Mitofusin 2 is required to maintain mitochondrial coenzyme Q levels

Arnaud Mourier,1 Elisa Motori,1 Tobias Brandt,2 Marie Lagouge,1 Ilian Atanassov,1 Anne Galinier,3 Gunter Rappl,4,5 Susanne Brodesser,6 Kjell Hultenby,7 Christoph Dieterich,1 and Nils-Göran Larsson1,7

1Max Planck Institute for Biology of Ageing, 50931 Cologne, Germany
2Max Planck Institute of Biophysics, 60438 Frankfurt, Germany
3STROMALab, UMR Université Paul Sabatier/Centre National de la Recherche Scientifique 5273, Institut National de la Santé et de la Recherche Médicale U1031, BP 84 225–F-31 432, Toulouse, France
4Department of Internal Medicine, University Hospital Cologne, and 5Center for Molecular Medicine Cologne (CMMC), University of Cologne, 50931 Cologne, Germany
5CECAD Research Center, Lipidomics Facility, University of Cologne, 50931 Cologne, Germany
6Department of Laboratory Medicine, Karolinska Institutet, Stockholm, Sweden
7Max Planck Institute for Biology of Ageing, 50931 Cologne, Germany

Mitochondria form a dynamic network within the cell as a result of balanced fusion and fission. Despite the established role of mitofusins (MFN1 and MFN2) in mitochondrial fusion, only MFN2 has been associated with metabolic and neurodegenerative diseases, which suggests that MFN2 is needed to maintain mitochondrial energy metabolism. The molecular basis for the mitochondrial dysfunction encountered in the absence of MFN2 is not understood. Here we show that loss of MFN2 leads to impaired mitochondrial respiration and reduced ATP production, and that this defective oxidative phosphorylation process unexpectedly originates from a depletion of the mitochondrial coenzyme Q pool. Our study unravels an unexpected and novel role for MFN2 in maintenance of the terpenoid biosynthesis pathway, which is necessary for mitochondrial coenzyme Q biosynthesis. The reduced respiratory chain function in cells lacking MFN2 can be partially rescued by coenzyme Q10 supplementation, which suggests a possible therapeutic strategy for patients with diseases caused by mutations in the Mfn2 gene.

Introduction

Mammalian Mitofusin 1 (MFN1) and Mitofusin 2 (MFN2) belong to the GTPase family of proteins (Bourne et al., 1990) and are required for mitochondrial outer membrane fusion (Rojo et al., 2002). Ubiquitous knockout of the Mfn1 or Mfn2 gene results in embryonic lethality during midgestation (Chen et al., 2003) due to placental dysfunction (Chen et al., 2003, 2007). Remarkably, by using a conditional knockout allele in conjunction with cre-recombinase expression only in the embryo, Mfn1 or Mfn2 knockout mice are born alive (Chen et al., 2007). Mice ubiquitously lacking MFN1 are apparently healthy, whereas mice without MFN2 die in the early postnatal period and show severe defects in movement and balance (Chen et al., 2007). Experiments performed on immortalized Mfn1 and Mfn2 knockout mouse embryonic fibroblasts (MEFs) have shown that heterooligomeric complexes between MFN1 and MFN2 are important for control of mitochondrial fusion (Detmer and Chan, 2007). Despite its well-established role in mitochondrial fusion, a growing body of evidence suggests that MFN2 has additional functions, such as ER–mitochondria tethering (de Brito and Scorrano, 2008) and control of mitochondrial transport in axons (Misko et al., 2010), which may explain some of the disease manifestations in humans with Mfn2 mutations (Züchner et al., 2004; Detmer et al., 2008). The different functions of MFN2 are supported by in vivo studies in mice, showing that MFN2 is required for normal glucose homeostasis (Sebastián et al., 2012), and is essential for axonal projections of dopaminergic neurons (Lee et al., 2012; Pham et al., 2012). However, the molecular basis for the mitochondrial dysfunction, caused by loss of MFN2, remains to be established. Here we report the unexpected finding that loss of MFN2 causes depletion of mitochondrial coenzyme Q, which,
oxidative phosphorylation (OXPHOS) in heart, a typical highly energy-demanding tissue. Heart conditional knockouts for \(\text{Mfn1}\) or \(\text{Mfn2}\) were born at Mendelian proportions and had no enlargement of the heart when assessed at the age of 20 wk (Fig. S1, A and B), which is consistent with results from independent knockout strains (Papanicolaou et al., 2011, 2012). Furthermore, mitochondrial DNA (mtDNA) levels (Fig. S1 C) and mitochondrial mass were not affected in \(\text{Mfn1}\) and \(\text{Mfn2}\) knockout hearts, as determined by electron microscopy analysis (Fig. S1 D).

The absence of changes in mitochondrial content prompted us to analyze mitochondrial activity at the single-cell level. To this end, we performed retrograde perfusion to isolate adult cardiomyocytes from control and \(\text{Mfn1}\) and \(\text{Mfn2}\) knockout hearts expressing a mitochondrial targeted YFP. Confocal microscopy analysis of cardiomyocytes isolated from \(\text{Mfn1}\) knockout hearts showed that the mitochondria were homogeneous in shape and organized in a classical linear arrangement between myofibrils, in turn, leads to respiratory chain dysfunction. A global reduction in the terpenoid synthesis pathway is present in both \(\text{Mfn2}\) and \(\text{Mfn1}\) conditional knockout mice, but the defect is more severe and impacts respiratory chain function only in \(\text{Mfn2}\) knockout mice. Moreover, addition of coenzyme Q to immortalized \(\text{Mfn2}\) knockout MEFs or isolated mitochondria partially rescues the respiratory chain dysfunction, which suggests a therapeutic strategy for intervention in affected patients.

**Results**

**MFN2 is required to maintain mitochondrial morphology and function in cardiomyocytes**

To further investigate the role of mitofusins in energy metabolism, we performed a detailed characterization of how loss of MFN1 and MFN2 affects mitochondrial morphology and oxidative phosphorylation (OXPHOS) in heart, a typical highly energy-demanding tissue. Heart conditional knockouts for \(\text{Mfn1}\) or \(\text{Mfn2}\) were born at Mendelian proportions and had no enlargement of the heart when assessed at the age of 20 wk (Fig. S1, A and B), which is consistent with results from independent knockout strains (Papanicolaou et al., 2011, 2012). Furthermore, mitochondrial DNA (mtDNA) levels (Fig. S1 C) and mitochondrial mass were not affected in \(\text{Mfn1}\) and \(\text{Mfn2}\) knockout hearts, as determined by electron microscopy analysis (Fig. S1 D).

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Mfn2 knockout hearts exhibited atypical morphology with bud formation (Fig. 2 A). In addition, significant fractions of abnormally small (diameter <300 nm) or large (diameter >1.5 µm) mitochondria were observed in Mfn2 knockout hearts (Figs. 2 A and S2 A). The functional defects observed in single cells were further quantified by FACS analysis of isolated heart mitochondria after double staining with MitoTracker green and TMRM, which revealed a twofold increase of Δψ-negative mitochondria in Mfn2 knockout in comparison with Mfn1 knockout or control hearts (Fig. 2, B and C).

Loss of MFN2 impairs the mitochondrial respiratory chain capacity

The increase of Δψ-negative mitochondria in hearts lacking MFN2 prompted us to characterize the OXPHOS system properties. To this end, isolated heart mitochondria were incubated with respiratory substrates whose metabolism results in delivery of electrons at the level of complex I (pyruvate, glutamate, malate) or complex II (succinate + rotenone), and the oxygen consumption rate was recorded in the phosphorylating (ADP and Pi), nonphosphorylating (oligomycin), and uncoupled state (carbonyl cyanide m-chlorophenyl hydrazone [CCCP]).

![Figure 2](image_url)

Figure 2. Loss of MFN2 results in mitochondrial heterogeneity and dysfunction. (A) Electron cryo-tomography analysis of isolated mitochondria. Slices through typical tomographic volumes and example segmented 3D representations from control, Mfn1, and Mfn2 knockout heart mitochondria are shown (blue, cristae; transparent gray, IMS; outer membrane omitted for clarity). Bars, 250 nm. (B) Representative analysis out of 14 FACS analyses of isolated control heart mitochondria incubated with succinate and rotenone, and stained with MitoTracker green FM and TMRM. FACS analyses were performed using the FlowJo software, applying a biexponential transformation to digital data. (C) FACS analysis of control (white bars, n = 5 independent experiments), Mfn1 knockout (gray bars, n = 4 independent experiments), and Mfn2 knockout (black bars, n = 5 independent experiments) heart mitochondria to determine the percentage of membrane potential-negative (green) mitochondria in the presence of succinate and rotenone. Error bars indicate ±SEM. ***, P < 0.001.
were severely decreased in Mfn2 knockout heart mitochondria in comparison with controls (Fig. 4 B). The coenzyme Q isoforms are organic antioxidant compounds present in all cellular membranes and are specifically required in the mitochondrial inner membrane to transfer electrons from complex I and II to complex III of the respiratory chain. To gain further insight into the potential consequences of the observed coenzyme Q deficiency, we measured the NADH–cytochrome c reductase (complex I-III) and succinate-cytochrome c reductase (complex II-III) activities, as they are dependent on the endogenous coenzyme Q pool. Interestingly, both of these respiratory chain enzyme activities were impaired in Mfn2 knockout heart mitochondria (Fig. 4 C). Moreover, the succinate cytochrome c reductase deficiency observed in isolated mitochondria from Mfn2 knockout hearts could be partially rescued by the addition of the artificial quinone decylubiquinone (Fig. 4 D). The rescue of the succinate cytochrome c reductase deficiency was comparable to the rescue seen after decylubiquinone treatment of cell lines from patients with genetic defects impairing the synthesis of coenzyme Q10 (Mollet et al., 2007). Reactive oxygen species (ROS) production was assessed by measuring H₂O₂.

We observed a clear decrease in respiration capacity under phosphoryllating and uncoupled conditions in Mfn2 knockout heart mitochondria at 20 (Fig. 3 A) and 60 wk of age (Fig. S2 B). In agreement with the reduced respiratory chain capacity, the ATP production rate was also affected in the absence of MFN2 (Fig. 3 B). To decipher the molecular origin of the respiratory chain dysfunction, we assessed the enzyme activities and steady-state levels of the respiratory chain complexes. Surprisingly, the respiratory chain complex enzyme activities were not decreased (Fig. 3 C) and the levels of different cytochromes, such as a-a₃ (complex IV), bH-bL (complex III), and c, were normal in Mfn2 knockout heart mitochondria (Fig. S2 C). Also, the levels of different subunits of the OXPHOS system complexes as determined by Western blot analyses were unaffected (Fig. 3 D).

**Mfn2 heart knockout mitochondria display a severe coenzyme Q deficiency**

Because the lipid environment has been shown to impact the respiratory activity, we determined the mitochondrial lipid composition and found no alterations in phospholipid content (Fig. 4 A and S2 D). However, the levels of coenzyme Q9 and Q10 were severely decreased in Mfn2 knockout heart mitochondria in comparison with controls (Fig. 4 B). The coenzyme Q isoforms are organic antioxidant compounds present in all cellular membranes and are specifically required in the mitochondrial inner membrane to transfer electrons from complex I and II to complex III of the respiratory chain. To gain further insight into the potential consequences of the observed coenzyme Q deficiency, we measured the NADH–cytochrome c reductase (complex I-III) and succinate-cytochrome c reductase (complex II-III) activities, as they are dependent on the endogenous coenzyme Q pool. Interestingly, both of these respiratory chain enzyme activities were impaired in Mfn2 knockout heart mitochondria (Fig. 4 C). Moreover, the succinate cytochrome c reductase deficiency observed in isolated mitochondria from Mfn2 knockout hearts could be partially rescued by the addition of the artificial quinone decylubiquinone (Fig. 4 D). The rescue of the succinate cytochrome c reductase deficiency was comparable to the rescue seen after decylubiquinone treatment of cell lines from patients with genetic defects impairing the synthesis of coenzyme Q10 (Mollet et al., 2007). Reactive oxygen species (ROS) production was assessed by measuring H₂O₂.

![Figure 3](image-url)
The coenzyme Q deficiency is caused by multiple terpenoid synthesis defects

To decipher the molecular origin of the coenzyme Q deficiency, we characterized the proteome changes associated with loss of the MFN1 or MFN2 proteins, respectively. To this end, we performed stable isotope labeling by amino acids in cell culture (SILAC) experiments by quantitatively comparing the proteomes of MFN1 and MFN2 knockout MEFs (Fig. 5, A and B; and Fig. S3, B–D) with control MEFs, whose proteome had been saturated with heavy isotope labeled arginine and lysine ($^{13}$C$_6$, $^{15}$N$_2$, l-arginine, $^{13}$C$_6$, $^{15}$N$_4$, l-lysine). Surprisingly, the terpenoid biosynthesis pathway clearly stood out as the unique metabolic process significantly down-regulated in both MFN1 and MFN2 knockout MEFs. The levels of all detected enzymes in the terpenoid biosynthesis pathway, except one of the two acetyl-CoA acetyltransferase isoforms (Acat1), were decreased in comparison with controls (Fig. 5, A and B; and Fig. S3, B–D). In contrast, none of the downstream enzymes acting in the coenzyme Q biosynthesis pathway (Coq2, Coq9) or in the cholesterol and steriodogenesis pathways were significantly affected (Fig. 5, A and B; and Fig. S3 D). Furthermore, qRT-PCR analyses showed that the levels of transcripts encoding enzymes involved in the terpenoid (Fdhps, ggps1, Hmgcr, Hmgcs1, Hmgcs2) and coenzyme Q synthesis pathways (coq2, coq3, coq5, coq7, pdss1, pdss2) were not affected in MFN2 knockout hearts (Fig. 5 C). Thus, the decreased levels of enzymes involved in the terpenoid biosynthesis pathway could not be attributed to changes in their mRNA expression.

To investigate how the global down-regulation of terpenoid synthesis enzyme levels impacted terpenoid biosynthesis, we quantified terpenoid metabolites in MFN1 and MFN2 knockout hearts. The levels of isopentenyl-pyrophosphate (IPP) and p-HBA (unpublished data) were unaltered in MFN1 and MFN2 knockout hearts. However, the levels of dimethylallyl pyrophosphate (DMAP), geranyl diphosphate (GPP), farnesyl diphosphate (FPP), and geranyl-geranyldiphosphate (GGPP) were severely reduced in the MFN2 knockout hearts (Fig. 5 D) and were normal (IPP; DMAP and GPP) or slightly reduced (FPP and GGPP) in the MFN1 knockout hearts (Fig. 4 E). This experiment was performed on heart mitochondria from control (white bars, n = 5 independent experiments), MFN1 (gray bars, n = 5 independent experiments), and MFN2 conditional knockout (black bars, n = 5 independent experiments) animals. (D) Succinate-cytochrome c reductase activity determined in the absence (white and black bars) or presence (gray bars) of an exogenous coenzyme Q analogue (decylubiquinone). The experiment was performed on mitochondria from control (white bars, n = 5 independent experiments) and MFN2 knockout (black bars, n = 5 independent experiments) hearts. [E] Assessment of the hydrogen peroxide production rate in nonphosphorylating conditions in the presence of succinate, rotenone sensitive) was clearly reduced in MFN1 knockout hearts. The proteomic and metabolomic analyses showed that the levels of transcripts encoding enzymes involved in the terpenoid (Fdhps, ggps1, Hmgcr, Hmgcs1, Hmgcs2) and coenzyme Q synthesis pathways (coq2, coq3, coq5, coq7, pdss1, pdss2) were not affected in MFN2 knockout hearts (Fig. 5 C). Thus, the decreased levels of enzymes involved in the terpenoid biosynthesis pathway could not be attributed to changes in their mRNA expression.

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in turn, originates from a global defect of the terpenoid synthesis pathway, as determined by a combined proteomic and metabolomic approach. It should be noted that cultured cells lacking MFN1 or MFN2 typically have a fragmented mitochondrial network. It is thus formally possible that the coenzyme Coenzyme Q10 supplementation partially rescues mitochondrial morphology and function in Mfn2 knockout MEFs.

The loss of MFN2 in cardiomyocytes leads to impaired respiratory chain activity that is caused by CoQ deficiency, which, in turn, originates from a global defect of the terpenoid synthesis pathway, as determined by a combined proteomic and metabolomic approach. It should be noted that cultured cells lacking MFN1 or MFN2 typically have a fragmented mitochondrial network. It is thus formally possible that the coenzyme

![Diagram](image-url)

**Figure 5. Terpenoid synthesis pathway defects in mitofusin knockouts.** (A) Scheme of the mevalonate pathway in mammals. Red, proteins down-regulated according to SILAC-based protein quantification performed with Mfn1 and Mfn2 knockout MEFs; blue, not affected; gray, not detected. *, COQ2 protein level was determined by Western blot analysis. (B) Quantitative proteomic analysis of two independent batches of control and Mfn2 knockout MEFs using SILAC analysis. Negative values indicate down-regulation of the protein in Mfn2 knockout relative to control MEFs. (C) Transcript levels of different enzymes involved in the terpenoid and coenzyme Q synthesis pathways determined by qRT-PCR with total RNA extracted from control (white bars) and Mfn2 knockout (black bars) hearts (n = 5 pairs of mice). (D) Terpenoid metabolites content of whole heart extracts isolated from control (white bars, n = 12 samples), Mfn1 knockout (gray bars, n = 7 samples), and Mfn2 knockout (black bars, n = 6 samples). Error bars indicate ±SEM. *, P < 0.05; **, P < 0.01; ***, P < 0.001.
Q deficiency originates from impaired mitochondrial fusion as such rather than from a specific role for MFN2. To distinguish between these possibilities, we investigated the connection between coenzyme Q homeostasis and mitochondrial network morphology in Mfn1 and Mfn2 knockout MEFs. We first grew MEFs in glycolytic (glucose) and oxidative (galactose) medium and observed a growth defect of Mfn2 knockout MEFs, but not control or Mfn1 knockout MEFs, in galactose medium (Fig. S4, A–D). The cell growth phenotype suggests that loss of MFN2 causes a respiratory chain deficiency and, consistent with this observation, we observed a pronounced respiration defect in Mfn2 knockout MEFs grown on galactose (Fig. 6, A and B). The defective mitochondrial respiration in permeabilized Mfn2 knockout MEFs was associated with a deficiency of coenzyme Q9 and Q10 in both isolated mitochondria (Fig. 7 C) and whole cells (Fig. 6 D). To investigate how impaired terpenoid biosynthesis impacts respiratory chain function and mitochondrial morphology, we treated control MEFs with mevastatin (MEV), a well-known inhibitor of the terpenoid biosynthetic pathway. MEV treatment of wild-type MEFs lead to a profound reduction of coenzyme Q levels and impaired mitochondrial respiration, thus closely mimicking the bioenergetic phenotype of Mfn2 knockout MEFs (Fig. 6, A and D). However, the MEV treatment did not affect the mitochondrial network morphology of control cells (Fig. 7, A, C, and D), thus showing that the coenzyme Q deficiency as such does not induce profound alterations of mitochondrial dynamics. As the opposite might still be true, we investigated whether the coenzyme Q deficiency could be rescued by restoring a tubular mitochondrial network in MEFs lacking MFN2. To this end, we knocked down Drp1, which is involved in mitochondrial fission, in control MEFs and found a slight increase in mitochondrial elongation in comparison with MEFs transfected with a scrambled siRNA (Fig. S5 A), whereas the fragmented mitochondrial network of Mfn2 knockout MEFs

Figure 6. OXPHOS dysfunction in the absence of Mfn2 is due to coenzyme Q deficiency. (A) Assessment of cellular respiration in control (white bars, n = 5 independent experiments), Mfn1 knockout (gray bars, n = 5 independent experiments), Mfn2 knockout (black bars, n = 5 independent experiments), and MEV-treated control (light gray bars, n = 5 independent experiments) MEFs that were grown in DMEM glucose, permeabilized, and analyzed in the presence of succinate and glycerol-3-phosphate. (B) Assessment of cellular respiration in control (white bars, n = 5 independent experiments), Mfn1 knockout (gray bars, n = 5 independent experiments), and Mfn2 knockout (black bars, n = 5 independent experiments) MEFs that were grown in DMEM galactose, permeabilized, and analyzed in the presence of succinate and glycerol-3-phosphate. Dashed bars indicate cells of the same genotypes grown in the presence of coenzyme Q10 (n = 5 per genotype). (C) Quinone quantification performed on mitochondria isolated from control (white bars, n = 5 independent experiments), Mfn1 knockout (gray bars, n = 5 independent experiments), and Mfn2 knockout (black bars, n = 5 independent experiments) MEFs. (D) Quinone quantification performed on whole cell extracts isolated from control (white bars, n = 5 independent experiments), Mfn1 knockout (gray bars, n = 5 independent experiments), Mfn2 knockout (black bars, n = 5 independent experiments), and MEV-treated control (black bars with white dots, n = 5 independent experiments) MEFs grown in DMEM glucose. (E) Quinone quantification performed on whole cell extracts isolated from control (white bars, n = 5 independent experiments), Mfn1 knockout (gray bars, n = 5 independent experiments), and Mfn2 knockout (black bars, n = 5 independent experiments) MEFs grown in DMEM galactose. Dotted bars indicate cells of the same genotypes grown in the presence of coenzyme Q10 (n = 5 independent experiments). Error bars indicate ±SEM. ***, P < 0.001.
cellular respiration (Fig. 6 B) and the quinone deficiency (Fig. 6 E) in MEFs lacking MFN2. As previously reported (Chen et al., 2003), the absence of MFN2 causes a stochastic loss of $\Delta \psi$, as depicted by the strong variation in the TMRM staining (Fig. 7 B). Strikingly, we observed that supplementation of the culture medium with coenzyme Q could partially rescue the TMRM heterogeneity (Fig. 7, B and C; and Fig. S5, E and F). This partial rescue was also accompanied by a partial rescue of mitochondrial network morphology induced by Drp1 knockdown in Mfn2 knockout MEFs causes no changes of coenzyme Q content (Fig. S5, B and D), thus demonstrating that the coenzyme Q deficiency is not directly related to the mitochondrial network morphology. Interestingly, supplementation of the culture medium with coenzyme Q10 partially rescued the cellular respiration (Fig. 6 B) and the quinone deficiency (Fig. 6 E) in MEFs lacking MFN2. As previously reported (Chen et al., 2003), the absence of MFN2 causes a stochastic loss of $\Delta \psi$, as depicted by the strong variation in the TMRM staining (Fig. 7 B). Strikingly, we observed that supplementation of the culture medium with coenzyme Q could partially rescue the TMRM heterogeneity (Fig. 7, B and C; and Fig. S5, E and F). This partial rescue in $\Delta \psi$ was also accompanied by a partial rescue of
the mitochondrial network morphology, as indicated by the appearance of more elongated mitochondria (Fig. 7, B and D). Importantly, this rescue in mitochondrial morphology was not observed when Mfn1 knockout MEFs were supplemented with coenzyme Q10, which additionally supports the finding that Mfn2 has a distinct role in coenzyme Q pool maintenance.

**Discussion**

Mammalian Mfn1 and Mfn2 have originally been described as playing a central role in mitochondrial outer membrane fusion to control mitochondrial network morphology and allow the exchange of content between adjacent mitochondria. The role for Mfn2 in mammalian mitochondria has attracted much interest because mutations in Mfn2 cause the Charcot Marie Tooth type 2A (CMT2A) axonopathy (Züchner et al., 2004; Detmer et al., 2008). Recently, Mfn2 has been shown to be essential for axonal mitochondrial transport (Misko et al., 2010) and to maintain the axonal projections of dopaminergic neurons (Lee et al., 2012; Pham et al., 2012). Beyond its role in mitochondrial fusion, Mfn2 has also been implicated in ER–mitochondria tethering (de Brito and Scorrano, 2008) and in ER stress caused by mitochondrial dysfunction (Sebastián et al., 2012). However, the molecular basis for the mitochondrial dysfunction caused by loss of Mfn2 has been unclear. We report here that mouse hearts or MEFs lacking Mfn2 are respiratory chain deficient despite having normal levels and normal enzyme activities of individual respiratory chain complexes, which is consistent with previous studies of fibroblasts from CMT2A patients (Loiseau et al., 2007; Guillet et al., 2010). Unexpectedly, we show here that the deficient respiratory chain function in the absence of Mfn2 is explained by a deficiency of coenzyme Q. Loss of Mfn2 is associated with a fragmentation of the mitochondrial network, but this alteration is likely not the cause of the coenzyme Q deficiency because normalization of the mitochondrial network morphology in Mfn2 knockout MEFs does not normalize coenzyme Q levels (Fig. S5 G).

A continuous cellular supply of coenzyme Q requires de novo synthesis from the terpenoid pathway, a multistep process that has been reported to involve many subcellular compartments (Szkopinska, 2000). Whereas the condensation of the long isoprenoid side chain to the para-hydroxybenzoic acid (PHB) can occur in mitochondria (Tran and Clarke, 2007; Wang and Hekimi, 2013), the exact subcellular localization of the terpenoid and coenzyme Q biosynthetic pathways as well as the mechanisms that govern the intracellular transport of their metabolites remain to be defined (Szkopinska, 2000). Our analyses shed important new light on these molecular pathways, and we show here that loss of Mfn1 and Mfn2 is associated with a down-regulation of both levels of enzymes and metabolites of the terpenoid biosynthesis pathway. The enzymes of the terpenoid biosynthesis pathway have been found in different organelles such as mitochondria, ER, and peroxisomes (Keller et al., 1986; Olivier and Kribs, 2000; Kovacs et al., 2001). Interestingly, Mfn2 was previously reported to be key in high-level steroid production of trophoblast and Leydig cells (Duarte et al., 2012, 2014; Wasilewski et al., 2012). Our results suggest that the previously observed steroidogenesis defects of trophoblast and Leydig cells lacking Mfn2 are likely to originate from a deficiency in the terpenoid synthesis pathway.

In summary, we report here the novel and unexpected finding that Mfn2, an outer mitochondrial membrane protein, has an essential role in regulating the terpenoid biosynthesis pathway, which, in turn, is required to maintain mitochondrial coenzyme Q levels for optimal function of the respiratory chain. We show that the coenzyme Q deficiency and the respiratory chain deficiency can be partially rescued by coenzyme Q10 supplementation. This finding may well be of direct clinical importance because mutations in Mfn2 are known to cause human disease. Indeed, a recently published case study indicates that coenzyme Q supplementation may have a positive effect in patients with pathogenic Mfn2 mutations (Takahashi et al., 2012). Our results suggest that clinical trials to strainently evaluate the effects of coenzyme Q10 supplementation in patients with diseases caused by Mfn2 mutations may be warranted.

**Materials and methods**

**Creation of Mfn1 and Mfn2 knockout mice**

Mice with a loxP-flanked exon 3 of the Mfn1 gene or a loxP-flanked exon 5 of the Mfn2 gene were generated in the C57BL/6N background as described previously (Lee et al., 2012) and crossed with mice expressing cre recombinase in heart and skeletal muscle (+/Ckmm-cre). The resulting double heterozygous offspring (+/Mfn1lox-/+; +/Ckmm-cre and +/Mfn2lox-/+; +/Ckmm-cre) were crossed with mice having homozygous loxP-flanked alleles (Mfn1lox+/Mfn1lox- and Mfn2lox+/Mfn2lox-) to generate tissue-specific knockouts for Mfn1 (Mfn1lox+/Mfn1lox-; +/+Ckmm-cre) and Mfn2 (Mfn2lox+/Mfn2lox-; +/+Ckmm-cre). In some crosses, an allele allowing cell-type-specific expression of mitochondria-targeted YFP from the ROSA26 locus (Gt(ROSA26Sor<+>laczDepressor-P)2FLmpy, Sterky et al., 2011) was introduced.

**Electron cryo-microscopy**

For analysis by electron cryo-microscopy, mitochondria were washed twice with 320 mM trehalose, 20 mM Tris, pH 7.3, and 1 mM EGTA. Samples were mixed 1:1 with fiducial gold markers (10 nm gold particles conjugated to protein A; Aurion), blotted, and immediately plunged-frozen in liquid ethane on holey carbon grids (Quantifoil Micro Tools). Single tilt series (±60°, step size 1.5°) were collected on a Polara microscope (300 kV; FEI) using an Ultrascan 4 × 4k CCD (Gatan, Inc.) and a post-column Quantum energy filter (Gatan, Inc.) at −90 µm defocus. The nominal magnification was 34,000×, resulting in a pixel size of 6 Å. A total dose of ~130 e⁻/Å² was used. Tilt series alignment using the gold fiducial markers and tomogram reconstruction by back-projection were performed using the IMOD software package (Kremer et al., 1996). To increase contrast, a final filtering step applying nonlinear anisotropic diffusion (Franzak and Hegerl, 2001) was performed. Manual segmentation was performed with the program Amira (FEI).

**Isolation of cardiomyocytes**

To isolate adult cardiomyocytes, 15-wk-old mice were sacrificed by cervical dislocation, and the aorta was quickly clamped to the perfusion cannula. Using a Radnoti heart perfusion apparatus, the heart was first washed (1.5 mL/min) for 3 min with the perfusion buffer (120 mM NaCl, 4.8 mM KCl, 1.25 mM KH₂PO₄, 1.25 mM MgSO₄, 11 mM glucose, 20 mM Hepes, and 30 mA, pH 7.2). Then the heart was perfused for 3 min with the calcium-free perfusion buffer, and 240 U/mL collagenase type II (EMD Millipore) was added for ~30 min digestion. During digestion, the calcium concentration was progressively increased to 250 µM. To stop the digestion, the heart, which was flaccid and looked pale, was disrupted in perfusion buffer (250 µM calcium, 0.5 mg/ml BSA). After several rounds of washing by sedimentation, the isolated cardiomyocytes, which had a mostly rod-shaped morphology, were seeded at 70–80% density in culture medium (DMEM F12 without phenol red [Invitrogen], 1 µg/mL insulin, 0.55 µg/mL transferrin, 0.65 µg/ml selenium, 2 mM glutamine, 10 mM Hepes, 1% penicillin/streptomycin, and 0.05% BSA). Plating was...
performed on 23-mm glass-bottomed dishes (World Precision Instruments) and cardiomyocytes were left to attach for 2 h at room temperature before confocal imaging. To avoid spontaneous contraction of cells, 25 µM blebbistatin was added.

Live imaging of adult cardiomyocytes and MEFs

An inverted laser-scanning confocal microscope (Leica) equipped with a white light laser. Adult cardiomyocytes derived from mice expressing a mitochondrially targeted YFP were first equilibrated 30 min at 37°C, 5% CO₂, then loaded with 50 nM TMRM (Invitrogen) in culture medium for 30 min at 37°C. Fluorescence images were acquired with a 100x objective lens (HCX Plan-Apochromat CS 100x oil, 1.46 NA), using the LAS AF software. The following parameters in sequential line scan were used: YFP (excitation, 488 nm; emission, 512–552 nm; image size 1,024 x 1,024 pixels, scan speed 200 Hz) and TMRM (excitation, 550 nm; emission, 595–670 nm; image size 1,024 x 1,024 pixels, scan speed 200 Hz). For TMRM labeling in MEFs, cells were first loaded with 100 nM Mitotracker Green (Invitrogen) for 15 min at 37°C, 5% CO₂, then washed twice with culture medium to remove the probe in excess. Finally, MEFs were loaded with 20 nM TMRM for 30 min at 37°C before imaging. Images were acquired using the following parameters: Mitotracker Green (excitation, 488 nm; emission, 504–543 nm; image size 1,024 x 1,024 pixels, scan speed 400 Hz) and TMRM (excitation, 550 nm; emission, 590–668 nm; image size 1,024 x 1,024 pixels, scan speed 400 Hz). During acquisition, the laser power was kept as low as possible to reduce photo-toxicity; a 2x averaging of the optical plane was also applied to increase the signal-to-noise ratio. Transmission light images were acquired with a scan-BF detector and a 488-nm excitation laser line.

Imaging and analysis of mitochondrial potential in adult cardiomyocytes. The fraction of inactive mitochondria was quantified using the JC10pl plugin (Bole and Conibares, 2006) from ImageJ by calculating analysis of YFP signal (corresponding to the overall mitochondrial population) and TMRM signal (corresponding to the active mitochondria). The fraction of YFP signal overlapping the TMRM signal was expressed with the Manders coefficient. The intensity profile analysis was obtained by measuring the maximum fluorescence intensity of a segmented line drawn along mitochondria using ImageJ.

Analysis of mitochondrial potential in MEFs. The heterogeneity in TMRM and Mitotracker Green channels were calculated for their difference by using the equation signal = |TMRM-MitoTracker Green|. The pixel area in the resulting difference mask was then quantified and subsequently normalized over the corresponding thresholded single-stack image of the Mitotracker Green channel in order to obtain the fraction of mitochondria displaying heterogeneous membrane potential.

Analysis of mitochondrial morphology in MEFs. Classification of the mitochondrial morphology was performed according to the following criteria: cells displaying an intact tubular mitochondrial network were classified as tubular; cells with globular or rod-like mitochondria were classified as fragmented; cells showing equal presence of both tubular and fragmented mitochondria were instead classified as intermediate.

Image processing. Image panels were assembled with Photoshop (Adobe); no digital manipulation was applied except for adjustment of brightness and contrast.

Isolation of mitochondria

To isolate intact heart mitochondria, mice were sacrificed by cervical dislocation and hearts were quickly collected in ice-cold DPBS (Gibco) and homogenized with a few strokes of a Potter S homogenizer (Sartorius) in 5 ml of ice-cold mitochondria isolation buffer (MIB; 310 mM sucrose, 20 mM Tris-HCl, and 1 mM EGTA, pH 7.2). Mitochondria were purified by differential centrifugation (1,200 g for 10 min) and supernatants were then centrifuged at 10,000 g for 10 min. The crude mitochondrial pellet was resuspended in an appropriate volume of MIB. The mitochondrial protein concentration was determined using the Protein DC Lowry based assay (Bio-Rad Laboratories, Inc.).

Transmission electron microscopy

Electron micrographs of mitochondria were obtained as previously described (Hansson et al., 2004). In brief, small pieces from the left myocardium were fixed in 2% glutaraldehyde, 0.5% paraformaldehyde, 0.1 M sodium cacodylate, 0.1 M sucrose, and 3 mM CaCl₂, pH 7.4, for 30 min, followed by 24 h at 4°C. Specimens were rinsed in a buffer containing 0.15 M sodium cacodylate and 3 mM CaCl₂, pH 7.4; postfixed in 2% osmium tetroxide, 0.07 M sodium cacodylate, and 1.5 mM CaCl₂, pH 7.4, at 4°C for 2 h; dehydrated in ethanol followed by acetone; and embedded in LX112 (Ladd Research Industries). Ultrathin sections (40–50 nm) from longitudinal parts were cut and examined in a transmission electron microscope (Tecnai 10; FEI) at 80 kV. Digital images at a final magnification of 5,200× were randomly taken on myofibrils from sections of the myocardium. Printed digital images were used and the volume density (Vv) of mitochondria was calculated by point counting using a 2-cm square lattice (Weibel, 1979). A pilot study was performed to determine the number of blocks and images needed for an appropriate sample using a cumulative mean plot for evaluation (Weibel, 1979). Two different methods were used. YC sections were sectioned and 10 random images were collected from each block.

Measurement of mitochondrial respiration

The mitochondrial oxygen consumption flux was measured as described previously (Mourier et al., 2014) at 37°C using 65–125 µg of crude mitochondria diluted in 2.1 ml of mitochondrial respiration buffer (120 mM sucrose, 50 mM KCl, 20 mM Tris-HCl, 2 mM KH₂PO₄, 2 mM MgCl₂, and 1 mM EGTA, pH 7.2) in an Oxygen-graph-2k (Orbostars Instruments). The oxygen consumption rate was measured using either 10 mM pyruvate, 5 mM glutamate, and 5 mM malate or 10 mM succinate and 10 mM rotenone. Oxygen consumption was assessed in the phosphorylating state with 1 mM ADP or in the nonphosphorylating state by adding 2.5 µg/ml oligomycin. In the control mitochondria, the respiratory control ratio (RCR) values were >10 with pyruvate/glutamate/malate and >5 with succinate/rotenone. Respiration was uncoupled by successive addition of CCCP up to 3 µM to reach maximal respiration.

Measurement of respiratory chain enzyme activities

Mitochondrial proteins (15–50 µg) were diluted in phosphate buffer (50 mM KH₂PO₄, pH 7.4) followed by spectrophotometric analysis of isolated respiratory chain complex activities at 37°C using a spectrophotometer (UV-3600, Hitachi). Citrate synthase activity was measured at 412 nm (ε = 13,600 M⁻¹ cm⁻¹) after the addition of 0.1 mM acetyl-CoA, 0.5 mM oxaloacetate, and 1 mM 5,5-dithiobis-2-nitrobenzoic acid (DTNB). NADH dehydrogenase activity was determined at 340 nm (ε = 6220 M⁻¹ cm⁻¹) after the addition of 0.25 mM NADH, 0.25 mM decylubiquinone, and 1 mM KCN, controlling for rotenone sensitivity. Succinate dehydrogenase (SDH) activity was measured at 600 nm (ε = 21,000 M⁻¹ cm⁻¹) after the addition of 40 mM succinate, 35 µM dichloro phenolindophenol (DCIP), and 1 mM KCN. The complex III activity was measured after the reduction of cytochrome c at 550 nm (ε = 18,000 M⁻¹ cm⁻¹) in the presence of 40 mM succinate or 5 mM NADH, 80 µM cytochrome c, and 0.1 mM of sodium azide. The specific complex III activity was defined as the flux difference before and after the addition of 1 µM antimycin A. The NADH and succinate cytochrome c reductase activity was also followed at 540 nm (ε = 18,000 M⁻¹ cm⁻¹) in the presence of 40 mM succinate or 5 mM NADH, 80 µM cytochrome c, and 0.1 mM of sodium azide. SODH II activities were defined as the flux difference before and after the addition of 1 µM antimycin A. When used, decylubiquinone was added at 0.02 mg/ml. The COX activity was assessed using a classical TMPD/ascorbate assay. In brief, isolated mitochondria (65 µg/ml) were suspended in mitochondrial respiration buffer (see “Measurement of mitochondrial respiration”). Oxygen consumption was assessed in the presence of TMPD (0.2 mM), ascorbate (1 mM), and antimycin A (0.5 µM). After a few minutes of stationary respiration, 2 mM KCN was injected into the chamber. The COX activity corresponds to the KCN-sensitive respiration. All chemicals were obtained from Sigma-Aldrich.

Measurement of mitochondrial ATP production

To assess the ATP production rate, isolated mitochondria (65 µg/ml) were suspended in the mitochondrial respiration buffer (see “Measurement of mitochondrial respiration”). After addition of ADP (1 mM), succinate (2 mM), and antimycin (10 nM) or addition of ADP (1 mM), pyruvate (10 mM), glutamate (5 mM), and malate (5 mM), the oxygen consumption and ATP synthesis rates were both measured. Aliquots were collected every 20 s and precipitated in 7% HClO₄/25 mM EDTA, centrifuged at 16,000 g for 10 min, and then neutralized with 2 M KOH and 0.3 M MOPS. The ATP content in these samples was determined with the ATPLite 1 step (PerkinElmer). In a parallel experiment, oligomycin (2.5 µg/ml protein) was added to the mitochondrial suspension to determine the nonoxidative ATP synthesis rate.
Measurement of hydrogen peroxide production

The rate of H$_2$O$_2$ production was determined by monitoring the oxidation of the fluorogenic indicator Amplex red (Life Technologies) in the presence of horseradish peroxidase. The concentrations of horseradish peroxidase and Amplex red in the incubation medium were 5 U/ml and 1 µM, respectively. Fluorescence was recorded at the following wavelengths: excitation, 560 nm; emission, 590 nm. A standard curve was obtained by adding known amounts of H$_2$O$_2$ to the assay medium in the presence of the reagents. Mitochondria (65 µg protein/ml) were incubated in the respiratory medium (see “Measurement of mitochondrial respiration”) at 37°C and the H$_2$O$_2$ production rate measurements were initiated by substrate addition. The H$_2$O$_2$ production rate was determined from the slope of a plot of the fluorogenic indicator versus time.

FACS analysis of mitochondria

FACS analyses were performed using freshly isolated mitochondria (10–20 µg/ml) incubated in respiration buffer supplemented with 10 mM pyruvate, 5 mM glutamate, and 5 mM malate or 10 mM succinate and 10 mM rotenone. After a few minutes of preincubation at 37°C with 250 nM Mitotracker Green FM and 400 nM TMRRM, the mitochondrial potential positive and negative populations were analyzed by quadrant and polygonal region analysis. The cytotoxic fluorometric analysis was performed by using a FACSaria III cell sorter (BD) equipped with 488 nm and 633 nm lasers, and fluorescence was measured using the FL1 and FL2 channels. Data were acquired using DIVA 6.0 software (BD). To analyze the membrane potential, optimum concentration of oligomycin and CCCP defined in the respiratory assay were successively added. FACS analyses were performed using the Flowjo software applying a biexponential transformation to digital data.

Determination of coenzyme Q and terpenoids intermediates content

Mitochondria were additionally purified on a 20% Percoll gradient to separate mitochondria from mitochondrial-associated membranes (MAM). Quinones were extracted by suspending pellets of isolated mitochondria or MEFs in 200 µl of 1 mM cupric sulfate followed by addition of 200 µl ethanol. After sonication, the homogenate was vortexed in the presence of 400 µl hexane. After collection of the upper phase containing hexane, it was dried using a SpeedVac apparatus (Thermo Fisher Scientific). Next, samples were suspended in 50–200 µl ethanol/methanol (9:1), vortexed, incubated for 2 min in an ultrasonication bath, and filtered through a 0.2-µm modified nylon centrifugal filter (VWR) using a centrifuge [5424R; Eppendorf] set to 6°C and 15,000 rpm.

For absolute quantification of Q9 and Q10 in positive ESI multiple reaction monitoring (MRM) mode, an Acquity UPLC system (Waters) was connected to a Xevo TQ (Waters). An Acquity UPLC (Waters) BEH C18 1.7 µm, 2.1 x 50 mm column was used at 40°C. Solvent A was 90% methanol + 10% propanol + 0.1% FA, and solvent B was 45% acetonitrile/acetone + 10% propanol + 0.1% FA. A linear gradient of solvent A ranging from 100% to 0% in 5 min at a flow rate of 0.45 ml/min was used. Between 2 and 6 µl of the samples were injected. The sample manager was set to 6°C for the standards and the samples were defrosted and directly injected. The MRM transitions used for quantification are presented in the Fig. S5 H. Terpenoid intermediates were extracted as described previously (Henneman et al., 2008). In brief, half of a heart was homogenized in 2 ml of 2-propanol:100 mM HEPES, pH 7.8 (1:1 vol/vol), and sonicated for 5 min. After 10 min of centrifugation (20,000 g at 4°C), supernatant was collected and evaporated for 30 min using a SpeedVac apparatus. Samples were then suspended in 100 µl ethanol/methanol (9:1), sonicated for 1 min, and filtrated successively through a 0.45 µm and 0.2 µm modified nylon centrifugal filter (VWR). The absolute quantification of DMAP, FPP, GPP, GGPP, and AMPP was determined using an Acquity UPLC (Waters) connected to a Xevo TQ (Waters). An Acquity UPLC BEH C18 1.7 µm, 2.1 x 50 mm column was used at 40°C. Solvent A was 20 mM ammonium bicarbonate + 0.1% FA and B acetonitrile/water (3:1) + 0.1% FA. A linear gradient from 95% solvent A to 0% in 5 min at a flow rate of 0.35 ml/min was used. 7.5 µl was injected. The sample manager was set to 6°C for the standards and the samples were defrosted and directly injected. The MRM transitions used for quantification are presented in the Table S5 H. Determination of Q9 and Q10 in negative ESI MRM was determined using an Acquity UPLC (Waters) connected to a Xevo TQ (Waters). An Acquity UPLC BEH C18 1.7 µm, 2.1 x 50 mm column was used at 40°C. Solvent A was 90% methanol + 10% propanol + 0.1% FA, and solvent B was 45% acetonitrile/acetone + 10% propanol + 0.1% FA. A linear gradient of solvent A ranging from 100% to 0% in 5 min at a flow rate of 0.45 ml/min was used. Between 2 and 6 µl of the samples were injected. The sample manager was set to 6°C for the standards and the samples were defrosted and directly injected. The MRM transitions used for quantification are presented in the Fig. S5 H. Terpenoid intermediates were extracted as described previously (Henneman et al., 2008). In brief, half of a heart was homogenized in 2 ml of 2-propanol:100 mM HEPES, pH 7.8 (1:1 vol/vol), and sonicated for 5 min. After 10 min of centrifugation (20,000 g at 4°C), supernatant was collected and evaporated for 30 min using a SpeedVac apparatus. Samples were then suspended in 100 µl ethanol/methanol (9:1), sonicated for 1 min, and filtrated successively through a 0.45 µm and 0.2 µm modified nylon centrifugal filter (VWR). The absolute quantification of DMAP, FPP, GPP, GGPP, and AMPP was determined using an Acquity UPLC (Waters) connected to a Xevo TQ (Waters). An Acquity UPLC BEH C18 1.7 µm, 2.1 x 50 mm column was used at 40°C. Solvent A was 20 mM ammonium bicarbonate + 0.1% FA and B acetonitrile/water (3:1) + 0.1% FA. A linear gradient from 95% solvent A to 0% in 5 min at a flow rate of 0.35 ml/min was used. 7.5 µl was injected. The sample manager was set to 6°C for the standards and the samples were defrosted and directly injected. The MRM transitions used for quantification are presented in the Table S5 H.

RNAi

For the knockdown of Drp1, previously published constructs expressing either a GFP-encoding scramble or a GFP-encoding Drp1 microRNA were used (Motori et al., 2013). Cloning of constructs was performed by using the BLOCK-iT Lentiviral Pol II miR RNAi Expression System with EmGFP (Invitrogen). The following sequences were inserted into a pcDNA 6.2-GW/EmGFP-miR vector: 5'-GAAAGTACTGCCGCTGGAGACGTTTTGGCAGCTGACTCGTACGTACGGATCATT3'-[ scramble], 5' -CCGTAAACCT CTACAGGGTCAGTCAGTGGCCAAAACGCCTACCTAGGTAAGAATTTAC-3' [Drp1 microRNA]. 10^6 cells were electroporated with 1.5 µg of plasmid by using the Amaxa 4D Nucleofector (Lonza) according to the manufacturer’s instructions. Cells were then harvested at day 2 after electroporation and transfection efficiency was ≥60% in all analyzed samples.

Immunocytochemistry and confocal microscopy

Cells were fixed with 4% PFA in PBS and processed for immunocytochemistry as described previously (Motori et al., 2013). In brief, cells were fixed for 10 min at room temperature with 4% PFA in PBS, then permeabilized for 5 min at room temperature with 0.1% Triton X-100 in PBS. Blocking of unspecified sites as well as incubation with primary and secondary antibodies was performed in 3% BSA in PBS. Incubation with primary antibodies was carried at +4°C overnight by using chicken anti-GFP (1:1,000; Aves Labs) and rabbit anti-TOM20 (1:1,000; Santa Cruz Biotechnology, Inc.). After washing in blocking buffer, the secondary antibodies donkey anti-chicken conjugated to Alexa Fluor 488 and donkey anti-rabbit conjugated to Alexa Fluor 546 (1:1,000; Jackson ImmunoResearch Laboratories, Inc.) were applied for 2 h at room temperature. Slides were extensively washed in PBS and nuclei were counterstained with DAPI before mounting.

Fluorescence images were acquired using a laser-scanning confocal microscope (TCS SP5X; Leica), equipped with a white light laser, a 405-nm diode UV laser, a 100x objective lens (HCX Plan-Apochromat CS 100x oil, 1.46 NA). Individual stacks were taken in sequential mode by applying the following parameters: DAPI (excitation, 405 nm; emission, 428–467 nm; image size 1,024 × 1,024 pixels, scan speed 200 Hz), Alexa Fluor 488 (excitation, 488 nm; emission, 501–542 nm; image size 1,024 × 1,024 pixels, scan speed 200 Hz), Alexa Fluor 546 (excitation, 549 nm; emission, 563–638 nm; image size 1,024 × 1,024 pixels, scan speed 200 Hz).

Proteome analysis by SILAC

Stable isotope labeling by amino acids in culture was performed as described previously (Ong and Mann, 2006). In brief, control MEFs were grown in DMEM medium supplemented with dialyzed serum in the presence of either light or heavy arginine and lysine (^13C, ^15N-arginine, ^13C, ^15N-lysine; Thermo Fisher Scientific) to allow full incorporation of the amino acids (>98%). After 7 days in culture, labeled and control MEFs were collected and lysed in buffer (6 mM guanidinium chloride, 10 mM Tris (2-carboxyethyl) phosphine hydrochloride, 40 mM chloroacetamide, and 100 mM Tris-HCl; Kulak et al., 2014). After complete lysis, 100 µg of protein sample (50 µg labeled control [heavy] mixed with 50 µg nonlabeled samples [light]) was mixed with 2 µg of Trypsin gold (Promega) and incubated overnight at 37°C to achieve complete digestion. Peptides were cleaned with home-made StageTip.
extracted with an additional 2 ml of chloroform. The combined organic and then centrifuged (4,000 g, 10 min). Data were collected on a mass spectrometer (QExactive Plus Orbitrap; Thermo Fisher Scientific) using a Top10 method. For MS mode, the following parameters were used: resolution 70,000, maximum injection time 100 ms, scan range 300–1,800 m/z. For MSMS mode: resolution 17,500, maximum injection time 120 ms, scan range 100–2,000 m/z.

Mass spectrometry data analysis
The raw data were analyzed with MaxQuant (Cox and Mann, 2008) version 1.4.1.2 using the integrated Andromeda search engine (Cox et al., 2011). The data were searched against the canonical and isomeric sequences of the mouse reference proteome (proteome ID UP000000589, downloaded June 2014) from UniProt. The database was automatically complemented with 247 sequences of contaminating proteins using the “include contaminants” option of MaxQuant. For data analysis, cysteine carboxymethylation was set as “fixed modification” and methionine oxidation and protein Nterminal acetylation as “variable modification.” The digestion parameters were set to “specific” and “Trypsin/P,” allowing for cleavage after lysine and also arginine when followed by proline. The minimum number of peptides and razor peptides for protein identification was 1; the minimum number of unique peptides was 0. The minimum peptide length was 7 amino acids. Protein identification was reversed at a peptide spectrum matches and protein false discovery rate (FDR) of 0.01. The “second peptide” option was on in order to identify co-fragmented peptides. For protein quantification, the heavy labels were set to Arg10 and Lys8. The “requantify” option was on and a minimum of two ratio counts was required for reporting the protein ratio. Unique plus razor peptides were used for protein quantification.

Protein quantification significance analysis
Downstream data analysis was performed using the Perseus statistical framework (https://www.perseus-framework.org/) version 1.5.0.0. Protein contaminants and reverse identifications were removed and protein ratios were reversed. Next, protein ratios and intensities were transformed [log 2 and log 10, respectively]. Proteins that were significantly down-regulated in the knockout versus the wild type were identified using significance B (Cox and Mann, 2008). A p-value of 0.05 was used as a threshold for significance. Only protein ratios that were significantly lower in the down-regulated proteins were used while “specific” and “Trypsin/P,” allowing for cleavage after lysine and also arginine when followed by proline. The minimum number of peptides and razor peptides for protein identification was 1; the minimum number of unique peptides was 0. The minimum peptide length was 7 amino acids. Protein identification was reversed at a peptide spectrum matches and protein false discovery rate (FDR) of 0.01. The “second peptide” option was on in order to identify co-fragmented peptides. For protein quantification, the heavy labels were set to Arg10 and Lys8. The “requantify” option was on and a minimum of two ratio counts was required for reporting the protein ratio. Unique plus razor peptides were used for protein quantification.

Western blot analyses
Proteins from mitochondrial and whole cell lysates were separated by SDS-PAGE and blotted onto PVDF membranes (GE Healthcare). The following primary antibodies were used: mouse anti–β-actin (1:5,000, A5316; Sigma-Aldrich), rabbit anti-Mitofusin-1 (1:1,000. ab104274; Abcam), mouse anti–MFN2 (1:1,000, ab56889; Abcam), mouse anti–VDAC (1:5,000, MABN504; EMD Millipore), and mouse anti–OXPHOS antibody cocktail (1:1,000, ab110413; Abcam). The following secondary antibodies were used: donkey anti–rabbit IgG (NA9340V; GE Healthcare) and sheep anti–mouse (NA9X451; GE Healthcare). Detection was done by incubation with HRP-conjugated secondary antibodies and conversion to chemiluminescence with ECL (GE Healthcare).

miDNA quantification
DNA was extracted using the DNeasy Tissue and Blood kit (QIAGEN) according to the manufacturer’s instructions. 5 ng of total DNA from the heart was used for detection of miDNA and genomic DNA by qPCR with the following
References


