Na⁺/H⁺ antiporter activity by respiratory complex I controls mitochondrial Δψ and is impaired in LHON disease

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The mitochondrial electron transport chain (mETC) converts the energy of substrate oxidation into a H⁺ electrochemical gradient (Δp), which is composed by an inner mitochondrial membrane (IMM) potential (ΔΨmt) and a pH gradient (ΔpH). So far, ΔΨmt has been assumed to be composed exclusively by H⁺. Mitochondrial Ca²⁺ and Na⁺ homeostasis, which are essential for cellular function, are controlled by exchangers and antiporters in the inner mitochondrial membrane (IMM). In the last few years, some of them have been identified, except for the mitochondrial Na⁺/H⁺ exchanger (mNHE). Here, using a rainbow of mitochondrial and nuclear genetic models, we have identified it as, specifically, the P-module of complex I (CI). In turn, its activity creates a Na⁺ gradient across the IMM, parallel to ΔpH, which accounts for half of the ΔΨmt in coupled respiring mitochondria. We have also found that a deregulation of this mNHE function in CI, without affecting its enzymatic activity, occurs in Leber hereditary optic neuropathy (LHON), which has profound consequences in ΔΨmt and mitochondrial Ca²⁺ homeostasis and explains the previously unknown molecular pathogenesis of this neurodegenerative disease.
All living organisms rely on the formation of transmembrane potentials to support energy balance, and the maintenance and regulation of these transmembrane potentials are crucial determinants of cell outcomes and homeostasis. Eukaryotes rely on a plasma membrane potential and a mitochondrial inner membrane potential (ΔΨmt), the latter being particularly important for energy production and cell fate determination. The mETC is composed of several complexes and supercomplexes. Mitochondrial complexes I (CI) and II (CII) respectively oxidize NADH and succinate to reduce ubiquinone (CoQ) to ubiquinol. Complex III (CIII) uses ubiquinol to reduce cytochrome c (cyt c), and complex IV (CIV) oxidizes cyt c to reduce O2 to H2O. This series of reactions is coupled to the translocation of H+ by CI, CIII, and CIV across the inner mitochondrial membrane (IMM) to form a H+-motive force (Δp). Δp, in turn, activates the phosphorylation of ADP to ATP, which is coupled to the electrophoretic entry of H+ through a fifth complex (CV). Δp is composed of an electrical component (ΔΨmt) and a chemical component (ΔpH), with ΔΨmt accounting for approximately 80% of the total Δp.

Mitochondria also contain a panoply of exchangers and antiporters that allow them to maintain respiration, osmolarity, and volume and permit the entry and extrusion of substrates and metabolites. For instance, the mitochondrial Ca2+ uniporter (MCU) introduces Ca2+ into the mitochondrial matrix and Ca2+ is, in turn, extruded by the mitochondrial Na+/Ca2+ exchanger (NCLX) in an electrogenic exchange for 3 Na+ ions. Mitochondrial Na+ exit is mediated by a highly active, electroneutral Na+/H+ exchanger (NHE) of unknown molecular identity.

In the last few years, the relationship between mitochondrial Ca2+ imbalance and neurological diseases is becoming clearer. LHON is a mitochondrial disorder causing central vision loss at early age by degeneration of the optic nerve and it is produced by mutations in mitochondrial DNA (mtDNA)-encoded CI subunits. Particularly, G11778A, the most common mutation producing LHON, do not show a detectable decrease in CI activity (Figure 1a). Thus, we wondered whether, instead, the molecular pathogenesis could be related to an affection in mitochondrial Ca2+. We observed that basal mitochondrial Ca2+ levels were increased and mitochondrial Ca2+ exit rate was lowered altered in LHON cybrids, and that this was not due to lower levels in NCLX amount (Figure 1b-e). Thus, as NCLX activity is coupled to NHE function, we tested whether these defects may be due alterations in the latter. Indeed, we observed that NHE activity in LHON cybrids was abnormally low (Figure 1f). This prompted to question whether other mtDNA mutations or the presence/absence of OXPHOS complexes may also cause a decrease in NHE activity. To note, as decreases in CIII or CIV amounts produce defects in CI assembly, we tested this in cells expressing Emericella nidulans alternative oxidase (AOX) to maintain CI assembly despite the lack of CIII or CIV. We found that only the mutations in CI subunits promoted a decrease in NHE activity, proportionally to the level of mutated DNA (mutation in ND6 is in a 50% heteroplasmy with mtDNAWT; Figure 1g).

CI is composed of three structural modules. The N-module mediates NADH oxidation and transfers the electrons to the Q-module, also called the CoQ-reducing module. The energy released by the NADH-CoQ oxidoreduction is transferred to the subunits in the H+ pumping module, or P-module, which is evolutionarily related to the Na+/H+ antiporters of halophilic and halophilic bacteria, and it is composed by the mtDNA-encoded subunits. Knockout models of different CI subunits produce similar fold decrease in CI activity, regardless of whether they affect the N-module or the P-module (Extended Data Fig. 1a). In contrast, in the literature contradictory findings can be found regarding their bioenergetic impact. We wondered whether quantitative differences in NHE activity could lie behind such contrasting behavior and, surprisingly, we found that the knock-out for the N-module subunit NDUFS4 not only not...
showed a decrease as the other CI deficiency models, but an increase in NHE activity (Figure 1h-k).

**Figure 1. Characterization of CI-dependent NHE activity.** (a) Rotenone-sensitive NADH-decylubiquinone oxidoreductase activity in LHON vs control mitochondrial membranes. (b) Mitochondrial Ca$^{2+}$ levels in LHON vs control cells before and after addition of ATP. (c) Mitochondrial Ca$^{2+}$ basal levels in LHON vs control cells. (d) Mitochondrial Ca$^{2+}$ influx rate in LHON vs control cells. Inset: western blot of mitochondrial extracts showing the mitochondrial calcium uniporter (MCU) levels and a loading control (SDHA). (e) Mitochondrial Ca$^{2+}$ efflux rate in LHON vs control cells. Inset: western blot of mitochondrial extracts showing NCLX levels, a loading control (SDHA) and the ladder. (f) Passive NHE activity in LHON vs control mitochondria. (g) Passive NHE activity in control, ρ0, Cytb$^+$+AOX, Cox10$^{WT}$+AOX, Cox10$^{K0}$+AOX, ND6$^{K0}$ and ND4$^{K0}$ mitochondria. (h) Mitochondrial matrix Na$^+$ extrusion measured with SBFI-AM in mtDNA$^{WT}$, ND6$^{K0}$ and ND4$^{K0}$ intact mitochondria respiring on succinate. (i) Mitochondrial matrix Na$^+$ extrusion measured with SBFI-AM in NDUSF4$^{WT}$ and NDUSF4$^{K0}$ intact mitochondria respiring on succinate. (j) Mitochondrial matrix H$^+$ extrusion measured with BCECF-AM in mtDNA$^{WT}$, ND6$^{K0}$ and ND4$^{K0}$ intact mitochondria respiring on succinate. (k) Mitochondrial matrix H$^+$ extrusion measured with BCECF-AM in NDUSF4$^{WT}$ and NDUSF4$^{K0}$ intact mitochondria respiring on succinate.

Consistent with ND6 mutant mtDNA being in heteroplasmy14-20, ND6$^{K0}$ cells showed residual CI assembly with respect to control isogenic cybrid cells (WT), whereas ND4$^{K0}$ cells contained no detectable levels of assembled CI (Figure 2a). The remaining mitochondrial complexes showed a redistributed pattern on blue-native gel electrophoresis (BNGE) consistent with reduced levels of CI-containing supercomplexes (Figure 2a). In contrast, NDUSF4$^{K0}$ MAFs showed slightly faster migration of CI-containing supercomplexes than NDUSF4$^{WT}$ MAFs, reflecting the assembly of CI with only the Q- and P-modules (Figure 2b). Thus, NDUSF4$^{K0}$ has a higher NHE activity with its CI P-module conserved. In contrast to other CI defects in the P-module (ND4$^{K0}$ and ND6$^{K0}$ models) and the LHON causing point mutation in the mtDNA encoded ND4 protein, produces a specific decrease in NHE activity. This, together with the fact that the mtDNA-encoded CI subunits are related to bacterial NHE and that the *E. coli* CI may act as a NHE22-24, strongly support that CI P-module is a bona-fide NHE in mitochondria. Indeed, this is supported by *in vitro* data using isolated bovine CI reconstituted in liposomes25.
As we now had a loss- and gain-in-function models of NHE, we wondered what the bioenergetic consequences would be, if any, in these cell models (ND4KO, ND6KO vs NDUFS4KO). In line with a loss of Cl activity, all cell models, including wild type cells chronically exposed to low-dose of the Q-module-binding inhibitor rotenone (Chronic Rot), showed an increase in combined CII+III activity compared with untreated or vehicle-treated mtDNAWT cells (Figure 2c-f and Extended Data Figure 1b, summarized in Table S1). Increased CII+III activity is a well-known consequence of Fgr kinase activity in these models.\(^{15}\) In contrast, isolated CIV activity was elevated only in ND4KO and NDUFS4KO cells (Figure 2g-j and Extended Data Figure 1c), while CV activity, in forward or reverse mode, was higher only in NDUFS4KO cells (Extended Data Figure 1d-e, summarized in Table S1). Analysis of oxygen consumption by permeabilized cells respiring on CI, CII, or CIV substrates revealed that Cl-dependent respiration was much lower in ND4KO, ND6KO, and Chronic Rot cells with respect to their wild type or vehicle-treated counterparts but was surprisingly unaltered in NDUFS4KO cells (CI in Figure 3a-c and Extended Data Figure 2a). In the presence of CII substrates, independently of the presence of ADP, respiration rates were higher in all cell models except for ND6KO (CII and CII+ADP in Figure 3a-c and Extended Data Figure 2a). CIV-dependent

**Figure 2. Characterization of CI-deficiency models.** (A) Blue-native electrophoresis (BN-PAGE) of mitochondria from mtDNAWT, ND6KO, and NDUFS4KO hybrid cell lines (n=3 per genotype) immunoblotted to detect subunits of CI (Anti-NDUF9), CIV (Anti-Col), CII (Anti-Core2), CII (Anti-SDHA), and CV (Anti-ATPβ). (B) BN-PAGE of mitochondria from NDUFS4WT and NDUFS4KO MAFs (n=2) blotted against the same epitopes as in (A). The lateral labels in (A) and (B) indicate the migrated positions of all detected complexes and supercomplexes. (C) Antimycin A (AA)-sensitive succinate-cyt c oxidoreductase activity in mtDNAWT and ND6KO mitochondrial membranes. (D) AA-sensitive succinate-cyt c oxidoreductase activity in mtDNAWT and ND4KO mitochondrial membranes. (E) AA-sensitive succinate-cyt c oxidoreductase activity in NDUFS4WT and NDUFS4KO mitochondrial membranes. (F) AA-sensitive succinate-cyt c oxidoreductase activity in mtDNAWT mitochondrial membranes from cells treated for 6 h with either vehicle or 250 nM rotenone (Chronic Rot). (G) Potassium cyanide (KCN)-sensitive cyt c oxidase activity in mtDNAWT and ND6KO mitochondrial membranes. (H) KCN-sensitive cyt c oxidase activity in mtDNAWT and ND4KO mitochondrial membranes. (I) KCN-sensitive cyt c oxidase activity in NDUFS4WT and NDUFS4KO mitochondrial membranes. (J) KCN-sensitive cyt c oxidase activity in mtDNAWT mitochondrial membranes from cells treated for 6 h with either vehicle or 250 nM rotenone (Chronic Rot).
respiration was similar in mtDNA\(^{WT}\) and ND6\(^{KO}\) cells and elevated in ND4\(^{KO}\) and NDUFS4\(^{KO}\) cells (CIV in Figure 3a-c), resembling isolated CIV activity (Figure 2g-j and Extended Data Figure 2c).

Analysis of ΔΨ\textsubscript{mt} revealed that ND6\(^{KO}\) mitochondria were depolarized relative to isogenic controls resiping on any substrate (Figure 3d). In contrast, ND4\(^{KO}\) mitochondria were depolarized only when resiping on CI substrates, showing normal mitochondrial membrane polarization when resiping on CII and CIV substrates (Figure 3e). NDUFS4\(^{KO}\) mitochondria were not depolarized by CI substrates and were hyperpolarized by CII and CIV substrates (Figure 3f). The mitochondria of Chronic-Rot–treated cells were depolarized by CI substrates and hyperpolarized by CII substrates (Extended Data Figure 2b-d). The different cell models thus show apparent contradictory behaviors. What most called our attention was the discrepancy between elevated CII+III and CIV activities and weaker or unaffected mitochondrial membrane polarization in ND6\(^{KO}\) and ND4\(^{KO}\) cells supplied with CII and CIV substrates (Figure 2c, d, g and h, CII and CIV insets in Figure 3d-e). This contrasted with NDUFS4\(^{KO}\) and Chronic Rot

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**Figure 3. Bioenergetic characterization of the ND6\(^{KO}\), ND4\(^{KO}\), and NDUFS4\(^{KO}\) cell lines.** (a) Oxygen consumption rates in mtDNA\(^{WT}\) and ND6\(^{KO}\) permeabilized cells (n=5). (b) Oxygen consumption rates in mtDNA\(^{WT}\) and ND4\(^{KO}\) permeabilized cells (n=7). (c) Oxygen consumption rates in NDUFS4\(^{WT}\) and NDUFS4\(^{KO}\) permeabilized cells (n=4). GM, glutamate/malate; Rot, 1 μM rotenone; Succ, succinate; ADP, adenosine diphosphate; AA, 1 μM antimycin A; TMPD+Asc, N,N,N′,N′-tetramethyl-p-phenylenediamine + ascorbate; Azide, 50 mM, sodium azide. CI denotes CI-dependent respiration, CII denotes CII-dependent respiration, CII+ADP denotes CII+ADP-dependent respiration, and CIV denotes CIV-dependent respiration. (d-f) Calibrated TMRM signal in mtDNA\(^{WT}\), ND6\(^{KO}\) (d), ND4\(^{KO}\) (e) and NDUFS4\(^{KO}\) cells (f), providing the CI-dependent ΔΨ\textsubscript{mt} (Panel CI in d-f), CII-dependent ΔΨ\textsubscript{mt} (Panel CII in d-f), CII+ADP-dependent ΔΨ\textsubscript{mt} (Panel CII+ADP in d-f), and CIV-dependent ΔΨ\textsubscript{mt} (Panel CIV in d-f).
cells, which also showed elevated CII+III activity (and also elevated CIV activity in the case of NDUFS4KO cells), but in this case accompanied by CII-dependent mitochondrial hyperpolarization (and CIV-dependent hyperpolarization in NDUFS4KO cells; Figure 2e, i and Figure 3f, Figure 2f-j and Extended Data Figure 2b-d), despite an even higher forward CV activity in the case of NDUFS4KO cells (Extended Data Figure 1d). To exclude an effect of artifacts arising from the use of permeabilized cells, we repeated the same experiments with isolated mitochondria. The results were the same in all cases except for ND6KO mitochondria, in which respiration rates and ΔΨmt were like those in mitochondria isolated from isogenic wild type cells (Extended Data Figure 3, summarized in Table S1). These results show that knockout models for different CI subunits have distinct bioenergetic footprints under non-CI substrates.

This variable behavior has several possible explanations. We first pondered the possible involvement of mitochondrial volume, H+ leak, or mitochondrial transition pore (MTP) opening. However, none of these parameters was changed in ND6KO or ND4KO mitochondria with respect to their isogenic control (Extended Data Figure 4). In line with previous reports, a proteomics analysis of isolated mitochondria from ND6KO, ND4KO, and NDUFS4KO cells showed a pronounced reduction in most CI subunits in ND4KO and ND6KO cells, contrasting with unaltered levels of P-module subunits in NDUFS4KO cells (Extended Data Figure 5a and e). All the cell models had a similar protein content of other mitochondrial complex subunits (Extended Data Fig 5b-d and Fig. 5f-h). In a false discovery rate analysis, the only difference detected between NDUFS4KO MAFs and the ND6KO and ND4KO lines was the presence in the cybrid lines of mitochondrial medium-chain specific acyl-CoA dehydrogenase (ACADM) (Extended Data Figure 5i-k). However, this fatty acid oxidation enzyme locates in the mitochondrial matrix, whereas the discrepancies we observed are related to the IMM ΔΨmt provoked by CII and CIV substrates. We therefore discarded an involvement of ACADM in the generation and maintenance of ΔΨmt.

Strikingly, frozen-thawed-permeabilized ND6KO and ND4KO mitochondria supplied with CII and CIV substrates had markedly higher oxygen consumption rates relative to isogenic controls than observed with intact ND6KO and ND4KO mitochondria (compare Fig. 4a and b with Extended Data Figure 3a and b). This finding suggested that the distinct bioenergetic footprints of the different CI subunit knockout models might be determined by IMM integrity. We first tested whether the same sample treated with different CI inhibitors may affect equally NHE activity. Only high doses of rotenone were able to decrease NHE activity, in contrast to the N-module inhibitor DPI (Extended Data Figure 6 a-b), which is compatible with the fact that rotenone is also able to bind ND4 at high doses, potentially inhibiting CII NHE. Accordingly, CI activation state only increased NHE function, as occurred with NDUFS4KO, and in contrast to the two other CI deficiency models (Extended Data Figure 6 c-e), which is also in agreement with previous in vitro data.

As ND6KO and ND4KO did not show the expected hyperpolarization under CII and/or CIV substrates, we decided to measure H+ pumping in the same conditions, in all CI deficiency models, as it could be a result of lower H+ ejected by CIII and/or CIV. As expected, H+ pumping was lower in all three of the knockout lines when they were reliant on CI substrates (Extended Data Figure 6f, g, i and j). However, with CI substrates, ND6KO and ND4KO mitochondria showed higher H+ pumping than isogenic controls, resembling NDUFS4KO mitochondria (Extended Data Figure 6f, h, i and k). This finding fits well with our data showing higher CII+III activity in all CI-deficiency models (compare Extended Data Figure 6h and k with Fig. 2c-f). However, the absence of hyperpolarization in ND6KO and ND4KO mitochondria occurred despite
their showing higher $\text{H}^+$ pumping. This intriguing contradiction opens the possibility that CI P-module-dependent NHE activity influences $\Delta \Psi_{\text{mt}}$, so that it is governed not only by a $\text{H}^+$ gradient, but also by a $\text{Na}^+$ gradient.

To evaluate this hypothesis, we tackled this by three different approaches. First, we recorded $\Delta \Psi_{\text{mt}}$ in wild type and ND4$^{\text{KO}}$ mitochondria, supplied with CII substrates and upon the addition of the chemical Na$^+$/$\text{H}^+$ exchanger monensin to chemically restore the NHE function in isolated mitochondria. Whereas nigericin, a chemical K$^+$/H$^+$ exchanger, promoted hyperpolarization in isolated wild type mitochondria (Extended Data Figure 6l), monensin did not (Extended Data Figure 6m and n). In contrast, both drugs supported hyperpolarization of ND4$^{\text{KO}}$ mitochondria (Extended Data Figure 6l and m), indicating that the restoration of a Na$^+$ gradient by monensin addition contributed to the establishment of $\Delta \Psi_{\text{mt}}$ in ND4$^{\text{KO}}$ mitochondria. No similar phenomenon was observed with NDUFS4$^{\text{KO}}$ mitochondria (Extended Data Figure 6n).

Second, we measured respiration rates and $\Delta \Psi_{\text{mt}}$ in intact wild type mitochondria maintained in an osmotically compensated, Na$^+$-free buffer. The absence of Na$^+$ slightly decreased the respiration rate and promoted mitochondrial depolarization by a third or a half, depending on the substrate conditions (Figure 4c and Extended Data Figure 6o). Mouse heart mitochondria behaved similarly (Extended Data Figure 6q-r). In parallel, the absence of Na$^+$ promoted apparently higher CI- and CII-dependent $\text{H}^+$ pumping, which we interpreted as the consequence of an inoperative NHE, unable to dissipate $\Delta \text{pH}$ (Figure 4d and e). None of these phenomena was observed in ND4$^{\text{KO}}$ mitochondria (Figure 4f-h and Extended Data Figure 6p), pointing to a role of the CI NHE-dependent formation of $\Delta \Psi_{\text{mt}}$, possibly reliant on a Na$^+$ gradient. Within this approach we also compared respiration rates and $\Delta \Psi_{\text{mt}}$ between wild type and ND4$^{\text{KO}}$ mitochondria in the absence of Na$^+$. Under these conditions, ND4$^{\text{KO}}$ mitochondria showed even higher respiration (compare Extended Data Figure 6s and Extended Data Figure 3b) and, more importantly, hyperpolarization with respect to mtDNA$^{\text{WT}}$ under CII and CIV substrates (Figure 4t-w). This outcome resembled the bioenergetic footprint of NDUFS4$^{\text{KO}}$ and Chronic Rot mitochondria maintained in Na$^+$-containing buffer (Table S1), indicating that the build-up of a Na$^+$ gradient-dependent $\Delta \Psi_{\text{mt}}$ specifically in wild type mitochondria was masking the effect of higher CII+CIII activity and $\text{H}^+$ pumping on respiration and $\Delta \Psi_{\text{mt}}$ in the ND6$^{\text{KO}}$ and ND4$^{\text{KO}}$ cell models, which may be unable to form a Na$^+$ gradient-dependent $\Delta \Psi_{\text{mt}}$. 


In a third approach, we directly measured the mitochondrial Na$^+$ gradient in isolated mitochondria. For respiration driven by CI or CII substrates, the Na$^+$ gradient in ND6$^{KO}$ and ND4$^{KO}$ mitochondria was below-control in proportion to the level of assembled CI (Figure 4i and j). In contrast, the Na$^+$ gradient in NDUFS4$^{KO}$ mitochondria was unaffected (Figure 4k and l).

Figure 4. Mitochondrial Na$^+$ gradient controls ΔΨmt and is lowered in LHON. (a) Oxygen consumption rates in mtDNA$^{WT}$ and ND6$^{KO}$ frozen-thawed-permeabilized mitochondria in the presence of cyt c (n=3). (b) Oxygen consumption rates in mtDNA$^{WT}$ and ND4$^{KO}$ frozen-thawed-permeabilized mitochondria in the presence of cyt c (n=3). (c) Relative contribution of Na$^+$ to ΔΨmt in mtDNA$^{WT}$ mitochondria respiring on different substrates, calculated as the ratio of the calibrated TMRM signal in Na$^+$-containing buffer to that in Na$^+$-free buffer. (d-e) Mitochondrial matrix H$^+$ pumping measured from the BCEFC-AM signal in mtDNA$^{WT}$ isolated mitochondria respiring on CI (d) or CII (e) substrates in Na$^+$-containing and Na$^+$-free buffer. (f) Relative contribution of Na$^+$ to ΔΨmt in ND4$^{KO}$ mitochondria respiring on CI, CII, CI+ADP, or CIV substrates, calculated as the ratio of the calibrated TMRM signal in Na$^+$-containing to that in Na$^+$-free buffer. (g-h) Mitochondrial matrix H$^+$ pumping measured from the BCEFC-AM signal in ND4$^{KO}$ isolated mitochondria respiring on CI (g) or CII (h) substrates in Na$^+$-containing and Na$^+$-free buffer. (i-j) Na$^+$ gradient in mtDNA$^{WT}$, ND6$^{KO}$, and ND4$^{KO}$ isolated mitochondria incubated with SBF1-AM and respiring on GM (i) or succinate (j), calculated as the ratio of the matrix Na$^+$ concentrations determined after and before the addition of rotenone (i) or antimycin A (j). (k-l) Na$^+$ gradient in NDUFS4$^{WT}$ and NDUFS4$^{KO}$ isolated mitochondria incubated with SBF1-AM and respiring on GM (k) or succinate (l), calculated as the ratio of the matrix Na$^+$ concentrations determined after and before the addition of rotenone (k) or antimycin A (l). (m-n) Mitochondrial matrix H$^+$ pumping measured from the BCEFC-AM signal in control vs LHON isolated mitochondria respiring on CI (m) or CII (n) substrates in Na$^+$-containing buffer. (o) Antimycin A-sensitive succinate-cytochrome c oxidoreductase activity in LHON vs control mitochondrial membranes. (p-q) Na$^+$ gradient in control versus LHON isolated mitochondria incubated with SBF1-AM and respiring on GM (p) or succinate (q), calculated as the ratio of the matrix Na$^+$ concentrations determined after and before the addition of rotenone (p) or antimycin A (q). (r-s) Calibrated TMRM signal in control versus LHON respiring with CI (r) or CI+ADP (s) substrates.
l), indicating that formation of the Na\(^+\) gradient requires a fully assembled CI P-module. Measurement of the cytosolic and mitochondrial Na\(^+\) content in intact cells confirmed that (1) there are different concentrations of Na\(^+\) in the cytosol and mitochondria (Extended Data Figure 7a), that (2) total Na\(^+\) content was similar in all the cell models (Extended Data Figure 7b) and (3) that the transmembrane Na\(^+\) gradient exists not only in isolated mitochondria, but also in a similar magnitude in intact cells (Extended Data Figure 7b).

From the measurements taken throughout this study, we calculate that for respiration driven by CI or CII substrates the Na\(^+\) gradient contributes around one third of the total ΔΨ\(_{mt}\) (Table S2). However, when CV is activated by the presence of ADP (state 3) and partially dissipates ΔpH, the electrical contribution of the Na\(^+\) gradient reaches approximately half of the ΔΨ\(_{mt}\) (Table S2). The mitochondrial Na\(^+\) gradient thus either parallels the mitochondrial H\(^+\) gradient (compare mtDNA\(^{WT}\) in Extended Data Figure 7d and e with mtDNA\(^{WT}\) in Figure 4i and j) or even surpasses it (compare NDUFS4\(^{WT}\) in Extended Data Figure 7f and g with NDUFS4\(^{WT}\) in Figure 4k and l). This near-equal contribution of the Na\(^+\) gradient and the H\(^+\) gradient to ΔΨ\(_{mt}\) in coupled-respiring mitochondria predicts that mutations of key CI P-module residues, altering CI NHE activity may alter ΔΨ\(_{mt}\), independently of their CI NADH-CoQ oxidoreductase activity and H\(^+\) pumping capacity. Indeed, LHON mitochondria showed no differences in H\(^+\) pumping capacity under CI substrates and higher values for CI-dependent respiration (Figure 4m and n), which fits well with CI (Figure 1a) and CII+III activities (Figure 4o). In accordance with a lesser NHE activity and its contribution to the build-up of ΔΨ\(_{mt}\), LHON mitochondria showed lower Na\(^+\) gradient under CI and CII substrates (Figure 4o and p), depolarized mitochondria under CI substrates and absence of hyperpolarization with CII substrates (Figure 4q and r). This was confirmed by measuring ΔΨ\(_{mt}\) in LHON intact cybrids (Extended Data Figure 7h).

The results reported here demonstrate that CI operates as an NHE in isolated mitochondria and intact cells, regardless if CI is in its active conformation, and that doing that, CI builds up a Na\(^+\) gradient across the IMM that significantly contributes to the ΔΨ\(_{mt}\). CI-NHE activity has a H\(^+\)-dissipating effect (ΔpH becomes partially substituted by a Na\(^+\) gradient), and this effect allows higher respiration rates at lower levels of CI NADH-CoQ oxidoreductase activity. Indeed, NDUFS4\(^{KO}\) mitochondria supplied with CI substrates were able to respire and build up ΔΨ\(_{mt}\) to a similar extent as their wild type counterparts (CI panel in Figure 3f and l and Extended Data Figure 3f) despite their lower CI activity (Extended Data Figure 1a), highlighting that higher CI-NHE activity (Figure 1i and k, and Extended Data Figure 6e) allows the formation of a mitochondrial Na\(^+\) gradient (Figure 4k) at a lower available H\(^+\) gradient (Extended Data Figure 7f). In parallel, because ND6\(^{KO}\) and ND4\(^{KO}\) cells were incapable of forming a mitochondrial Na\(^+\) gradient (Figure 4i-j), they were unable to increase mitochondrial respiration or hyperpolarize (Figure 3a-e and Extended Data Figure 3a-e) despite having higher CII-III activity (Figure 2c-d). In other words, the presence of CI-NHE activity and the concomitant formation of a mitochondrial Na\(^+\) gradient enables the control of ΔΨ\(_{mt}\), in addition to ΔpH, and a more efficient use of substrates in terms of oxygen consumption. Indeed, a specific deficiency in the CI-NHE function, independently of CI NADH-CoQ oxidoreductase activity, contributes to the molecular mechanism of the m.11778G>A mutation at ND4, responsible for the mtDNA-linked Leber hereditary optic neuropathy, through deregulation of mitochondrial Ca\(^{2+}\) homeostasis. The discovery that the Na\(^+\) gradient controls ΔΨ\(_{mt}\), together with its tight regulation by a non-canonical NHE function of CI, introduces a new and unexpected layer of regulation to mitochondrial bioenergetics, with extensive implications for neurological physiology and disease.
Main references


2. Mitchell, P. Coupling of phosphorylation to electron and hydrogen transfer by a chemiosmotic type of mechanism. *Nature* **191**, 144-148 (1961). [https://doi.org/10.1038/191144a0](https://doi.org/10.1038/191144a0)


24 Stolpe, S. & Friedrich, T. The *Escherichia coli* NADH:ubiquinone oxidoreductase (complex I) is a primary proton pump but may be capable of secondary sodium antiport. *J Biol Chem* **279**, 18377-18383 (2004). [https://doi.org/10.1074/jbc.M311242200](https://doi.org/10.1074/jbc.M311242200)


Acknowledgments

We thank Dr. Sara Cogliati (CMBSO, UAM-CSIC) for helping with TEM imaging; Andrea Curtabbi, Carmen Morales Vidal, Dr. Demetrio Julián Santiago Castillo, Dr. Silvia Priori (CNIC), Dr. Eduardo Rial (CIB-CSIC) for fruitful discussion, and M. M. Muñoz-Hernandez, R. Martínez de Mena, E.R. Martínez Jiménez, and C. Jiménez for technical assistance.

Author contributions

Conceptualization: PHA and JAE. Methodology: PHA, YMM (Opa1 western blot) and EC (Proteomics). Investigation: PHA. Funding acquisition: JAE and JV. Project administration: JAE. Writing: PHA and JAE. Writing – review & editing: PHA, YMM, EC, JV and JAE.

Competing interests’ declaration

Authors declare that they have no competing interests.

Additional information
**Funding:** This study was supported by competitive grants from the *Ministerio de Ciencia e Innovación* (MCIN) RTI2018-099357-B-100, and CIBERFES (CB16/10/00282), Human Frontier Science Program (grant RGP0016/2018), and Leducq Transatlantic Networks (17CVD04) to JAE. PHA is supported by a JdC IJC2020-042679-I, YMM is supported by a FPI-SO fellowship PRE2018-083478. The CNIC is supported by the ISCIII, the MCIN and the Pro CNIC Foundation, and is a Severo Ochoa Center of Excellence

**Data and materials availability:** All data are available in the main text or the supplementary materials. Cell lines will be available under a materials transfer agreement (MTAs).

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**Extended Data**

**Materials and methods**

**Cell culture and treatment**

mtDNAWT cybrids, ND6KO cybrids, ND4KO cybrids, hmtDNAWT cybrids, LHON cybrids, NDUFS4WT mouse adult fibroblasts (MAFs), NDUFS4KO MAFs, and human control and LHON cybrids were cultured in DMEM supplemented with 5% heat-inactivated FBS, 100 U/mL penicillin, and 100 µg/mL streptomycin. All cells were correctly recognized from proteomic analysis, CI assembly assessed by Blue Native polyacrylamide gel electrophoresis (BN-PAGE), and CI spectrophotometric activity.

Chronic rotenone (Chronic Rot) treatment consisted of incubating mtDNAWT cybrids with 250 nM rotenone for 4-6 hours. This rotenone concentration was maintained throughout the detachment procedure in all buffers (trypsin and PBS) until the cell pellet was placed on ice, since removing rotenone or increasing its content during detachment was found to uncouple mitochondria.

All cultures were routinely checked for mycoplasma contamination and tested negative.

**Mitochondria isolation**

Mitochondria were isolated from cybrids, MAFs, or mouse heart using a protocol adapted for either cell culture or mouse tissues [28]. Briefly, hearts and liver were cleaned of blood by rinsing in ice-cold phosphate buffered solution (PBS), and the tissue was disrupted with a blade on ice. Disrupted tissue, cybrids, or MAFs were resuspended in a buffer containing 0.32 M sucrose, 10 mM Tris-base and 1 mM EDTA (pH 7.4), placed in a glass Elvehjem potter, and homogenized by up and down strokes using a motor-driven Teflon pestle. Successive homogenization–centrifugation steps yielded the mitochondria-containing fraction, which was quantified with the Bradford protein assay.

**Mitochondrial membrane isolation and measurement of complex activities**

Mitochondrial membranes from cultured cells or liver were obtained after freeze–thawing isolated mitochondria, and OXPHOS enzyme activity was measured as previously described [29-30], using between 0.5 and 20 µg protein per sample, depending on the activity to be measured. Rotenone-sensitive NADH-ubiquinone decylubiquinone (DQ) oxidoreduction (CI activity) was measured by changes in absorbance at 340 nm. Antimycin A-sensitive succinate-cyt c oxidoreductase activity (CII + CIII activity) was calculated after measuring changes in absorbance at 550 nm. Potassium cyanide (KCN)-sensitive cyt c-oxygen oxidoreductase activity (CIV activity) was calculated after measuring changes in absorbance at 550 nm. Oligomycin-
sensitive ATPase activity (CV activity) was calculated after measuring changes in absorbance at 340 nm driven by the pyruvate kinase reaction coupled to ADP phosphorylation by CV.

**Measurement of O2 consumption and mitochondrial membrane potential**

Measurements were made with an O2k Oxygraph instrument (Oroboros Instruments) with the attached fluorescent module. Tetramethylrhodamine methyl ester (TMRM) in quenching mode (2 μM) was used for the detection of mitochondrial membrane potential. Oxygen and TMRM calibration were performed before every experiment. Cells (2x10^6) or isolated mitochondria or mitochondrial membranes (50 μg) were loaded in the O2k Oxygraph chambers with Mir05 buffer (Oroboros Instruments) and 20 μg digitonin (only for cells); substrates and inhibitors were added sequentially. Glutamate (10 mM) and malate (0.5 mM) were added to activate CI-dependent respiration and rotenone (0.5 μM) was added to suppress it. Succinate (10 mM) was then added to activate CII-dependent respiration, and subsequently 1 mM ADP was added to activate state 3 respiration; both of which were inhibited by adding 2.5 μM antimycin A. Lastly, 0.5 mM N, N', N', N'-tetramethyl-p-phenylenediamine (TMPD) and 2 mM ascorbate were added to activate CIV-dependent respiration, which was later suppressed by addition of 100 mM sodium azide. In some experiments, 10 μM cyt c was included in the sequence. In some other experiments, 100 nM nigericin or 100 nM monensin was added after succinate.

Na^+-containing respiration buffer was prepared by replacing potassium phosphate with sodium phosphate in the Mir05 formula (Oroboros instruments). Na^+-free respiration buffer was made with phosphoric acid and buffered to physiological pH with N-methyl D-glucamine (NMDG). Both buffers were compensated in osmolality and ionic strength using NMDG. In addition, in these experiments the acidic forms of substrates, appropriately resuspended in Na^+-free respiration solution and buffered to reach respiration solution pH, were used to avoid the introduction of Na^+ into the Na^+-free respiration buffer or additional Na^+ into the Na^+-containing buffer. Mitochondrial membrane potential was calibrated according to the manufacturer’s instructions.

**Blue native gel electrophoresis**

Complex and supercomplex levels, compositions, and distributions were analyzed in isolated mitochondria by BNGE. Mitochondrial proteins were solubilized with 10% digitonin (4 g/g; Sigma-Aldrich D5628) and separated on 3–13%-gradient Blue native gels. Gradient gels (1.5-mm thick) were prepared using a gradient former connected to a peristaltic pump.

**SDS gel electrophoresis**

Mitochondria were resuspended in RIPA buffer (1% Triton-X-100, 50 mM Tris-HCl pH 7.4, 50 mM sodium chloride, 0.5% sodium deoxycholate, and 5 mM EDTA) supplemented with a protease inhibitors cocktail mix (Sigma P8340). Samples were then incubated at 4°C on a rotating wheel for 15 minutes and centrifuged for 15 minutes at 13,000 x g at 4°C. Supernatants were collected and transferred to fresh 1.5 ml Eppendorf tubes. Protein concentration was quantified by the Bradford assay.

The levels and distribution of Opa1 were analyzed by sodium dodecyl sulphate (SDS) gel electrophoresis (SDSGE) on 7.5% gels. Loading buffer (50 mM Tris-HCl pH 6.8, 2% SDS, 10% glycerol, 1% 2-mercaptoethanol, 0.02% bromophenol blue) was added to samples, which were incubated for 5 minutes at 95°C.

**Immunodetection of single proteins, complexes, and supercomplexes**
BNGE or SDSGE proteins were electrobotted using a Mini Trans-Blot Cell system (Bio-Rad) onto polyvinylidene difluoride (PVDF) transfer membrane (Immobilon-FL, 0.45 μm; Merck Millipore, IPFL00010) for 1 hour at 100 V in transfer buffer (48 mM Tris, 39 mM glycine, and 20% ethanol). Non-specific binding sites were blocked by incubating membranes with phosphate-buffered saline (PBS) containing 5% bovine serum albumin (BSA) for 1 hour at RT. For protein detection, membranes were incubated in blocking buffer containing primary antibody overnight at 4°C. After three washes with PBS 0.1% Tween-20 (PBS-T) for 30 min, membranes were incubated with appropriate secondary antibodies for 1 hour at RT. Membranes were then washed twice with PBS-T and once with PBS. To study complex and supercomplex assembly, the PVDF membrane was sequentially probed with antibodies against Cl (anti-NDUFA9, Abcam ab14713), CIII (anti-UQCR2, Proteintech, 14742-1 AP) and CII (anti-Fp70, Invitrogen, 459200). Parallel BNGE western blots were performed to probe CIV (anti-COI, Invitrogen, MTCO1 459600) and CV (anti-ATPβ, Abcam, Ab 14730). To study Opa1 processing, the PVDF membrane was sequentially probed with antibodies against Opa1 (anti-Opa1, Abcam, ab42364) and Fp70 (anti-Fp70, Invitrogen, 459200). Fiji software was used for Opa1 quantification. Antibody binding was detected by fluorescence as previously described.

Measurement of mitochondrial ATP synthesis

ATP synthesis was assessed in isolated mitochondria (1.5 to 15 μg mitochondrial protein) using succinate as substrate (plus rotenone) in the presence of ADP in a kinetic luminescence assay based on the luciferin/luciferase reaction.

Measurements of mitochondrial matrix Na+ and H+

The probe-loading protocol was adapted from. Isolated mitochondria from cells were incubated for 20 min at 37 °C with 10 μM 2′,7′-bis(2-carboxyethyl)-5-(and-6)-carboxyfluorescein-acetoxyethyl ester (BCECF-AM; for matrix H+) or with 1,3-benzenedicarboxylic acid, 4,4′-[1,4,10-trioxa-7,13-diazacyclopentadecane-7,13-diylbis(5-methoxy-6,12-benzofurandiy)]bis-,tetakis [(acetylxy)methyl] ester (SBFI-AM; for matrix Na+). This was followed by two rounds of centrifugation (12000 g, 4°C, 5 min) and resuspension in sucrose buffer. The last resuspension was in Na+-free respiration buffer.

Probe-loaded mitochondria were distributed in p96 wells in Na+-free buffer, unless otherwise indicated. Measurements were made with a Fluoroskan Ascent microplate reader (Thermo Fisher Scientific) at 390/485 nm excitation pair for BCECF-AM and 355/390 nm excitation pair for SBFI-AM. Emission was recorded at 530 nm emission for BCECF-AM and SBFI-AM. Substrates and inhibitors were subsequently added as in the O2 consumption experiments, without ADP, TMPD, ascorbate, or sodium azide. The H+ pumping rate was obtained from the slope after adding respiratory substrates; the slope was flat after the addition of the corresponding inhibitors. For the measurement of reverse NHE activity using BCECF-AM, 10 mM NaCl was added after succinate. For measurement of forward NHE activity and Na+ gradient using SBFI-AM, 10 mM NaCl was added at the beginning of the experiment.

Calibration was performed by adding equal amounts of mitochondria from the same cell type to several wells containing a graded series of pH or Na+ concentration in the presence of 1 μM nigericin, 1 μM monensin, and 1 μM gramicidin.

Spectrophotometric measurement of NHE activity in isolated mitochondria

The protocol was adapted to the analysis of cell-culture-derived mitochondria from. Briefly, passive mitochondrial swelling was recorded in 50 μg of mitochondria resuspended in 133 mM
sodium acetate, 0.2 mM Tris-EGTA, pH 7.0 at room temperature. Absorbance was recorded at 550 nm in a UV/VIS JASCO spectrophotometer (Thermo Fisher Scientific).

In some experiments, sodium acetate was exchanged for sodium chloride as a swelling control. In others, before addition of mitochondria to the sodium acetate buffer, mitochondrial CI was activated by incubating the sample with glutamate/malate in Miro05 buffer for 5 min at 37ºC. Alternatively, mitochondrial CI was deactivated by incubating the mitochondria without substrates in Miro05 buffer for 15 min at 37ºC.

**Measurement of Na⁺ gradient in whole cells by confocal microscopy**

Cytosolic and mitochondrial Na⁺ were detected as in29. Briefly, cybrids were plated the day before experiments, washed three times with Hank’s balanced salt solution with Ca²⁺/Mg²⁺/glucose (HBSS + Ca/Mg + glucose), and incubated with 5 μM Asante NaTRIUM Green-2-Acteoxymethyl ester (ANG2-AM) for 30 min at 37 ºC in the dark. ANG2-AM was washed out, and new HBSS + Ca/Mg + glucose was added, including 1 μM CoroNa Red. Cells were further incubated for 30 min at 37 ºC in the dark. After this, the medium was changed again, cells were washed once with HBSS + Ca/Mg + glucose, and the plate was placed on the automated stage of a Leica SP-5 confocal microscope for live imaging. The planes were focused for image capture, and images were taken with a ×63 objective. Experiments started and ended at 20% O₂ and 5% CO₂. Loaded cells were excited with an argon/krypton laser using the 496-nm line for ANG2-AM and the 514-nm line for CoroNa Red. Fluorescence emission of ANG2-AM was detected in the 515–550-nm range, whereas the emission of CoroNa Red was detected in the 555–575-nm range.

*In situ* calibration of the same cells was performed after a two-wash step with Na⁺-free HBSS + Ca/Mg + glucose. Increasing Na⁺ concentrations, starting from 0 mM Na⁺, were then applied in the presence of 1 μM nigericin, 1 μM monensin, and 1 μM gramicidin, and images were taken with the excitation/emission wavelengths indicated above. Calibration solutions were equilibrated for at least 5 min.

**Measurement of mitochondrial Ca²⁺ in whole cells by confocal microscopy**

The protocol followed was exactly as in29. Briefly, cells were transiently transfected with Cepia2mt and mitochondrial Ca²⁺ imaged under a Leica SP5 microscope. Mitochondrial Ca²⁺ entry was stimulated by addition of ATP 40 μM and calibrated using ionomycin 1 μM and EDTA 2.5 mM.

**Measurement of ΔΨmt in whole cells by confocal microscopy**

Cells were seeded the day prior experimentation. Then, cells were incubated with TMRM 50 nM, for non-quenching mode, for 30 min before imaging in a Leica SP5 microscope. Samples were excited with an argon/krypton laser using the 543-nm line and emission was detected in the 555–595-nm range. Calibration was performed following guidelines and indications from PMID: 22495585***.

**Measurement of mitochondrial volume**

For the determination of mitochondrial volume 50 μg of mitochondria and 1 mL Mir05 respiration buffer were placed in a cuvette, and the absorbance at 550 nm was recorded with a UV/VIS JASCO spectrophotometer (Thermo Fisher Scientific). Parallel measurements were taken at steady state and in mitochondria respiring from different substrates for 5 min at 37 ºC. A
valinomycin technical control was included to determine whether this technique was suitable for the measurement of mitochondrial volume changes, with positive results.

Transmission electron microscopy

Cells were analyzed by electron microscopy as previously described. Thin sections including mitochondria were imaged with a JEM1010 electron microscope (Jeol).

Proteomic analysis

Protein extracts of isolated mitochondria from mtDNAWT, ND6KO, ND4KO, NDUFS4WT, and NDUFS4KO samples were obtained by homogenization with ceramic beads (MagNa Lyser Green Beads, Roche, Germany) in CS buffer (Pipes pH 6.8, MgCl2, NaCl, EDTA, sucrose, SDS, sodium orthovanadate; Biochain Institute, Inc. #K3013010-5) freshly supplemented with protease and phosphatase inhibitors. Extracted proteins (around 200 μg) from atrial biopsies were subjected to in-filter reduction and alkylation using iodoacetamide followed by trypsin digestion (Nanosep Centrifugal Devices with Omega Membrane-10K, PALL), and the resulting peptides were TMT-labeled according to the manufacturer’s instructions. Labeled peptides were subjected to LC-MS analysis using a Proxeon Easy nano-flow HPLC system (Thermo Fisher Scientific, Bremen, Germany) coupled via a nanoelectrospray ion source (Thermo Fisher Scientific) to an Orbitrap Fusion mass spectrometer (Thermo Fisher) using with a 2-cm trap column and a 50-cm analytical column (75 μm I.D, 2 μm particle size, Acclaim PepMap RSLC, 100 C18; Thermo Fisher Scientific) in a continuous acetonitrile gradient consisting of 0–30% A for 60 min, 50–90% B for 3 min (A= 0.1% formic acid; B= 100% acetonitrile, 0.1% formic acid) at a flow rate of 200 nL/min. Mass spectra were acquired in a data-dependent manner, with an automatic switch between MS and MS/MS using a top-speed method and dynamic exclusion. MS spectra were collected in the Orbitrap analyzer using a mass range of 400–1500 m/z at 60,000 resolution. HCD fragmentation was performed at 33 eV of normalized collision energy and MS/MS spectra were analyzed at 30,000 resolution in the Orbitrap.

Proteins were identified with the SEQUEST HT algorithm integrated in Proteome Discoverer 2.5 (Thermo Scientific). MS/MS scans were searched against a pig reference proteome database (human_pig_202105_pro-sw-tr.target-decoy.fasta), (296316 sequences in total). For database searching, parameters were selected as follows: trypsin digestion with 2 maximum missed cleavage sites, precursor mass tolerance of 2 Da, and a fragment mass tolerance of 0.03 Da. Methionine oxidation (+15.994915 Da) and asparagine and glutamine deamidation (+0.984016 Da) were set as variable modifications, whereas cysteine carbamidomethylation (+57.021464 Da) and TMT labeling (+229.162932 Da) at peptide N-terminal ends and Lys residues were considered fixed modifications. False discovery rates (FDR) for peptide identifications were calculated by the refined method with an additional filter for a precursor mass tolerance of 10 ppm. A 1% FDR was used as the criterion used for peptide identification.

Quantitative information from TMT reporter intensities was integrated from the spectrum level to the peptide level and then to the protein level based on the WSPP model, using the GIA integration.

Statistical analysis

Statistical analyses and graphics were produced with GraphPad Prism 8 software. Datasets were compared by t test, analysis of variance (ANOVA), or nonparametric analysis as appropriate and with P values adjusted for multiple tests. P values from each comparison are shown in every graph. All results are presented as mean ± SD or mean ± SEM.
Extended Data Figure 1. Isolated complex activities in CI-deficiency models. (a) Rotenone (Rot)-sensitive NADH-CoQ oxidoreductase activity in mtDNAWT, ND6KO and ND4KO, and NDUFS4WT and NDUFS4KO mitochondrial membranes from MAFs and Liver. (b) Antimycin A (AA)-sensitive succinate-cyt c oxidoreductase activity in NDUFS4WT and NDUFS4KO mouse liver mitochondrial membranes. (c) KCN-sensitive cyt c oxidase activity in NDUFS4WT and NDUFS4KO mouse liver mitochondrial membranes. (d) Oligomycin (Olig)-sensitive ATPase activity in mtDNAWT, ND6KO, ND4KO, Vehicle-treated, Chronic rotenone-treated and NDUFS4WT and NDUFS4KO mitochondrial membranes from MAFs and Liver. (e) Olig-sensitive ATP synthase activity in mtDNAWT, ND6KO, ND4KO, Vehicle-treated, Chronic rotenone-treated and NDUFS4WT and NDUFS4KO isolated mitochondria in the presence of 1 µM Rot and respiring on succinate.
Extended Data Figure 2. Bioenergetics of the Chronic Rot CI-deficiency model. (a) Oxygen consumption rates were measured in permeabilized mtDNA WT cells that had been treated with vehicle or Chronic Rot (n=4). Insets and acronyms are the same as those in Fig. 2; FCCP, carbonyl cyanide-p-trifluoromethoxyphenylhydrazone. (b-d) Calibrated TMRM signal in permeabilized mtDNA WT cells that had been treated with vehicle or Chronic Rot, respiring on CI substrates (b), CII substrates (C), or CII substrates +ADP (d).
Extended Data Figure 3. Bioenergetics analysis of isolated mitochondria from the CI deficiency models. (a) Oxygen consumption rates in isolated mitochondria from mtDNA\(^{\text{WT}}\) and \(\text{ND6}^{\text{KO}}\) (n=3) cells. (b) Oxygen consumption rates in isolated mitochondria from mtDNA\(^{\text{WT}}\) and \(\text{ND4}^{\text{KO}}\) (n=3) cells. (c) Oxygen consumption rates in isolated mitochondria from \(\text{NDUFS4}^{\text{WT}}\) and \(\text{NDUFS4}^{\text{KO}}\) (n=3) cells. Abbreviations are the same as in Fig. 2. (d) Calibrated TMRM signal in \(\text{ND6}^{\text{KO}}\), \(\text{ND4}^{\text{KO}}\) (e) and \(\text{NDUFS4}^{\text{KO}}\) (f) mitochondria respiring on CI substrates (CI inset), CII substrates (CII inset), CII substrates +ADP (CII+ADP inset), and CIV substrates (CIV inset).
Extended Data Figure 4. ND6KO and ND4KO mitochondria have similar size, H⁺ leak, and MTP permeability to mtDNAWT mitochondria. (a) Mitochondrial volume measured by absorbance at 550 nm in the presence of CI substrates (Glut/Mal), CI substrates + ADP (Glut/Mal+ADP), or CII substrates + 1 µM rotenone (Rot/Succ+ADP) in mtDNAWT, ND6KO and ND4KO mitochondria. (b) mtDNAWT and ND6KO mitochondrial volume in the absence or presence of valinomycin. (c) Transmission electron microscopy (TEM) imaging of mitochondria in mtDNAWT, ND6KO, and ND4KO cells (n=2). (d) H⁺ leak contribution to baseline CII respiration +/- ADP in Vehicle and Chronic Rot-treated, as well as in mtDNAWT and ND4KO permeabilized cells, calculated as the ratio of baseline CII respiration (without ADP) to H⁺ leak. (e) ATP synthesis contribution to CII respiration in Vehicle and Chronic Rot-treated, as well as in mtDNAWT and ND4KO permeabilized cells, calculated as the ratio of coupled CII respiration (with ADP) to ATP synthesis. (f) Oxygen consumption rates in mtDNAWT, ND6KO, and ND4KO permeabilized cells respiring on Succ + Rot + ADP in the absence or presence of cyt c. (g) Calibrated TMRM signal in mtDNAWT, ND6KO, and ND4KO permeabilized cells respiring on Succ + Rot + ADP in the absence or presence of cyt c. (h) Western blot analysis of OPA1 expression in mtDNAWT, ND6KO, and ND4KO isolated mitochondrial fractions; anti-Fp70 was used as a loading control. Inset: shows quantification of three independent experiments.
Extended Data Figure 5. Proteomics analysis of mtDNA<sup>WT</sup>, ND6<sup>KO</sup>, ND4<sup>KO</sup>, NDUFS4<sup>WT</sup>, and NDUFS4<sup>KO</sup> isolated mitochondria. Relative mean abundances of (a) CI subunits in mtDNA<sup>WT</sup>, ND6<sup>KO</sup>, and ND4<sup>KO</sup> mitochondria; (b) CII subunits in mtDNA<sup>WT</sup>, ND6<sup>KO</sup>, and ND4<sup>KO</sup> mitochondria; (c) CIII subunits in mtDNA<sup>WT</sup>, ND6<sup>KO</sup>, and ND4<sup>KO</sup> mitochondria; (d) CIV subunits in mtDNA<sup>WT</sup>, ND6<sup>KO</sup>, and ND4<sup>KO</sup> mitochondria; (e) of CI subunits in NDUFS4<sup>WT</sup> and NDUFS4<sup>KO</sup> mitochondria; (f) of CII subunits in NDUFS4<sup>WT</sup> and NDUFS4<sup>KO</sup> mitochondria; (g) of CIII subunits in NDUFS4<sup>WT</sup> and NDUFS4<sup>KO</sup> mitochondria and (h) of CIV subunits in NDUFS4<sup>WT</sup> and NDUFS4<sup>KO</sup> mitochondria. (i-k) Volcano plot showing relative abundances of all mitochondrial proteins detected in (i) mtDNA<sup>WT</sup> vs ND6<sup>KO</sup>; (j) in mtDNA<sup>WT</sup> vs ND4<sup>KO</sup>; (k) in NDUFS4<sup>WT</sup> vs NDUFS4<sup>KO</sup>. For all experiments, n=3 three independent biological replicates.
Extended Data Figure 6. CI-NHE activity-dependent Na⁺ gradient contributes to total ΔΨmt. (a) Dose-response of mouse liver mitochondrial NHE activity to rotenone. (b) Dose-response of mouse liver mitochondrial NHE activity to DPI. (c) Mouse liver mitochondrial NHE activity in different CI activation states. (d) NHE activity in mtDNAWT, ND6KO and ND4KO mitochondria. (e) NHE activity in NDUSF4WT and NDUSF4KO mitochondria. (f) Calibrated BCECF-AM signals of mtDNAWT, ND6KO and ND4KO isolated mitochondria under CI (Inset CI-CIII-CIV) and CII substrates (Inset CII-CIII-CIV). (g) Calibrated BCECF-AM signals of NDUSF4WT and NDUSF4KO isolated mitochondria under CI (Inset CI-CIII-CIV) and CII substrates (Inset CII-CIII-CIV). (h) Calibrated TMRM signal before and after the addition of nigericin (100 nM), monensin (100 nM) in mtDNAWT and ND4KO or monensin (100 nM) in NDUSF4WT and NDUSF4KO isolated mitochondria respiring in Rot/Succ + ADP. (i-k) Oxygen consumption rates measured in mtDNAWT (i), ND4KO (j) and C57BL/6N heart (k) mitochondria in Na⁺-containing buffer (black or reddish lines) and Na⁺-free buffer (gray or brownish lines; n=4). (l) Relative contribution of Na⁺ to ΔΨmt in C57BL/6N heart mitochondria respiring on CI, CII, CII+ADP, or CIV substrates, calculated as the ratio of the calibrated TMRM signal in Na⁺-containing to that in Na⁺-free buffer. (m) Oxygen consumption rates measured in mtDNAWT and ND4KO mitochondria in Na⁺-free buffer (n=4). (n) Calibrated TMRM signal in mtDNAWT and ND4KO mitochondria respiring on CI (CI inset), CII (CII inset), CII+ADP (CII+ADP inset), or
CIV (CIV inset) substrates in a Na⁺-free buffer. Abbreviations in f, g, i-k and m are the same as those in Fig. 2
Extended Data Figure 7. The mitochondrial Na\(^+\) gradient parallels the H\(^+\) gradient and contributes to \(\Delta\psi_{\text{mt}}\) in intact cells. (a) Cytosolic and mitochondrial Na\(^+\) concentrations in mtDNA\(^{\text{WT}}\), ND6\(^{\text{KO}}\), and ND4\(^{\text{KO}}\) intact cells measured by confocal microscopy. (b) Total Na\(^+\) concentrations calculated as the sum of cytosolic and mitochondrial Na\(^+\) concentrations, measured by confocal microscopy. (c) Mitochondrial Na\(^+\) gradient calculated as the ratio between cytosolic and mitochondrial Na\(^+\) concentrations measured by confocal microscopy. (d) H\(^+\) gradient in mtDNA\(^{\text{WT}}\), ND6\(^{\text{KO}}\) and ND4\(^{\text{KO}}\) isolated mitochondria incubated with BCECF-AM and respiring on GM, calculated as the ratio of matrix H\(^+\) concentrations determined after and before addition of rotenone. (e) H\(^+\) gradient in mtDNA\(^{\text{WT}}\), ND6\(^{\text{KO}}\), and ND4\(^{\text{KO}}\) isolated mitochondria incubated with BCECF-AM and respiring on succinate + rotenone 1µM, calculated as the ratio of matrix H\(^+\) concentrations determined after and before addition of AA. (f) H\(^+\) gradient in NDUFS4\(^{\text{WT}}\) and NDUFS4\(^{\text{KO}}\) isolated mitochondria incubated with BCECF-AM and respiring on GM, calculated as the ratio of matrix H\(^+\) concentrations determined after and before addition of rotenone. (g) H\(^+\) gradient in NDUFS4\(^{\text{WT}}\) and NDUFS4\(^{\text{KO}}\) isolated mitochondria incubated with BCECF-AM and respiring on succinate + rotenone 1µM, calculated as the ratio of matrix H\(^+\) concentrations determined after and before addition of AA. (h) Calibrated TMRM signal measured by confocal microscopy in intact control vs LHON cybrids.
Extended Data Table 1. Summary of bioenergetics characterization of CI-deficiency models.

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<tr>
<th>Comparisons vs Control</th>
<th>CII+III activity</th>
<th>CIV activity</th>
<th>CV activity</th>
<th>Respiration</th>
<th>( \Delta \Psi_{mt} )</th>
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<td>ND6&lt;sup&gt;KO&lt;/sup&gt;</td>
<td>Higher (Fig. 2c)</td>
<td>Equal (Fig. 2g)</td>
<td>Equal (ED Fig. 1d-e)</td>
<td>Equal (Fig. 3a and ED Fig. 3a)</td>
<td>Equal (Fig. 3d and ED Fig. 3d)</td>
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<tr>
<td>ND4&lt;sup&gt;KO&lt;/sup&gt;</td>
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<td>Higher (Fig. 2h)</td>
<td>Equal (ED Fig. 1d-e)</td>
<td>Higher (Fig. 3b and ED Fig. 3b)</td>
<td>Equal (Fig. 3e and ED Fig. 3e)</td>
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<tr>
<td>NDUFS4&lt;sup&gt;KO&lt;/sup&gt;</td>
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<td>Higher (Fig. 2i)</td>
<td>Higher (ED Fig. 1d-e)</td>
<td>Higher (Fig. 3c and ED Fig. 3c)</td>
<td>Hyperpolarized (Fig. 3f and ED Fig. 3f)</td>
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<tr>
<td>Chronic Rot</td>
<td>Higher (Fig. 2f)</td>
<td>Equal (Fig. 2)</td>
<td>Equal (ED Fig. 1d-e)</td>
<td>Higher (ED Fig. 2e)</td>
<td>Hyperpolarized (ED Fig. 2c and d)</td>
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Extended Data Table 2. Summary of total and relative contributions of the Na\textsuperscript{+} gradient to ΔΨ\textsubscript{mt} in the presence of different respiratory substrates and oxidizing conditions.

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<th>Estimation approach</th>
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<th>Contribution to ΔΨ\textsubscript{mt} (%)</th>
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<td></td>
<td>ΔΨ\textsubscript{mt} measurement in mtDNA\textsuperscript{WT± Na\textsuperscript{+}}</td>
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