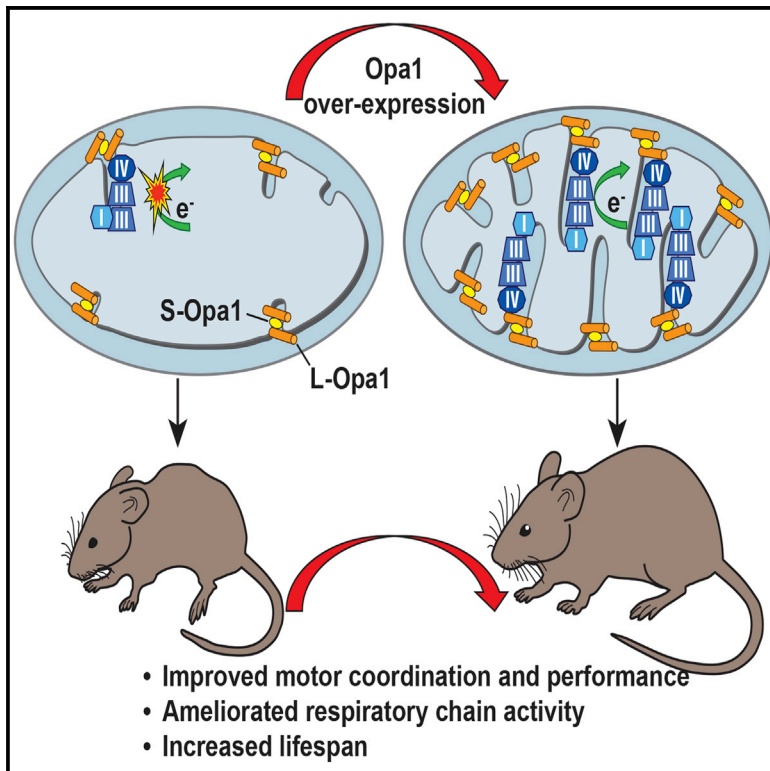


Cell Metabolism

Opa1 Overexpression Ameliorates the Phenotype of Two Mitochondrial Disease Mouse Models

Graphical Abstract



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In Brief

Although mitochondrial diseases have diverse causes, they are all characterized by defective oxidative phosphorylation. Civiletto et al. show that overexpression of the mitochondria-shaping protein OPA1, which improves respiratory chain efficiency, improves the phenotypes of two pre-clinical models of defective mitochondrial bioenergetics.

Highlights

- *Opa1* overexpression improves the phenotype of *Ndufs4*^{-/-} and *Cox15*^{sm/sm} mice
- O₂ consumption rate and respiratory chain activities are increased in double mutants
- Mitochondrial ultrastructure is corrected by *Opa1* overexpression in *Cox15*^{sm/sm}
- Respiratory complexes and supercomplexes are stabilized in *Cox15*^{sm/sm}::*Opa1*^{tg}



Opa1 Overexpression Ameliorates the Phenotype of Two Mitochondrial Disease Mouse Models

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SUMMARY

Increased levels of the mitochondria-shaping protein Opa1 improve respiratory chain efficiency and protect from tissue damage, suggesting that it could be an attractive target to counteract mitochondrial dysfunction. Here we show that *Opa1* overexpression ameliorates two mouse models of defective mitochondrial bioenergetics. The offspring from crosses of a constitutive knockout for the structural complex I component *Ndufs4* (*Ndufs4*^{-/-}), and of a muscle-specific conditional knockout for the complex IV assembly factor *Cox15* (*Cox15*^{sm/sm}), with *Opa1* transgenic (*Opa1*^{tg}) mice showed improved motor skills and respiratory chain activities compared to the naive, non-*Opa1*-overexpressing, models. While the amelioration was modest in *Ndufs4*^{-/-}::*Opa1*^{tg} mice, correction of cristae ultrastructure and mitochondrial respiration, improvement of motor performance and prolongation of lifespan were remarkable in *Cox15*^{sm/sm}::*Opa1*^{tg} mice. Mechanistically, respiratory chain supercomplexes were increased in *Cox15*^{sm/sm}::*Opa1*^{tg} mice, and residual monomeric complex IV was stabilized. In conclusion, cristae shape amelioration by controlled *Opa1* overexpression improves two mouse models of mitochondrial disease.

INTRODUCTION

Mutations in mitochondrial DNA (mtDNA), and in the vast repertoire of nuclear genes that converge on the formation and function of the mitochondrial respiratory chain (MRC), are responsible for primary “mitochondrial disorders,” a group of highly heterogeneous conditions, hallmarked by faulty oxidative phosphorylation (OXPHOS), that can affect any organ, at any age, and by any mode of transmission (Koopman et al., 2012). When taken as a whole, mitochondrial disorders are among the most frequent genetic diseases, affecting >1 in 5,000 individuals in

the European population (Elliott et al., 2008). Despite substantial progress in mitochondrial medicine, the complexity of mitochondrial biology and genetics still constitutes a major challenge for understanding the mechanistic basis of mitochondrial disorders and explains, at least in part, their huge clinical and biochemical variability, which is also a major hurdle toward effective treatment. However, the development of “general” therapeutic strategies, extendable to diverse disease-associated OXPHOS defects, is now a realistic goal, based on rapidly expanding knowledge of the molecular mechanisms underpinning mitochondrial biogenesis, quality control, and signaling pathways. Several of these mechanisms are related to the control of mitochondrial shape and organization of the mitochondrial network.

Mitochondria are highly dynamic organelles that fuse and divide to adapt their structure and morphology to the energetic needs of the cell (Kasahara and Scorrano, 2014). Dynamin-related GTPases located on the inner and outer mitochondrial membranes (IMMs and OMMs) control the fission and fusion processes (Griparic and van der Bliek, 2001). Mitochondrial fission and fusion regulate a number of cellular processes, including organelle distribution during cell proliferation, bioenergetics proficiency, mitochondrial calcium flux, mitochondrial apoptosis, autophagy, and even complex morphogenetic processes such as the formation of dendritic spines (Kasahara and Scorrano, 2014). Fission is regulated by Dynamin-Related Protein 1 (DRP1) and by its OMM partners Fission 1 (FIS1), Mitochondrial Fission Factor (MFF), and Mitochondrial Division (MID) 49 and 51; fusion is controlled by the OMM proteins Mitofusins 1 and 2 (MFN1 and 2) and by the IMM protein Optic Atrophy 1 (OPA1). In humans, eight OPA1 isoforms are produced by alternative splicing of a single gene (Delettre et al., 2001), which are further processed by at least three proteases to form several long (L) and short (S) forms of OPA1 (Anand et al., 2014; Ehses et al., 2009; Griparic et al., 2007; Head et al., 2009; Ishihara et al., 2006; Song et al., 2007), which oligomerize to form functionally active quaternary structures (Frezza et al., 2006). In addition to its role in mitochondrial fusion, OPA1 controls the cristae remodelling arm of mitochondrial apoptosis (Frezza et al., 2006) and the physical and functional organization of the MRC complexes in MRC supercomplexes (Cogliati et al., 2013). These functional quaternary structures increase electron flow channeling through MRC complexes during respiration, thus minimizing



electron leaks (Acín-Pérez and Enriquez, 2014; Enriquez and Le-naz, 2014), and stabilize individual MRC complexes such as Complex III, CIII (Acín-Pérez et al., 2004). Consequently, OPA1 levels directly affect mitochondrial respiratory efficiency (Cogliati et al., 2013) and proteolytic inactivation of OPA1 has been shown to occur in cells from mitochondrial disease patients (Duvezin-Caubet et al., 2007). However, the role of OPA1 in vivo and the potential of its stabilization as a strategy to combat mitochondrial dysfunction and disease remains unclear. Genetic ablation of *Opa1* is lethal during mouse embryonic development (Rahn et al., 2013) and causes massive dysfunction also in postmitotic tissues (L.S., unpublished data). On the other hand, high levels of OPA1 overexpression are toxic in cells, leading to paradoxical mitochondrial fragmentation (Cipolat et al., 2004). In order to avoid these effects, we have used X-chromosome-targeted *Opa1* transgenesis (*Opa1^{tg}*) to obtain moderate, ubiquitous overexpression in a mouse line (Cogliati et al., 2013). Remarkably, *Opa1^{tg}* mice are fertile and viable, have normal lifespan, and are protected from a panoply of tissue insults, including denervation-induced skeletal muscle atrophy, brain and heart ischemia-reperfusion damage, and liver failure caused by Fas-ligand-induced apoptosis (see the accompanying paper by Varanita et al., 2015). We therefore reasoned that *Opa1* overexpression could be exploited to correct the biochemical impairment and mitigate the pathology associated with genetically determined OXPHOS failure in available mouse models of mitochondrial disease. To this aim, we crossed the *Opa1^{tg}* mouse with (i) a constitutive knockout mouse for *Ndufs4* (*Ndufs4^{-/-}*) (Karamanlidis et al., 2013; Kruse et al., 2008; Quintana et al., 2010; Quintana et al., 2012), encoding the 18 kDa subunit of complex I (CI), and (ii) a skeletal-muscle-specific knockout mouse for *Cox15*, in which the *Cre*-recombinase-driven ablation of *Cox15* is dependent upon the muscle-specific actin gene promoter (*Cox15^{sm/sm}*) (Viscomi et al., 2011). *Cox15* encodes a key enzyme in the biosynthesis of heme *a*, an essential catalytic redox component of complex IV ([CIV]; cytochrome c oxidase [COX]) (Bareth et al., 2013; Barros et al., 2001; Barros and Tzagoloff, 2002; Glerum et al., 1997; Petruzzella et al., 1998; Wang et al., 2009). In humans, mutations in *NDUFS4* are associated with early-onset fatal Leigh syndrome due to severe Complex I (CI) deficiency (Petruzzella et al., 2001; Scacco et al., 2003; van den Heuvel et al., 1998). Mutations in *COX15* have been reported in children with severe isolated cardiomyopathy, encephalopathy, or cardioencephalomyopathy (Alfadhel et al., 2011; Antonicka et al., 2003; Bugiani et al., 2005). Here we report that the mild *Opa1* overexpression achieved in the *Opa1^{tg}* mice corrects these two models of CI and CIV deficiency, the latter to a greater extent.

RESULTS

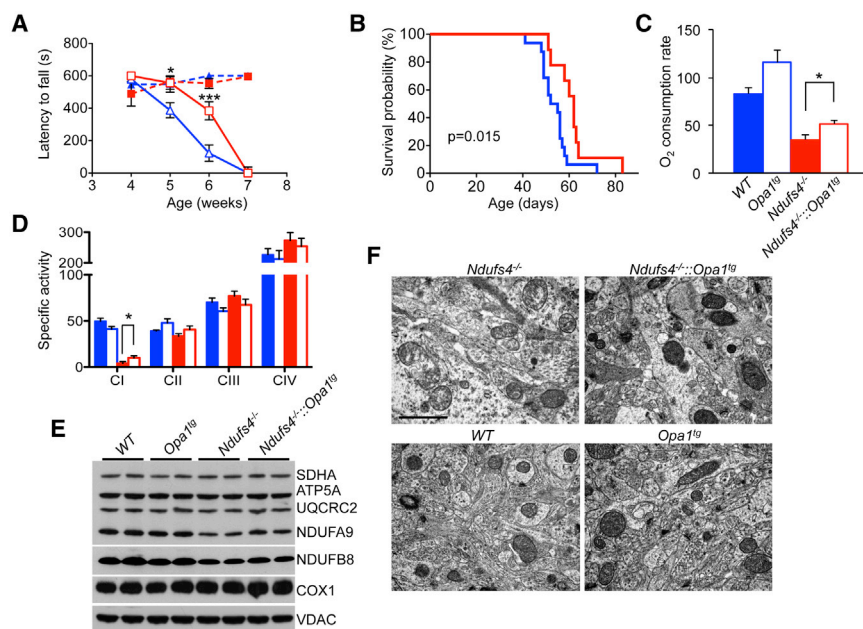
Generation of *Ndufs4^{-/-}::Opa1^{tg}* and *Cox15^{sm/sm}::Opa1^{tg}* Mouse Lines

All procedures on animals were conducted under the UK Animals (Scientific Procedures) Act, 1986, and approved by the local ethical review committee. To obtain transgenic overexpression of *Opa1* in OXPHOS defective mouse lines, we crossed the *Opa1^{tg}* mouse with either *Ndufs4^{-/-}* or *Cox15^{sm/sm}* mice. The constitutive *Ndufs4^{-/-}* mouse shows severe CI deficiency, and

the accumulation of a catalytically inactive 830 kDa CI assembly intermediate, associated with the onset of a rapidly progressive syndrome, dominated by neurological impairment starting approximately 40 days after birth (Calvaruso et al., 2012; Kruse et al., 2008). The *Cox15^{sm/sm}* mouse shows profound skeletal muscle COX deficiency, leading to muscle wasting, severe motor performance impairment, and markedly reduced survival (Viscomi et al., 2011).

Ndufs4^{-/-}::Opa1^{tg} Mice Live Longer and Show Better Motor Coordination than *Ndufs4^{-/-}* Mice

Western blot analysis on two animals for each genotype showed tissue-specific increases in the levels of long and short forms of OPA1 in brain (Figure S1A), skeletal muscle (Figure S1B), and heart (Figure S1C) in *Opa1^{tg}* and *Ndufs4^{-/-}::Opa1^{tg}* relative to their *Ndufs4^{-/-}* and wild-type (WT) littermates. Motor coordination, measured as latency to fall by weekly performed rotarod test, showed steady downhill in both *Ndufs4^{-/-}::Opa1^{tg}* and *Ndufs4^{-/-}* groups, each composed of six animals. However, the scores, which were identical in the two groups at the beginning of the observation (week 4 after birth), showed significant differences in subsequent measurements, with the *Ndufs4^{-/-}::Opa1^{tg}* group performing consistently better than the *Ndufs4^{-/-}* group in weeks 5 (latency to fall: 556 ± 39 versus 388 ± 46 s) and 6 (latency to fall: 384 ± 56 versus 123 ± 50 s). At week 7, both groups were virtually unable to perform the test at any rate, due to collapse of neurological conditions or death (Figure 1A). Furthermore, Kaplan-Meier analysis showed that survival probability was moderately, but significantly, prolonged in *Ndufs4^{-/-}::Opa1^{tg}* versus *Ndufs4^{-/-}* littermates (survival median: 62.0 versus 53.5 days; log-rank test *p* = 0.015) (Figure 1B). The amelioration in motor coordination and prolongation of the lifespan was associated with a ~1.5-fold increase in glutamate-malate-dependent state-3 oxygen consumption rate of isolated brain mitochondria (reflecting CI-driven respiration), which nevertheless remained well below the values obtained in brain mitochondria of WT and *Opa1^{tg}* littermates (Figure 1C). Accordingly, CI activity, measured by spectrophotometric assay (Figure 1D), was ~2.5-fold increased (*p* < 0.05) in *Ndufs4^{-/-}::Opa1^{tg}* versus *Ndufs4^{-/-}* isolated brain mitochondria. Several MRC complex subunits were analyzed by western blotting on SDS-PAGE of isolated brain mitochondria. Notably, the two tested CI subunits (NDUFA9 and NDUF8), as well as the UQCRC2 subunit of CIII, were increased ~1.3-fold in *Ndufs4^{-/-}::Opa1^{tg}* versus *Ndufs4^{-/-}* brain mitochondria (Figure 1E). The *p* values of two-tail, unpaired Student's *t* test of densitometric analysis obtained from four independent samples/genotype was *p* = 0.02 for NDUFA9, *p* = 0.04 for NDUF8, and *p* = 0.01 for UQCRC2. Contrariwise, no significant change was observed for SDHA (CII), COX1 (CIV), and ATP5A (CV) subunits. Blue native gel electrophoresis (BNGE)-based in-gel activity of CI and CI-containing supercomplexes (sc) showed hardly any reactive band in both *Ndufs4^{-/-}::Opa1^{tg}* double mutants and *Ndufs4^{-/-}* littermates (Figure S1D). Western blot analysis using antibodies against NDUF8 (specific to CI), UQCRC1 (specific to CIII), and COX1 (specific to CIV) showed, in the same genotypes, a catalytically active 200 kDa assembly intermediate as well as the catalytically inactive 830 kDa CI and 830 kDa+CIII₂ assembly intermediates previously reported in the *Ndufs4^{-/-}* mouse model (Calvaruso et al., 2012), but neither



the significance levels calculate by unpaired, two-tail Student's t test *p < 0.05.

(E) Western blot analysis of subunits from CI (NDUFA9 and NDUFB8), CII (SDHA), CIII (UQCRC2), CIV (COX1), and CV (ATP5A) in brain isolated mitochondria. VDAC is taken as an internal control.

(F) Electron microscopy of brain specimens from the different genotypes. Note that mitochondrial morphology is well preserved in *Ndufs4^{-/-}::Opa1^{tg}* versus *Ndufs4^{-/-}*. Scale bar 1 μ m.

CI holocomplex nor CI supercomplexes containing CI+CIII₂ or CI+CIII₂+CIV (Figure S1E). However, electron microscopy analysis of forebrain (Figure 1F) and cerebellum (not shown) suggested amelioration in cristae architecture, with no change in mitochondrial shape and dimensions.

These findings suggest that moderate *Opa1* overexpression partially rescues the phenotype of a CI-deficient mouse model by improving cristae morphology.

Survival and Functional Improvement in the Double Mutant *Cox15^{sm/sm}::Opa1^{tg}* Mice

Similar to *Cox15^{sm/sm}::Opa1^{tg}*, male and female *Cox15^{sm/sm}::Opa1^{tg}* mice were smaller than WT or *Opa1^{tg}* littermates (p < 0.005; n = 4 for each genotype; data not shown). To test muscle performance, we monthly monitored motor endurance by a standard treadmill test starting at 2 months of age. Albeit still significantly lower than WT (n = 8) and *Opa1^{tg}* (n = 9) littermates, the scores of the *Cox15^{sm/sm}::Opa1^{tg}* double mutant group (n = 8) were ~2- to 3-fold higher than those of the *Cox15^{sm/sm}* group (n = 10) (p < 0.005 at each time point; Figure 2A). On average, the distance covered by *Cox15^{sm/sm}::Opa1^{tg}* animals was three times that of the *Cox15^{sm/sm}* littermates (169 \pm 20 versus 58 \pm 6 m, p = 2.59 \times 10⁻⁶) (Figure S2A). This improvement was recorded in both homozygous females and hemizygous males, with no significant gender difference (Figure S2B). Interestingly, *Opa1^{tg}* performed better than WT littermates (890 \pm 50 versus 803 \pm 27 m, p < 0.001; Figure S2A). We followed the animals up to 6 months of age. Notably, *Cox15^{sm/sm}::Opa1^{tg}* double mutants showed a remarkable increase in survival probability compared to their *Cox15^{sm/sm}* littermates (log-rank test p = 0.0003). Only 1/17 (7%) *Cox15^{sm/sm}::Opa1^{tg}* animal died during

our observation period, versus 12/23 (50%) *Cox15^{sm/sm}* littermates (Figure 2B).

These findings indicate that controlled *Opa1* overexpression ameliorates life expectancy and motor function of a mouse model of muscle CIV deficiency.

Amelioration of Mitochondrial Morphology and Biochemistry in *Cox15^{sm/sm}::Opa1^{tg}* Muscle

H&E staining of the mid-portion of gastrocnemius muscle (Figure 3A) showed numerous centralized nuclei in both *Cox15^{sm/sm}* and *Cox15^{sm/sm}::Opa1^{tg}*, the proportion of which did not significantly differ between the two groups (Figure 3B). However, the average cross-sectional area (CSA) of muscle fibers was doubled in the *Cox15^{sm/sm}::Opa1^{tg}* relative to *Cox15^{sm/sm}* muscles (p < 0.05), although still smaller (~50%) than the areas measured in WT and *Opa1^{tg}* muscles (p < 0.05) (Figure 3C). We also tested whether the muscle hypotrophic and dystrophic features of the *Cox15^{sm/sm}* mice were associated with increased apoptosis or altered autophagy. TUNEL staining in muscle sections did not reveal hardly any positive fiber in either *Cox15^{sm/sm}* or *Cox15^{sm/sm}::Opa1^{tg}*, similar to WT and *Opa1^{tg}* (not shown), suggesting that apoptosis is not a major player in the pathogenesis of *Cox15^{sm/sm}* myopathy. Immunostaining of muscle sections with Lamp1 and P62 antibodies revealed increased reactions in *Cox15^{sm/sm}* samples compared to WT and *Opa1^{tg}*, which, however, did not change in *Cox15^{sm/sm}::Opa1^{tg}* (not shown). These data indicate that apoptosis and autophagy are unlikely to contribute to the amelioration observed in *Cox15^{sm/sm}::Opa1^{tg}* mice.

Next, we performed ultrastructural analysis of muscle fibers (Figure 4). Examination of tibialis anterior muscle revealed

Figure 1. Improved Motor Function and Extended Lifespan in *Ndufs4^{-/-}::Opa1^{tg}* Double Mutants

(A) Rotarod analysis (n = 6/group). Dashed blue: WT; dashed red: *Opa1^{tg}*; solid blue: *Ndufs4^{-/-}*; solid red: *Ndufs4^{-/-}::Opa1^{tg}*. Error bars represent SEM. The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test, *p < 0.05, ***p < 0.005, and refer to the comparison between *Ndufs4^{-/-}::Opa1^{tg}* and *Ndufs4^{-/-}*.

(B) Kaplan-Meier survival probability. Significance was assessed by the log rank test. Solid blue: *Ndufs4^{-/-}* (n = 16); solid red: *Ndufs4^{-/-}::Opa1^{tg}* (n = 9).

(C) Oxygen consumption measurements (nmol O₂/min/mg of protein). Solid blue: WT (n = 5); blue outline: *Opa1^{tg}* (n = 5); solid red: *Ndufs4^{-/-}* (n = 5); red outline: *Ndufs4^{-/-}::Opa1^{tg}* (n = 6). The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test *p < 0.05.

(D) MRC activities (nmol/min/mg of protein) in isolated brain mitochondria. Solid blue: WT (n = 8); blue outline: *Opa1^{tg}* (n = 5); solid red: *Ndufs4^{-/-}* (n = 7); red outline: *Ndufs4^{-/-}::Opa1^{tg}* (n = 8). Error bars represent SEM. The asterisks represent

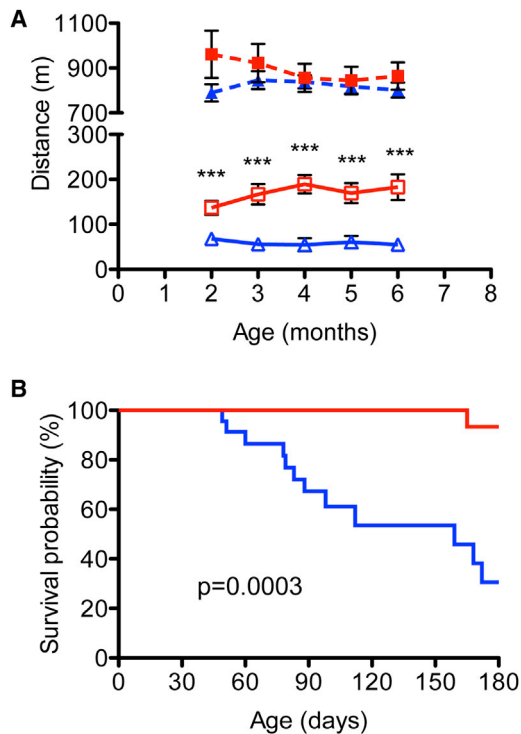


Figure 2. Improved Motor Function and Extended Lifespan in *Cox15^{sm/sm}::Opa1^{Tg}* Animals

(A) Treadmill analysis of motor performance. Dashed blue: WT (n = 8); dashed red: *Opa1^{Tg}* (n = 9); solid blue: *Cox15^{sm/sm}* (n = 10); solid red *Cox15^{sm/sm}::Opa1^{Tg}* (n = 8). Error bars represent SEM. The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test, ***p < 0.005, and refer to the comparison between *Cox15^{sm/sm}::Opa1^{Tg}* and *Cox15^{sm/sm}*.

(B) Kaplan–Meier survival probability. Significance was assessed by the log rank test. Solid blue: *Cox15^{sm/sm}* (n = 23); solid red: *Cox15^{sm/sm}::Opa1^{Tg}* (n = 17).

marked improvement of mitochondrial and myofibrillar morphology in *Cox15^{sm/sm}::Opa1^{Tg}* compared to *Cox15^{sm/sm}*. While *Cox15^{sm/sm}* muscle mitochondria showed profound cristae disorganization and prominent vacuolar degeneration of the inner mitochondrial compartment, the *Cox15^{sm/sm}::Opa1^{Tg}* cristae were organized in an ordered and parallel array and were tightly folded, similar to those observed in WT and *Opa1^{Tg}* littermate specimens (Figure 4A). No hyperfused organelles were observed in *Opa1^{Tg}* muscle. Likewise, myofibrils, which in *Cox15^{sm/sm}* appeared disorganized and separated from each other by the accumulation of membranous and granular material, showed a compact orderly array in *Cox15^{sm/sm}::Opa1^{Tg}*, again resembling those of WT and *Opa1^{Tg}* specimens (Figure 4A). Quantification of area, perimeter, length, and width of muscle mitochondria showed significantly increased values (1.5- to 5-fold) in *Cox15^{sm/sm}*, relative to WT and *Opa1^{Tg}* specimens, indicating mitochondrial ballooning. These parameters were partially corrected in *Cox15^{sm/sm}::Opa1^{Tg}* samples (1.3- to 1.8-fold versus WT) (Figure 4B).

We then tested whether the improvements in motor performance and muscle morphology observed in *Cox15^{sm/sm}::Opa1^{Tg}* animals were associated with ameliorated biochemical parameters. Glutamate-malate dependent state-3 oxygen consumption

rate in isolated muscle mitochondria (reflecting CI-driven respiration) was decreased in *Cox15^{sm/sm}* animals (128 ± 37 nmol/min/mg protein), but increased significantly in *Cox15^{sm/sm}::Opa1^{Tg}* (190 ± 48 nmol/min/mg; p < 0.05), up to lower normal values, not significantly different from those obtained in WT (250 ± 84 nmol/min/mg) and *Opa1^{Tg}* (276 ± 101 nmol/min/mg) samples (Figure 5A). Similar results were obtained by measuring succinate- and TMPD/ascorbate-dependent state-3 oxygen consumption rate (reflecting CII- and CIV-driven respiration, respectively) (Figure 5A). We also analyzed the activity of the single respiratory complexes by spectrophotometric assays. The specific activity of CIV normalized to that of citrate synthase (CS) was significantly increased (p < 0.05) in *Cox15^{sm/sm}::Opa1^{Tg}* (16 ± 3) versus *Cox15^{sm/sm}* (10 ± 4) muscle homogenates, albeit still much lower than that of WT (47 ± 18) or *Opa1^{Tg}* (44 ± 7) muscles (Figure 5B). These results were qualitatively confirmed by histochemical COX staining, which showed reduced COX-negative fibers in COX/SDH double reaction in *Cox15^{sm/sm}::Opa1^{Tg}* versus *Cox15^{sm/sm}* muscle fibers (Figure 5C).

Overall, these data indicate that the ameliorated *Cox15^{sm/sm}::Opa1^{Tg}* mitochondrial ultrastructure is accompanied by improved respiration and CIV activity.

Mitochondrial Protein and Transcriptional Analysis in *Cox15^{sm/sm}::Opa1^{Tg}* Mice

In order to investigate the molecular bases of the observed correction of mitochondrial respiration in *Cox15^{sm/sm}::Opa1^{Tg}* animals, we first analyzed the amount of subunits specific to each MRC complex by western blotting analysis of muscle homogenates separated by SDS-PAGE (Figure 6A). COX1, the mtDNA-encoded largest CIV subunit, which contains the two heme-a moieties and the Cu_B center, and COX4 (isoform 1), a nucleus-encoded early-assembled CIV subunit, were both reduced in *Cox15^{sm/sm}* compared to WT and *Opa1^{Tg}* samples, but did increase in those from *Cox15^{sm/sm}::Opa1^{Tg}* animals (Figure 6B). In both *Cox15^{sm/sm}* and *Cox15^{sm/sm}::Opa1^{Tg}* samples, subunits of other MRC complexes were generally increased, particularly the UQRCR2 subunit of CIII, suggesting compensatory activation of MRC biogenesis induced by CIV deficiency. Interestingly, analysis of mitochondrial and nuclear transcripts of CIV subunits revealed a specific, marked reduction of *mt-Cox1* mRNA in *Cox15^{sm/sm}* samples, which was fully rescued in *Cox15^{sm/sm}::Opa1^{Tg}* samples (Figure 6C). Conversely, other mtDNA- (*mt-Cox2*) and nucleus-encoded (*Cox411* and *Cox5a*) transcripts were unchanged in *Cox15^{sm/sm}* versus WT mice and were not affected by *Opa1* overexpression (Figure 6C). We detected no quantitative change in any group of animals for transcripts *Ppargc1a* and *Tfam*, encoding PGC-1 α and TFAM, respectively (Figure 6C). The mtDNA copy number, which was increased in *Cox15^{sm/sm}*, returned to WT levels in *Cox15^{sm/sm}::Opa1^{Tg}* (Figure S3A), suggesting normalization of the homeostatic control on mtDNA maintenance (Scarpulla et al., 2012). Importantly, the *Cox15* transcript level measured in skeletal muscle of *Cox15^{sm/sm}* was ~9% of that present in WT and *Opa1^{Tg}* littermates and did not change significantly in *Cox15^{sm/sm}::Opa1^{Tg}* (Figure S3B), ruling out a direct effect of *Opa1* overexpression on *Cox15* transcription.

In order to analyze the composition and amount of OPA1, we performed a western blot experiment using an anti-OPA1

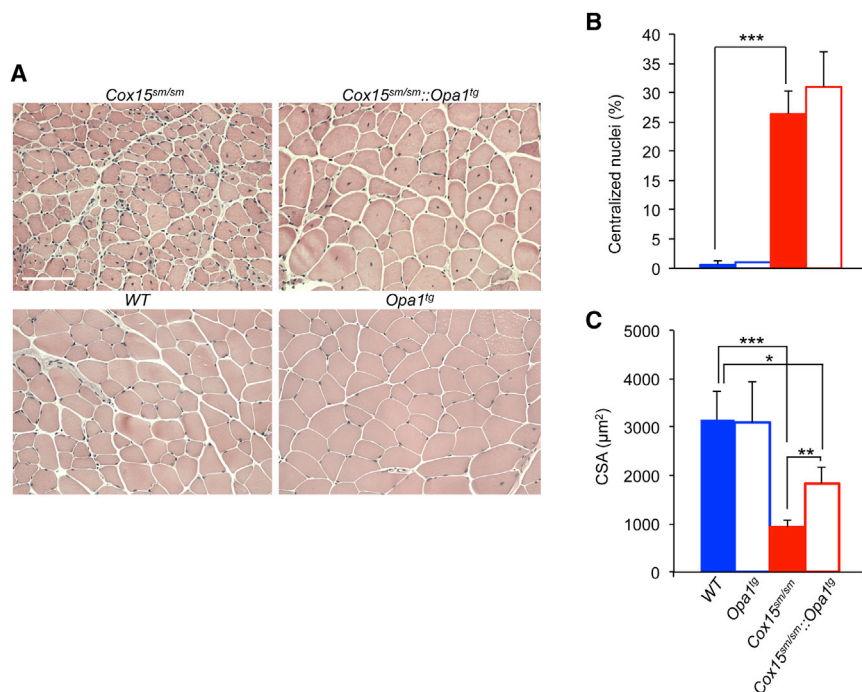


Figure 3. Improved Morphological Features of *Cox15^{sm/sm}::Opa1^{Tg}* Skeletal Muscle

(A) H&E staining of skeletal muscles from mice of the different genotypes. Scale bar 100 μm.

(B) Quantification of the number of centralized nuclei in the different genotypes. 600 fibers/sample were analyzed (n = 4/genotype). Solid blue: WT; blue outline: *Opa1^{Tg}*; solid red: *Cox15^{sm/sm}*; red outline: *Cox15^{sm/sm}::Opa1^{Tg}*. Statistical significance was calculated by unpaired, two-tail Student's t test: ***p < 0.005. Error bars represent SEM.

(C) Cross-sectional area of muscle fibers of gastrocnemius. 600 fibers/sample were analyzed (n = 4/genotype). Solid blue: WT; blue outline: *Opa1^{Tg}*; solid red: *Cox15^{sm/sm}*; red outline: *Cox15^{sm/sm}::Opa1^{Tg}*. Statistical significance was calculated by unpaired, two-tail Student's t test: *p < 0.05; **p < 0.01; ***p < 0.005. Error bars represent SEM.

antibody that specifically recognizes five OPA1 species in skeletal muscle, namely isoforms 1 and 7, and short forms 1–3, resulting from proteolytic cleavage (Figure 6A). Densitometric quantification showed a significant increase (~1.4-fold) in the amount of all five detected OPA1 species in *Opa1^{Tg}* muscle samples relative to WT. The amount of OPA1 species was further increased (~1.3- to 1.8-fold) in *Cox15^{sm/sm}* versus WT and significantly more so in *Cox15^{sm/sm}::Opa1^{Tg}* (~1.5- to 2-fold), particularly for the short forms 1–3 (Figure S3C).

These data suggest that *Opa1* overexpression leads to stabilization in CIV protein and transcript levels.

MRC Supercomplexes Are Increased in *Cox15^{sm/sm}::Opa1^{Tg}* Mice

Individual stability of MRC complexes depends on their assembly in respiratory chain supercomplexes (sc) (Acín-Pérez et al., 2004). We therefore analyzed CI and CIV BNGE in-gel activities in digitonin-treated isolated muscle mitochondria. In *Cox15^{sm/sm}* samples, the intensity of the band corresponding to individual CIV holocomplex was markedly reduced, and hardly any reaction was detected in the high-molecular-weight area harboring sc. However, individual CIV in-gel activity was increased, and CIV reactive high-molecular-weight bands were clearly detected in *Cox15^{sm/sm}::Opa1^{Tg}* mitochondria (Figure 7A), indicating stabilization of CIV holocomplex and CIV-containing sc. Likewise, several high-molecular-weight CI-reactive bands, corresponding to CI- and CIV-containing sc, were virtually absent in the same *Cox15^{sm/sm}* samples but were clearly, albeit weakly, visible in the *Cox15^{sm/sm}::Opa1^{Tg}* samples (Figure 7B). Western blot analysis on BNGE samples using an anti-COX1-specific antibody confirmed the results of in-gel activity: a band corresponding to CIV holocomplex was increased in *Cox15^{sm/sm}::Opa1^{Tg}* compared to *Cox15^{sm/sm}* and high-molecular-weight Cox1-reactive bands, which were detected in WT and

an anti-Ndufa9 antibody showed slight increased Ndufa9 amount in *Cox15^{sm/sm}::Opa1^{Tg}* versus *Cox15^{sm/sm}* (Figure 7D). These results indicate that increased OPA1 levels stabilize RCS and residual CIV in *Cox15^{sm/sm}* mice.

DISCUSSION

The moderate overexpression of *Opa1* achieved in the *Opa1^{Tg}* mouse model is well tolerated and compatible with normal development, fertility, and lifespan. No obvious detrimental effects were noted during the 6-month period of our experimental observation; although, as mentioned in the accompanying paper by Varanita et al. (2015), prolonged *Opa1* overexpression may lead to increased strain-specific prevalence of cancer, a somehow expected result given the antiapoptotic role of OPA1. However, increased OPA1 amount in key tissues, including skeletal and cardiac muscle, brain, and liver, confer remarkable protection against a wide spectrum of experimental tissue damage models (accompanying paper by Varanita et al., 2015). Here, we demonstrate that this effect can be exploited to improve the motor performance, as well as biochemical and molecular phenotypes of two mouse models of genetically determined OXPHOS failure.

The *Ndufs4^{-/-}* mouse is characterized by the complete absence of a structural component of CI. In the current map of mammalian CI, the NDUFS4 18 kDa subunit has been modeled into a density in a cleft between the 75 kDa subunit and the 49 kDa, 30 kDa, and NDUFB8 (TYKY) subunits (Vinothkumar et al., 2014). This localization may explain why disease mutations in the 18 kDa subunit lead to accumulation of late-stage interrupted-assembly intermediates lacking the NADH-dehydrogenase module (Calvaruso et al., 2012). The indispensable role of NDUFS4 in preserving the CI structure and redox activity can also explain the limited improvement in motor coordination,

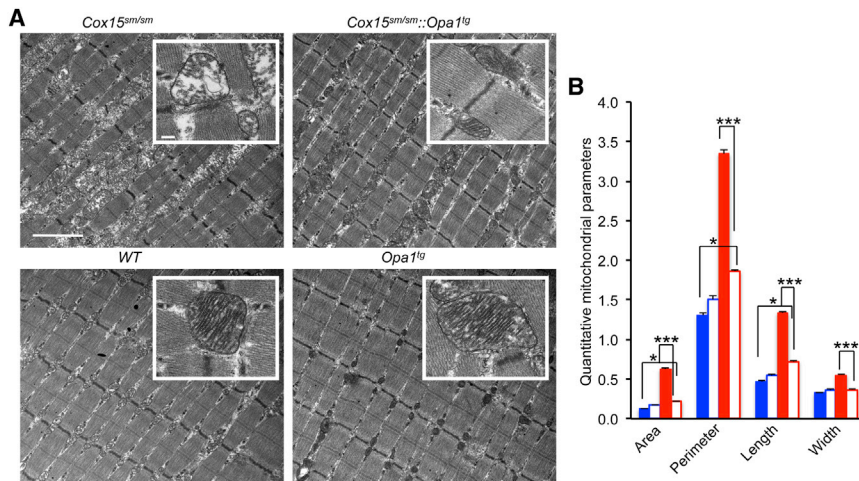


Figure 4. Amelioration of Mitochondrial Ultrastructure in *Cox15^{sm/sm}::Opa1^{Tg}* Skeletal Muscle

(A) EM microphotographs of specimens from the different genotypes. Note that while there is accumulation of inter-myofibrillar amorphous material and disorganization of the mitochondrial cristae (inset) in the *Cox15^{sm/sm}*, the myofibrillary array and cristae structure (inset) are both well preserved in the *Cox15^{sm/sm}::Opa1^{Tg}*. Mitochondria appear larger and with denser cristae in *Opa1^{Tg}* than in WT. Scale bar 2 μ m; scale bar of the inset: 200 nm.

(B) Morphometric quantification of the area, perimeter, length and width of mitochondria from the different genotypes. A total of 200 mitochondria from WT and *Opa1^{Tg}* mice ($n = 2$) were analyzed, and 500 from *Cox15^{sm/sm}* and *Cox15^{sm/sm}::Opa1^{Tg}* ($n = 2$).

measured by rotarod, and biochemical phenotypes observed in *Ndufs4^{-/-}::Opa1^{Tg}* double mutants. Rapamycin-induced inhibition of mTOR, which was recently shown to alleviate the phenotype, also failed to correct the CI defect of *Ndufs4^{-/-}* mice (Johnson et al., 2013). However, the cristae-centric approach used here to treat CI deficiency improved survival to levels comparable to those achieved by every-other-day rapamycin administration (Johnson et al., 2013). It would be interesting to verify if the combination of *Opa1* overexpression and mTOR inhibition is additive and can achieve even better amelioration. We also cannot exclude that OPA1-driven correction of OXPHOS failure may be less amenable in brain than in other tissues, e.g., skeletal muscle.

The *Cox15^{sm/sm}* mouse model is based on muscle-specific ablation of *Cox15*, dependent on the activity of the *Cre* recombinase under the control of the skeletal-muscle-specific actin promoter. Although *Cox15^{sm/sm}* animals display early-onset progressive mitochondrial myopathy, characterized by profound muscle weakness and wasting leading to early death, the genetic lesion underpinning this phenotype can be considered as a hypomorphic mutant allele, the effects of which on COX activity and MRC proficiency are severe but partial. Accordingly, *Cox15* transcript was drastically decreased, but not absent, in *Cox15^{sm/sm}* and *Cox15^{sm/sm}::Opa1^{Tg}* muscles, explaining the presence of some residual COX activity in muscle of adult animals and the “mosaic”-like distribution of the histochemical reaction to COX, which is preserved in scattered muscle fibers (Figure 5C). The molecular *Cox15^{sm/sm}* defect resembles that of the majority of mitochondrial disorders (i.e., partial rather than complete biochemical impairment) due to decreased but not abolished function of the mutant gene product. Incidentally, this category of mitochondrial diseases also includes the few *COX15* mutant patients reported in the literature (Alfadhel et al., 2011; Antonicka et al., 2003; Bugiani et al., 2005). These considerations may explain the marked and persistent improvement of the motor performance observed in *Cox15^{sm/sm}::Opa1^{Tg}* double mutants and suggest that OXPHOS impairment due to decreased, but not abolished, function of the mutant gene product is amenable to correction by potentiating the effects of OPA1 on mitochondrial cristae shape and MRC function.

The *Cox15^{sm/sm}::Opa1^{Tg}* animals displayed not only a dramatic increase in motor endurance but also a remarkable extension of survival up to, and in most cases well beyond, the 6-month period of observation of our experimental protocol. These effects were associated with robust increase of muscle mitochondrial respiration accompanied by a milder but significant increase of muscle COX activity. The discrepancy between the nearly complete reversion to normal in the respiratory rate and the modest increment of COX activity may be explained by increased stabilization of CIV holocomplex and CIV-associated supercomplexes, as indicated by results of BNGE in-gel activity and immunoblotting. The recovery in the motor performance and biochemical proficiency of the *Cox15^{sm/sm}::Opa1^{Tg}* double mutants was associated with marked correction of the profound ultrastructural alteration in cristae morphology associated with *Cox15* ablation. This effect is likely to be directly connected to the role of OPA1 in cristae shape and may be correlated to partial stabilization of the CIV holocomplex and supercomplex organization in the cristae membrane. Accordingly, both nucleus- and mitochondria-encoded MRC subunits were increased in *Cox15^{sm/sm}::Opa1^{Tg}* versus *Cox15^{sm/sm}* muscles, while the corresponding transcripts were unchanged, with the notable exception of *mt-Cox1* mRNA. COX1 is the mtDNA-encoded catalytic CIV subunit that contains the heme-a moieties essential for COX1 stabilization (Bareth et al., 2013). The concordant decrease of both COX1 protein and *mt-Cox1* mRNA in *Cox15^{sm/sm}* was completely reverted in *Cox15^{sm/sm}::Opa1^{Tg}*. Taken together, these results substantiate the possibility that in mammals, as in *Saccharomyces cerevisiae*, COX1 biosynthesis is regulated by a feedback mechanism linking translation and post-translation processing to transcription (Herrmann et al., 2013). In yeast, a key role in this homeostatic loop is carried out by the *mt-Cox1*-specific translation activators MSS51 and PET309 (Herrmann et al., 2013). The mammalian ortholog of PET309 has been identified as the leucine-rich pentatricopeptide repeat-containing (LRPPRC) protein, an RNA binding factor that is mutated in French-Canadian, COX defective Leigh-like syndrome in humans (Mootha et al., 2003). Interestingly, LRPPRC interacts with OPA1, at least in *D. melanogaster* (Banerjee and Chinthapalli, 2014). However, as there is no obvious mammalian

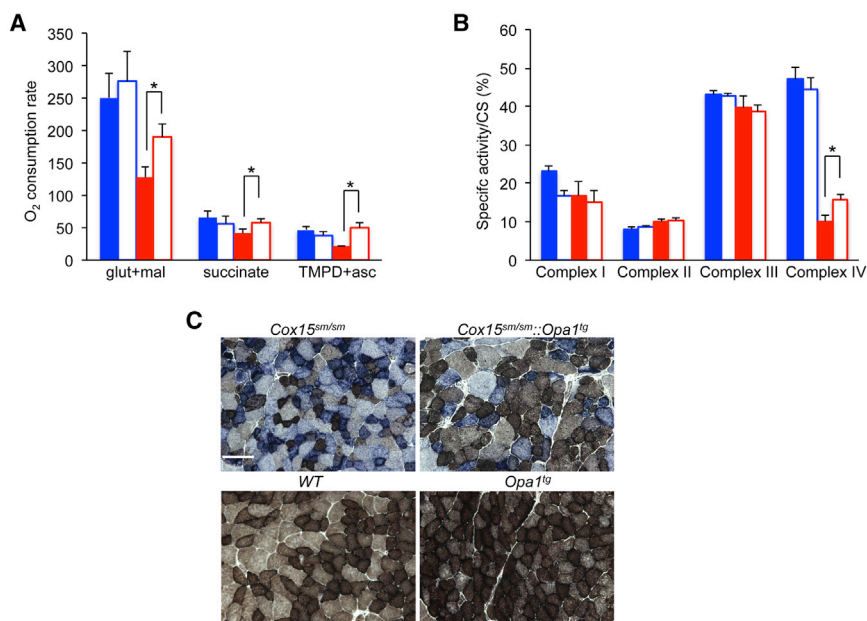


Figure 5. Increased Respiration and Mrc activities in *Cox15^{sm/sm}::Opa1^{Tg}* Skeletal Muscle

(A) Oxygen consumption rate (nmol O₂/min/mg of protein). Solid blue: WT (n = 5); blue outline: *Opa1^{Tg}* (n = 5); solid red: *Cox15^{sm/sm}* (n = 5); red outline: *Cox15^{sm/sm}::Opa1^{Tg}* (n = 6). Error bars represent SEM. The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test, *p < 0.05.

(B) MRC complex activities (nmol/min/mg of protein) in skeletal muscle mitochondria shown as a percentage of CS activity. Solid blue: WT (n = 5); blue outline: *Opa1^{Tg}* (n = 5); solid red: *Cox15^{sm/sm}* (n = 5); red outline: *Cox15^{sm/sm}::Opa1^{Tg}* (n = 6). Error bars represent SD. The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test, *p < 0.05.

(C) COX/SDH histochemical double staining. Scale bar: 50 μm.

ortholog of MSS51, and the 5' UTR sequence of the yeast *Cox1* mRNA, to which MSS51 binds, is virtually absent in the mammalian transcript, these feedback mechanisms are probably different in their molecular details between yeast and mammalian systems.

Finally, while apoptosis seems not to play a significant role in *Cox15^{sm/sm}* myopathy and is not increased in *Cox15^{sm/sm}::Opa1^{Tg}* samples, immunoanalysis suggested dysregulated autophagy in response to *Cox15* ablation, but whether this is relevant

to the pathogenesis of the disease remains to be further investigated.

Our study demonstrates that OPA1-dependent mitochondrial cristae and sc stabilization is effective in correcting mitochondrial disease conditions characterized by partial, however severe, MRC defects. OPA1 can even partially improve defects caused by complete lack of mitochondrial CI via an unknown mechanism. Our results indicate OPA1 as a new target for effective therapy of primary mitochondrial disorders. From a

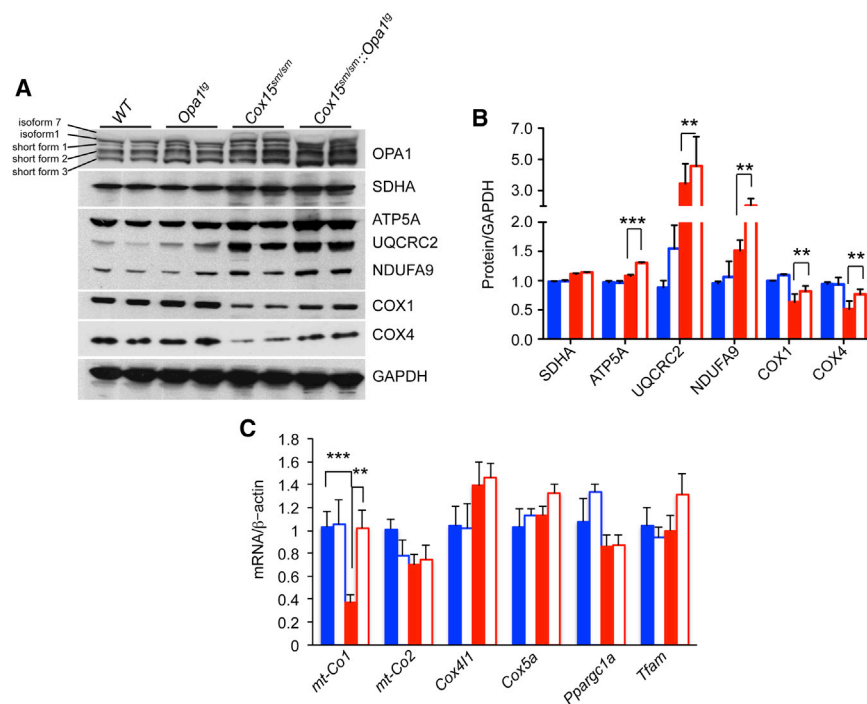


Figure 6. Molecular Analysis of *Cox15^{sm/sm}::Opa1^{Tg}* Skeletal Muscle

(A) Western blot analysis of OPA1, SDHA (CII), ATP5A (CV), UQCRC2 (CIII), NDUFA9 (CI), COX1, and COX4 (CIV); GAPDH is taken as a loading reference.

(B) Densitometric analysis of MRC subunits. Solid blue: WT (n = 4); blue outline: *Opa1^{Tg}* (n = 4); solid red: *Cox15^{sm/sm}* (n = 4); red outline: *Cox15^{sm/sm}::Opa1^{Tg}* (n = 4). Error bars represent SEM. The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test. **p < 0.01; ***p < 0.005.

(C) mRNA expression analysis of CIV- and mitochondrial-biogenesis-related genes. Gene transcripts, retrotranscribed into cDNA, were quantified by RT-PCR; normalized to the amount of the β-Actin gene transcript; taken as a standard; and expressed as time-fold variation relative to the WT. Error bars represent SEM. The asterisks represent the significance levels calculate by unpaired, two-tail Student's t test. **p < 0.01; ***p < 0.005.

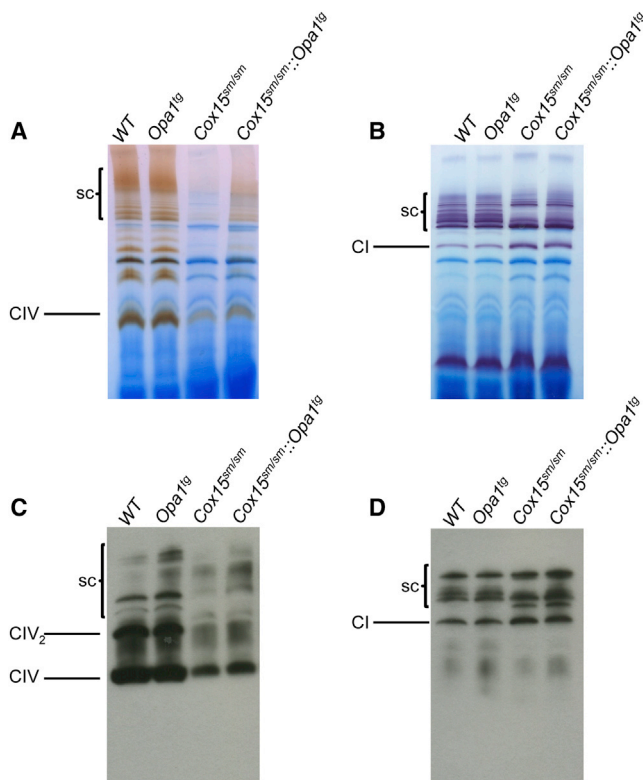


Figure 7. Stabilization of Respiratory Complexes and Supercomplexes in *Cox15^{sm/sm};;Opa1^{Tg}* Skeletal Muscle

(A) CIV-specific BNGE in-gel activity. Note the increased staining of the bands corresponding to CIV-holoenzyme and CIV-containing sc in *Cox15^{sm/sm};;Opa1^{Tg}* versus *Cox15^{sm/sm}*.

(B) CI-specific BNGE in-gel activity. Note the increased staining of the bands corresponding to the sc in *Cox15^{sm/sm};;Opa1^{Tg}* versus *Cox15^{sm/sm}*.

(C) Western blot analysis of MRC CIV from BNGE of digitonin-treated isolated mitochondria. The blot was immunodecorated with an anti-COX1 antibody.

(D) Western blot analysis of MRC CI- and CIV-containing sc's from BNGE of digitonin-treated isolated mitochondria. The blot was immunodecorated with an anti-NDUFB8 antibody.

translational point of view, future work is warranted to elucidate the molecular details underpinning these remarkable effects, extend the experimental observation to additional OXPHOS-defective models, and eventually transfer this set of proof-of-principle observations into effective therapeutic approaches by, for example, selecting compounds or bioreactors able to control the expression or stability of OPA1 in suitable mammalian cell systems and animal models.

EXPERIMENTAL PROCEDURES

Reagents and Materials

Antibodies (COX1, COX4, UQCRC2, NDUFA9, ATP5a, and SDHA) were from Mitoscience, GAPDH were from Millipore, P62 was from Sigma, LC3-I/II was from Cell Signaling, Lamp1 was from Sigma, and OPA1 was from BD Biosciences.

Animal Work

All procedures were conducted under the UK Animals (Scientific Procedures) Act, 1986, approved by Home Office license (PPL: 7538) and local ethical review. The mice were kept on a C57Bl6/129Sv mixed background, and WT

littermates were used as controls. The animals were maintained in a temperature- and humidity-controlled animal-care facility with a 12 hr light/dark cycle and free access to water and food and were sacrificed by cervical dislocation.

Behavioral Analysis

A treadmill apparatus (Columbus Instruments) was used to measure motor exercise endurance according to the number of falls in the motivational grid during a gradually accelerating program with speed initially at 6.5 m/min and increasing by 0.5 m/min every 3 min. The test was terminated by exhaustion, defined as >10 falls/min into the motivational grid.

A rotarod apparatus (Ugo Basile) was used to assess coordination skills. After two acclimation sessions, the mice underwent three trial sessions at least 20 min apart, using a standard acceleration protocol pre-set by the constructor.

Oxygen Consumption Studies

Mouse brains were homogenized in 0.075 M sucrose, 0.225 M mannitol, 1 mM EGTA, and 0.01% fatty-acids-free BSA (pH 7.4); skeletal muscle were homogenized in 150 mM sucrose, 50 mM Tris-HCl, 50 mM KCl, 10 mM EDTA, 0.2% BSA (pH 7.4), and subtilisin 1 mg/g of muscle (Frezza et al., 2007). Mitochondria isolated by differential centrifugation and resuspended 25 mM sucrose, 75 mM sorbitol, 100 mM KCl, 0.05 mM EDTA, 5 mM MgCl₂, 10 mM Tris-HCl (pH 7.4), and 10 mM H₃PO₄ (pH 7.4) (Fernández-Vizarrá et al., 2002).

For oxygraphic measurements, 250–500 μg of mitochondrial proteins were incubated in a buffer containing 225 mM sucrose, 75 mM mannitol, 10 mM Tris-HCl (pH 7.4), 10 mM KCl, 10 mM KH₂PO₄, 5 mM MgCl₂, and 1 mg/ml fatty-acids-free BSA (pH 7.4). Oxygen consumption was evaluated by a Clark oxygen electrode (Hansatech, Instruments), using the following substrates and inhibitors concentrations: 5 mM glutamate and 2.5 mM malate for ci-dependent respiration, 5 mM succinate and 2 μM rotenone for cii-dependent respiration, 6 mM ascorbate, and 300 μM TMPD and antimycin A 0.25 μg/ml for civ-dependent respiration. 100 μM ADP was added to stimulate ATP-coupled oxygen consumption. 100 μM NaCN was added to completely inhibit respiration (Frezza et al., 2007).

BNGE

For BNGE analysis, 250 μg of mitochondria isolated as described above were resuspended in native page buffer (Invitrogen), protease inhibitors, and 4% digitonin and incubated for 1 hr on ice before centrifuging at 20,000 × g at 4°C. 5% Coomassie G250 was added to the supernatant. 30 μg were separated by 3%–12% gradient BNGE and either stained with for in-gel activities or electroblotted on PVDF membranes for immunodetection (Nijtmans et al., 2002).

Morphological Analysis

For histochemical analysis, tissues were frozen in liquid-nitrogen-pre-cooled isopentane. 8-μm-thick sections were stained for COX and SDH, as described Sciacco and Bonilla (1996). Analysis of centralized nuclei and analysis was performed on H&E-stained sections using ImageJ on four samples/genotype (600 fibers/sample). For ultrastructural studies, samples were fixed with 2.5% glutaraldehyde in 10 mM phosphate buffer (pH 7.4). Quantification of mitochondrial morphometry (area, perimeter, long axis, and short axis) in electron micrographs was performed using ImageJ on 200 mitochondria from WT and *Opa1^{Tg}* muscles and 500 mitochondria from *Cox15^{sm/sm}* and *Cox15^{sm/sm};;Opa1^{Tg}* muscles.

Biochemical Analysis of MRC Complexes

Brain and skeletal muscle samples were snap-frozen in liquid nitrogen and homogenized in 10 mM phosphate buffer (pH 7.4). The spectrophotometric activity of CI, CII, CIII, and CIV, as well as CS, was measured as described in Bugiani et al. (2004).

Real-Time PCR

MtDNA content and transcripts analysis was carried out by SYBR Green real-time PCR, as described in Viscomi et al. (2011).

Western Blot Analysis

Mouse tissues were homogenized in ten volumes of 10 mM potassium phosphate buffer (pH 7.4). Mitochondrial-enriched fractions were collected after centrifugation at 800 × g for 10 min in the presence of protease inhibitors,

and frozen and thawed three times in liquid nitrogen. Protein concentration was determined by the Lowry method. Aliquots, 70 μ g each, were run through a 4%–12% SDS-PAGE and electroblotted onto a nitrocellulose membrane, which was then immunodecorated with different antibodies.

Statistical Analysis

All numerical data are expressed as mean \pm SEM. Student's unpaired two-tail t test and Kaplan-Meier distribution were used for statistical analysis. Differences were considered statistically significant for $p < 0.05$.

SUPPLEMENTAL INFORMATION

Supplemental Information includes three figures and can be found with this article online at <http://dx.doi.org/10.1016/j.cmet.2015.04.016>.

AUTHOR CONTRIBUTIONS

G.C. carried out the *in vivo* phenotype analysis and performed biochemical measurements and molecular analysis and wrote the manuscript; T.V. generated mouse colonies, carried out BNGE in-gel activity, western blot, and ultrastructural analysis and wrote the manuscript; T.G. and S.B. helped to carry out *in vivo* phenotype analysis; R.C., S.M., and C.L. carried out histological, histochemical, and morphometric analysis; C.V. supervised and participated in the experimental design, supervised the experimental work, and contributed to write the manuscript; L.S. and M.Z. designed the study and wrote the manuscript.

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