se genera un subcomplejo de ATPasa, indicando que DAPIT promueve el mantenimiento de la estructura de este complejo.

Por otro lado, en esta tesis se han desarrollado herramientas celulares para estudiar la capacidad de síntesis de ATP de forma independiente al transporte de electrones. Para ello, se han utilizado herramientas optogenéticas. Estas técnicas se han utilizado tradicionalmente en estudios neurales, para modificar y controlar el flujo a través de canales modificando su actividad. En esta tesis, se han desarrollado células capaces de transformar la energía de la luz en energía aprovechable para crecer. Gracias a la expresión de una deltarodopsina en células de mamífero, éstas son capaces de rescatar deficiencias energéticas usando la luz como fuente de energía.



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# Abbreviations



Å	Ångström
A/BA	Acrylamide/bis-acrylamide
ADP	Adenosine diphosphate
ATP	Adenosine triphosphate
ATPase	ATP synthase
BN	Blue native
bp	(DNA) base pairs
BSA	Bovine serum albumin
CaCl <sub>2</sub>	Calcium chloride
CoQ	Ubiquinone
CsA	Cyclosporine A
Cyt c	Cytochrome c
DDM	n-Dodecyl--D-Maltoside
DNA	Deoxyribonucleic acid
DP5	Maltopentose
MgAMP-PNP	Adenosine 5'-[β, γ-imido]triphosphate magnesium
MgADP	Adenosin 5' (Trihydrogen Pyrophosphate)
EDTA	Ethylenediaminetetraacetic acid
EMMA	European Mouse Mutant Archive
ES	Embryonic stem
EuMMCR	European Mouse Mutant Cell Repository
FBS	Foetal bovine serum
FCCP	Carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone
FRT	Flippase recognition target
GTT	Glucose tolerance test
IF <sub>1</sub>	Inhibitor factor 1
kDa	kilo Dalton
IMM	Inner mitochondrial membrane
IMS	Intermembrane space
ITT	Insulin tolerance test
IVF	<i>In vitro</i> fertilisation
IMS	Intermembrane space
nDNA	Nuclear DNA
mETC	mitochondrial electron transport chain
MOI	Multiplicity of infection
MOPS	(3-(N-morpholino)propanesulfonic acid)
mPTP	Mitochondrial permeability transition pore

MRI	Magnetic resonance imaging
mtDNA	Mitochondrial DNA
NADH	Reduced nicotinamide adenine nucleotide
ORF	Open reading frame
OXPHOS	Oxidative phosphorylation system
o/n	Overnight
PBS	Phosphate-buffered saline
Pi	Inorganic phosphate
rev/s	Revolutions per second
ROS	Reactive oxygen species
RT	Room temperature (25°C)
SDS	Sodium dodecyl sulphate
STZ	Streptozotocin
TCA	Tricarboxylic acid cycle
Tris	Tris(hydroxymethyl)aminomethane
$\mu\text{M}$	micro Molar
V	Vol





# Introduction

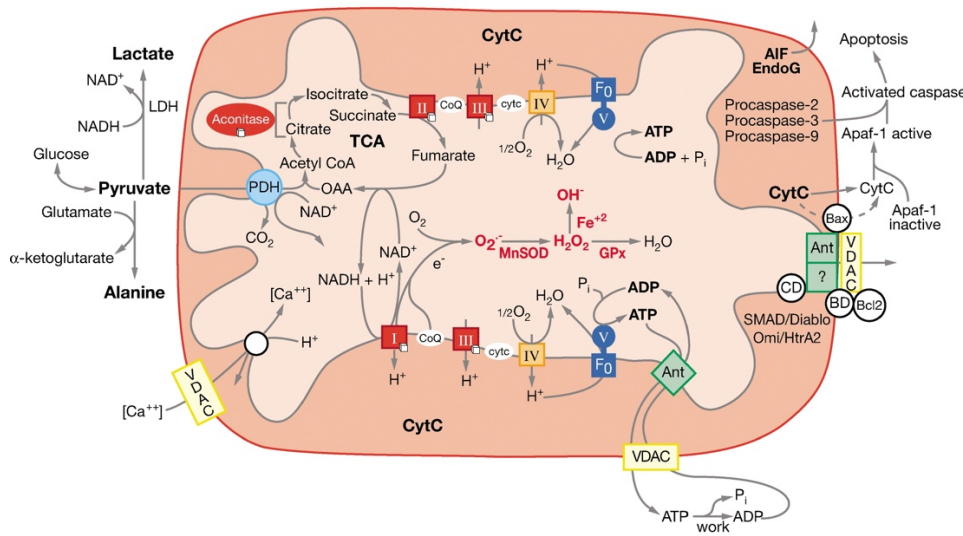


## I. MITOCHONDRIA BIOLOGY

### I.1 Mitochondrial function

Mitochondria are endosymbiotic organelles that harbour different critical functions, provide energy to cells in the form of ATP, and also harbour the TCA cycle and beta oxidation of fatty acids and the metabolism of certain amino acids. They also generate reactive oxygen species (ROS), buffer calcium from the cytosol and regulate cell death. Mitochondria have their own genetic information in mitochondrial DNA (mtDNA), discovered in the 1960s (Nass and Nass, 1963; Schatz, Haslbrunner and Tuppy, 1964; Sagan, 1967; Schwartz and Dayhoff, 1978). mtDNA not only provides genetic information, but is also important for cell viability. Mitochondria are not only important for their role in energy production, but other roles, such as controlling stem cell function and fate are being progressively recognised for mitochondria (Folmes *et al.*, 2012). In addition, a metabolic change from glycolytic to oxidative, as well as the regulation of the mitochondria permeability transition pore (mPTP), are both necessary to achieve embryonic cardiomyocyte maturation (Hom *et al.*, 2011). It has been shown that mutation in mtDNA causes diseases (Holt, Harding and Morgan-Hughes, 1988; Wallace *et al.*, 1988), triggers carcinogenesis (Carew and Huang, 2002; Petros *et al.*, 2005) and is related with the apoptosis process (Kroemer and Reed, 2000). The accumulation of these mutations results in premature aging in mice (Trifunovic *et al.*, 2004).

In mitochondria, energy is produced through mitochondrial oxidative phosphorylation (OXPHOS). Glucose is converted into two pyruvates and thus reduces two cytosolic  $\text{NAD}^+$  to NADH and produces two ATP molecules. Next pyruvate enters mitochondria and is transformed into Acetyl-CoA and  $\text{CO}_2$  to produce  $\text{NADH}+\text{H}^+$  and  $\text{FADH}_2$ . Fatty acids are also oxidised by beta oxidation to produce acetyl-CoA,  $\text{NADH}+\text{H}^+$  and  $\text{FADH}_2$ . In both cases, acetyl-CoA enters the tricarboxylic acid cycle (TCA) and generates more  $\text{NADH}+\text{H}^+$ . Both  $\text{NADH}+\text{H}^+$  and  $\text{FADH}_2$  are oxidised by the mitochondrial electron transport chain (mETC). NADH releases electrons to complex I by reducing ubiquinone (CoQ) to ubiquinol. The different enzymes that use FAD as a cofactor (like Complex II) release electrons directly to CoQ to form ubiquinol. Ubiquinol is reoxidised by complex III, which sends electrons downhill to cytochrome c (cyt c) and complex IV, and finally to oxygen to produce  $\text{H}_2\text{O}$  (Fig. 1) (Wallace, Fan and Procaccio, 2010; Enríquez, 2016).



**Figure 1. Mitochondrial oxidative phosphorylation diagram** (adapted from Wallace, 2005). Glucose is converted into pyruvates which enter into the mitochondria and is transformed into Acetyl-CoA producing  $\text{NADH} + \text{H}^+$  and  $\text{FADH}_2$ . These two molecules are oxidised by the mitochondrial electron transport chain.

Complexes I, III and IV use the energy released by the electron flux to translocate protons ( $\text{H}^+$ ) from the inner mitochondrial matrix to the intermembrane space (IS). In this way they generate an electrochemical gradient, which is also known as proton motive force (p.m.f.) and drives ATP synthesis from ADP and  $\text{P}_i$  by  $\text{H}^+$ -ATPase.

## 1.2 Mitochondrial morphology

Mitochondria are delimited by two membranes: the inner (IMM) and the outer (OMM). The space between both membranes is known as the inner mitochondrial space (IMS), while the matrix is located inside the IMM. This morphology was observed for the first time by electron microscopy (Palade, 1952; Sjostrand, 1953). The IMM and OMM form different sub-compartments that perform specific functions. Thus the OMM forms specialised structures that interact with the endoplasmic reticulum (ER), named mitochondria-associated membranes (MAMs) (Vance, 2014) whose composition differs from other OMM areas (Giorgi *et al.*, 2015). In this way there are defined specialised regions in the IMM very much enriched in OXPHOS components located in the cristae edge.

Mitochondrial morphology can switch from an elongated form to a more fragmented status due to fusion and fission events and, as a result, the cristae structures in the IMM change. These cristae-remodelling processes occur during apoptosis, for example, when cytochrome c is released from mitochondria (Scorrano *et al.*, 2002). The morphology of these cristae also changes under physiological conditions, in different organs and

developmental stages. Mitochondrial fission occurs during the process of eliminating non-functional mitochondria by mitophagy (Twig *et al.*, 2008), or in the G2/M phase of the cell cycle to secure mitochondria segregation during cell division (Smirnova *et al.*, 2001). The opposite phenomenon, the fusion process, increases in a different cell cycle phase, G1/S, (Mitra *et al.*, 2009), during a starvation period (Rambold *et al.*, 2011) or during embryonic development (Chen *et al.*, 2000).

During development, mitochondria take a simple structure with fragmented cristae and a small mtDNA copy number (Folmes *et al.*, 2012; Folmes and Terzic, 2016). Cardiomyocytes from E9.5 embryos have fragmented and perinuclear mitochondria, but on day E13.5 they extend more into the cytoplasm, and become organised and filamentous (Hom *et al.*, 2011). Taken together, all these changes facilitate energy production.

The morphology of mitochondrial cristae is important from a bioenergetic point of view (Cogliati *et al.*, 2013). Cristae increase the IMM surface and the amount of mETC (Scheffler, 1999), and it has been proposed that this increases ATP production capacity by oxidative phosphorylation (Palade, 1953). Apart from this incremented surface, it is believed that cristae also provide a benefit in energy production terms (Cogliati, Enriquez and Scorrano, 2016), based on mathematical models (Demongeot *et al.*, 2007), by minimising the distance between ADP/ATP translocation and the ATP synthesis site.

Cardiolipin is an important lipid that determines mitochondrial shape. It is an acidic phospholipid found in mitochondrial membranes and its reduction in the IMM has been shown to severely alter the mitochondria ultrastructure, as in Barth syndrome (Xu *et al.*, 2006; Chicco and Sparagna, 2007). Cardiolipin has also been described to establish the supercomplexes of mETC (Pfeiffer *et al.*, 2003; McKenzie *et al.*, 2006), and the III<sub>2</sub>+IV<sub>2</sub> association is unstable in its absence (Mileykovskaya and Dowhan, 2014).

Another protein involved in cristae remodelling is Opa1, optic atrophy type 1 protein, a dynamic-like GTPase whose mutations are associated with autosomal dominant optic atrophy type I disease (Alexander *et al.*, 2000; Delettre *et al.*, 2000). Processing large Opa1 isoforms is essential for mitochondrial fusion (Duvezin-Caubet *et al.*, 2006). There are different proteases that control Opa1 cleavage, including Yme1L or Oma1 (MacVicar and Langer, 2016).

The ATPase complex is an important modulator of cristae formation. In *Paramecium multimicronucleatum* mitochondria, ATPase is on the edge of cristae (Allen, Schroeder and Fok, 1989). In yeast, ATPase subunits e and g have been reported as non-essential for ATP synthesis, but their absence confers a characteristic structure to cristae in the form of concentric circles (Paumard *et al.*, 2002). Other subunits, such as ATP6, have been described as modulators of ATPase conformation, but are not absolutely necessary for cristae formation (Rak *et al.*, 2007; Habersetzer *et al.*, 2013). Another hypothesis that could

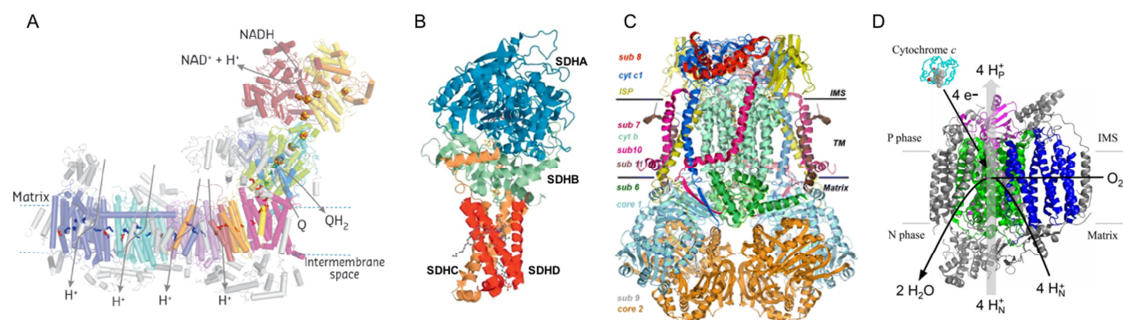
explain the role of ATPase is that it forms a pH gradient that could lead to a membrane curvature that gives rise to cristae formation (Rieger, Junge and Busch, 2014).

### I.3 Electron transport chain complexes and their organisation

The OXPHOS system constitutes the most important pathway to produce energy in the cell. It is formed by ATPase and four complexes: complexes I, II, III and IV.

#### Complex I

The NADH dehydrogenase complex transfers two electrons from NADH to ubiquinone, and is coupled with the translocation of four protons from the matrix to the IMS. It is composed of 44 proteins with a weight of 960kDa, of which seven are codified by mtDNA. Recently, its structure has been resolved by cryoelectronmicroscopy with a resolution of 4.2 Å in bovine mitochondria (Fiedorczuk *et al.*, 2016; Zhu, Vinothkumar and Hirst, 2016). Complex I is formed by three different modules: N, that links NADH and oxidises it; Q, that transfers electrons to CoQ; and P, the hydrophobic module in the IMM involved in proton pumping (Fig. 2A).



**Figure 2. Complexes from the mitochondrial electron transport chain. A.** Mammalian complex I structure. **B.** Complex II structure (modified from (Bezawork-Geleta *et al.*, 2017). subunits of SDH: SDHA (blue), SDHB (green), SDHC (brown), and SDHD (red). **C.** Complex II structure, modified from (Xia *et al.*, 2007). **D.** Structure and cofactor of mammalian cytochrome c oxidase. The three subunits codified in mtDNA are in green (Cox1), pink (Cox2) and blue (Cox3) (modified from: Rich, 2017).

#### Complex II

Complex II, or succinate dehydrogenase, is also an enzyme of the tricarboxylic acid cycle that catalyses the oxidation of succinate to fumarate. It transfers electrons from  $\text{FADH}_2$  to ubiquinone. It is located in the IMM, faces the intermembrane space and is composed of four subunits: SDHA, SDHB (both make up the catalytic site), SDHC and SDHD (allow its location in the IMM) (Hägerhäll, 1997) (Fig. 2B). It is a special complex as it does not pump protons, and is the only complex that does not contain any subunits codified by mtDNA.

### Complex III

Complex III, or cytochrome c reductase, transfers electrons from ubiquinone to cytochrome c (Fig. 2C). This process is coupled with the pumping of four protons to the IMS. It is composed of 11 subunits, of which only one, mtCytb, is codified by mtDNA (Iwata, 1998).

### Complex IV

Complex IV, or cytochrome c oxidase, is the last of the complexes of the mETC. Its function is to transfer electrons from cytochrome c to molecular oxygen by generating H<sub>2</sub>O. Four protons are translocated during this reaction. This complex is formed by 14 subunits, three of which form the catalytic core, and are encoded by mtDNA, Cox1, Cox2 and Cox3 (Fig. 2D).

### Complexes organisation

The organisation of the mETC in the IMM has been a matter of debate for decades. In the 1950s the “solid model” was proposed by Keilin and Hartree to explain the organisation of mitochondrial complexes (Keilin and Hartree, 1947). This model described the complexes joined as one to act as static entities to secure communication among them (Slater, 2003). The “solid model” was questioned after the isolation of active individual complexes took place (Hatefi *et al.*, 1962). Moreover, the description of the mobility of coenzyme Q between CII and CIII went against the proposed “solid model”.

A new model, namely the “fluid model”, was described in 1986 by Hackenbrock and colleagues (Hackenbrock, Chazotte and Gupte, 1986). This model stated that complexes could diffuse into the IMM in an aleatory manner. Electron transport was described as a random collision between complexes that depended on the velocity of coenzyme Q and of cytochrome c diffusion

In 2000, the “supercomplexes” structures were discovered by native electrophoresis, Blue Native gel (BN-PAGE) (Schägger and Pfeiffer, 2000). BN-PAGE allowed the discovery of the stable associations among OXPHOS complexes. The largest form of supercomplexes is formed by the I+III<sub>2</sub>+IV association, known as the respirasome. Other frequent associations are the I+III<sub>2</sub> supercomplex and III<sub>2</sub>+IV. Although these associations also exist in plants, they can vary in the stoichiometry (Eubel, Heinemeyer and Braun, 2004). As example, in *Saccharomyces cerevisiae*, since they lack complex I, the only supercomplex observed was III<sub>2</sub>+IV<sub>2</sub> (Schägger and Pfeiffer, 2000). The largest amounts of these supercomplexes are found in mammals such as mouse, bovine or human.

Based on evidence that favours the two previous models, (Boekema and Braun, 2007; Lenaz and Genova, 2007), a new model was proposed, known as the “plasticity model”

(Acín-Pérez *et al.*, 2008), which combines the possibility of finding complexes, and not only those associated among them, but also individually.

The I+III<sub>2</sub> supercomplex from *Arabidopsis thaliana* was the first to be investigated by cryoelectronmicroscopy techniques (Dudkina *et al.*, 2005). In 2016, three independent groups simultaneously provided respirasome structures with significantly improved resolution (Gu *et al.*, 2016; Letts, Fiedorczuk and Sazanov, 2016; Wu *et al.*, 2016). Sazanov (Letts, Fiedorczuk and Sazanov, 2016), and the Kühlbrandt groups (Sousa *et al.*, 2016) provided a resolution of 5.8 Å or 9 Å, respectively, from bovine heart. The Yang group provided a resolution of up to 4 Å from porcine mitochondria and human mitochondria. These groups also provided details of two different respirasome conformations, “tight” and “loose”. In a tight conformation, complex IV comes into contact with both complex I and III, whereas in a loose conformation complex IV comes into contact only with complex I (Letts, Fiedorczuk and Sazanov, 2016).

The stability of supercomplexes is influenced by different proteins, and also by other non-protein factors. The first proposed proteins were Rcf1 and Rfc2 in *S. cerevisiae* (Chen *et al.*, 2012). The deletion of these proteins suppresses supercomplexes formation because it impairs the association between CIII and CIV. Currently, it seems that these factors are in fact directly involved in the assembly of CIV, and indirectly impact supercomplex formation through CIV assembly alterations (Garlich *et al.*, 2017). Therefore, they cannot be considered specific factors for supercomplex assembly.

The only *bona fide* assembly factor described to date is COX7A2L, or SCAF1, which denotes the name of “Supercomplex Assembly factor 1”. SCAF1 is a protein of 113 amino acids, including the mitochondrial import sequence. A mutated version of SCAF1 has been described in a subset of inbred mouse strains, notably all investigated C57B/L6-derived substrains (Lapiente-Brun *et al.*, 2013). This mutated form of 111 amino acids has an intramolecular microdeletion which causes the loss of two amino acids that render it unable to interact with complex IV (Cogliati *et al.* 2016). The mechanism of interaction between CIII and CIV is determined by the interchange of the structural subunits of CIV that determine if this complex associates with CIII<sub>2</sub>, dimerise-forming CIV<sub>2</sub> or remains as monomer CIV. These proteins are COX7A1, which favours CIV dimer formation, COX7A2 that favours CIV monomers, and COX7A2L or SCAF1 that allows the physical superassembly between complexes III<sub>2</sub> and IV. Additional tissue-specific isoforms modulate CIV dimerisation: COX6A2 or COX6A1 (Cogliati *et al.*, 2016).

Apart from these protein factors, other components, such as cardiolipin, also regulate the supercomplexes assembly (Pfeiffer *et al.*, 2003). This phospholipid is important for membrane potential maintenance (Gohil *et al.*, 2004). Cardiolipin-deficient cells show the instability of the III<sub>2</sub>+IV<sub>2</sub> supercomplex (Pfeiffer *et al.*, 2003).

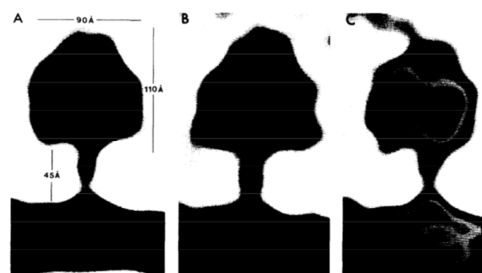
The regulation of supercomplexes formation and their plasticity is important to substrate adaptation and optimised mitochondria metabolism. The assembly of CIII and CI generates two independent pools of coenzyme Q for the electrons from both NADH and FADH<sub>2</sub> by avoiding substrate competition in the mETC (Lapiente-Brun et al, 2003; Guaras et al, 2006).

The organisation of complexes in supercomplexes is an important issue during embryo development in mice. The assembly of supercomplexes around stage E11.5 of embryo development has been shown to correspond to a cardiac development step (Beutner, Eliseev and Porter, 2014), which indicates that mitochondrial regulation is also important during cardiac maturity (Menendez-Montes *et al.*, 2016).

## II. ATP SYNTHASE

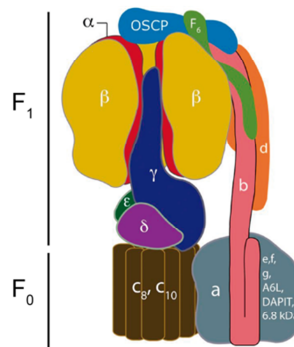
### II.1 F<sub>1</sub>F<sub>0</sub>-ATPase structure

The F<sub>1</sub>F<sub>0</sub>-ATPase complex is formed by 16 proteins (Collinson *et al.*, 1994) and is located in the membranes of different structures, including mitochondria, chloroplasts or eubacteria. The general shape of the complex is similar in all three. F<sub>1</sub>F<sub>0</sub>-ATPase is composed of three domains, the F<sub>1</sub>, F<sub>0</sub> and the slender stalk that links all the domains together. The F<sub>1</sub> domain is extrinsic to the membrane, which is where ATP is produced and hydrolysed. The F<sub>0</sub> domain is embedded in the membrane and forms the channel that allows protons to pass through the membrane and to couple it with ATP synthesis. Initially the slender stalk was considered an artifact, until it was defined by electron microscopy in *Escherichia coli* (Gogol, Lücken and Capaldi, 1987) (Fig. 3). Later, it was demonstrated that if this stalk was disrupted, the F<sub>1</sub> domain would be released as a globular complex.



**Figure 3. Analysis of the electron microscopy images of ATPase from *Escherichia coli*.** The average of the different side views of F<sub>1</sub> and F<sub>0</sub> computed from (A) 30 particles aligned by correlation methods; (B) 10 particles with a cleft in the F<sub>1</sub> domain; (C) 10 particles with a solid F<sub>1</sub> domain.

The  $F_1$  domain is comprised of five globular proteins,  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$  and  $\epsilon$  in a specific stoichiometry  $\alpha_3\beta_3\gamma_1\delta_1\epsilon_1$  with a weight of 350kDa (J.E.Walker *et al.*, 1985; Abrahams *et al.*, 1994; Collinson *et al.*, 1994). Subunits  $\alpha$  and  $\beta$  are arranged in an alternating conformation, which generates a spherical structure with a diameter of 100 Å. This structure envelops subunits  $\gamma$  that are localised in the central block by fixing these subunits to the  $F_0$  domain. Subunits  $\delta$  and  $\epsilon$  contribute to this association by binding to the end of the cylinder form by c subunits (Fig. 4).



**Figure 4. Organisation of protein subunits in mitochondria ATP synthase (from (Walker, 2013)).**

The  $F_0$  domain's name is attributed to oligomycin because is the domain that confers sensitivity to this antibiotic (Racker, 1963; J. Symersky *et al.*, 2012).  $F_0$  is composed of subunits a, b, c, d, e, f, g, and A6L in the membrane domain. The c-subunits form a ring with different numbers of subunits depending on the species. In bovine mitochondria, the number of c-subunits is 8, and is 10 in yeast (Jindrich Symersky *et al.*, 2012).

Other ATPase subunits form the central stalk, or stator, whose function is to fix  $F_1$  during rotation in order to provide structural sustenance. It is composed of the oligomycin-sensitive conferring protein (OCP), and subunits b, d and F6. The structure of these proteins is of low-degree conservation between species, but their function is very well conserved.

There are other subunits that form part of ATPase, but their role is not yet well-defined. The f subunit, essential for ATP synthesis, produces the phenotype of uncoupled ATP synthase when it is lost (Spannagel *et al.*, 1997). Subunit i has been described only in yeast ATPase, and it produces the dissociation of the ATPase complex when it is absent. Although subunits e, k, and g are involved in dimer formation, the only essential subunit for this process is e (no protein homologous to subunit k in yeast has been described in mammals).

Two new peptides have been recently discovered, proteins MLQ and AGP. MLQ is a 6.8kDa mitochondrial proteolipid, MP68 (Terzi *et al.*, 1990; Chen *et al.*, 2007; Meyer *et al.*,

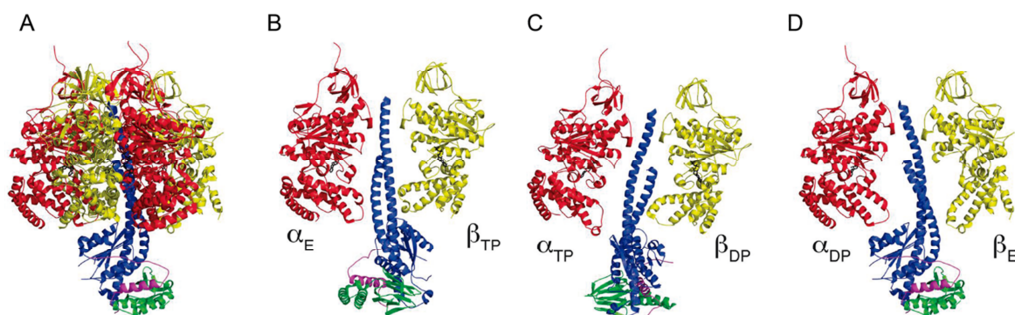


In physiological contexts, ATPase dimerisation is also important during ageing. In mature cells, the proportion of dysfunctional mitochondria is higher than in young cells. These mitochondria also display a vesicular morphology that lacks cristae, and ATPase dimers dissociate into monomers (Daum et al., 2013).

As described in Section 1.2, ATPase plays an important role in cristae formation, promoting cristae morphology maturation. In line with this, ATPase is important during germ cell differentiation in *Drosophila*, and not because of its role in energy production, but because of the function it performs in cristae formation (Teixeira et al., 2015).

### II.3 ATP synthesis

The catalytic ATPase site has been described based on X-ray images from bovine (Abrahams et al., 1994) and yeast  $F_1F_0$ -ATPase. As the  $\gamma$  subunit assembly takes place asymmetrically, ATPase takes an asymmetric structure and can be observed in different conformations. While the central stalk rotates, ATPase can be found in three varying conformations of the  $\beta$ -subunit, which confers different catalytic properties (Fig. 6). Although conformations  $\beta_{TP}$  and  $\beta_{DP}$  take similar structures, they exhibit differential preference in binding nucleotides. While  $\beta_{TP}$  can bind MgAMP-PNP (a non-hydrolysable analogue of ATP),  $\beta_{DP}$  binds MgADP. Conformation  $\beta_E$ , also known as “open” or “empty”, takes a different structure due to the  $\gamma$ -subunit, and is unable to bind any nucleotide. This latter conformation is the active one, which releases the ATP molecule.



**Figure 6. Structure of the F<sub>1</sub> domain of bovine ATPase** (modified from (Walker, 2013)). **A.** Catalytic F<sub>1</sub> with the three different conformations. After each rotation of 120°, a conformation change occurs. **B.** State represented by conformation  $\beta_{TP}$  with affinity to Mg-AMP-PNP. **C.** Conformation represented by the presence of  $\beta_{DP}$ , with affinity to Mg-ADP. **D.** State without any affinity to nucleotides, characterised by  $\beta_E$ . It is the active conformation.

During ATP synthesis, the direction of c-ring cylinder movement is right-handed and this rotation is generated by a p.m.f. at a speed of 100-150 rev/s. During ATP hydrolysis,

however, the rotation direction reverses. This rotation is produced by the interaction with protein a, and the 360° rotation prompts the synthesis of the 3 ATP molecules.

#### II.4 F<sub>1</sub>F<sub>0</sub>-ATPase regulation

F<sub>1</sub>F<sub>0</sub>-ATPase activity depends on the energy available in cells. When the proton gradient is low, the β-subunit binds MgADP by assuming a conformation that render the complex inactive. Other molecules inhibit ATP synthesis, such as ATPase inhibitory factor 1 (IF<sub>1</sub>). At a pH of 6.7, it forms a complex with ATPase that both inhibits ATP synthesis and prevents ATP hydrolysis. IF<sub>1</sub> is also regulated by post-translational modifications, specifically by phosphorylation, which controls the ATPase function in different contexts depending on substrate availability or the energy demanded by the cell (García-Bermúdez, Cuezva and Bernardi, 2016; Esparza-Moltó, Nuevo-Tapióles and Cuezva, 2017).

#### II.5 Role of ATP synthase in apoptosis

The apoptosis process occurs when cells die in a programmed manner. It is an energy-consuming process that takes place in events like cell turnover, embryo development or cell death caused by toxic agents (Elmore, 2007). During apoptosis, the permeabilisation of the IMM occurs and apoptotic factors are released from the mitochondria to the cytoplasm. The structure that would lead to the release of these factors is known as the mitochondrial permeability transition pore (mPTP) (Rasola and Bernardi, 2007; Kwong and Molkenin, 2015). The mPTP is a structure located in the IMM and is permeable to solutes and ions and, specifically, the release of calcium and cytochrome c triggers the apoptosis process (Eskes *et al.*, 1998).

The “permeability transition” concept was proposed for the first time by Haworth and Hunter (Hunter and Haworth, 1979), but its physiological role and composition have been debated for some time. Initially it was suggested that the adenine nucleotide translocator (ANT) is involved in mPTP formation because of the inhibition of the mPTP by adenine nucleotides. The use of genetic tools to knockout ANT has demonstrated that this hypothesis is incorrect, and that ANT is not essential for mPTP formation. A similar situation occurs with the Vdac complex; nor does it participate in mPTP regulation (Halestrap, 2009).

It is believed that ATP synthase is required for mPTP formation (Giorgio *et al.*, 2013) and ATPase dimers have been proposed as the mPTP. The purification of these dimers retains mPTP activity (Giorgio *et al.*, 2013). Moreover, their role in calcium release in *Drosophila* (Von Stockum *et al.*, 2015) has been demonstrated and suggests that the process is not exclusively mammalian (Azzolin *et al.*, 2010). Alternatively, it has been proposed other

mPTP identity independently of ATPase dimerisation. In this hypothesis, it is described the c-ring subunit of ATPase as the mPTP (Alavian *et al.*, 2014).

Currently, the composition of the mPTP remains unclear. Different studies have raised doubts about the proposed models. The idea that c-ring forms the mPTP has been ruled out using atom molecular dynamic simulations (Zhou *et al.*, 2017). More recently, the belief that the peripheral stalk and c-subunits are required to retain mPTP activity has been excluded (He, Carroll, *et al.*, 2017; He, Ford, *et al.*, 2017).

## II.6 New peptides associated with ATPase

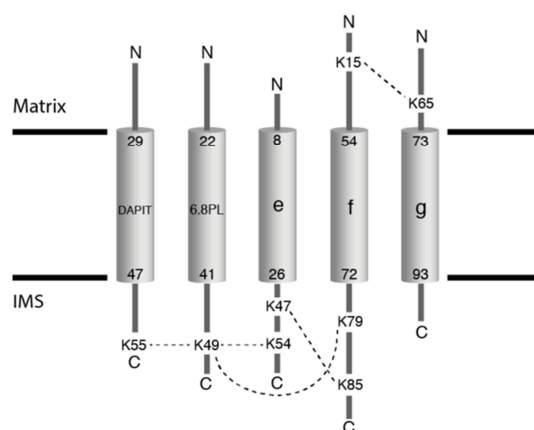
DAPIT and MP68 have remained undetected for a long time. Experimental conditions were important to detect them. When exogenous phospholipids are absent, the small proteins are lost during chromatography, but if the preparation includes some phospholipids, DAPIT and MP68 are found in the purified ATPase complex. Peptides DAPIT and MP68 have been identified with sequences AGPEADAQFHFTGIKKYFN and MLQSLIKKVVWIPMKPYTQAYQEI, respectively (Fig. 6). Both have been detected in mass spectrometry experiments (Rexroth *et al.*, 2004; Chen *et al.*, 2007), in the monomeric and dimeric forms of the bovine ATPase complex (Meyer *et al.*, 2007; Nůsková *et al.*, 2015), and as individual proteins. The same study found no differences in the hydrolysis capacity of ATPase when MP68 was dissociated. Both share a common transmembrane region (Fig. 7) and several phosphorylation sites which have not been confirmed by a proteomic analysis (Carroll, Fearnley and Walker, 2006).



**Figure 7. Alignment and structural analysis of proteins MLQ and AGP.** The transmembrane region is marked in red, phosphorylation sites are marked in blue, and green and yellow denote the unmarked amino acids, which are identical, similar and non-similar. **A.** Alignment of the DAPIT sequence from *R. norvegicus* (R.n.), *Bos taurus* (B.t.) (gi91207977), *Homo sapiens* (H.s.) (gi 14249376), and *S. bullata* (S.b.) (gi23505740). **B.** Alignment of the MP68 sequence from *R. norvegicus* (R.n.), *Bos taurus* (B.t.) (gi91207977) and *S. bullata* (S.b.) (gi23505740) (modified from (Meyer *et al.*, 2007).

These two peptides were initially attributed to peripheral functions, but then more important roles were ascribed to them when they appeared in a screening of regulators of oxidative phosphorylation (Baughman *et al.*, 2009). In the screening, for which shRNAs were used, more than 85% of mRNA levels were depleted and the oxygen consumption rate (OCR) was measured. No alterations in the OCR were observed in either the shRNA of the *Dapit* and *Mp68* cells, or in the expression levels of other proteins of the mETC, including NDUFB8, SDHB or ATP5a. Although DAPIT and MP68 were proposed as the principal candidates to interact with mETC genes, changes in the oxidative phenotype in these mutants was not observed.

The interactions between the subunits of the ATPase complex, more specifically those involving DAPIT and MP68 have been analysed in covalent cross-linking assays. DAPIT and MP68 protein, as well as other peptides including ATP8, e, f, or g, have been described as supernumerary subunits. Nonetheless, this has not been confirmed by gel staining intensity where all the ATPase subunits have been resolved (Lee *et al.*, 2015). Lysine K49, in the carboxyl terminal of MP68, is linked to the same region of DAPIT K55, and also to different lysines of supernumerary subunits e, f or g (Fig. 8).

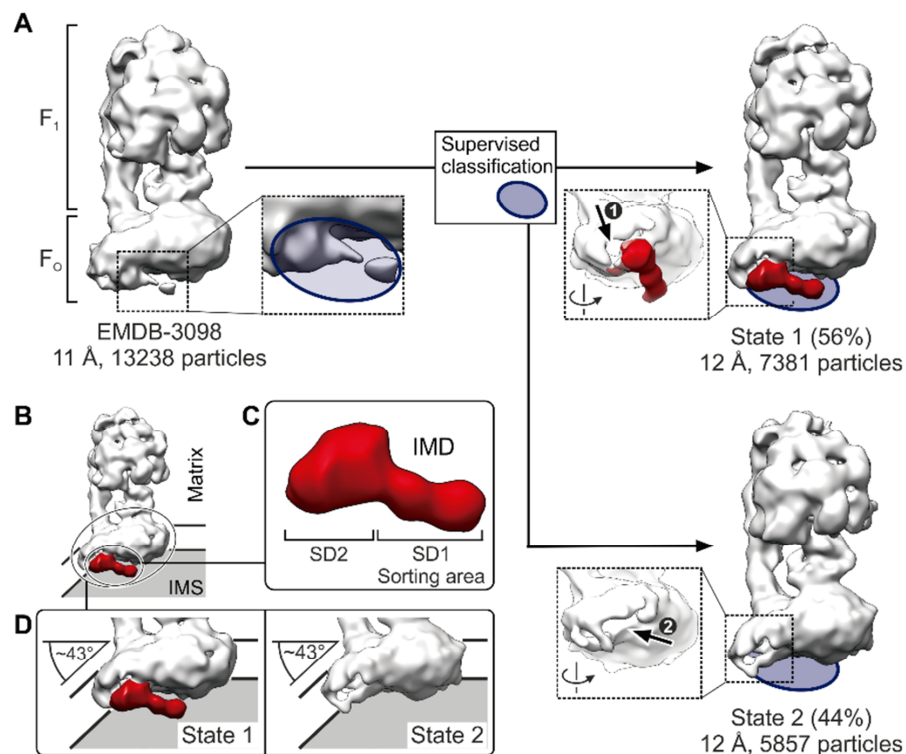


**Figure 8. The cross-links detected between some ATPase subunits.** The residues that cross-link between the supernumerary subunits of ATPase are shown in the figure (from (Lee *et al.*, 2015)).

Recently, the three-dimensional structure of a new ATPase domain has been observed in cryo-electron microscopy (cryo-EM) images and by a computational analysis (Fischer *et al.*, 2017). It is named the intermembrane space domain (IMD), and is described as a new domain associated with the traditional  $F_0$  ATPase complex, but is clearly separated.

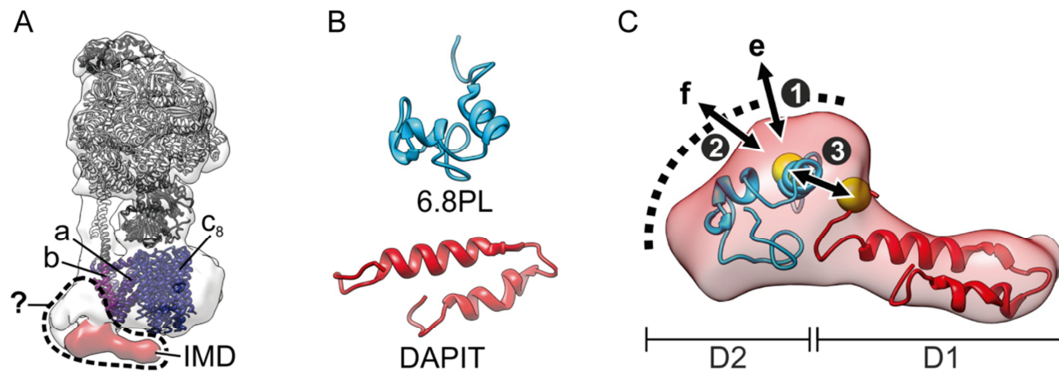
The presence or absence of the IMD determines state 1 or state 2 of the ATPase structure, respectively (Fig. 9A). The IMD was identified in individual particles associated with ATPase that come into contact with the rotor region (Fig. 9B), and its structure is divided

into a tubular domain, SD1, and a globular domain, SD2 (Fig. 9C). In the particle where the IMD is absent, the transmembrane region remains unaffected, which indicates that the IMD does not play a role in membrane bending (Fig. 9D).



**Figure 9. Description of FoF1 ATP synthase with and without an intermembrane space domain (IMD).** **A.** After a computational analysis of the cryo-EM images, an IMD was described to be slightly associated with bovine ATPase. Two groups of structures are separated, state 1 where an IMD is present, and state 2 where an IMD is absent. **B.** The IMD is located in the inner mitochondrial space (IMS). **C.** The IMD structure with two domains: SD1 (tubular) and SD2 (globular). **D.** The IMD is not essential for membrane bending. From (Fischer *et al.*, 2017).

The absence of DAPIT and MP68 in stage 2 led to the conclusion that the IMD is formed by them, supported by the fact that strong interactions with other ATPase components were lacking, and also by their absence in preparations with small amounts of phospholipids. Furthermore, different structural models were proposed using the IMD structure and the previous cross-linking analysis, which concluded that these two proteins are the only candidates that occupy the IMD space (Fig. 10A-C).

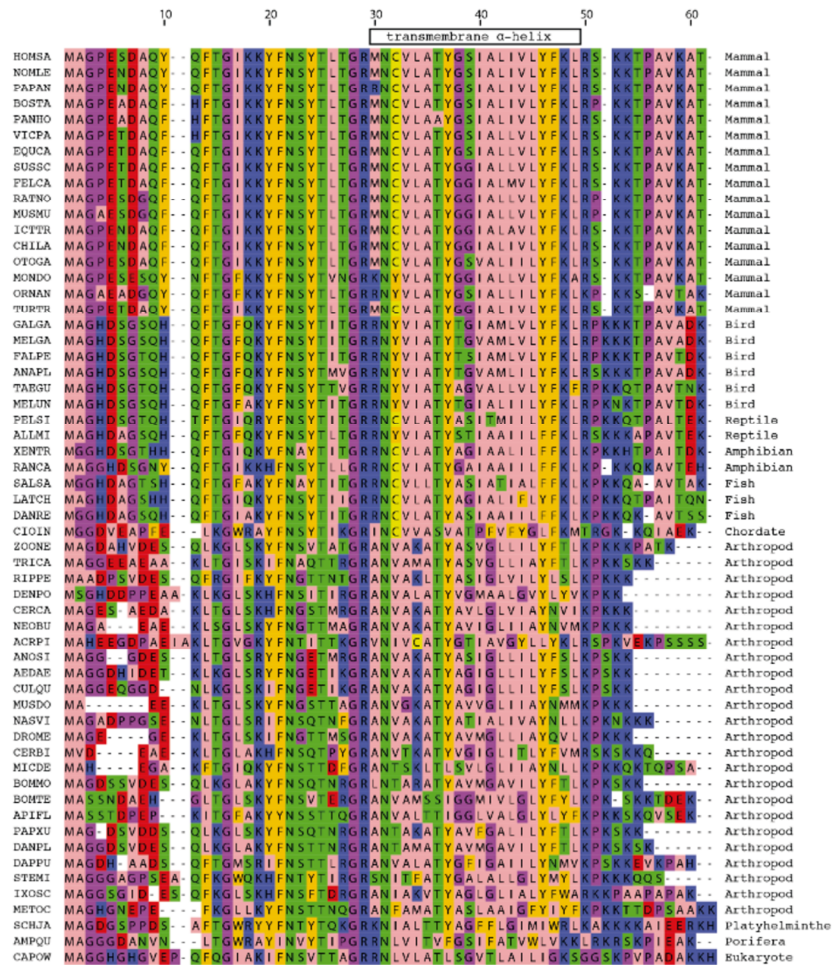


**Figure 10. Structural composition of the mammalian IMD.** **A.** Representation of state 1 of the IMD. The  $F_1$  domain is depicted in grey, the c-ring rotor of ATPase is shown in blue, subunit a in purple and subunit b in magenta. **B.** Model of subunits DAPIT MP68 obtained by iTasser. **C.** Modelling DAPIT and MP68 in the IMD. Yellow spheres indicate the residues involved in cross-linking. From (Fischer *et al.*, 2017).

Similar IMD structures were found in other ATP synthases from yeast or ciliates, but their shape differed. The common aspect of these three cases was that the IMD always came into contact with the rotor region, which indicates its importance in regulating its function or its structure. The interaction between the IMD and the c subunits could affect the function of ATPase; in this scenario, the IMD would act as an inhibitor of  $F_0$  (similarly to IF1 in  $F_1$ ). Nonetheless, the function performed by the IMD did not come over very clearly, and one suggestion is that it may be involved in the regulation of the mPTP opening monitored by either a lipid plug or dimer formation.

### DAPIT (AGP)

DAPIT (diabetes-associated protein in insulin-sensitive tissue) is a basic hydrophobic protein that is 58 amino acids long with a single transmembrane helix. Its C-terminal domain is located in the intermembrane space (IMS) (Lee *et al.*, 2015) (Fig. 11).



**Figure 11. Conservation of the DAPIT protein sequence in metazoans.** The sequence of DAPIT proteins of several organisms appears with this colour code: pink, hydrophobic; orange, aromatic; blue, basic; red, acidic; green, hydrophilic; magenta, proline and glycine. Yellow denotes cysteine residues. From (Lee *et al.*, 2015).

This peptide has been described in a streptozotocin-induced diabetes rat model (Päivärinne and Kainulainen, 2001). Streptozotocin (STZ) is an antibiotic, an analogue of glucose molecules that induces diabetes type I. STZ enters the beta cells of the pancreas by specific transporter GLUT2, which is enriched in this cell type. Therefore in this model, a high DAPIT mRNA expression was detected in tissues such as the brain, skeletal or cardiac muscle, but levels were lower in other tissues like the lung, liver or kidney. Following STZ treatment, Dapit mRNA decreased in the diabetic rat compared to the control ones. For this reason, the detected ORF was called DAPIT, diabetes-associated protein in insulin tissues. The DAPIT protein has been highly conserved during evolution. The human protein has a 58-98% similarity to proteins *Fugu* and *Xenopus*, and shares 40% identity with *Drosophila melanogaster* homologous protein Neb-cGP. In all these species, it is noteworthy that 13 amino acids are conserved, which indicates their potential importance in the DAPIT structure or function (Päivärinne and Kainulainen, 2001).

Given its general expression pattern, it was initially considered to perform a general function. In particular, insect homologue Neb-cGP has been described as a stimulator of cAMP. According to these data, DAPIT may be involved in this process under diabetic conditions (Päivärinne and Kainulainen, 2001).

HeLa cells with low DAPIT levels, by shRNAs, show loss of the ATPase population in mitochondria at the protein level, but not mRNA (Ohsakaya *et al.*, 2011). They also present impaired ATP synthesis and cell growth. In these cells, these authors did not observe any disassembly of ATPase or supercomplexes.

With the human cells that overexpress the DAPIT protein, glucose metabolism modulation has been observed (Kontro *et al.*, 2015). In these cells, the mtDNA content and mitochondrial mass lower, while the mitochondrial membrane potential increases. A tendency towards glycolytic metabolism has been detected, which promotes the activation of HIF1alpha in Wnt/beta catenin signalling.

### **6.8kDa protein (MP68)**

This protein has been described simultaneously to the DAPIT protein (Terzi *et al.*, 1990; Chen *et al.*, 2007; Meyer *et al.*, 2007). Although the interference of the *Mp68* gene in cells reduces both the ATPase population and ATP production (Fujikawa *et al.*, 2014), the mRNA levels of the main ATPase subunits remain unaffected.

### III. METABOLIC CONTROL BY OPTOGENETICS

#### III.1 Optogenetics systems

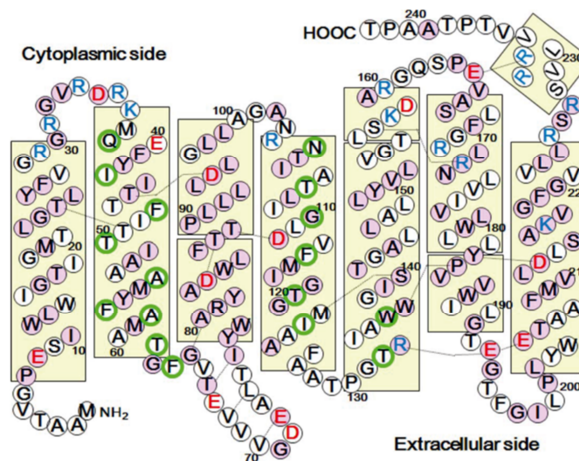
Optogenetic tools were traditionally used to modulate signalling in cells, whereby light induces conformational changes to proteins by triggering a cellular response. Initially, light-sensitive channels were developed to modulate the polarity of neurons to thus initiate a signalling pathway (Zemelman *et al.*, 2002). Beyond the neuroscience field, optogenetic applications are broader, and optogenetics is also used to manipulate gene expression (Yazawa *et al.*, 2009; Kennedy *et al.*, 2010) and to modulate the behavior of individual cells for example, by modifying cell polarity. The commonest method to do this is by modulating protein dimerisation with light (Levskaia *et al.*, 2009; Guglielmi, Falk and De Renzi, 2016). By way of example, changing the conformational status of proteins to produce an active protein is used to photoactivate Rac and to control cell motility (Wu *et al.*, 2009).

#### III.2 Optogenetics in metabolism

These tools have also been used in the metabolism field to modulate gene expression and to modify and study metabolism (Idevall-Hagren *et al.*, 2012; Inglés-Prieto *et al.*, 2015). All these examples use light as a trigger to induce conformational change. However, the possibility of using optogenetic systems to convert light energy into energy that can be used by cells to grow has been explored only minimally.

#### III.3 Deltarhodopsin

The first described microbial rhodopsin was in *Halobacterium salinarum* (Grote and O'Malley, 2011), but later others were found in several species. They all share a structure determined by seven transmembrane helices and they perform different functions as light-driven protons, ions or photosensors (Lanyi, 1998) (Fig. 12).



**Figure 12. Diagram of the rhodopsin structure of *Haloterrigena thermotolerans*.** Transmembrane helices are represented in rectangles, the amino acids with a positive and negative charge are depicted in blue and red, respectively. The green circles include the amino acids that interact between helices, and the pink circles show the most conserved residues between rhodopsins (from (Zhang *et al.*, 2013))

More specifically, Deltarhodopsin (HtdR) is a light-driven proton ( $H^+$ ) pump present in the membrane of *Haloterrigena turkmenica* (Kamo *et al.*, 2006) and others species. Its structure in different organisms is similar.

The functionality of HtdR has been demonstrated in artificial vesicles, which expresses HtdR together with  $F_0F_1$ -ATP synthase. HtdR translocates  $H^+$  into vesicles and ATP synthase can use the  $H^+$  gradient to generate ATP from ADP in the extra-vesicle medium. This protein has also been used in cells to partially rescue membrane damage following neurotoxin 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP) treatment (Hara *et al.*, 2013).

#### III.4 Detaching ATP synthesis from the mitochondrial electron transport chain by optogenetics

Deltarhodopsin expression in cells is a way to separate ATP synthesis from electron transport in the mETC. Previously, this effect was achieved by the expression of other proteins, such as Ndi1 from *Saccharomyces cerevisiae* and Aox from fungi (Perales-Clemente *et al.*, 2008). Ndi1 is a NADH dehydrogenase with NADH DH/CoQ reductase activity that can substitute the function of complex I in yeast, but without  $H^+$  pumping. Aox is an alternative oxidase present in plants and fungi that is able to oxidase the CoQ. With the expression of both these proteins in mitochondrial-deficient cells, it is possible to restore the capacity of these mutant cells so they can grow in the absence of pyruvate and uridine because they are able to restore the electron flux. Although these two proteins can

restore the electron flux, they cannot restore proton pumping, which means that these cells are unable to grow in galactose. Expressing deltarhodopsin in these cells would allow the ETC function to be fully restored.





# Objectives



The role attributed to ATP synthase as the main energy producer of the cell has been maintained for years. Nonetheless, new functions have been defined, such as a regulator, apoptosis or as a determinant of mitochondrial cristae morphology. For this reason, the structure of this complex has been studied in detail, which led to the discovery of new subunits associated with it. We are interested in characterising these new subunits and their role in the ATPase function.

Optogenetics is a technique that has been traditionally used in the neural field to modulate cell behavior. We aim to extend its applications to the metabolism field.

Taken together, the specific objectives of this doctoral thesis are the:

1. Characterisation of proteins DAPIT and MP68 at the structural and functional levels.
2. Modulation of survival in the absence of the functional mETC.



# Methods



## I. CELL CULTURE

### III.1 Mammalian cells

The following cells were used in this work (Table 1)

**Table 1. The cells used in this work.**

Line	nDNA	mtDNA	Mutation	ETC	Origin	Reference
<b>L929<sup>mtC57</sup></b>	L929	C57BL/6	-	All complexes are stable	Cybrids generated by transferring the platelet of cells C57BL/6 to $\rho^{0929}$	Moreno-Loshuertos <i>et al.</i> 2006
<b>ND4<sup>KO</sup></b>	L929	3T3	Mt-nd4 10227delA (homoplasmic)	Complex I absent	Random mutation in 3T3 cells and transfer to L929 cells	Perales-Clemente <i>et al.</i> 2010
<b>Cox10<sup>KO</sup></b>	129/SvJ	C57BL/6	Cox10 knockout (homozygosis)	Complexes I and IV absent	Cre protein expression in the Cox10 <sup>fllox/fllox</sup> fibroblasts	Diaz <i>et al.</i> 2006
<b>F3T3</b>	L929	3T3	-	All complexes are stable	Cybrids generated by the fusion of mitochondria from 3T3 cells with $\rho^{0929}$ cells	R. Moreno

### III.2 Culture medium

The culture medium used to grow the cell lines was DMEM medium that contained L-glutamine and glucose at a concentration of 25 mM. The following were also added: 5% of FBS, 1mM sodium pyruvate, 100 units/mL penicillin and 100 ug/mL streptomycin. For the culture of some mutant cells with deficiencies in the ETC, uridine was added at a concentration of 50 ug/mL.

The medium used in the growth curve (Section I.5) was modified to ensure that other components did not interfere with growth, only with the carbon source of glucose or galactose. This prepared medium consisted of DMEM (Sigma D5030) in water supplemented with 2 mM L-glutamine, 1 mM sodium pyruvate, 5% dialysed FBS, 100 units/mL penicillin, 100 ug/mL streptomycin and phenol red. Then depending on the experiment, the medium was supplemented with 5 mM or 25 mM of glucose or galactose.

### III.3 Culture conditions

Cells were grown at 37°C at a constant level of 5% CO<sub>2</sub>. The medium was frequently changed every 48 h, except in the growth curve experiment where the medium was changed daily.

### III.4 Ectopic gene expression in mammalian cells

To get a stable expression of an exogenous protein a lentiviral infection was used.

To obtain viral particles, 293T cells were used and transfected with transfection reagent X-treme-GENE-HO (Roche) following the manufacturer's instructions. Firstly cells were placed in a medium that contained 25 µM chloroquine to avoid degrading exogenous DNA. Next 7 µg of the DNA that coded the viral capsid (the psPAX2 vector), 3 µg of the DNA that coded the viral envelope (pMD2G) and 10 µg of plasmid pWXLD with the sequence of interest were added to the FBS-free medium (Table 2).

**Table 2 . Plasmids used to express exogenous proteins in mammalian cells.**

Plasmid	Bacterial resistance	Eukaryotes promotor	Eukaryotes selection	Origin	Use
<b>pWXLD</b>	ampicillin	EF1 alpha	puromycin	TronoLab (with some modifications)	Lentiviral vector to expression in eukaryotes
<b>psPAX2</b>	ampicillin	CMVeh	-	TronoLab	Lentiviral packaging vector (2nd generation)
<b>pMD2G</b>	ampicillin	CMV	-	TonoLab	Lentiviral envelop vector

The produced viral particles were released to the medium and were collected 48 h and 72 h post-transfection. The medium was filtered with a 0.45  $\mu$ M filter to eliminate cell debris before storing it at  $-80^{\circ}\text{C}$ . A small portion of it was kept to quantify the number of viral particles by qPCR (Scherr *et al.*, 2001). After determining the concentration, a particular MOI (multiplicity of infection) was selected (1, 5 or 10) as well as the infected cell line. Cells were incubated with viral particles for 6 h and were then cultured in normal medium. The antibiotic was added to the medium to select the infected cells after 48 h, which allowed enough time for the selection gene to be expressed, specifically the puromycin resistance gene.

### III.5 Growth curve

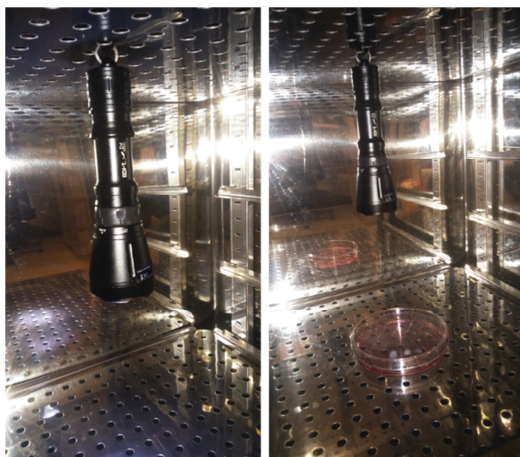
To determine the ability of the different cell types to grow, a growth curve in the presence of glucose or galactose as a carbon source was performed. The cells with mutations in the ETC protein cannot survive in galactose medium because its conversion into energy is very slow, and glycolysis does not suffice to produce all the ATP required for cells to grow.

Cells were cultured in p96 plates at a density of around 2000 cells/well (depending on cell type). Cell growth was monitored daily by the CyQUANT™ NF Cell Proliferation Assay Kit (Thermo Fisher C35006) to measure DNA content. The number of plates depended on the experiment time. Typically, 5 days sufficed to evaluate the growth changes in different substrates.

In some experiments, different inhibitors, such as Antimycin A (inhibit complex III), were used to examine cell growth in its presence.

### III.6 Cell growth in light conditions

To HtdR-expressing cell been an active protein which translocates protons, it is needed a source of light. Here, a standard cell incubation was transformed in an illuminated incubator thanks to the position of torch into it (Fig. 13). The torch used contains several LEDs that provides a whit light without warm up the place.



**Figure 13. Photography of the illuminated incubator assembly. The torch was ubicated at the top of the incubator and the light illuminates the plates.**

### III.7 Immunofluorescence

Immunofluorescence was completed to detect the presence of some proteins. Cells were grown in p12 plates on glass covers. Firstly, they were fixed with 4% paraformaldehyde in culture medium for 15 minutes at 37°C. Next they were washed several times with PBS. To permeabilise cells, covers were treated with PBS with triton X-100 for 5 min in the same plate. A blocking step was used by adding PBS with triton X-100 and 2% BSA for 10 min. Finally, the primary antibody was added to the blocking solution to detect the protein of interest (Table 3).

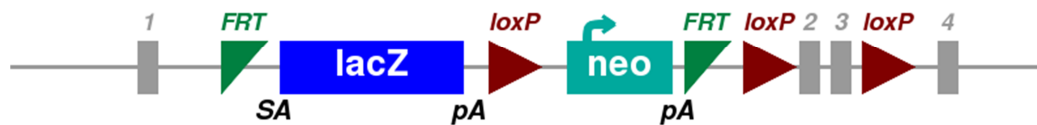
**Table 3. Antibodies used for immunofluorescence**

Detected antigen	Species	Catalogue number	Company
HA epitope	Rat IgG1	11867423001	Sigma
Tom20	Rabbit IgG	sc-11415	Santa Cruz
DAPIT	Rabbit IgG	17716-1-AP	Proteintech
Ndufa9	Mouse IgG	ab14713	Abcam

## IV. ANIMAL MODELS

### IV.1 Mouse strains

Two mouse strains were generated during this thesis. The first was a knockout for the DAPIT protein. The frozen sperm of a *Dapit* transgenic mouse C57BL/6N-*Usmg5*<sup>tm1a(EUCOMM)Wtsi/H</sup> was obtained from the European Mouse Mutant Archive (EMMA) with ID EM:07834 and the following construct (Fig. 14).



**Figure 14. Scheme of the construction of transgenic mice.** The *lacZ*- and neomycin-resistant genes lie between the FRT sites, and the main exons, 2 and 3, lie between the *loxP* sites (From EMMA).

This is a very versatile construct. Firstly, *Dapit* expression could be followed because of *lacZ* expression and its subsequent detection via X-gal staining. Secondly, since it is a conditional transgenic mouse, the DAPIT protein could be specifically deleted in different tissues depending on the particular cre line used.

*In vitro* fertilisation (IVF) of the C57BL/6 female mice was performed at the Transgenesis Unit of the CNIC. Descendants were genotyped to select the transgenic line of interest. Transgenic mice were then crossed with a flippase mouse line to thus eliminate the *neo* and *lacZ* cassettes found between the FRT sites. Lastly, in order to achieve full knockout, mice were crossed with the *Sox2:cre* mouse line (Hayashi, Tenzen and McMahon, 2003), which expresses cre recombinase under the control of the *Sox2* promoter during early development.

The second mouse strain generated was a knock-out of *Mp68*. In this case, embryonic stem (ES) cells were obtained from the European Mouse Mutant Cell Repository (EuMMCR). The transgenic ES for the 2010107E04Rik gene (*Mp68*) was performed in the JM8A1.N3 cell line. The basis of the structure of the used construct was the same as the transgenic *Dapit* one, the *lacZ* cassette, and the codifying exons placed between the *loxP* sites.

## IV.2 PCR for genotyping

To genotype mice, a piece of tail was cut from mice during weaning, and DNA was extracted using the REExtract-N-Amp™ Tissue PCR Kit following the datasheet indications. Taq DNA polymerase was used to genotype.

The primers used to genotype the different mouse lines are specified (Tables 4 and 5).

**Table 4. Primers to genotype the *Dap1t* transgenic mice**

To genotype the DAPIT mice	Sequence
Wild-type allele	TGCCAGTTCTTGGCTTACT
	AAGGCCAGCTAAAGTTGGAG
Transgenic allele	TGCCAGTTCTTGGCTTACT
	GAACCTCGGAATAGGAACTTCG
Floxed allele	AAGGCGCATAACGATACCAC
	CCGCCTACTGCGACTATAGAGA
Knockout allele	AAGGCGCATAACGATACCAC
	TGGGAAAGCACTTAACCCCA

**Table 5. Primers to genotype the *Mp68* transgenic mice**

To genotype the <i>Mp68</i> mice	Sequence
Wild-type allele	GGAATGTGGGTATCTAGTTTGGT
	AATGAAGACGGAGGCCTGTT
Transgenic allele	GGAATGTGGGTATCTAGTTTGGT
	ACTTCGGAATAGGAACTTCGGT
Floxed allele	AAGGCGCATAACGATACCAC
	CCGCCTACTGCGACTATAGAGA
Knockout allele	AAGGCGCATAACGATACCAC
	TAGACAGACAGCCAGTCCCG

## V. PROTEIN RESOLVING AND ANALYSIS

### V.1 Protein extraction and quantification

Proteins were extracted from cells with RIPA buffer (50 mM TrisHCl pH7.4, 50 mM NaCl, 1% Triton x-100, 0.5% sodium deoxycholate, 5 mM EDTA, and a protease inhibitor cocktail (11697498001 Sigma). Cells from a p100 plate were pelleted and washed with cold PBS twice. The pellet was then resuspended in 200  $\mu$ L of RIPA buffer, incubated at 4°C for 15 min and placed on a rotating wheel. Lastly, cells were centrifuged at 15,000g for 15 min at 4°C and the supernatant was collected.

The Bradford method was used to quantify the protein concentration in the samples (Bio-Rad 5000006) (Bradford, 1976). Next 1  $\mu$ L of sample was added to each well, which

already contained 200  $\mu\text{L}$  of water, and were quantified by triplicate readings. A standard protein curve using 0, 1, 2, 3, 4, 5, 10 and 15  $\mu\text{L}$  of 1 mg/mL stock of BSA was used to measure the absorbance of the samples. Finally, 50  $\mu\text{L}$  of Bradford reagent were added, mixed well and measured in a spectrophotometer at 595 nm.

## V.2 SDS-PAGE electrophoresis

Standard denaturing (SDS-PAGE) conditions were used to resolve proteins. A 30% percentage of acrylamide/bis-acrylamide (29:1 ratio) (A/BA) was used and they were run in gel with stacking and resolving parts (Table 6).

**Table 6. SDS-PAGE gel composition**

	Resolving 12.5%	Resolving 7%	Stacking
<b>Tris HCl 1M pH 8.8</b>	1.87 mL	1.87 mL	-
<b>Tris HCl 1M pH 6.8</b>	-	-	0.25 mL
<b>A/AB 30%</b>	2.08 mL	1.16 mL	0.33 mL
<b>Distilled water</b>	1.05 mL	1.94 mL	
<b>SDS 20%</b>	25 $\mu\text{L}$	25 $\mu\text{L}$	10 $\mu\text{L}$
<b>APS 20%</b>	15 $\mu\text{L}$	15 $\mu\text{L}$	10 $\mu\text{L}$
<b>TEMED</b>	5 $\mu\text{L}$	5 $\mu\text{L}$	5 $\mu\text{L}$

Samples were run in Tris-glycine buffer (Tris 25 mM, glycine 20 mM and SDS 0.5%) at 10 mA voltage during stacking, and voltage was raised to 20 mA once the samples reached the running gel (if the number of gels was more than 1, amperage was increased). This was done with a Mini-PROTEAN Tetra Cell (BioRad).

## V.3 Protein transference to a PVDF membrane and protein detection

After resolving proteins on gel, they were transferred to a PVDF membrane (Immobilon-FL, 0.45  $\mu\text{M}$ ) and were then transferred and run in transfer buffer (48 mM Tris, 39 mM glycine and 20% methanol) for 1 h and 15 min at 100 volts (V) by the Mini Trans-Blot Cell system (BioRad). The membrane was blocked to avoid non-specific unions of antibodies with 5% BSA in PBS or with Sea Blocking buffer (Thermo Scientific 37527) for 1 h at room temperature (RT) or o/n at 4°C.

To detect specific proteins, the primary antibody was diluted in blocking solution and incubated with the membrane for 2 h at RT (Table 7). Next the membrane was washed 3 times with PBS that contained 0.1% Tween-20 for 5 min. The secondary antibody diluted in blocking solution was also added and incubated for 1 h at RT (Table 8). Afterwards the membrane was washed once more (the last wash was done only with PBS).

To develop the signal, the Odyssey® Imaging System (from LI-COR Company) was used. By this technology, two longitudinal waves were detected at the same time, which allowed the simultaneous detection of two proteins of a similar size.

**Table 7. Primary antibodies used**

Detected antigen	Species	Code	Company
<b>ACTIN</b>	Rabbit IgG	A2066	Sigma
<b>COI</b>	Mouse IgG2	459600	Invitrogen
<b>DAPIT</b>	Rabbit	17716-1-AP	Proteintech
<b>NDUFA9</b>	Mouse IgG	ab14713	Abcam
<b>OPA1</b>	Mouse IgG	612606	BD Bioscience
<b>TOM20</b>	Rabbit IgG	sc-11415	Santa Cruz
<b>UQCRC1</b>	Mouse IgG1	Ab110252	Abcam
<b>UQCRC2</b>	Rabbit	17716-1-AP	Proteintech

**Table 8. Secondary antibodies used**

Detected antigen	Conjugation	Code	Company
<b>Mouse IgG (H+L)</b>	DyLight™ 800	610-145-121	Rockland
<b>Mouse IgG (H+L)</b>	Alexa Fluor 680	A21057	Life Technologies
<b>Rabbit IgG (H+L)</b>	DyLight™ 800	610-145-122	Rockland
<b>Rabbit IgG (H+L)</b>	Alexa Fluor 680	A210766	Life Technologies
<b>Rat IgG(H+L)</b>	IRDye800	926-32219	LI-COR
<b>Rat IgG(H+L)</b>	Alexa Fluor 680	A21096	Life Technologies

## VI. ANALYSIS OF MITOCHONDRIAL COMPLEXES AND SUPERCOMPLEXES

### VI.1 Mitochondria extraction from cells and tissues

Mitochondria isolation was done following the protocol of J.A. Enríquez's laboratory (Fernández-Vizarra, López-Pérez and Enriquez, 2002).

#### **Mitochondria from cells**

To isolate mitochondria from cells, 10 plates of 10 mm were required. Cells were trypsinised and then pelleted. The pellet was washed with PBS prior to extraction. At this point, the pellet can be stored at -80°C or directly extracted. In the next step, hypotonic medium (sucrose 83 mM, MOPS 10 mM, pH 7.2) was added to resuspend the pellet (seven volumes of the pellet volume). After 2 min on ice, the pellet was placed into a glass tissue potter and homogenised in a stirrer (Heidolph RZR 2041). Afterwards, the same volume was added to the hypertonic medium (250 mM sucrose, 30 mM MOPS pH 7.2). Then it was centrifuged at 1,000g for 5 min at 4°C. The supernatant was collected in several 1.5mL phials and centrifuged at 13,000g for 5 min at 4°C. Next pellets were collected and washed

with medium A (320 mM sucrose, 10 mM Tris, 1mM EDTA, pH 7.4) several times and placed in a single phial. Finally, samples were quantified by the Bradford method (see Section III.1) and stored at -80°C in 50 mM Bis-Tris, 1M aminocaproic acid at a concentration of 10 ug/uL.

### Mitochondria from tissues

Instead of using a hypo/hypertonic medium, the mitochondria from tissues were homogenised directly in medium A, except when extracting mitochondria from the heart, when fatty acid-free BSA was added.

#### VI.2 Mitochondria samples preparation

Fresh or defrosted mitochondria were lysed with digitonin, a soft detergent at a detergent:protein ratio of 4:1 (g/g). Each 10- $\mu$ L (100  $\mu$ g of protein) aliquot was treated with 4  $\mu$ L of 10% digitonin. The mix was incubated for 5 min on ice and centrifuged at 13,000g for 30 minutes at 4°C. The supernatant was then collected and applied to 3  $\mu$ L of loading buffer (5% Coomassie Blue G-250 in 1M aminocaproic acid).

#### VI.3 Blue-Native PAGE gel

Blue-Native Gel (BN) electrophoresis should be carried out to visualise mitochondrial complexes (Schägger, 1995; Wittig, Braun and Schägger, 2006). A gradient gel is required to resolve high-molecular-weight complexes (**Table 9**). Using the gradient, resolving gel was prepared in 1.5 mm-thick glass plates with the gradient former connected to a peristaltic pump.

**Table 9. BN gel composition**

Resolving gel component	Acrylamide solution, 3%	Acrylamide solution, 13%
<b>A/BA 48:1.5</b>	0.305 mL	0.866 mL
<b>Buffer 3x *</b>	1.667 mL	1.111 mL
<b>Distilled water</b>	3 mL	0.713 mL
<b>Glycerol 85%</b>	-	0.643 mL
<b>APS 20%</b>	20 $\mu$ L	7 $\mu$ L
<b>TEMED</b>	4 $\mu$ L	3 $\mu$ L

\* Buffer 3x: Bis-Tris 150 mM, aminocaproic acid 1.5 M, pH 7

See Section III.2 for the stacking gel composition

## VI.4 BN-PAGE electrophoresis

Mitochondria samples, already digitonised and containing the loading buffer, were loaded onto gels and run in a Mini-PROTEAN Tetra Cell apparatus (BioRad). Different buffers were used to run cathode A (50 mM tricine, 7.5 mM imidazole, 0.02% Coomassie blue G-250, pH 7), cathode B (50 mM tricine, 7.5 mM imidazole, pH 7), and anode (25 mM imidazole, pH 7). The first run was at 150V for 40 min using cathode A, with an extra 1 h at 300V with cathode B (the anode buffer was the same throughout the process).

To transfer the mitochondria complexes to a membrane, the same protein transfer procedure was followed (see Section III.3).

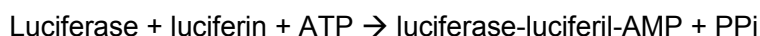
## VII. MITOCHONDRIAL ANALYSIS

### VII.1 Subcellular fractionation

Mitochondria isolation was done as previously described (see Section Methods IV.1). Fresh mitochondria were treated with trypsin to digest the proteins in the OMM, as previously described (Perales-Clemente *et al.*, 2011).

### VII.2 ATP synthesis

An ATP synthesis measurement was taken in fresh mitochondria. ATP was detected by the luciferin/luciferase fluorimetric method that follows this reaction:



The produced light was proportional to the amount of ATP, with maximal emission in 560 nm (Vives-Bauza, Yang and Manfredi, 2007).

Using flat-bottomed white plates (96-well Costar Assay Plates), 160  $\mu\text{L}$  of buffer A (50 mM KCl, 25 mM Tris-HCl, 2 mM EDTA, 0.1% BSA, 10 mM  $\text{KPO}_4$ , 0.1 mM  $\text{MgCl}_2$ , pH 7.4) were added. Next a standard ATP curve was done to relate the amount of light and the ATP concentration by adding 10  $\mu\text{L}$  at varying ATP concentrations (from 0 to 10 mM). To calculate ATP production in the experimental samples, 50  $\mu\text{g}$  of mitochondria were added (in the 160  $\mu\text{L}$  of buffer A) to each well. Later substrates were added (a 5  $\mu\text{L}$  mix of 6 mM DP5 + 5  $\mu\text{L}$  pyruvate/glutamate/malate, depending on the substrate to be evaluated, + 5  $\mu\text{L}$  ADP). Three replicates were needed for each sample measurement. Finally, 20  $\mu\text{L}$  were

added (0.5 M Tris-acetate, 0.4 mM luciferin, 10 ug/mL luciferase, pH 7.75). After mixing well, the plate was measured in a luminometer (Tubo Sirius, Berthold) for 5 min every 20 sec.

During the analysis, a regression line was done by correlating the ATP concentration and luminescence. Later ATP production was determined as the nmols of ATP produced in 1 minute per protein amount.

### VII.3 Ultrastructural mitochondrial morphology analysis by transmission electron microscopy

Apex hearts from mice were collected and fixed in 2.5% glutaraldehyde, 4% formaldehyde in 0.1 M HEPES buffer for 5 h at 4°C. After several washes with 0.4 M HEPES buffer, samples were postfixed with 1% osmium tetroxide and 3% potassium ferrocyanide in distilled water for 1 h at RT. Then samples were washed in distilled water 3 times for 10 minutes each time. Next they were dehydrated with different percentages of acetone and were embedded in a resin (Spurr Low-Viscosity Embedding).

After solidification, blocks were cut in 60-nm sections, mounted on a copper grid and lead citrate-stained. Samples were evaluated by JEOL 10-10 electron microscopy with different objectives.

### VII.4 Metabolic test in mice

#### **Glucose tolerance test (GTT)**

The main objective of the GTT experiment was to determine the capacity of the organism to clear glucose from blood (Harlass *et al.*, 1991). To carry this out, male mice were subjected to starvation conditions for 6 h before the test started (from 8 a.m. to 2 p.m.). Glucose was administered via intra-peritoneal injection. The amount of glucose used in this experiment was 1.8 g of glucose per kg of mouse weight. Next a small incision was made at the end of the mouse tail to obtain a sufficient amount of blood required to measure the glucose concentration. Blood glucose levels were measured prior to glucose administration at 15, 30, 60, 90 and 120 minutes using specific strips (Lite Blood Glucose Test Strips from FreeStyle).

### **Insulin tolerance test (ITT)**

The ITT measures glucose levels by monitoring the response to insulin administration. ITT was carried out in mice that had fasted for 2 h (from 12 p.m. to 2 p.m.). Insulin was then administered by injection at a concentration of 0.75 U/mL. As well as in GTT, blood glucose levels were measured with specific strips at 15, 30, 60 and 90 min after insulin administration.

### **VII.5 Mitochondrial swelling assay**

In isolated mitochondria, swelling occurs due to the large cytosolic amount of proteins and other solutes. This event occurs *in vitro* after opening the mPTP, which is why it is commonly used as a measure of mPTP opening.

Fresh mitochondria were used, 25 µg per sample in 100 µL of mPTP buffer (200 mM sucrose, 10 mM HEPES, 5 mM KH<sub>2</sub>PO<sub>4</sub>, 10 µM EGTA, pH 7.3). In 96-well glass plates, each sample was placed in a well and then measured by a spectrophotometer. A volume of 750 µM calcium chloride (CaCl<sub>2</sub>) was added to trigger mitochondrial swelling. Measurements were taken every 15 secs for 6 min in a spectrophotometer at A540. A 1 µM concentration of cyclosporine A (CsA) was added to half the wells as a negative control to avoid mitochondrial pore opening.

### **VII.6 Immunoprecipitation of and resolving ATP synthase proteins**

To isolate proteins from the ATP synthase complex, the ATP synthase Immunocapture kit from Abcam (ab109715) was used with some modifications. At least 1 mg of mitochondria at a concentration of 5.5 mg/mL in PBS was needed. This mixture was solubilised with 1% n-Dodecyl-D-Maltoside (DDM), a non-ionic detergent, to disrupt the external membrane. This mitochondrial suspension was kept on ice for 30 min before centrifuging at 13,000g for 30 minutes at 4°C. The supernatant was finally collected.

During centrifugation, agarose beads that contained specific immunocapture antibodies for the ATP synthase complex were prepared. They were cleaned with PBS and placed in a new phial (all the working steps with beads were done with blue tips cutting the end). The previously collected supernatant was added to beads, incubated at 4°C and left on a rotating wheel o/n.

The following day, samples were centrifuged at minimum velocity to allow beads to lower. Beads were resuspended again in PBS with protease inhibitors and 0.05% DDM, and were

placed in a 15-mL phial. Washing was done at 4°C by leaving the phial on the rotating wheel for 15 min. This last step was repeated 3 times, and in the last repetition beads were transferred to a 1.5-mL phial. To elute proteins from beads, sample buffer (50 mM Tris-HCl, 20% SDS, 10% glycerol, 1%  $\beta$ -mercaptoethanol, 0.02% bromophenol blue, pH 6.8) was added with no medium. The sample buffer was collected and stored at -80°C.

To resolve the proteins obtained from the immunocapture, a big gradient gel (10-15%) was done using the former gradient (Table 10) in a big glass. Runs were done with the same buffer as SDS-PAGE electrophoresis (Section III.2)

**Table 10. Big gradient gel composition**

Resolving gel component	Acrylamide solution, 10%	Acrylamide solution, 15%
<b>A/BA 29:1</b>	4 ml	3.5 mL
<b>Separated buffer 4x *</b>	3 ml	1.75 mL
<b>Distilled water</b>	4.9 ml	1.75 mL
<b>APS 20%</b>	15 $\mu$ L	10 $\mu$ L
<b>TEMED</b>	5.76 $\mu$ L	3.36 $\mu$ L

**\* Separated buffer 4x: Tris-HCl 1.5 M, SDS 0.4%, EDTA 8 mM pH 8.8**

**Stacking gel component**

<b>A/BA 29:1</b>	1.2 mL
<b>Concentrated buffer 4x *</b>	2 mL
<b>Distilled water</b>	4.7 mL
<b>APS 20%</b>	37 $\mu$ L
<b>TEMED</b>	20 $\mu$ L

**\* Concentrated buffer 4x: Tris-HCl 0.5 M, SDS 0.4%, EDTA 8 mM pH 6.8**

Then to stain proteins, silver staining (PlusOne Silver Staining Kit from GE Healthcare Life Science) was done.



# Results



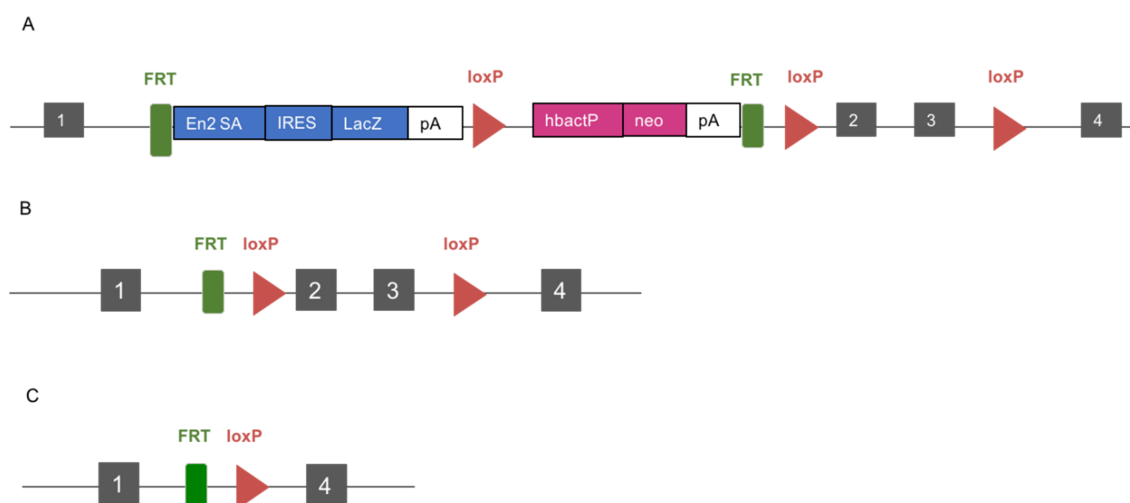
## I. PROTEINS ASSOCIATED WITH H<sup>+</sup>-ATP SYNTHASE

### I.1 Establishment of the mice DAPIT and MP68 knockout colonies

#### DAPIT knockout mice

DAPIT is a subunit of mitochondrial H<sup>+</sup>-ATP synthase, which has been identified very recently and whose role in the complex remains unclear. To study its expression pattern and role *in vivo*, a versatile knock-in mouse was established, which can be used as both lacZ reporter and conditional knock out from the European Mouse Mutant Archive (EMMA), ID EM:07834.

Following *in vitro* fertilisation, the obtained animals were crossed with C57BL/6 mice to verify the germinal transmission of the transgenic cassette (Fig. 15A). Next, the sequences between FRT sites, including neomycin resistance and LacZ genes were deleted by crossing the positive descendants with Rosa26:FLP mice, a transgenic mouse line that expresses the flipase gene (Fig. 15B). This was done to avoid interfering with gene expression levels. In this way, floxed mice were obtained with the codified exons of interest between loxP sites. Lastly, in order to obtain the knockout (KO) mice of interest, floxed mice were crossed with a Sox2:Cre line to delete the codified exons (Fig. 15C).

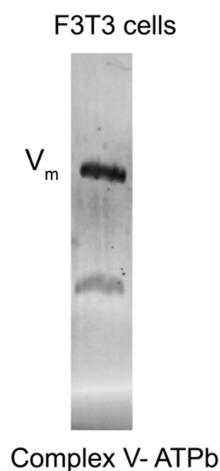


**Figure 15. Schematic representation of the different transgenic lines of proteins DAPIT and MP68. A.** Original construct of the transgenic Dapit mouse. **B.** Representation of an allelic construct after crossing transgenic Dapit with a flipase mouse. Neo-LacZ cassette deletion occurred. **C.** Construct was obtained after the crossing with a Cre protein expression mouse line. The exons between LoxP sites were deleted.

### MP68 knockout mice

To obtain the MP68 knockout mouse, the same strategy was employed as that for the DAPIT knockout with specific primers to the genotype (Fig. 15). In this case, ES cells were obtained from the European Mouse Mutant Cell Repository (EuMMCR). The transgenic ES for the 2010107E04Rik gene (*Mp68*) was used in the JM8A1.N3 cell line. ES cells were used to obtain a C57BL/6 chimera, which was crossed several times until the germinal transmission of the transgene was observed. To obtain the KO mice, floxed mice were crossed with a Sox2:Cre to generate total KO mice.

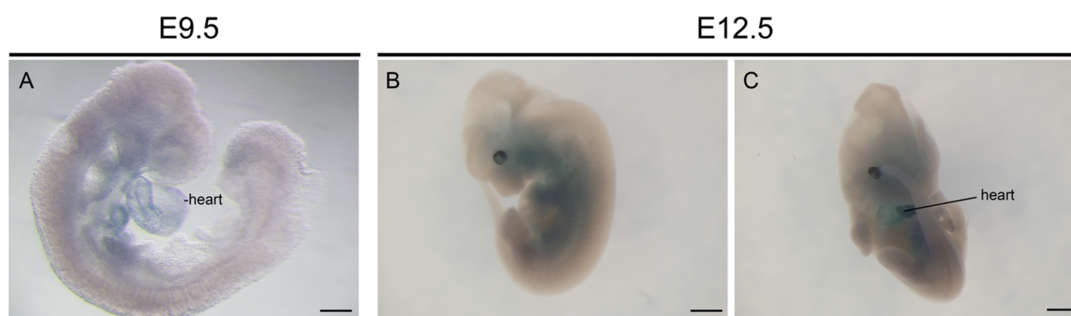
Previously to the establishment of mouse colonies, mitochondria isolated from fibroblast in culture were used to study the function of the ATPase associated proteins DAPIT and MP68. When we analysed the ATPase complex by staining with ATP beta antibody, the monomeric form of the ATPase was detected, but no multimeric forms were observed (Fig.16). This excludes the use of cells to analyse all the possible conformations of ATPase and highlights the importance of using an in vivo model to analyse ATPase supercomplexes. For this reason, the remaining experiments were performed with tissue-isolated mitochondria.



**Figure 16.** Blue native of cell mitochondria to analyse the ATPase organization by using ATP beta antibody.

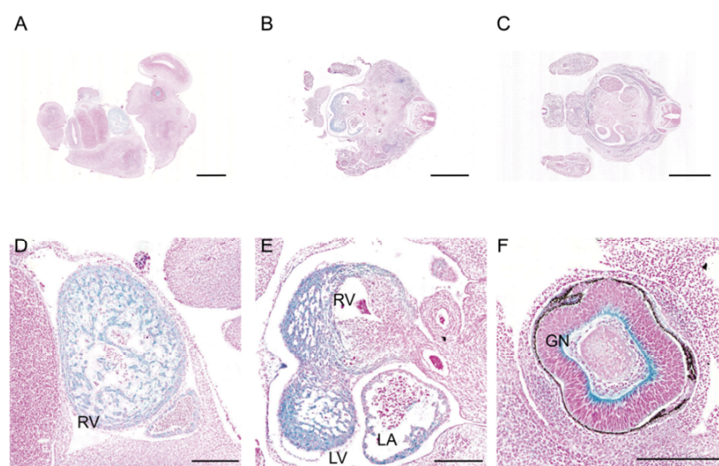
## I.2 Characterisation of DAPIT KO mice

The mice that harboured the original construct (Fig. 15A) constitutively expressed the  $\beta$ -galactosidase gene under the endogenous *Dapit* promoter. This allowed us to investigate the *Dapit* expression pattern during embryonic development by staining with X-gal. For this purpose, E9.5 and E12.5 embryos were chosen. In the E9.5 embryo, the signal was detected in heart. At E12.5, this signal became stronger and extended to other tissues from the ventral embryo part (Fig. 17).



**Figure 17. X-galactosidase staining of the whole embryo.** **A.** Mouse embryo in the E9.5 developmental stage staining to detect LacZ expression to follow *Dapit* gene expression. **B and C.** The mouse embryo that expresses LacZ in the ventral area, and more specifically in heart. Bars, 200  $\mu$ m (A), and 1 mm (B, C).

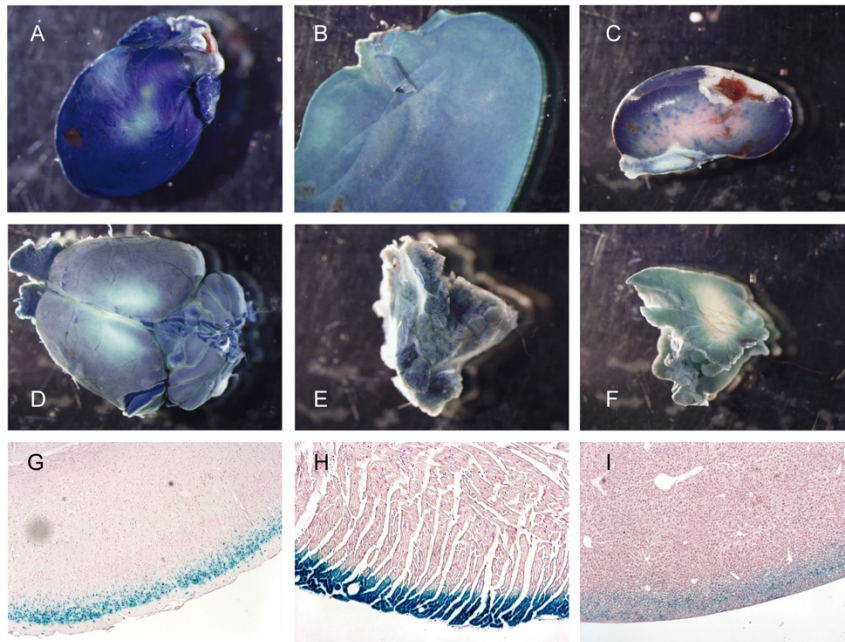
LacZ staining was then analysed in the E12.5 embryo sections. The signal was detected mainly in the heart and the neural system, including retina ganglion neurons (Fig. 18).



**Figure 18. X-galactosidase staining in the E12.5 embryo.** X-galactosidase staining (blue) represents the areas in which the *Dapit* gene is expressed in this stage. **A.** Longitudinal plane of the E12.5 embryo showing LacZ expression in blue. **B and C.** Transversal plane of the E12.5 embryo showing LacZ expression in blue. **D.** Longitudinal section of X-galactosidase staining of the embryonic heart. **E.** Transversal section of X-galactosidase staining of the embryonic heart. **F.** Transversal image of the eye embryo (RV: right ventricle, LV: left ventricle, LA: left atrium, GN: ganglion neuron. Scale A-C 1mm, D-F 250  $\mu$ m).

To determine whether the expression levels were maintained during development, adult organs were also analysed (Fig. 19). In addition to heart, lacZ staining was also detected in the liver, kidney, brain, skeletal muscle and spleen.

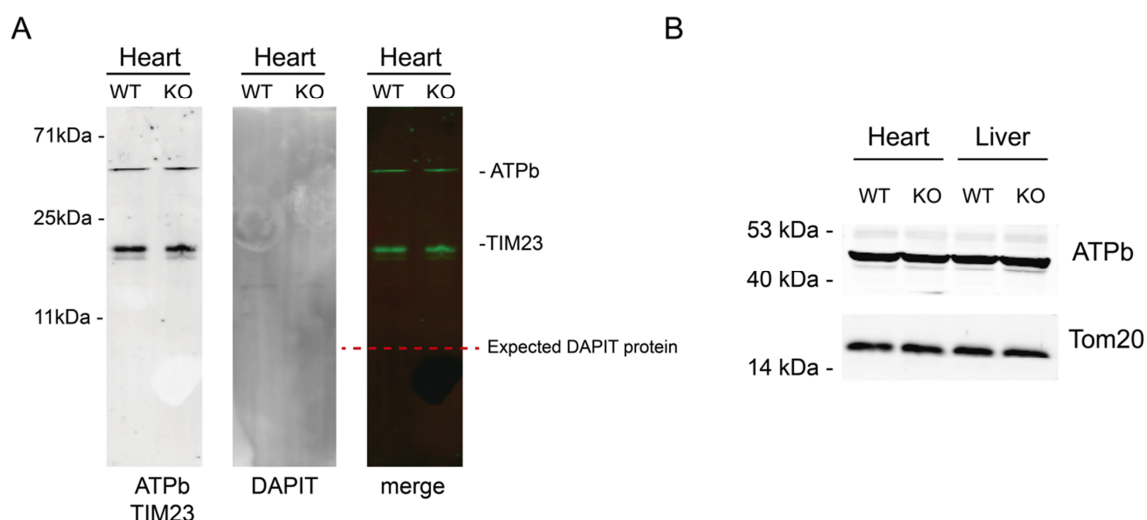
*Dapit* tg/tg



**Figure 19. X-galactosidase staining of the adult organs of the *Dapit* transgenic mice (*Dapit* tg/tg).** A-F. Whole mount X-galactosidase staining A. heart, B. liver, C. kidney D. brain, E. muscle, F. spleen. G-H. Cut organs after X-galactosidase staining. G. Brain slide after X-galactosidase staining. H. Heart slide after X-galactosidase staining. I. Liver slide after X-galactosidase staining.

### I.3 Characterisation of mitochondria and OXPHOS function in DAPIT knockout.

As DAPIT expression was detected at a high level in the heart in different developmental stages, DAPIT function was studied primarily in this tissue. To confirm the absence of the DAPIT protein in the total KO mice, mitochondrial protein from heart was analysed and resolved under denaturalising conditions by SDS gel. However, the available commercial antibody did not detect any signal at the expected location (Fig. 20A).



**Figure 20. The SDS gel of tissue mitochondria.** Mitochondria from wild-type (WT) and KO heart, liver, spleen and brain were analysed by Western blot. **A.** Using the DAPIIT antibody, no signal was detected at the expected size of 7kDa, which corresponds to the DAPIIT protein. **B.** Detection of the ATPb subunit to calculate the amount of the ATPase complex in mitochondria. Tom20 was used to check the quantity of total mitochondria in each sample.

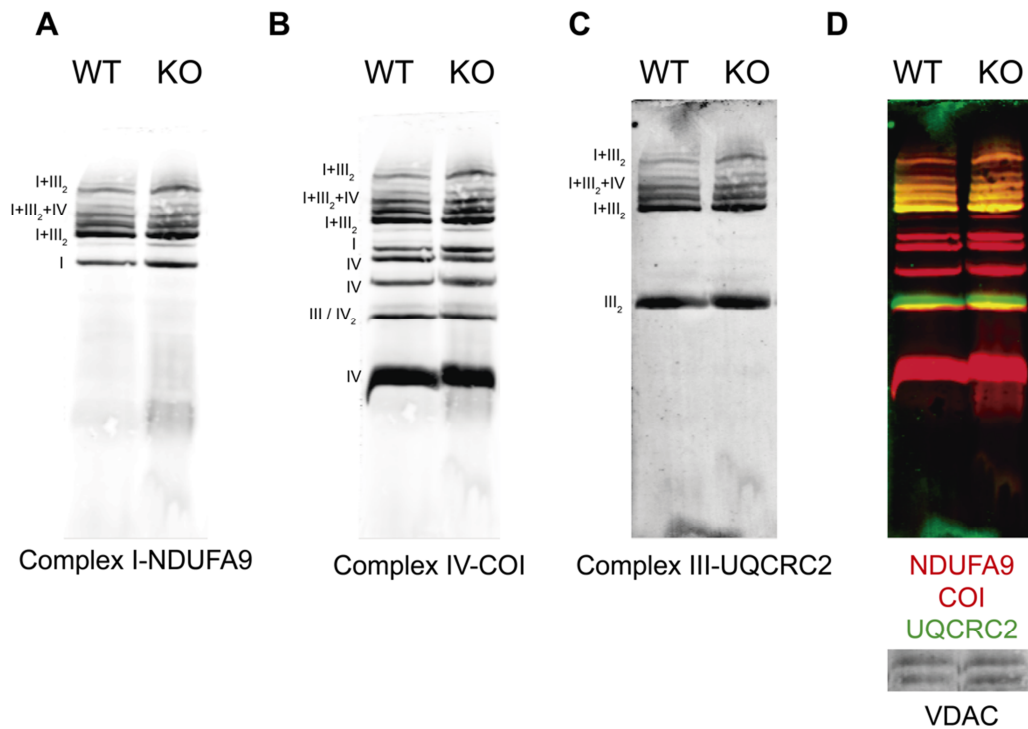
A reduction in the total quantity of the ATPase complex in the DAPIIT knockdown cell has been reported (Ohsakaya *et al.*, 2011). To analyse this, the WT and KO heart and liver mitochondria were analysed by Western blot (Fig. 20B). No reduction in ATPase was observed in the KO mitochondria compared with the WT, which indicates that lack of DAPIIT protein did not alter the amount of ATPase in the investigated tissues. This observation contrasts with previous reports which have shown that the H<sup>+</sup>-ATP synthase complex in DAPIIT knockdown cells reduces (Ohsakaya *et al.*, 2011).

Next the DAPIIT commercial antibody was tested under native conditions. Complex V was labelled in the WT sample, but not in the DAPIIT KO sample, which revealed an efficient deletion of the Dapit allele and the absence of the protein product (Fig. 21). Therefore, the commercial antibody only detected DAPIIT in its folded form.



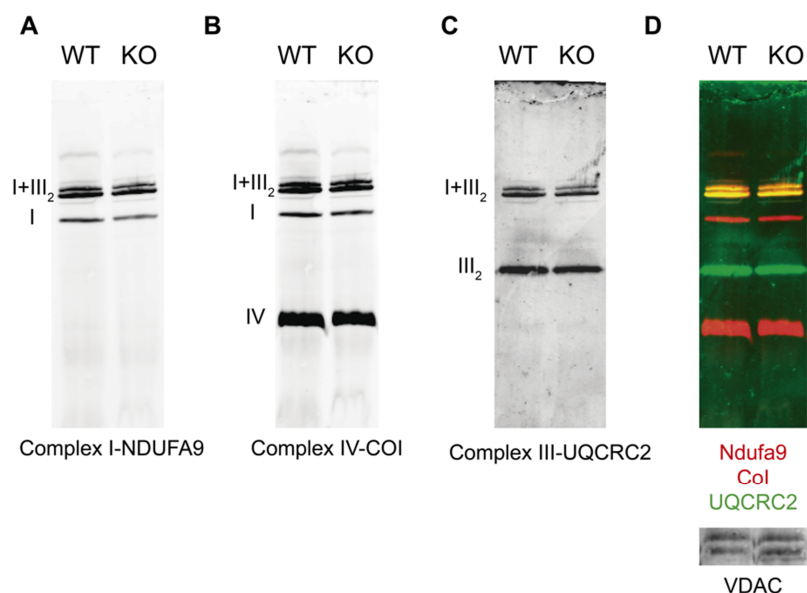
**Figure 21.** The blue native of the mitochondria from the WT and DAPIT KO heart mice. Western blotting revealed the absence of the DAPIT protein in the KO mice. Vdac was used as the loading control.

To analyse the effect of DAPIT loss of function on the different mETC complexes, the mitochondria from the DAPIT KO mice were analysed by BN (Fig. 22). No differences were observed in the organisation of complexes I, III and IV between the WT and KO heart mitochondria.



**Figure 22.** Analysis of the complexes of the heart mitochondria of the DAPIT WT and knockout by BN. Different complexes of digitonise mitochondria were analysed. **A.** The BN of the WT and KO mitochondria analysed with a specific complex I antibody that detects the Ndufa9 subunit. **B.** The previous membrane incubated with a specific antibody of complex IV that detected subunit Col. **C.** Analysis of complex III by the UQCRC2 antibody in the previous membrane. **D.** Merging of previous antibody signals.

These results are not heart-specific as they were confirmed in the liver mitochondria (Fig. 23).

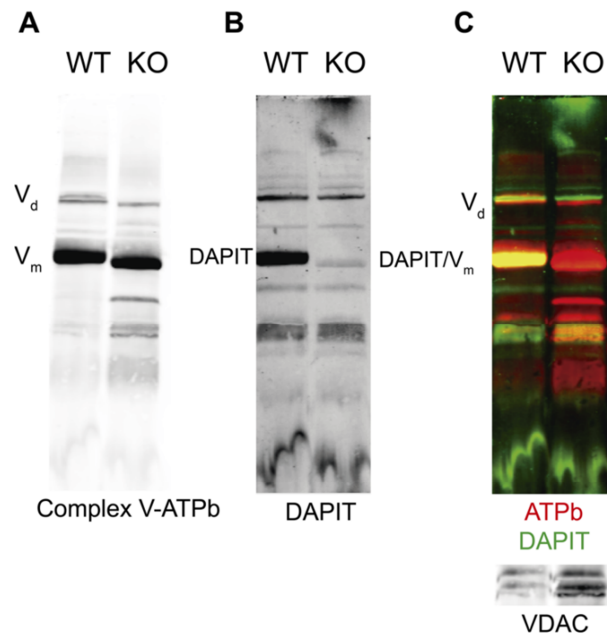


**Figure 23. Analysis of the complexes of the liver mitochondria of the DAPIT WT and knockout by BN.** Different complexes of digitonised mitochondria were analysed. **A.** The BN of the WT and KO mitochondria analysed with a specific complex I antibody that detected the Ndufa9 subunit. **B.** The previous membrane incubated with a specific antibody of complex IV that detected subunit Col. **C.** Analysis of complex III by an UQCRC2 antibody in the previous membrane. **D.** Merging of previous antibody signals.

Next ATPase organisation was analysed. In the WT mice, DAPIT co-migrated with the monomeric form of ATPase (Fig. 24). In the DAPIT knockout mitochondria, this complex migrated slightly more than in the WT, which suggests that the complex could have a reduced mass (Fig. 24A). As DAPIT has been described as a supernumerary subunit of ATPase, one possible explanation could be that its absence results in an observed reduced size of complex V. Other explanation could be the change in the conformation of ATPase complex that alter the migration pattern under native conditions.

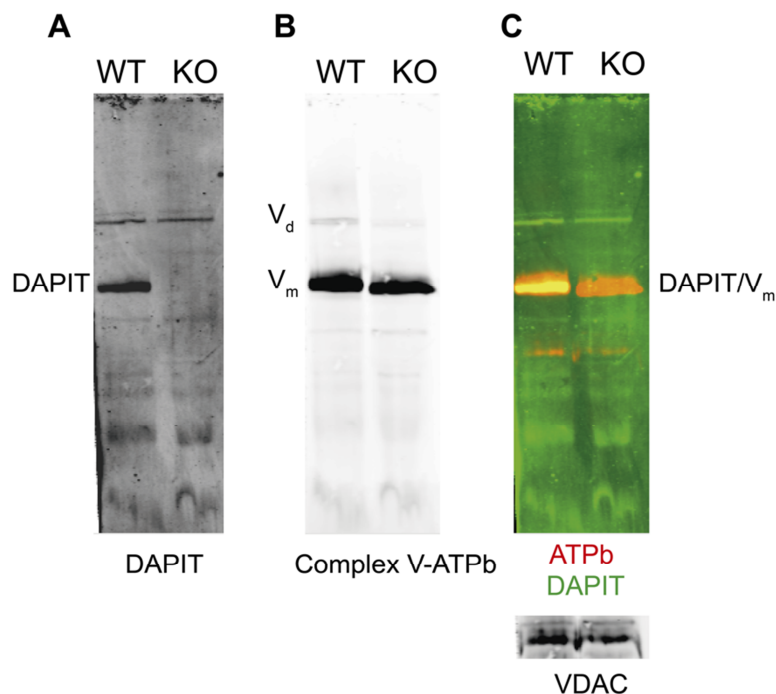
A change in ATPase organisation was observed in the DAPIT KO mice (Fig. 24B). In the WT mice, a high-molecular-weight band was observed that corresponded to ATPase in its dimeric form, but it was not observed in the DAPIT KO mice. This result indicates that DAPIT is necessary for ATPase dimer formation.

An additional lower-molecular-weight band was observed (Fig. 24A). This band was also stained with the anti-beta subunit antibody located in the ATPase F<sub>1</sub> domain. The molecular weight of this band was estimated to be between 400 and 500 kDa. This ruled out the possibility that this band corresponded to the F<sub>1</sub> domain, whose weight was estimated as 350 kDa.



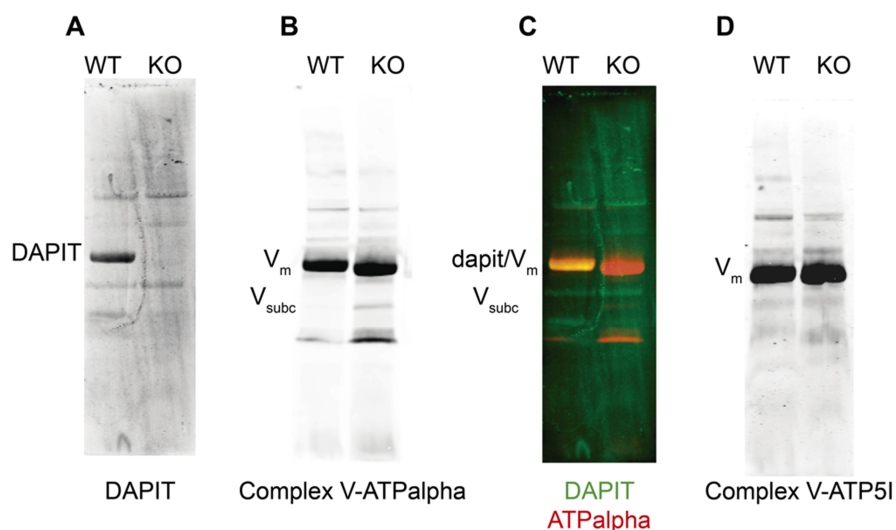
**Figure 24. Analysis of Complex V of the heart mitochondria of the DAPIT WT and KO by BN.** A. ATPase complex detection. B. DAPIT detection in the WT and KO heart mice mitochondria. C. Merging of the signal detected with the DAPIT and ATP subunit beta antibodies.

A similar result was detected in the DAPIT knockout mice liver. However, the ATPase dimers and the F<sub>1</sub> domain were not so clearly detected, which made it difficult to analyse this phenotype in the liver mitochondria (Fig. 25).



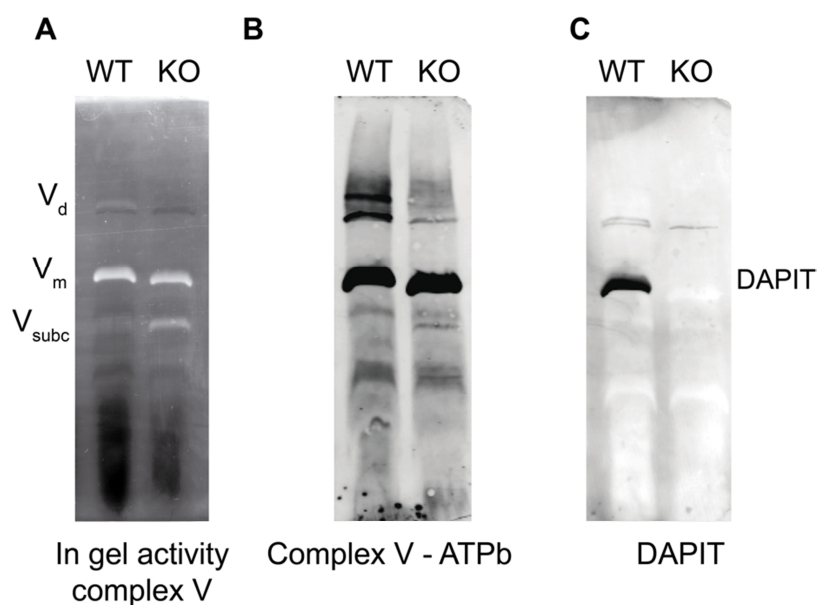
**Figure 25. Analysis of Complex V of the liver mitochondria of the DAPIT WT and knockout by BN.** A. DAPIT detection in the WT and KO heart mice mitochondria. B. ATPase complexes detection. C. Merging of the signal detected with the DAPIT and ATP subunit beta antibodies.

Further antibodies were used to confirm the identity of the ATPase subcomplex detected in the DAPIT KO mitochondria (Fig. 26). The alpha subunit, which is present in the F<sub>1</sub> domain of ATPase, was detected in that band (Fig. 26B). However, the ATP5I protein, which pertains to subunit F<sub>0</sub>, was not detected in the studied subcomplex (Fig. 26D).



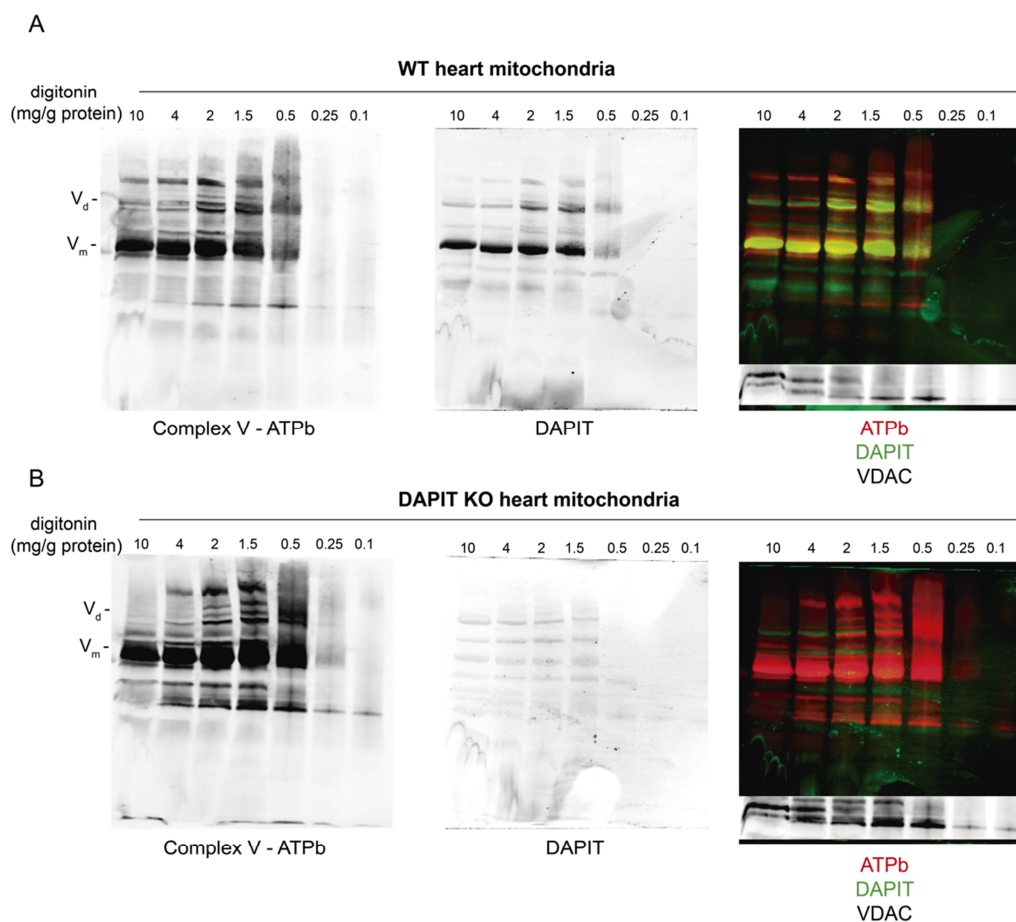
**Figure 26. Analysis of the heart mitochondria complexes by BN.** **A.** DAPIT protein detection in the heart mitochondria. **B.** Analysis of the ATPase complexes using an ATP alpha antibody. **C.** Merging of previous antibody signals. **D.** Another BN gel to analyse the ATP complex by an ATP5I antibody.

Then, the functionality of the ATPase subcomplex observed in DAPIT KO was evaluated by *in gel activity*. It was demonstrated that at least hydrolytic capacity remained in that subcomplex (Fig. 27).



**Figure 27. Functional analysis of the ATPase subcomplex in the DAPIT KO heart mitochondria.** **A.** In the gel activity of complex V using ATP as a substrate of the reaction. **B.** Western blot detection of the ATPb subunit in a parallel running gel. **C.** The previous membrane detected the DAPIT protein.

To exclude the possibility that these results were caused by the specific detergent concentration, a digitonin/protein proportion effect was evaluated. The ATPase subcomplex was detected independently of the amount of detergent used, which indicates that it is not a technical artifact, but is present in cells instead (Fig. 28).

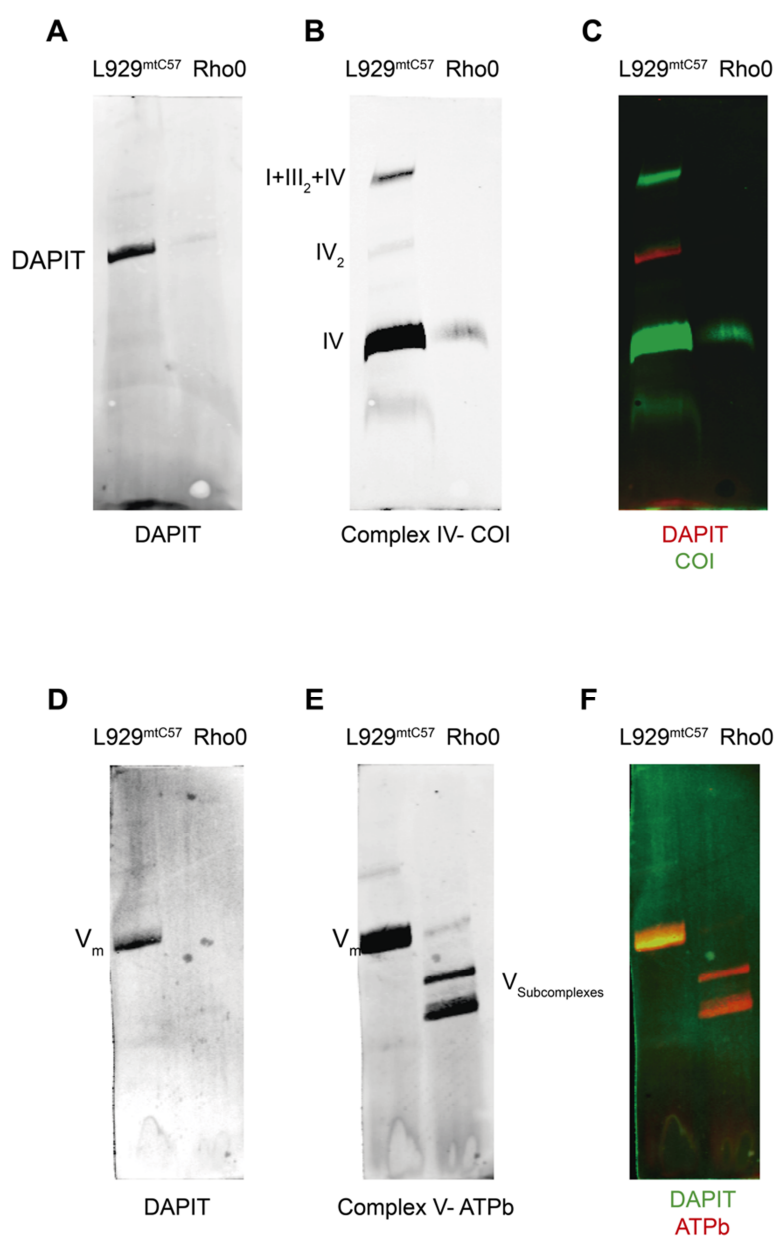


**Figure 28. Digitonine curve analysis in BN gel. A.** BN analysis of the heart mitochondria solubilised with different concentrations of digitonine. ATPase and the DAPIT protein were detected by antibodies. **B.** The same analysis using the DAPIT KO mitochondria.

Thus DAPIT, apart from being implicated in the dimerisation of ATPase, is necessary for ATPase monomers stability. In the absence of DAPIT, a 400-500 KDa subcomplex that contained the alpha and beta, but not the e, subunit was dissociated. All these results strongly suggest that it could correspond to the subcomplex identified in the Rho0 cells composed of at least subunits  $\alpha$ ,  $\beta$ , and  $\gamma_1$  from  $F_1$ , and subunit c from  $F_0$  (Nijtmans *et al.*, 1995).

The relationship between the different ATPase peripheral subunits is not clear. To better understand this, the ATPase structure in Rho0 cells was analysed. These cells have no

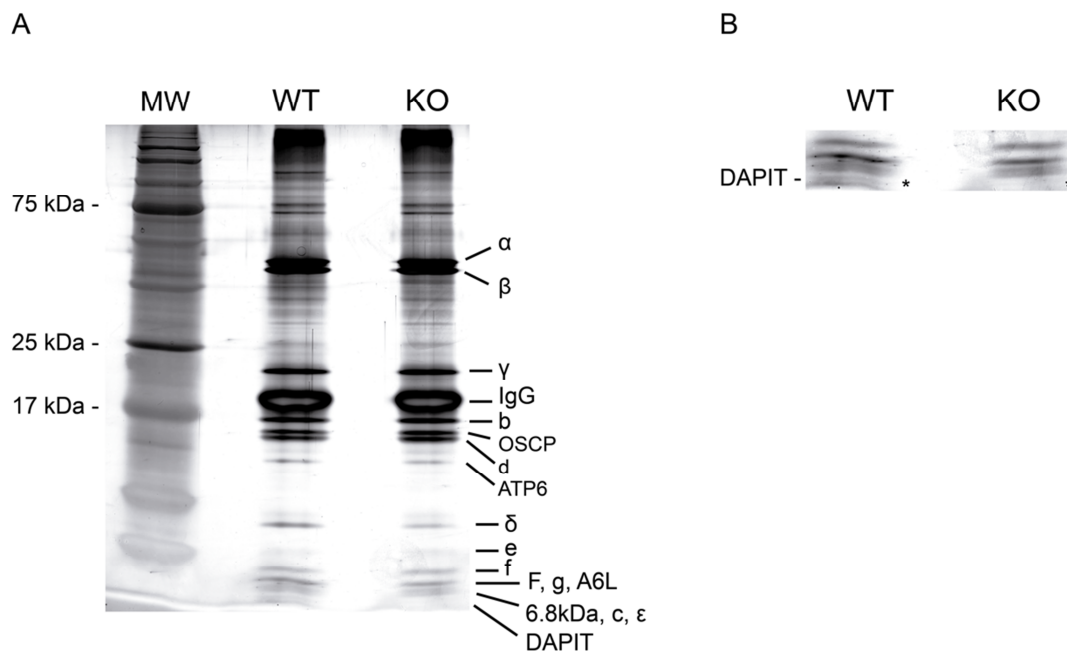
mtDNA, so the mitochondrial proteins coded by it are absent. Specifically, they lack subunits ATP6 and ATP8. The ATPase structure was altered in Rho0 cells given the absence of these subunits, and the size of the complex reduced. More importantly, the ATPase dimers were not found. Surprisingly, DAPIT was not detected in the ATPase of Rho0 cells (Fig. 29). Since Dapit is coded by the nuclear genome, this result indicates that DAPIT is bound to the rest of ATPase by the proteins coded by the mitochondrial genome. Moreover, the absence of DAPIT may be one of the reasons why the Rho0 ATPase complex is unable to form dimers.



**Figure 29. The Rho0 mitochondria analysis by BN gel. A-C.** In Rho0 cells, the CoI subunit is not present in this cell because it is codified by mtDNA. DAPIT is not detected, nor associated with the ATPase complex. **D-F.** In the Rho0 cells without the DAPIT protein, ATPase complex organisation differs and several subcomplexes appear.

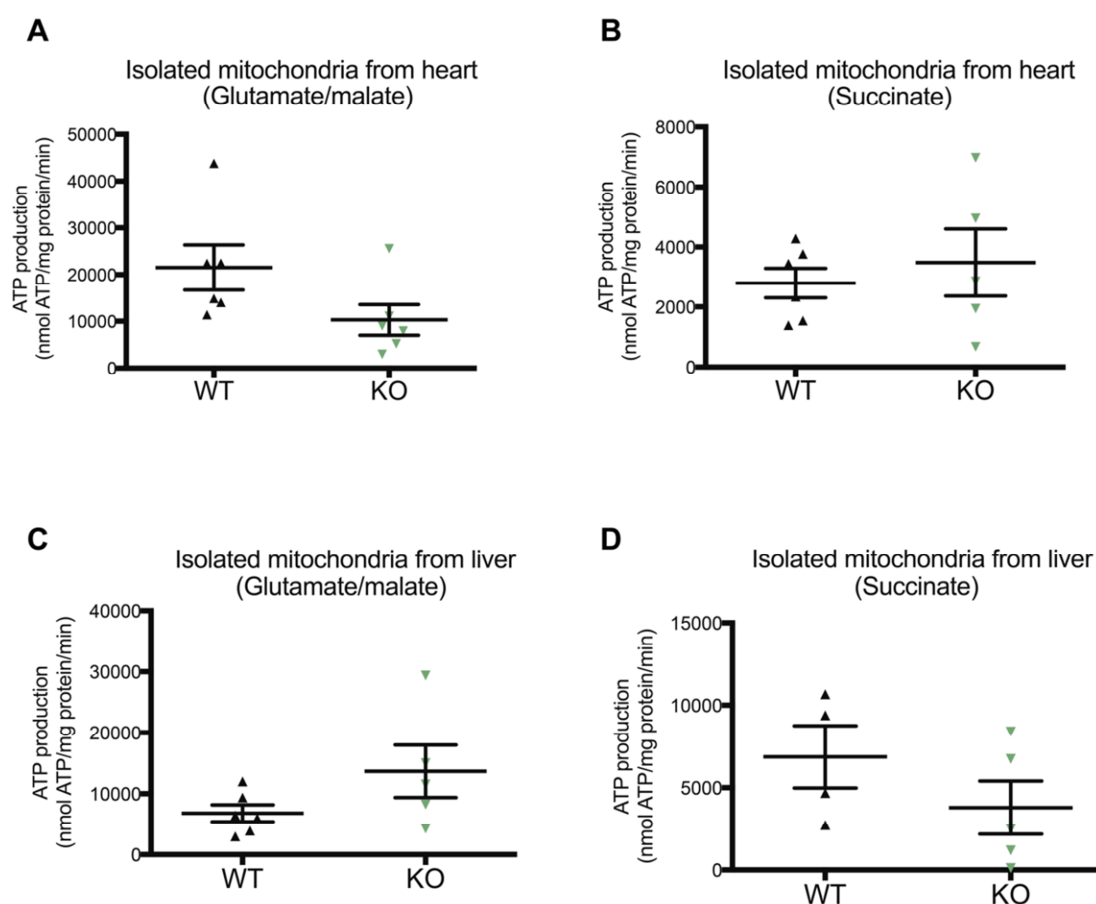
In order to analyse how the absence of DAPIT affected the rest of the ATPase complex, ATPase was immunoprecipitated from the tissue mitochondria. After resolving its subunits in gradient gel and staining proteins with silver, no differences were detected in the other ATPase subunits (Fig. 31). Only a small difference was detected at the bottom of the gel, which could correspond to the DAPIT peptide that was absent in the KO ATPase complex.

Taken together, these results indicate that while other subunits are necessary for DAPIT assembly in ATPase, DAPIT does not affect the assembly of these subunits.



**Figure 30. Immunoprecipitation of the ATPase complex.** **A.** The proteins that form the ATPase complex were isolated and resolved in a gradient acrylamide gel and stained with silver. In the WT and KO ATPase, the main proteins were identified and a difference was observed only in the band that contained the small-sized proteins due to the absence of DAPIT in the KO mitochondria. **B.** Zoom of the part of the gel where the DAPIT protein is. (\* is located, indicating the presence and absence of the DAPIT protein).

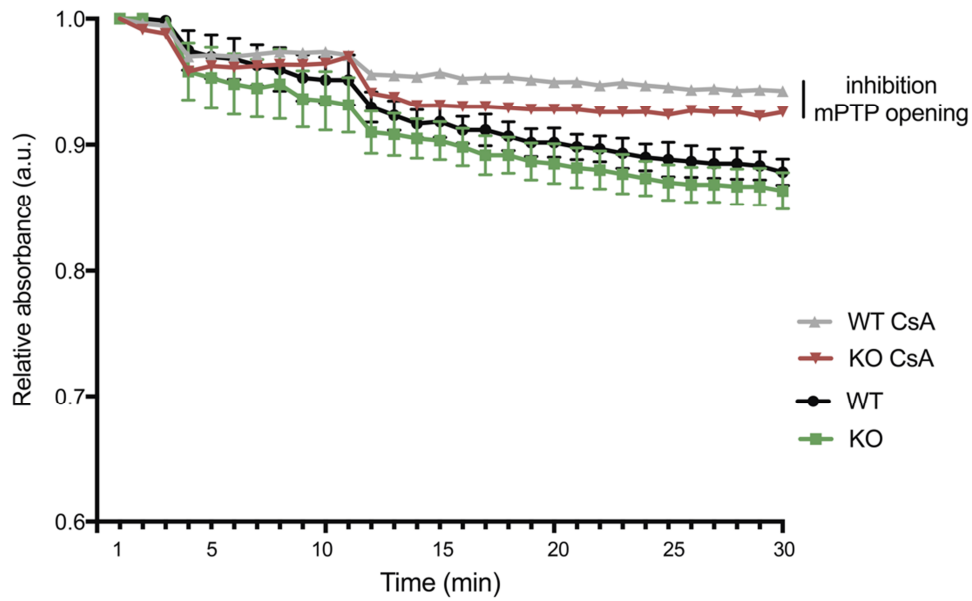
The results reported here indicated that ATPase was the only mitochondrial complex affected by the absence of DAPIT. The functionality of this complex was tested in the mitochondria isolated from heart and liver. ATP production was evaluated by adding different substrates, including glutamate/malate and succinate, to promote the electron transfer from complex I or II, respectively (Fig. 31). No differences in ATP synthesis were observed between the WT and DAPIT KO mitochondria. This result indicates that ATPase dimer formation is not essential for ATP production by mitochondrial ATPase.



**Figure 31. ATP synthesis in the tissue mitochondria.** Analysis of ATP production in the heart mitochondria with glutamate/malate (**A**) or succinate (**B**) as a substrate. Analysis of ATP production in the liver mitochondria with glutamate/malate (**C**) or succinate (**D**) as a substrate. (A-D represents the ATP production value and SEM) (A p-value=0.0841, B p-value=0.5651, C p-value=0.1339 and D p-value=0.2520 all by a two-tailed *t*-test).

All these results show that eliminating the DAPIT protein is a very specific way of inhibiting ATPase dimer formation without it affecting ATP production or other subunits of ATPase.

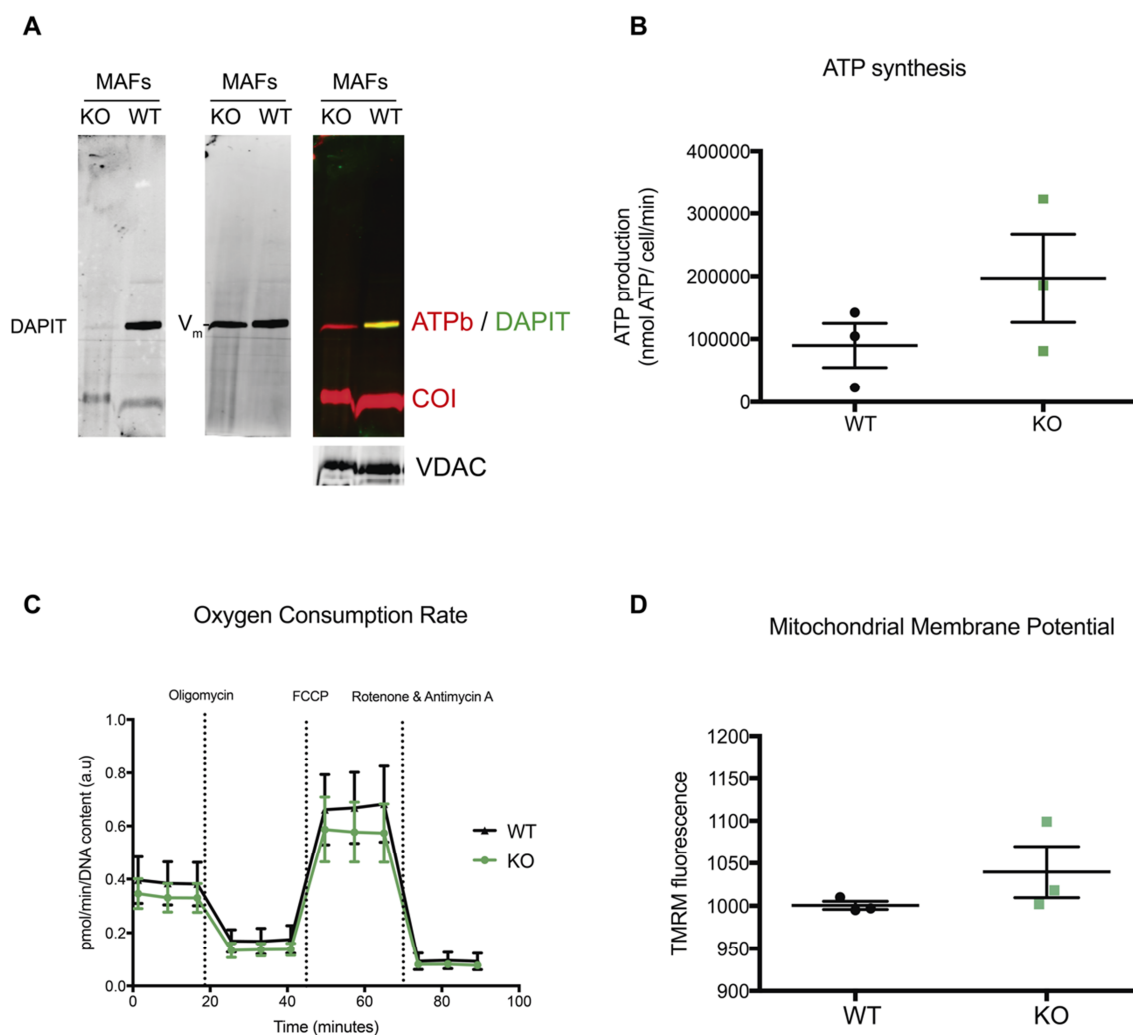
It has been suggested that ATPase dimers are implicated in mPTP opening (Giorgio *et al.*, 2013). To test this hypothesis, the aperture process was analysed in the isolated mitochondria. Addition of calcium to the medium with the heart mitochondria triggered pore aperture by altering mitochondrial morphology. This was detected by a change in the optic density of the medium, which can be evaluated by absorbance measurements. Taking into account the specificity of the signal, demonstrated by mPTP opening inhibition through cyclosporine A, no differences were observed between the WT and DAPIT KO mitochondria (Fig. 32). This result indicates that the disappearance of the ATPase dimers does not affect mPTP opening, and reveals that the ATPase dimers do not form the mitochondrial pore.



**Figure 32. Representation of absorbance variation with time after mPTP opening.** Calcium was added to trigger mPTP opening in presence or absence of CsA that acts as opening inhibitor.

#### I.4 Characterisation of DAPIT knockout mouse adult fibroblasts

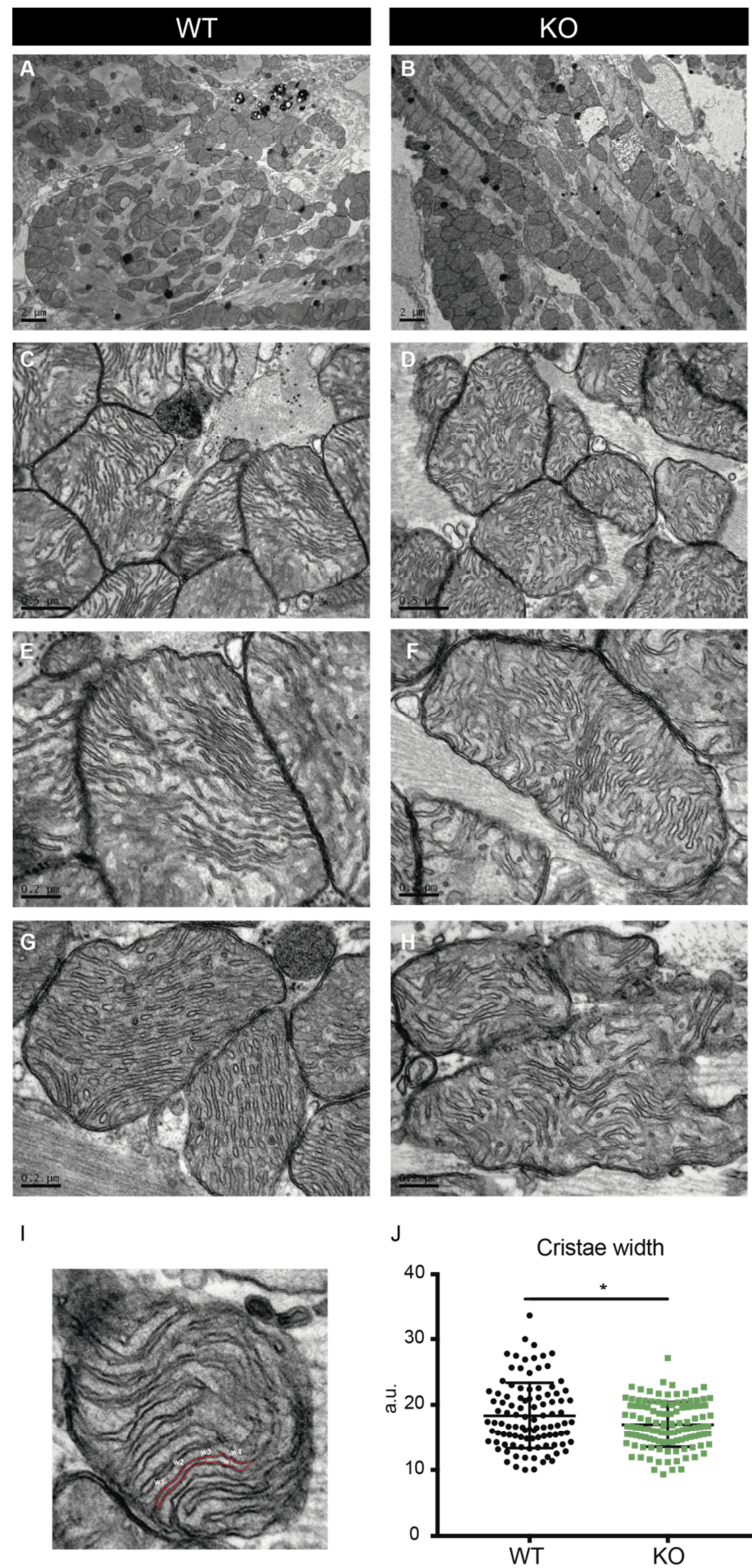
Next the phenotype of the immortalised mouse adult fibroblasts obtained from the WT and KO mice was analysed. After confirming lack of DAPIT by BN (Fig. 33A), ATP synthesis was measured in these cells, and no significant differences were detected (Fig. 33B). After treatment with different drugs, which allowed the basal and maxima respiratory capacity to be studied, no changes in the OCR were observed among the other parameters (Fig. 33C). Moreover, lack of DAPIT, associated with ATPase, did not bring about any change in the mitochondrial membrane potential (Fig. 33D). Altogether, these results support that DAPIT plays no major energetic role.



**Figure 33. Characterisation of mouse adult fibroblasts (MAFs).** Different measurements were taken in one clone of WT MAFs and in another clone of KO MAFs. **A.** BN of the WT and KO MAFs that detected DAPI, ATPb and VDAC proteins. **B.** Measuring ATP production in the WT and KO MAFs. The ATP production and SEM values are represented ( $p$ -value 0.244 by a two-tailed  $t$ -test). **C.** Time profile of the OCR of the WT and KO MAFs after oligomycin, FCCP, rotenone and antimycin A treatment. **D.** Detection of the mitochondrial membrane potential in the WT and KO MAFs. The value membrane potential and SEM are represented ( $p$ -value 0.2687 by a two-tailed  $t$ -test).

### I.5 Role of DAPI in mitochondria morphology

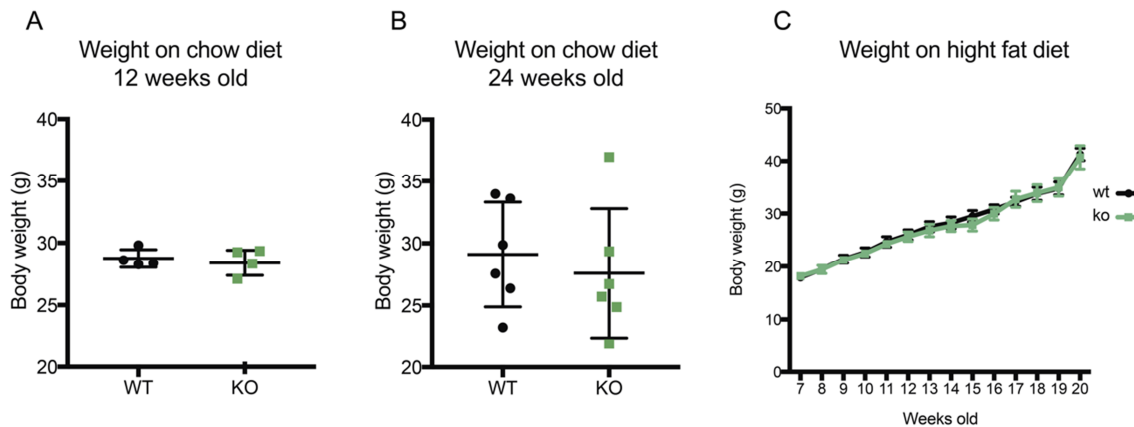
As ATPase dimers are located on the edge of cristae, their morphology was analysed in our mouse model in which ATPase dimers were absent. The cristae morphology was analysed by transmission electron microscopy (TEM) in the WT and KO mouse hearts. Slightly thinner mitochondrial cristae were detected in the KO mitochondria than in the WT hearts (Fig. 34).



**Figure 34. Mitochondria transmission electron microscopy analysis.** **A** and **B**. Heart mitochondria images from the WT and KO mice, respectively, at the 5000x magnification. **C** and **D**. Heart mitochondria images from the WT and KO mice, respectively, at the 40000x zoom. **E** and **F**. Heart mitochondria images from the WT and KO mice, respectively, at the 80000x zoom. **G** and **H**. Heart mitochondria images from the WT and KO mice, respectively, at the 80000x zoom. **I**. Representation of how cristae width was measured. A mean of four measurements ( $w_1$ ,  $w_2$ ,  $w_3$  and  $w_4$ ) was taken to refer to the width of each crista. **J**. Quantification of cristae width in the WT and KO mitochondria (\*  $p$ -value = 0.0192 by a two-tailed  $t$ -test, each dot represents one crista; two animals per group were analysed) (bars A-B 2  $\mu$ m, C-D 0.5  $\mu$ m and E-H 0.2  $\mu$ m).

## I.6 Characterisation of the DAPIT knockout mice phenotype

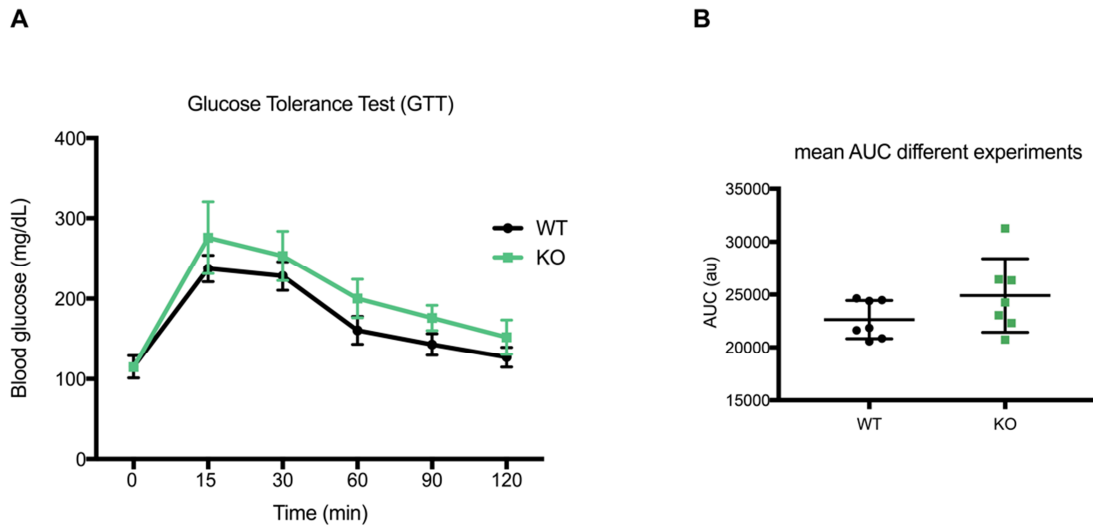
The WT and DAPIT KO mice looked similar when grown under standard conditions, and were fed chow diet (Fig. 35A-B). To test if some differences appeared under extreme conditions, animals were placed on a high-fat diet (HFD). However this diet, which contained a higher percentage of fat (60 kcal/g) did not show any difference (Fig. 35C).



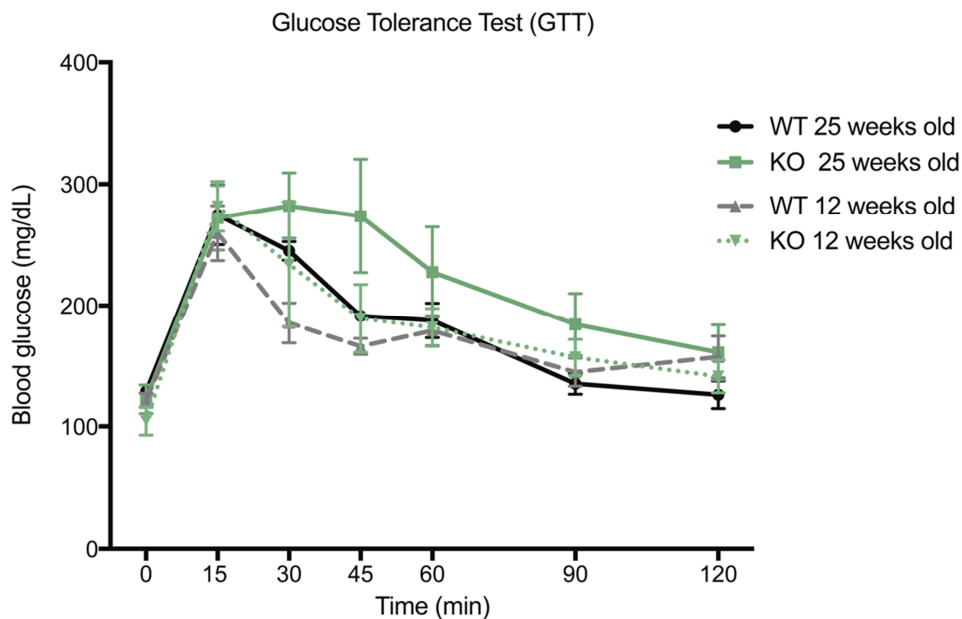
**Figure 35. Weight of mice.** **A.** Weight of the 12-week-old mice on the chow diet (p-value = 0.5805 by a two-tailed *t*-test). **B.** Weight of the 24-week-old mice on the chow diet (p-value = 0.5917 by a two-tailed *t*-test). **A** and **B**, each dot represents the weight of one mouse. **C.** Weight evolution of the high-fat diet. The weight of each group of animals was represented weekly (n=4).

DAPIT was initially identified as an upregulated protein in diabetes mellitus (Kontro *et al.*, 2012). To test its role in glucose metabolism, a glucose tolerance test was performed in the 12-week-old mice. Blood glucose levels were measured before and after glucose intraperitoneal administration (Fig. 36A), and a non-statistically significant tendency was observed for higher glucose levels in the KO vs. the WT group, although a tendency was found in all the seven experiments run (Fig. 36B).

To ensure that glucose regulation was not affected at any age range, the same experiment was repeated by comparing the 12- and the 25-week-old mice (Fig. 37). Once again, glucose levels were slightly higher in the KO than in the WT mice in both the 12- and 25-week-old mice. An age effect on glucose levels was detected as the glucose in the younger mice's blood was rapidly cleared.



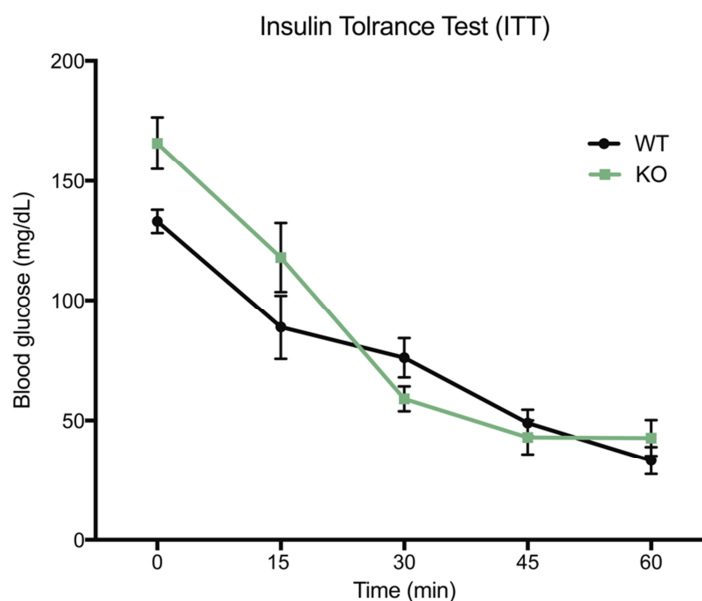
**Figure 36. Glucose tolerance test.** A. Blood glucose levels were measured in the WT (black line) and KO (green line) mice in 15- or 30-min intervals after glucose administration (n=5 mice, p-value = 0.2056 for the area under the curve (AUC) by a two-tailed *t*-test). A representative plot of the seven experiments is shown. B. The mean of several AUCs from seven different experiments (p-value = 0.1493 by a two-tailed *t*-test).



**Figure 37. Glucose tolerance test in mice of different ages.** Blood glucose levels were measured in the WT (black line) and KO (green line) mice, aged 12- and 25-weeks old in 15- or 30-min intervals after glucose administration (p-value = 0.5130 for the area under the curve when comparing the 12-week-old WT and the 12-week-old KO mice by a two-tailed *t*-test; p-value = 0.1658 for the area under the curve when comparing the 25-week-old WT and the 25-week-old mice KO by a two-tailed *t*-test; n=5).

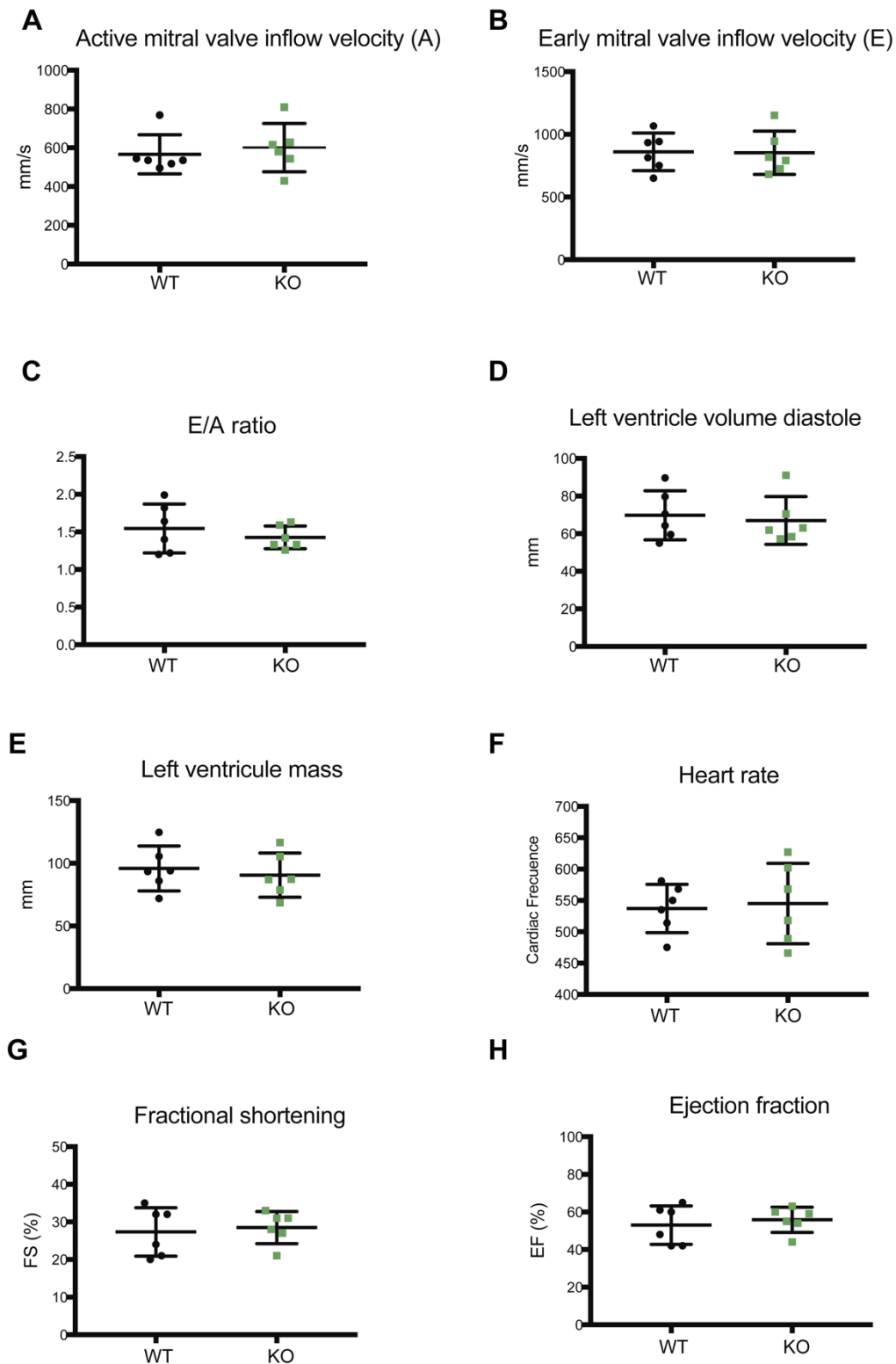
Next glucose clearance in response to insulin was analysed, but no differences between the WT and KO conditions were observed (Fig. 38).

Taken together, these results indicate that DAPIT does not play a major role in diabetes and glucose metabolism.



**Figure 38. Insulin tolerance test.** Blood glucose levels were measured in the 12-week-old WT (black line) and KO (green line) mice in 15-min intervals after insulin administration (n=4 mice, p-value = 0.2056 for the area under the curve by a two-tailed *t*-test). A representative plot of the three experiments is shown.

DAPIT is expressed at very high levels in the heart. To evaluate whether DAPIT affects organ function, an echocardiography was done on both the WT and KO mice (Fig. 39). Diverse parameters were evaluated, such as blood velocity in the mitral valve, and the E/A ratio, which represent the ratio between the top velocity flow in the early diastole and in the late diastole, to evaluate the function of the heart's left ventricle. Ventricle capacity was measured as the left ventricle volume in the diastole, the left ventricle mass or heart rate. Finally, the ejection fraction (EF) and fractional shortening (FS) values were calculated. No differences were observed in these parameters between the WT and DAPIT KO hearts.



**Figure 39. Echocardiography values representation.** The echocardiography values of the WT and KO mice are represented. **A.** Late mitral valve velocity (p-value = 0.6050 by a two-tailed *t*-test). **B.** Early mitral valve velocity (p-value = 0.9375 by a two-tailed *t*-test). **C.** E/A ratio to evaluate the left ventricle function (p-value = 0.4377 by a two-tailed *t*-test). **D.** Left ventricle volume in the diastole (p-value = 0.7180 by a two-tailed *t*-test). **E.** Left ventricle mass (p-value = 0.6141 by a two-tailed *t*-test). **F.** Cardiac frequency or heart rate (p-value = 0.8030 by a two-tailed *t*-test). **G.** Fractional shortening value to measure the fraction of diastolic dimension lost in the systole (p-value = 0.7193 by a two-tailed *t*-test). **H.** Ejection fraction value to evaluate the portion of blood ejected from the ventricle with each heartbeat (p-value = 0.5826 by a two-tailed *t*-test) (n= 6 mice per group, each dot represents a different mouse).

In the same echocardiographs, it was possible to detect some insufficiencies. They were divided into mitral insufficiency (when the mitral valve is unable to perfectly close, blood goes backwards from left ventricle), pulmonary insufficiency (the pulmonary valve does not work properly and blood goes back from the pulmonary artery to the right ventricle), tricuspid insufficiency (the tricuspid valve does not work properly and blood returns from the right ventricle to the atrium), and aortic insufficiency (blood goes back from the aorta to the left ventricle when the aortic valve does not close properly).

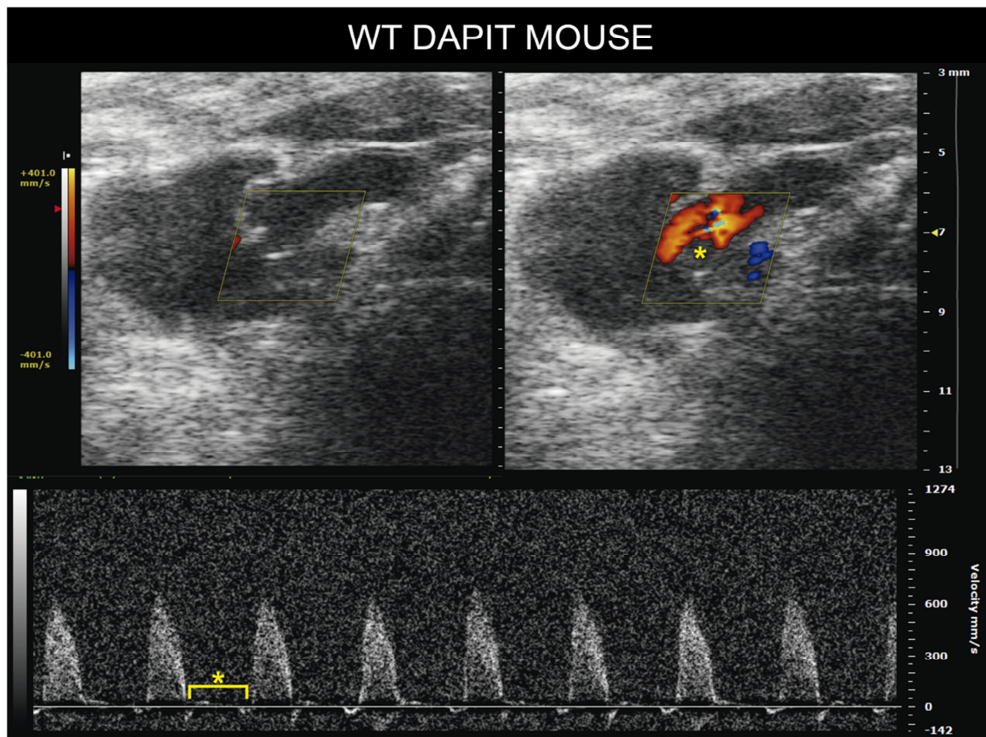
More types and amounts of valvular insufficiencies were detected in the KO mice compared to the WT mice (Fig. 40) ( $\chi^2 = 7.1106$ , p-value=0.007665).

insufficiencies	WT						KO					
	1	2	3	4	5	6	1	2	3	4	5	6
Mitral	Green	Green	Green	Green	Green	Green	Green	Green	Green	Green	Green	Green
Pulmonary	Green	Red	Green	Green	Green	Green	n.m	Green	Red	Green	Green	Green
Tricuspid	Green	Green	Green	Green	Green	Green	Green	Green	Red	Red	Red	Green
Aortic	Green	Green	Green	Green	Green	Green	Red	Green	Red	Red	Red	Green

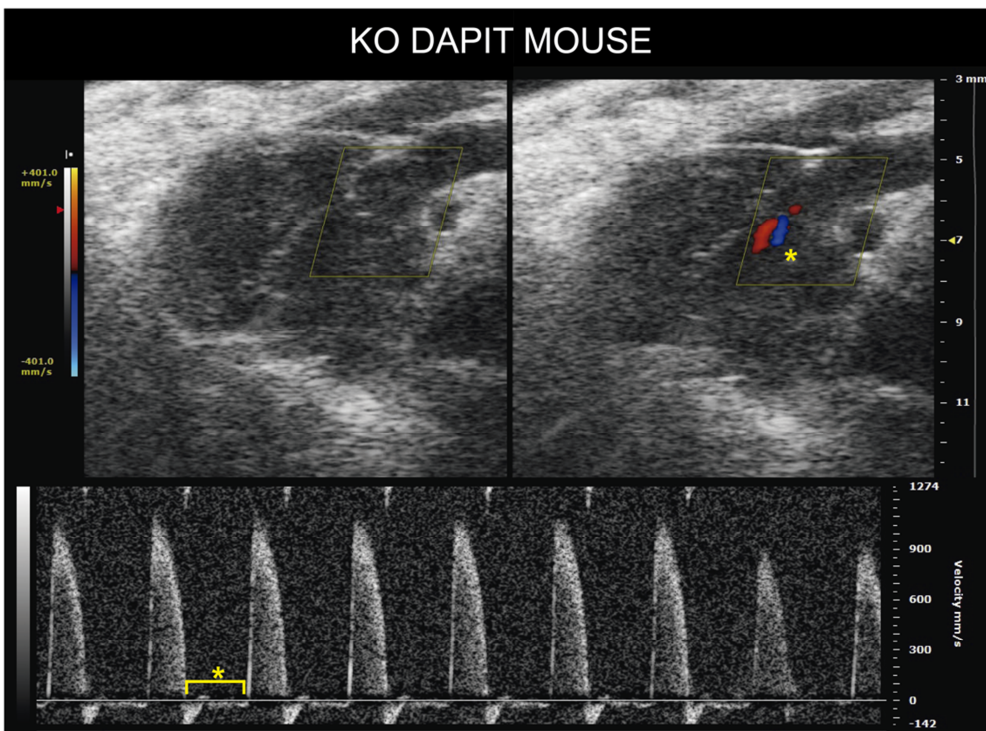
**Figure 40. Representation of the number of valve insufficiencies in the WT and DAPIT KO mice.** Six mice of each genotype were analysed. Green squares represent absence of insufficiency, and red ones indicate that one insufficiency type was detected (n.m: it was impossible to measure this parameter in this specific mouse). The difference was statistically significant in a chi-square test ( $\chi^2 = 7.1106$ , p-value=0.007665).

By way of example, aortic insufficiency was seen in the DAPIT KO mouse (Fig. 41). In the WT mouse (Fig. 41A), blood went from the left ventricle to the aorta, which was showed as positive velocity. A different situation was observed in the DAPIT KO mouse (Fig. 41B) as it was possible to distinguish negative velocity in the diastole, which indicates retrograde flux and aortic insufficiency.

A

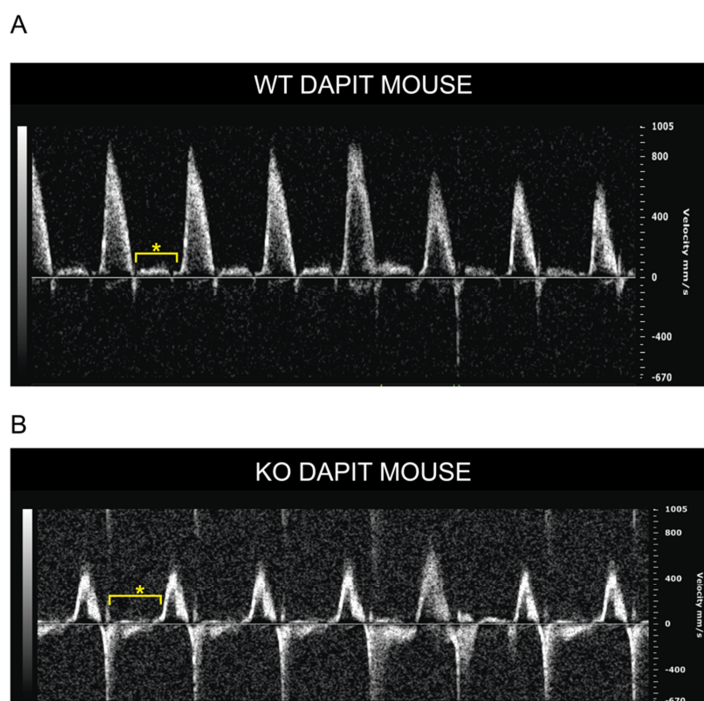


B

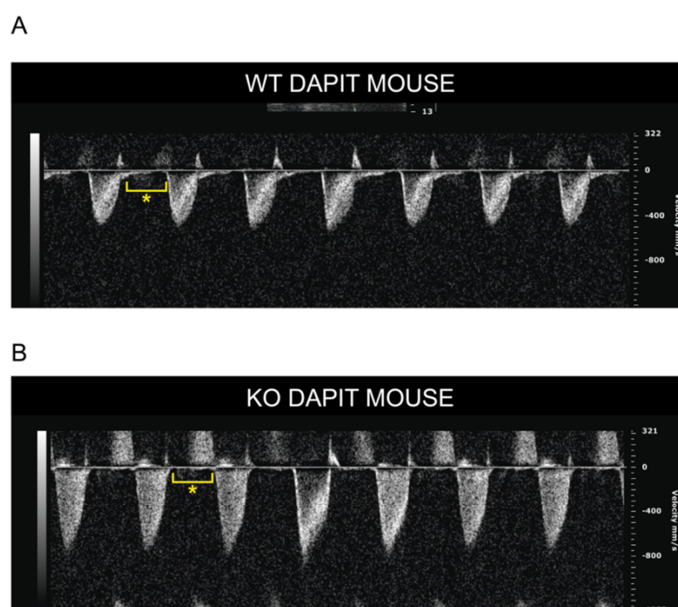


**Figure 41. Representative echocardiography image of aortic insufficiency. A.** An analysis of blood velocity in a representative WT mouse. **B.** An equal analysis done in the DAPIT KO mouse (a yellow star indicates differences in blood direction and velocity terms).

Other insufficiencies as tricuspid (Fig. 42) and pulmonary insufficiency (Fig. 43) were observed in DAPIT KO mice.

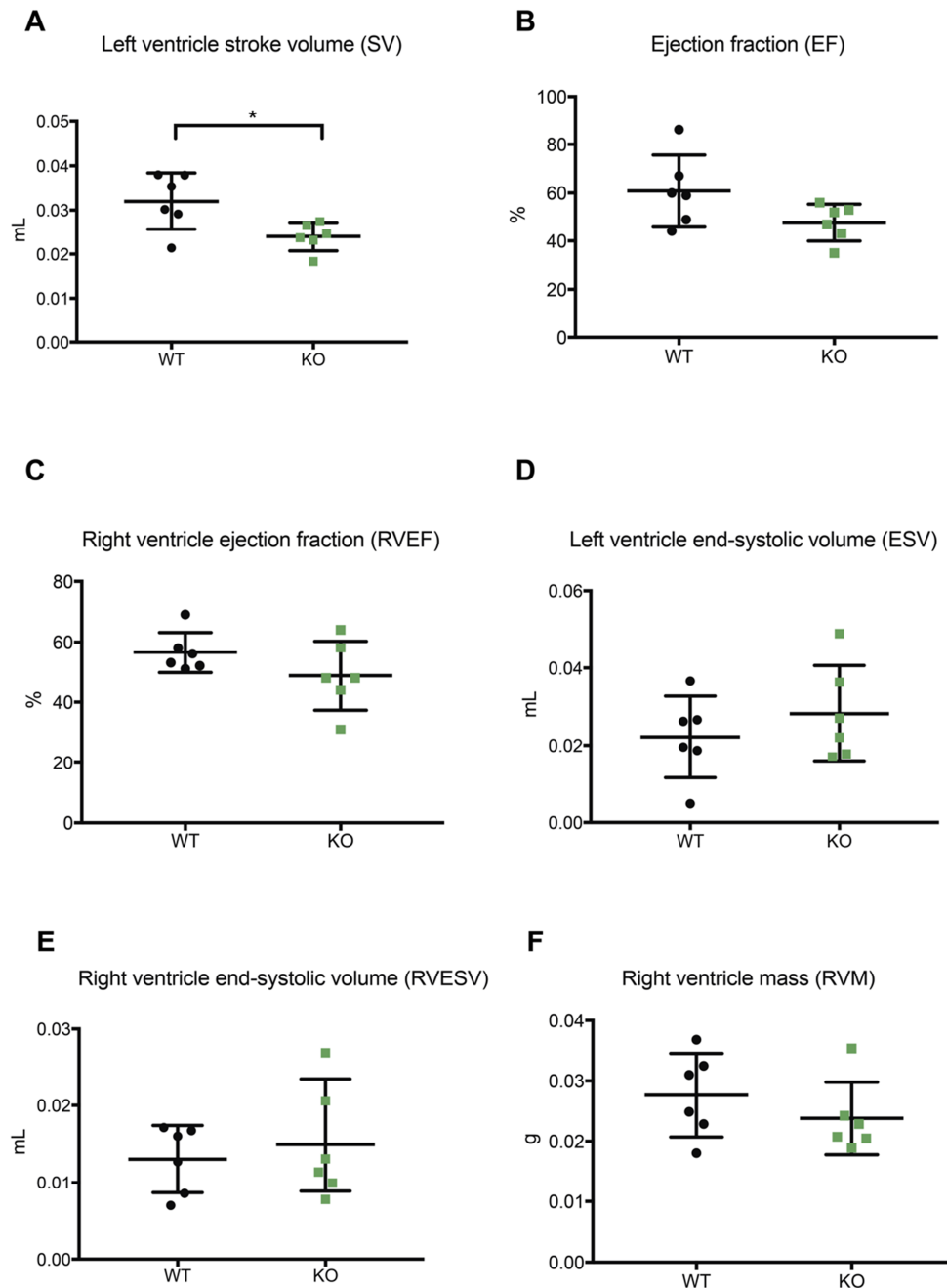


**Figure 42. Representative echocardiography image of tricuspid insufficiency.** A. An analysis of blood velocity in a representative WT mouse. B. An equal analysis done in the DAPIT KO mouse (a yellow star indicates differences in blood direction and velocity terms).



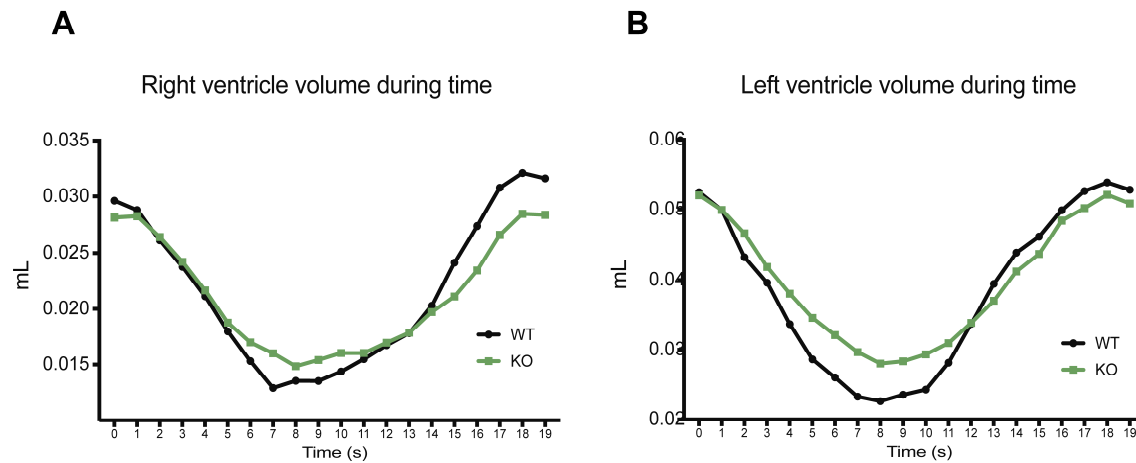
**Figure 43. Representative echocardiography image of pulmonary insufficiency.** A. An analysis of blood velocity in a representative WT mouse. B. An equal analysis done in the DAPIT KO mouse (a yellow star indicates differences in blood direction and velocity terms).

To analyse in more detail the heart functionality, a magnetic resonance imaging (MRI) was performed in WT and DAPIT KO mice. Different parameters were calculated (Fig. 44) showing a weak disffuntion in DAPIT KO hearts, as shown by reduced strock volume (Fig. 44A) and higher volume in the ventricles at the end of the systole (fig. 44C-D). All of these parameters together indicate a worse heart contraction in DAPIT KO compared with WT mice.



**Figure 44. Parameters calculated by analysis of magnetic resonance imaging.** **A.** Left ventricle stroke volume (p-value = 0.0205 by a two-tailed t-test). **B.** Ejection fraction (p-value = 0.0826 by a two-tailed t-test). **C.** Right ventricle ejection fraction (RVEF) (p-value = 0.1869 by a two-tailed t-test). **D.** Left ventricle end-systolic volume (ESV) (p-value = 0.3826 by a two-tailed t-test). **E.** Right ventricle end-systolic volume (RVESV) (p-value = 0.5923 by a two-tailed t-test). **F.** Right ventricle mass (RVM) (p-value = 0.3242 by a two-tailed t-test).

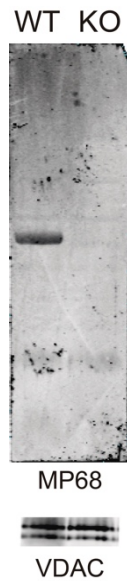
These results can be visualised representing the volume of both right and left ventricles during a pumping cycle. A lower diastolic volume and a higher systolic volume can be observed in the KO. This supports the previous data showing impaired contraction of the heart in the absence of DAPIT.



**Figure 45. Representation of end-systolic volume in right (A) and left (B) ventricle calculated using magnetic resonance imaging. The graph star to represent the ventricle volumen after systole.**

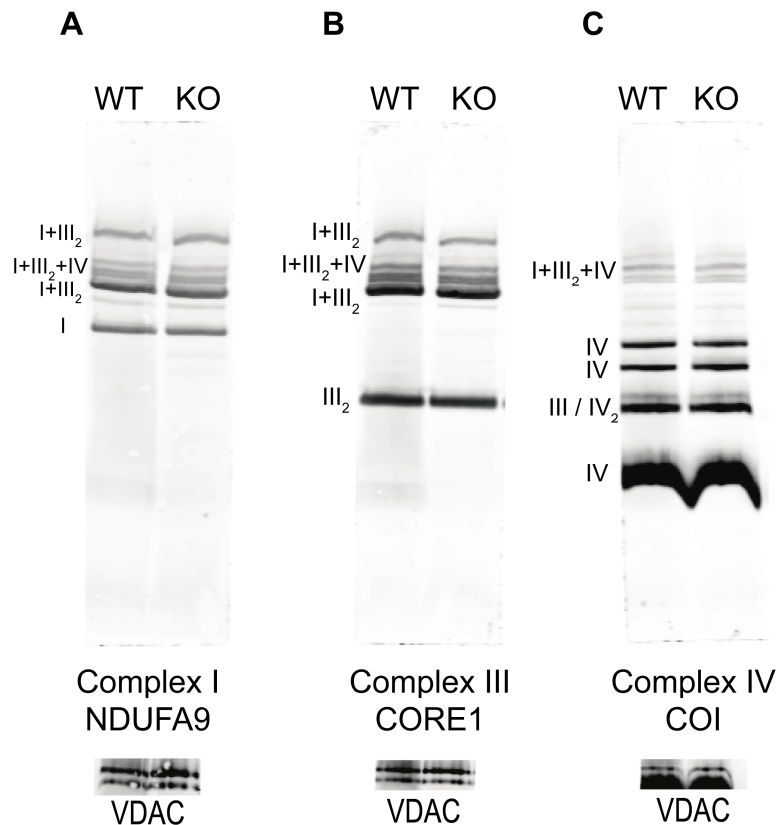
### I.7 Characterisation of mitochondria in MP68 knockout.

The MP68 function was studied primarily in heart and liver tissues. To confirm the absence of MP68 in the total KO mice, a mitochondrial protein was analysed under native conditions. An analysis of the KO heart mitochondria using a specific antibody confirmed that MP68 was lacking in the KO mice (Fig. 46).



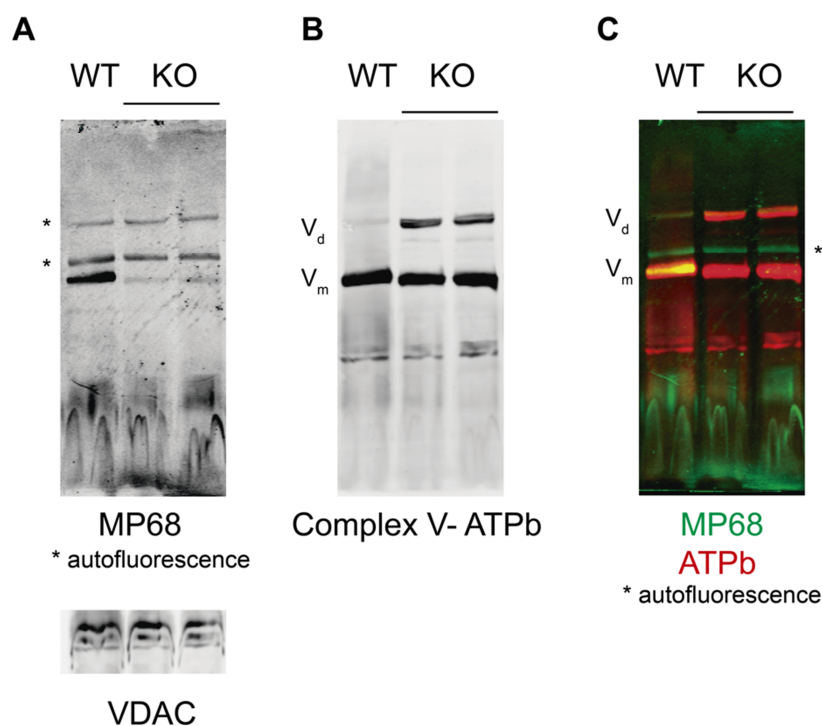
**Figure 46. BN to analyse the heart mitochondria.** MP68 was not detected in KO heart mitochondria. VDAC protein was used as loading control.

Then the status of mitochondrial complexes I, III and IV was evaluated (Fig. 47), where lack of MP68 did not affect the organisation of these complexes.



**Figure 47. Analysis of the heart mitochondria complexes of the MP68 WT and KO by BN.** Different complexes of the digitonised mitochondria were analysed. **A.** The BN of the WT and KO mitochondria analysed with a specific complex I antibody that detects the Ndufa9 subunit. **B.** The BN of the WT and KO mitochondria analysed with a specific antibody of complex IV that recognises subunit Col. **C.** Analysis of complex III by the UQCRC2 antibody. **D.** Analysis of complex V by the ATPb antibody.

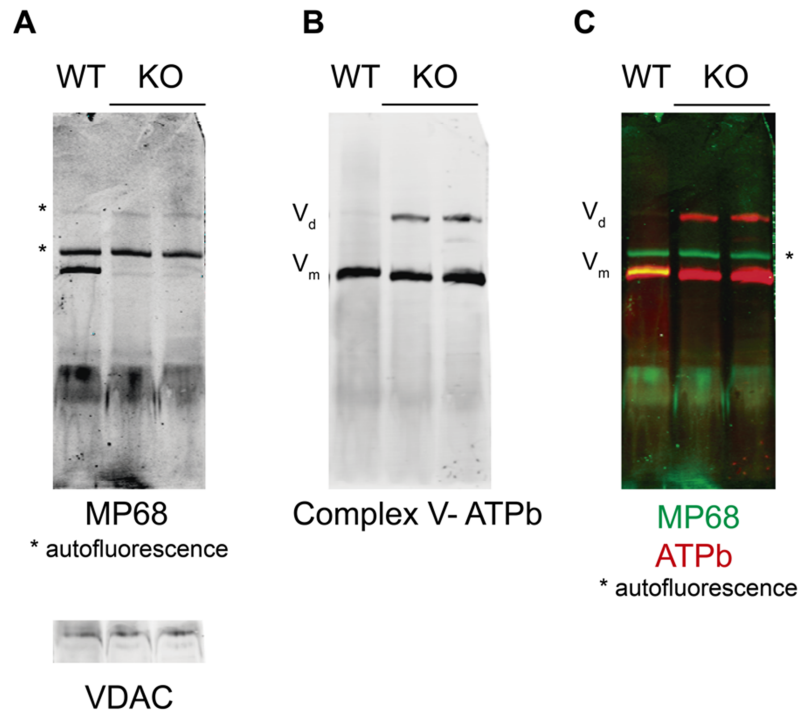
The heart mitochondria from one WT and two different KO mice were analysed (Fig. 48). It can be seen that the band that corresponds to MP68 is present in the WT, but not in the KO mice (Fig. 48A). In the same samples, the change observed in complex V is very interesting. This complex, detected by ATPb, presents a distinctive organisation in the WT and KO mitochondria. Interestingly, it was observed that the relation between complex V in the form of monomers and dimers is bigger in KO mitochondria than in the WT ones (Fig. 48B).



**Figure 48. Analysis of the heart mitochondria complexes of the MP68 WT and KO by BN.** **A.** The BN of the WT and KO mitochondria analysed with a specific MP68 antibody, and VDAC as the loading control. **B.** The previous membrane incubated with specific antibody ATPb that detects complex V. **C.** Merging of previous antibody signals.

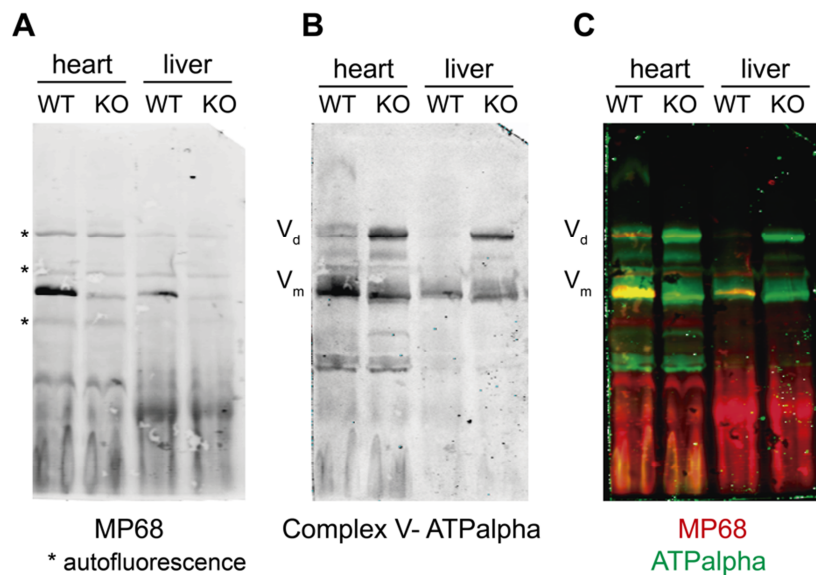
Interestingly, the monomeric form migrated slightly more in MP68 KO (Fig. 48B) samples, as occurs in the DAPIT KO ones.

The liver mitochondria were also analysed (Fig. 49), and the same phenotype was observed with both the absence of MP68 in KO and the differential organisation of complex V.



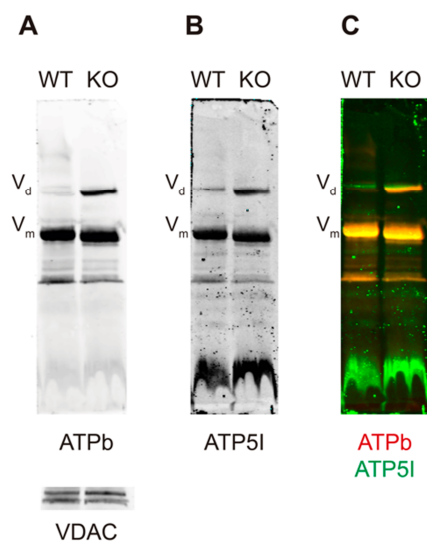
**Figure 49. Analysis of the liver mitochondria complexes of the MP68 WT and KO by BN.** **A.** The BN of the WT and KO mitochondria analysed with a specific MP68 antibody, and VDAC as the loading control. **B.** The previous membrane incubated with a specific antibody ATPb that detects complex V. **C.** Merging of previous antibody signals.

The different organisation of ATPase was confirmed by using a specific antibody for another subunit of ATPase, ATPalpha (Fig. 50).



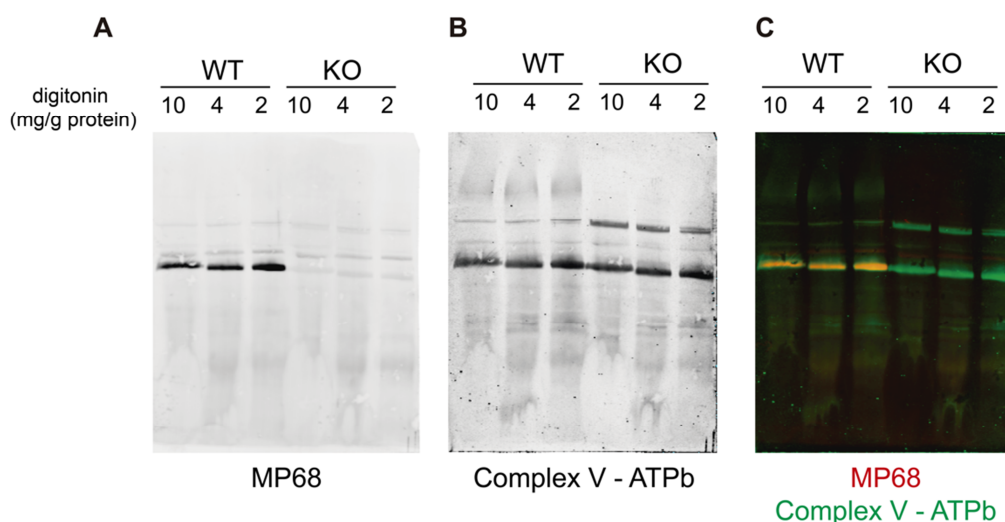
**Figure 50. Analysis of the heart and liver mitochondria complexes of the MP68 WT and KO by BN.** **A.** The BN of the WT and KO mitochondria analysed with a specific MP68 antibody. **B.** The previous membrane incubated with a specific antibody ATPalpha that detects complex V. **C.** Merging of previous antibody signals.

Another ATPase subunit, ATP5I, was described as being essential for dimerisation (Arselin *et al.*, 2003). This protein was detected in the monomeric and dimer form as ATPase of both WT and MP68 KO mitochondria (Fig. 51).



**Figure 51. Analysis of the heart mitochondria complexes of the MP68 WT and KO by BN.** **A.** The BN of the WT and KO mitochondria analysed with a specific ATPb antibody, and VDAC as the loading control. **B.** The previous membrane incubated with a specific antibody ATP5I that detects complex V. **C.** Merging of previous antibody signals.

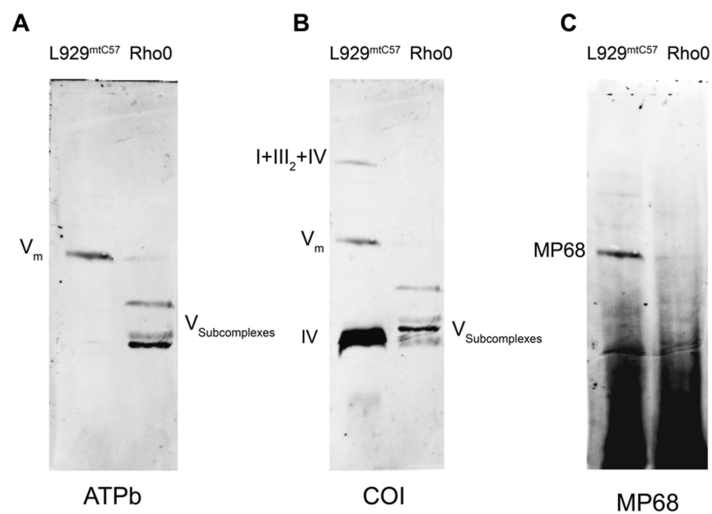
In order to see if the amount of used detergent affected the detection of ATPase organisation, the digitonin/protein ratio was progressively increased, and was visualised by BN gel (Fig. 52). It was possible to confirm the phenotype of the increased amounts of ATPase dimers independently on the amount of detergent.



**Figure 52. Blue Native analysis of the heart mitochondria solubilised at different digitonine concentrations.** **A.** MP68 was detected in the liver mitochondria. **B.** Complex V was detected by the ATPb antibody. **C.** Merging of both signals.

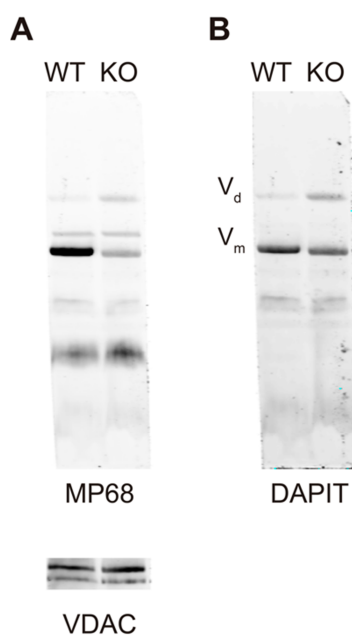
All these results together confirm that lack of MP68 affects ATPase organisation by promoting its dimerisation.

The relationship between mtDNA codified subunits of ATPase and DAPIT was previously described in this thesis. Now, the link between MP68 and these subunits is studied using also Rho0 cells. In these cells, the particular pattern of ATPase as well as the absence of MP68 in Rho0 cells was observed (Fig. 53). These results indicate that the presence of M68 in the ATPase complex depends on the presence of ATP6 and ATP8.



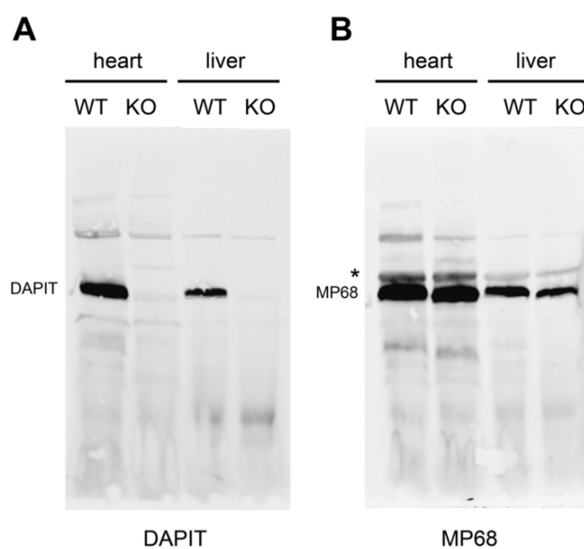
**Figure 53. The Rho0 mitochondria analysis by BN gel. A.** In Rho0 cells, ATPase complex organisation differs and several subcomplexes appear using ATPb antibody. **B.** mtDNA encoded subunit detected by using COI antibody. **C.** MP68 is not detected in Rho0 cells, nor associated with the ATPase complex

DAPIT and MP68 have been described to be of a similar small size, to form the IMD domain and to interact with one other. To understand how lack of MP68 could affect the interaction of DAPIT with the rest of ATPase, DAPIT was analysed in the MP68 KO hearts. While MP68 was not detected in the MP68 KO mitochondria, DAPIT was present in both the ATPase monomer and the dimers of the KO and WT mitochondria (Fig. 54). This means that MP68 is not necessary to make DAPIT form part of ATPase.



**Figure 54.** BN to analyse the DAPIT protein in the MP68 KO heart mitochondria. **A.** MP68 protein detection in the MP68 WT and KO heart mitochondria. **B.** DAPIT detection in MP68 WT and KO heart mitochondria.

To analyse the opposite, if DAPIT is necessary to correctly place of MP68 in ATPase complex, the presence of MP68 was analysed in absence of DAPIT (Fig. 55). In this situation no changes were observed, MP68 was located in ATPase complex even in absence of DAPIT in both heart a liver mitochondria.

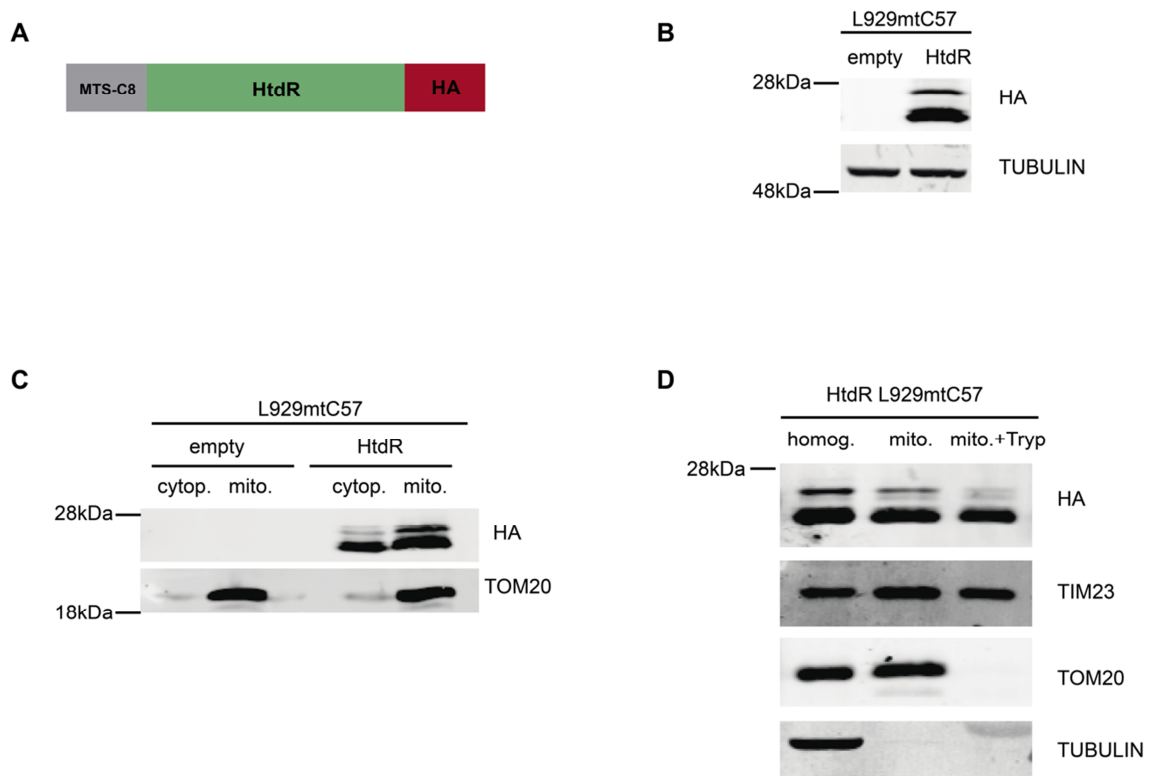


**Figure 55.** BN to analyse MP68 in the DAPIT KO heart mitochondria. **A.** DAPIT detection in the DAPIT WT and KO heart mitochondria. **B.** MP68 detection in the the DAPIT WT and KO heart mitochondria (\* autofluorescence).

## II. DELTARHODOPSIN

### II.1 Deltarhodopsin expression in mammalian cells

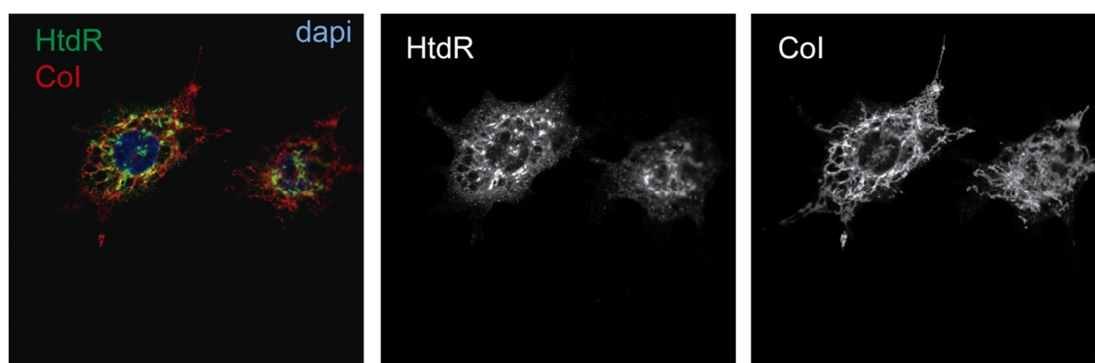
To target the HtdR protein in the mitochondria of mammalian cells, the HtdR sequence was cloned by adding a mitochondrial target signal (MTS) to the N-terminal domain, based on subunit 8 of complex IV, and an HA-tag to the C-terminal domain (Fig. 56A). Stable HtdR expression was performed by viral infection in the mammalian control cells, L929mtC57. At the total protein level two bands of about 28kDa in weight in HtdR-expressing cells were observed (Fig. 56B). The higher band corresponded to the total protein that was expressed, and the lower band correspond to HtdR without the MTS. This is the first evidence to demonstrate that HtdR is able to enter mitochondria. Next in order to observe the importing efficiency to mitochondria, the proportion of HtdR in both the cytoplasm and mitochondria was checked. Although the cytosol contained a portion of HtdR (Fig. 56C), the largest amount of the HtdR protein was located in mitochondria.



**Figure 56. Deltarhodopsin (HtdR) expression in mammalian cells.** **A.** Scheme of the HtdR construct, including MTS-C8 as the mitochondrial target signal (MTS), and HA tag. **B.** Immunodetection with the HA and tubulin antibodies of the total protein from the empty and HtdR-expressing cells. **C.** Immunodetection with the HA and Tom20 antibodies of the cytoplasm and mitochondria protein from the empty and HtdR-expressing cells. **D.** Immunodetection of mitochondrial proteins of the HtdR-expressing cells after treatment with trypsin with the HA, Tom20 and Tim23 antibodies.

To be functional, HtdR has to reach the inner mitochondrial membrane and be inserted into the right orientation. There the  $H^+$  gradient is generated and converted into chemical energy by ATP synthase. To test HtdR submitochondrial localisation, a protection assay was run. Following trypsin treatment, only IMM proteins Tim23 and HtdR remained undigested, while OMM protein Tom20 was digested. (Fig. 56D). This demonstrates that HtdR is targeted to the IMM, where it should be located to generate a proton gradient.

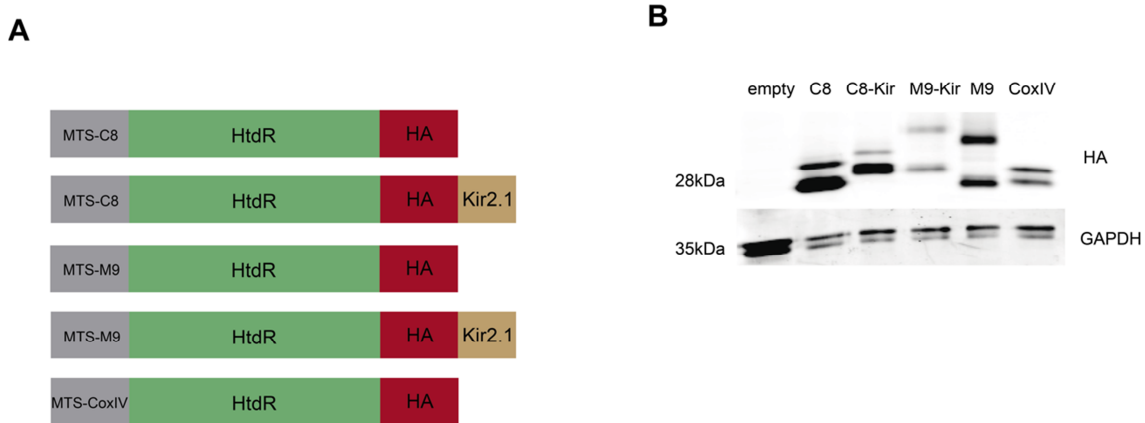
Next HtdR location was checked by immunofluorescence (Fig. 57). We observed partial co-localisation with the Col mitochondrial protein. This effect was due to the higher expression levels of the protein and its accumulation in the ER.



**Figure 57. Immunofluorescence of the HtdR-expressing cells.** Detection of HtdR with the HA antibody and mitochondria morphology with the Col antibody.

In order to improve mitochondrial expression, new proteins with different MTS were synthesised, including those from subunit 8 of cytochrome c oxidase (C8) and subunit 9 of ATPase (M9), and with sequence Kir2.1, an ER exporting signal (Fig. 58A).

From the analysis of the total protein from the cells that expressed the different constructs, it was concluded that HtdR expression existed in all cases. Comparing of the size of the different constructs, confirmed that the small band corresponded to HtdR after the MTS was removed, while the bigger band corresponded to HtdR with the MTS (Fig. 58B). The removal of the MTS further supports the notion that most HtdR is targeted to mitochondria. The C8-HtdR construct displayed the best performance in terms of expression level and efficiency of import judging by the cleavage of the MTS. It was, therefore, chosen for the following studies.



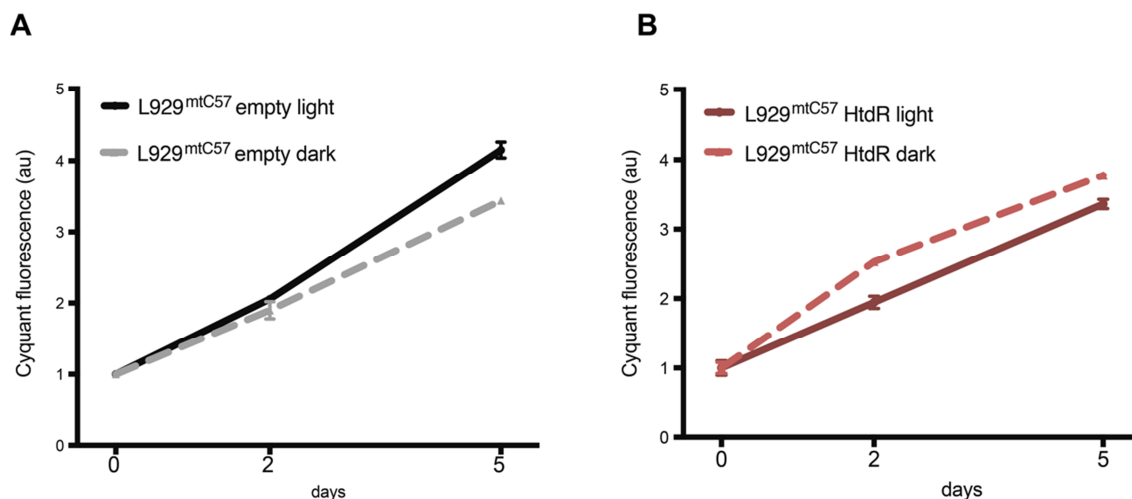
**Figure 58. Deltarhodopsin with different mitochondrial target signals.** **A.** Scheme of the HtdR construct with different mitochondrial target signals. **B.** Immunodetection of HtdR with different sequences that act as the MTS with the HA antibody.

## II.2 Cell growth without functional OXPHOS

To test the functionality and efficiency of HtdR, cell growth under conditions that force them to rely on mitochondrial ATP production rather than on glycolysis was tested by using different carbon sources and inhibitors.

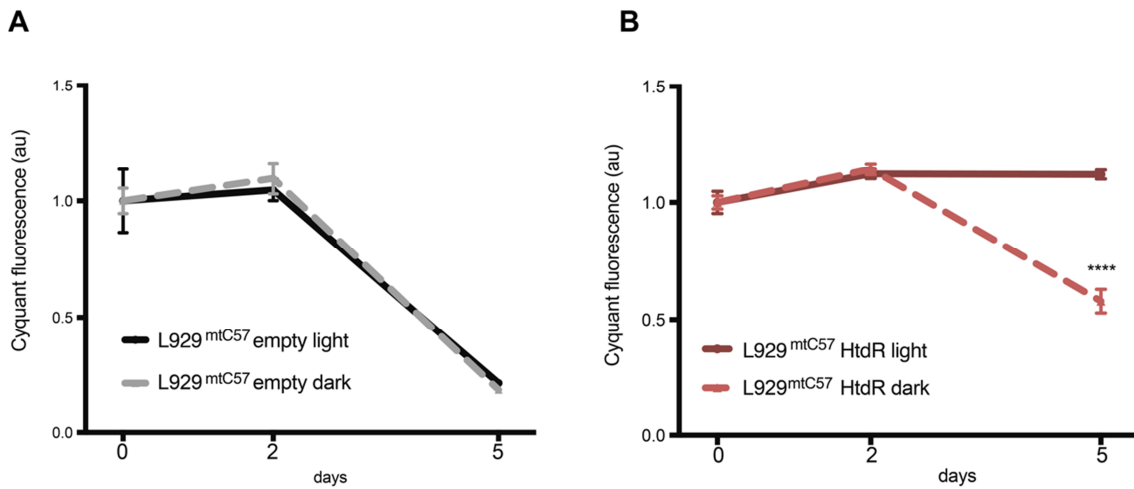
Unlike glucose, galactose does not allow cells to generate sufficient ATP by glycolysis to survive. However, it provides enough energy if generated by the ETC. Under these conditions, the ETC is the main source of energy (Bayona-Bafaluy, Fernández-Silva and Enríquez, 2002). In order to block the ETC, we used antimycin A (Aa), an inhibitor of complex III. Thus in galactose with Aa, cells cannot use glycolysis or ETC to produce ATP and cells die.

Growth of control and HtdR-expressing cells was evaluated in glucose-enriched medium that contained Aa for 5 days either in the dark or light. As expected, both the control (Fig. 59A) and HtdR-expressing cells (Fig. 59B) grew normally because they generated enough energy by glycolysis. Neither HtdR nor the used lighting conditions had any apparent toxic effect.



**Figure 59. Growth curve of the cells in the glucose-enriched medium. A.** Growth curve of empty vector expressing cells in glucose medium in the presence (black continuous line) and absence of light (grey discontinuous line). **B.** Growth curve of the HtdR cells in glucose medium in the presence (red continuous line) and absence of light (red discontinuous line).

When cells were shifted to galactose, the control cells were unable to survive in the presence of Aa either in the dark or light (Fig. 60A). On the contrary, the cells that expressed the HtdR protein survived in galactose plus Aa, but only if they were illuminated (Fig. 60B). This indicates that under light conditions HtdR generates sufficient p.m.f. to allow ATP synthesis by mitochondrial ATPase because, in this way, it rescues cells from death (Fig. 59B). The possibility that HtdR itself promotes cell survival by a different mechanism was ruled out by the fact that cells died in the absence of light. These results demonstrate for the first time that mammalian cells can convert light energy into electrochemical energy, which is used later by ATPase to produce ATP that can support cell growth.



**Figure 60. Growth curve of the cells in the galactose-enriched medium. A.** Growth curve of the control cells in galactose medium in the presence (black continuous line) and absence of light (grey discontinuous line). **B.** Growth curve of the HtdR cells in galactose medium in the presence (red continuous line) and absence of light (red discontinuous line) (in the last point: \*\*\*\* p-value < 0.0001 by a two-tailed *t*-test).

This cellular tool, in combination with the previously developed cell lines that express genes AOX and/or NDL1, will allow us to split the different function of the OXPHOS system into mammalian cells and to contribute to physiopathology.





# Discussion



## I. ASSOCIATED PROTEINS WITH ATP SYNTHASE

ATPase conformation is important for determining mitochondrial morphology and cristae formation. For the first time, we report that DAPIT is necessary for dimer formation. In the absence of the DAPIT protein, the capacity to form dimers disappears. The lack of DAPIT does not affect the loss of other ATPase subunits, indicating that our model is a very useful tool to analyse specifically how the loss of the ATPase dimers affects the cell. Moreover, DAPIT KO ATPase has the same capacity to produce ATP as the WT mitochondria, but without the presence of dimers. This indicates that ATPase function is independent of its dimeric conformation.

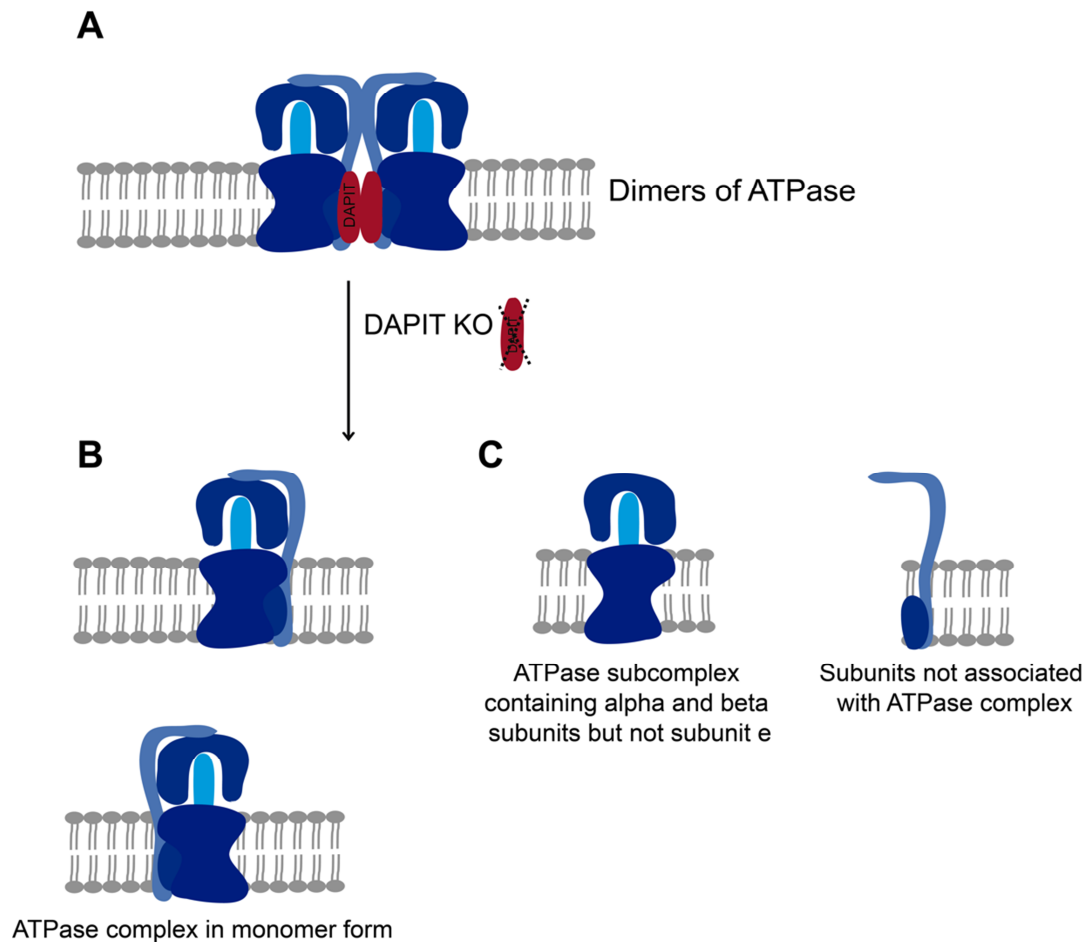
ATPase dimers have been proposed to form the mPTP (Giorgio *et al.*, 2013). Because the lack of DAPIT is a specific way to analyse the ATPase dimer function, we presently report that the absence of these dimers does not alter mPTP properties according to mitochondrial swelling measurements. The capacity of mPTP opening following calcium induction remained even in the mitochondrial DAPIT KO. In the future, these results could be verified by more specific experiments, including calcium retention capacity measurements to calculate the calcium amount threshold that triggers mPTP opening.

Interestingly, the ATPase monomers migrate differently under native conditions between WT and DAPIT KO mitochondria. Migration was higher in KO mitochondria than in WT one, indicating a small size or different conformation of ATPase monomer that alters the velocity of migration.

Apart from the absence of ATPase dimers in the DAPIT KO mitochondria, a new subcomplex was detected. We describe this complex as an association of subunits  $\alpha$ ,  $\beta$ , and  $\gamma_1$  from  $F_1$ , and subunits  $c$  from  $F_0$ . We defined the identity of this subcomplex based on its size, and also by the presence or absence of different subunits tested by Western blot, which indicates that the alpha and beta subunits are contained in it, but with no  $\epsilon$  subunit. Interestingly, a similar complex was described in the Rho0 cells. In these cells, without mtDNA, ATPase is unable to form dimers, but is found in a monomeric form and also in other subcomplexes (Nijtmans *et al.*, 1995). One of them, called  $V^*$ , has a similar weight as the subcomplex detected in the DAPIT KO mitochondria. Proteomic assays would be the best way to fully characterise the identity of this complex. Analysing its protein composition would be the determinant result to ensure how the subcomplex is generated through lack of DAPIT.

DAPIT was not detected in the proteomic analysis of ATPase from the Rho0 cells. This result was explained as a technical limitation (Wittig *et al.*, 2010). Nonetheless, we did not detect DAPIT when using the specific antibody, which could indicate that it is very likely to be absent in the ATPase of these cells. Lack of mtDNA that encodes subunits A6L and ATP8 plays a role as it affects the association of DAPIT with the rest of the ATPase. Interestingly, immunoprecipitation was carried out and demonstrated that the ATPase complex from the DAPIT KO mitochondria presented all its subunits, except DAPIT itself. This evidence led to affirm that the phenotype of absence of dimers observed in Rho0 cells could be caused by the absence of DAPIT. This reveals that DAPIT could be one of the last proteins to link to the ATPase complex, and that its association to ATPase depends on the presence of other subunits. This perfectly matches the fact that DAPIT is absolutely necessary for ATPase dimer formation.

The following model was proposed by considering all these results together (Fig. 61). In the presence of the DAPIT protein, ATPase is found in dimer association (Fig. 61A). However, this link is not possible when DAPIT is absent, and ATPase is found in a monomer form (Fig. 61B) and in addition, the monomeric ATPase is disestablished and generates a subcomplex (Fig. 61C).



**Figure 61. Schematic model showing the consequences of lack of the DAPIT protein. A.** ATPase dimers are formed in the presence of DAPIT. **B.** ATPase, in a monomer form, results from lack of DAPIT. **C.** The ATPase subcomplex forms in the absence of DAPIT that contains alpha and beta subunits, but not subunit e.

Recently, a new domain in ATPase has been described, the intermembrane space domain (IMD). The IMD may, or may not, be associated with the rest of ATPase, which determines two compositional states for IMD (Fischer *et al.*, 2017). These two states do not affect the 43° curvature of the IMM determined by ATPase. It has been suggested that the IMD is formed only by DAPIT and MP68. This indicates that DAPIT and MP68 could be essential for determining the ATPase state by acting either together or individually. Furthermore, the absence of these proteins would not affect the inner mitochondrial morphology.

Our results in an *in vivo* model were surprising if we consider previously published results in cells (Ohsakaya *et al.*, 2011; Kontro *et al.*, 2015). It has been reported that low DAPIT levels in cells affects the amount of ATPase in mitochondria. However, this was not the case in the DAPIT KO mice.

DAPIT signifies “diabetes-associated protein in insulin-sensitive tissues” and a role in glucose metabolism has been proposed. The generation of the DAPIT KO mouse colony was valuable to study the role of the DAPIT protein in diabetes, as previously described. DAPIT KO mice were submitted to a glucose tolerance test experiment, in which they did not show any defect in terms of clearing glucose from blood. Moreover, they responded in the same way as the WT mice responded to insulin administration. Taken together, these results suggest that lack of DAPIT does not affect glucose metabolism in the studied contexts. It would be interesting to induce diabetes in DAPIT KO mice, follow its evolution and determine through experimentation whether the DAPIT protein is required for adaptation in these circumstances.

In this thesis, several deficiencies in the cardiac function of DAPIT KO mice were found. A blood return was observed in the valves: pulmonary, aortic, tricuspid or mitral, with more frequency in DAPIT KO than in WT mice. Moreover, MRI revealed a ventricle stroke volume increased in the DAPIT KO hearts that generates impaired contraction of the heart in the DAPIT KO mice. Lack of the DAPIT protein somehow affects heart functionality, but how DAPIT affects this function is still unknown. In line with this, recently, it was published that the lack of DAPIT in zebrafish was associated with the development of dilated cardiomyopathy since the fishes with reduced DAPIT levels develop larger pericardial sac and atrial areas and reduced ventricular fractional shortening than the WT fishes (Nagata *et al.*, 2017). The relationship between DAPIT and cardiac defects was also suggested by a mouse model of congenital myopathy. This cardiomyopathy was caused by the lack of *Klh131*, a skeletal muscle specific gene, and it was found that *Dapit* gene is one the most upregulated genes (Papizan *et al.*, 2017). Similarly, but in skeletal muscle, DAPIT was observed as the protein that increased most its levels in a pig model of Duchenne muscular dystrophy (Fröhlich *et al.*, 2016).

The DAPIT KO mice are perfectly viable and fertile. These mice are able to live without ATPase dimers, which indicates that ATPase complex organisation is not essential for life. However, it could be relevant in other contexts not explored herein, such as exercise conditions or ageing. Moreover, the DAPIT protein is conserved in metazoan during evolution, which well indicates its relevant role. These suggest that other DAPIT functions remain hidden whose exploration is pending.

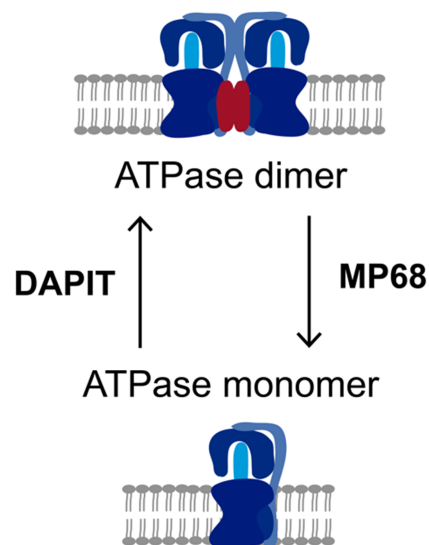
Considering the other subunit that forms the IMD, MP68, a mouse model to study the function of this protein was developed. Using this model, a key role in dimer formation regulation was demonstrated. The absence of MP68 promotes the dimerisation of ATPase in both the heart and liver mitochondria.

Moreover, the migration pattern of ATPase under native conditions was also different when comparing the MP68 WT and KO mitochondria. This fact could be explained by not only lack of MP68, but also by the change in ATPase complex morphology that occurred when MP68 was not present in ATPase.

As both proteins form the IMD, and given the faster migration pattern of monomeric ATPase in both mouse models, our results suggest that changes in IMD composition could alter ATPase conformation.

The lack of MP68 does not affect to the localisation of DAPIT in the ATPase complex. Interestingly the opposite was also true; in DAPIT KO mitochondria, MP68 remains in ATPase complex when DAPIT is absent. This fact indicates that our mouse models are very specific and this allows to study the lack of only one of these proteins.

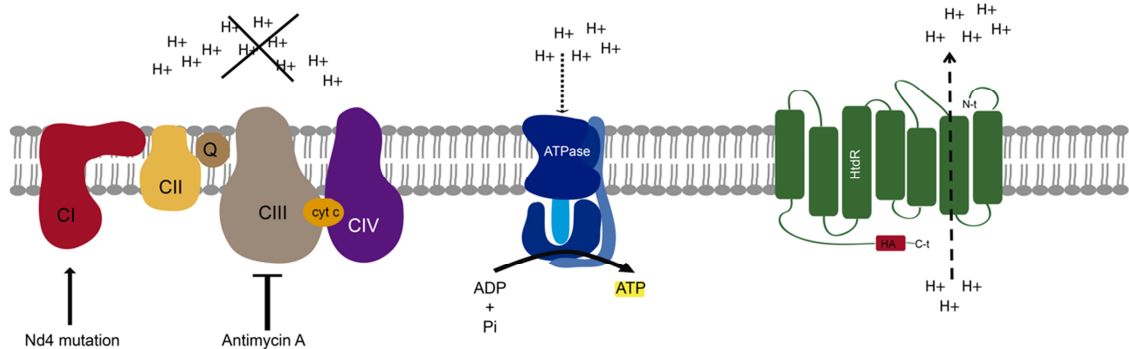
Although both proteins form part of the IMD, the absence of each one leads to an opposite phenotype regarding ATPase dimerisation (Fig. 62). A structural analysis in both mutants could clarify how the absence of these proteins affects the ATPase structure and its dimerisation.



**Figure 62. Schematic representation of the role of DAPIT and MP68 in ATP synthase dimerisation.**

## II. DELTARHODOPSINE

In this work, we demonstrated that light can be used by mammalian cells to obtain energy. This was achieved by the expression of the  $H^+$  proton-pumping protein HtdR in the IMM. This protein uses the energy of photons to translocate protons to the intermembrane space, which is used by ATPase to produce ATP from ADP in the absence of functional glycolysis and the ETC (Fig. 63). This is a proof of concept that mammalian cells can use light as a source of energy.



**Figure 63. Working model that describes the effect of HtdR expression on mammalian cells under electron transport chain (ETC) dysfunction.** ETC inhibition, due to an antimycin effect, is unable to pump protein to the mitochondrial IS. The HtdR is able to restore the proton gradient, which can be used by ATP synthase to produce ATP.

This is also a very interesting tool to characterise ETC deficiencies. There are some mitochondrial mutations in which it is unclear whether the phenotype is caused by lack of energy or by other secondary mutation effects (Kirches, 2011). This tool could be used in the future to determine the mechanisms of these diseases.

The use of cell-based robots that can be controlled by the use of light has been recently reported (Park *et al.*, 2016). However, this device needs glucose media as a source of energy. The possibility of combining this robot with HtdR herein reported could increase the independency of these devices by enabling them to obtain their own energy.

There is increasing evidence that metabolism is not only the way cells obtain energy, but can also play a role in cell specification and cellular differentiation (Chung *et al.*, 2007, 2010; Tormos *et al.*, 2011). Using this tool to manipulate the energy status of the cell can provide new insights into the role of metabolism.

In conclusion, this work represents the proof of concept that mammalian cells can use light energy to grow, and a powerful tool to manipulate metabolism.



# Conclusions



1. Absence of DAPIT does not affect the viability of mice
2. DAPIT determines dimerisation of the ATPase complex and lack of DAPIT does not allow the formation of ATPase dimers
3. Absence of DAPIT generates an ATPase subcomplex that probably contains the subunits  $\alpha$ ,  $\beta$ , and  $\gamma_1$  from  $F_1$ , and the c subunits from  $F_0$
4. The ATPase complex phenotype observed in Rho0 cells could be caused by lack of the DAPIT protein in these cells
5. DAPIT deficit does not affect mPTP opening
6. ATPase dimers are not the mPTP
7. DAPIT deficiency in mice does not affect glucose metabolism
8. Lack of MP68 promotes ATPase dimer formation
9. HtdR-expressing cells can use light energy to grow



# Conclusiones



1. La ausencia de DAPIT asociado a la ATPasa no afecta a la viabilidad del ratón.
2. DAPIT determina la dimerización del complejo ATPasa y su ausencia no permite la formación de dímeros de ATPasa.
3. La ausencia de DAPIT genera subcomplejos de ATPasa formados probablemente por las subunidades  $\alpha$ ,  $\beta$ , y  $\gamma_1$  del dominio  $F_1$ , y las subunidades c del dominio  $F_0$ .
4. El fenotipo de organización del complejo ATPasa observado en células Rho0 puede deberse a la ausencia de DAPIT en dichas células.
5. La falta de DAPIT en la célula no afecta a la apertura del mPTP
6. Los dímeros de ATPasa no forman el mPTP.
7. La ausencia de DAPIT en ratones no afecta al metabolismo de la glucosa.
8. La ausencia de MP68 promueve la formación de dímeros de ATPasa.
9. Las células que expresan HtdR son capaces de usar la luz para crecer.



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