Technical Communication

Cite

Author contributions
EG and TK collaborated closely with WGT in the development of the Q-Module, with guidance from ALM and PRR. TK and LHDC designed, carried out and analyzed the experiments. CD contributed to SUIT protocol development. TK, LHDC and EG co-wrote the manuscript. All authors commented on and approved the manuscript.

Conflicts of interest
EG is founder and CEO of Oroboros Instruments, Innsbruck, Austria;
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Abstract

Redox states of the mitochondrial coenzyme Q pool, which reacts with the electron transfer system, reflect the balance between (1) reducing capacities of electron flow from fuel substrates converging at the Q-junction, (2) oxidative capacities downstream of Q to O₂, and (3) the load on the OXPHOS system utilizing or dissipating the protonmotive force.

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**Coupling and pathway control of coenzyme Q redox state and respiration in isolated mitochondria**

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coupling control
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NS-pathway
additivity

A three-electrode sensor (Rich 1988; Moore et al 1988) was implemented into the NextGen-O2k to monitor continuously the redox state of CoQ2 added as a Q-mimetic simultaneously with O2 consumption. The Q-Module was optimized for high signal-to-noise ratio, minimum drift, and minimum oxygen diffusion. CoQ2 equilibrates in the same manner as Q at Complexes I, II and III. The CoQ2 redox state is monitored amperometrically with the working electrode, which is poised at CoQ2 redox peak potentials determined by cyclic voltammetry. The voltammogram also provides quality control of the Q-sensor and reveals chemical interferences.

The CoQ2 redox state and O2 consumption were measured simultaneously in isolated mouse cardiac and brain mitochondria. CoQ2 — and by implication mitochondrial Q — was more oxidized when O2 flux was stimulated by coupling control: when energy demand increased from LEAK to OXPHOS and electron transfer capacities in the succinate pathway. In contrast, CoQ2 was more reduced when O2 flux was stimulated by pathway-control of electron input capacities, increasing from the NADH (N)- to succinate (S)-linked pathway which converge at the Q-junction, with CI-Q-CIII and CII-Q-CIII segments, respectively. N- and S-respiratory pathway capacities were not completely additive, compatible with partitioning of Q intermediary between the solid-state and liquid-state models of supercomplex organization. The direct proportionality of CoQ2 reduction and electron input capacities through the CI-Q-CIII and CII-Q-CIII segments suggests that CoQ2 is accurately mimicking mitochondrial Q-redox changes.

1. Introduction

The redox state of mitochondrial metabolites plays a key role in mitochondrial respiratory control. Analysis of O2 consumption is one of the most established methods to study mitochondrial function in health and disease. High-resolution respirometry (HRR) is a state-of-the-art technique to measure mitochondrial respiration in a wide variety of sample preparations with the application of substrate-uncoupler-inhibitor titration (SUIT) protocols (Doerrier et al 2018). Extensions of HRR in the Oroboros Oxygraph-2k (O2k, Oroboros Instruments, Innsbruck, Austria) by fluorometric or potentiometric modules allow simultaneous measurements of respiration and additional
mitochondrial parameters (e.g. mitochondrial membrane potential, ATP synthesis, hydrogen peroxide production, Ca$^{2+}$, pH, NO, H$_2$S). The Oroboros NextGen-O2k is a novel all-in-one instrument which extends HRR with the amperometric measurement of the redox state of a Q-mimetic, such as CoQ2, added to the experimental chamber. The Q-mimetic equilibrates with the part of the mitochondrial coenzyme Q-pool which reacts with the electron transfer system (ETS-reactive Q). Thus, pivotal information is obtained on the relation between Q-redox state and electron transfer at controlled O$_2$ regimes in the range of hyperoxia to anoxia, whilst saving resources (time, biological sample, and reagents), and ensuring reproducibility and accuracy of the results. Importantly, mitochondrial coenzyme Q-redox states are not measured directly, but the signal is indirect and mediated by the Q-mimetic.

Mitochondrial coenzyme Q (mtCoQ; ubiquinone; 2,3-dimethoxy-5-methyl-6-polypropenyl-1,4-benzoquinone) was discovered in 1957 by Crane and colleagues. CoQ occurs in mitochondrial and other cellular membranes. It is a lipid composed of a benzoquinone ring with an isoprenoid side chain, two methoxy groups and one methyl group (Wolf et al 1958). The number N of isoprenoid units is indicated as CoQN; for example, CoQ8 occurs in Trypanosoma brucei, CoQ9 in Caenorhabditis elegans and rodents, CoQ10 in humans. Some species have more than one CoQ form, e.g. human and rodent mitochondria contain different proportions of CoQ9 and CoQ10 (Aberg et al 1992; Awad et al 2018; Hernández-Camacho et al 2018; Watts, Ristow 2017).

CoQ is widely distributed among non-mitochondrial compartments. In hepatocytes CoQ localizes in the Golgi apparatus (Crane et al 1985; Nyquist et al 1970), peroxisomes (Turunen et al 2004), microsomes (Seshadri Sastry et al 1961), and the plasma membrane electron transfer system (review: Morré, Morré 2011). >30 % of the membrane-bound CoQ is extramitochondrial in rat liver (Kalén et al 1987; Morré, Morré 1989). Additionally, lysosomes have an ETS comparable to mitochondria where CoQ acts as an electron carrier (Gille, Nohl 2000). Consequently, isolated mitochondria may be considered as the gold standard for selective measurement of the mitochondrial Q-redox state.

Q is a central component of the mtETS (Crane et al 1959; Hatefi et al 1959, Mitchell 1961) and is involved in antioxidant defense (Noh et al 2013), mitophagy (Rodríguez-Hernández et al 2009), and regulation of permeability transition (Balaban et al 2005; Bentinger et al 2007; Fontaine et al 1998; Lopez-Lluch et al 2010). Several branches of the ETS converge at the Q-junction: In mammalian mitochondria, ETS-reactive Q (redox active, Q$_{ra}$; Kröger and Klingenberg, 1973) is reduced by electron supply from (1) Complex I (CI), (2) CII, (3) electron-transferring flavoprotein Complex, (4) mglycerophosphate dehydrogenase Complex, (5) dihydro-orotate dehydrogenase Complex, and from other enzyme complexes (Enriquez, Lenaz 2014; Gnaiger 2020). Q$_{ra}$ is oxidized downstream through CIII. Electrons are subsequently transferred via cytochrome c to CIV and the terminal electron acceptor O$_2$.

The concept of the Q-cycle was proposed originally by Mitchell (1975) and was elaborated further in several modifications, describing how CIII translocates hydrogen ions against the protonmotive force pmF (Mitchell 1961, 1975; Rich 1984; Trumpower 1990; Trumpower, Gennis 1994; Crofts 2004). CoQ exists in different protonation states
and three different redox states: ubiquinone (oxidized), ubiquinol (CoQH₂, reduced), and an intermediate semiquinone. QH₂ binds to the Qₒ site of CIII, while ubiquinone binds to the Q₁ site of CIII. First, QH₂ reduces the iron-sulfur protein and loads cytochrome c₁ with one electron. The other electron is transferred to the b₁ heme and reduces the b₉h heme, which transfers the electron to ubiquinone at the Q₁ site, reducing it to a semiquinone. A second QH₂ — oxidized at the Qₒ site — is required to fully reduce this semiquinone to ubiquinol at Q₁ site. This results in two ubiquinols oxidized at the Qₒ site per one ubiquinone reduced at the Q₁ site. In a full Q-cycle, two H⁺ leave the mt-matrix, and four H⁺ enter the intermembrane space with a change of two net transmembrane charges. The reduced cytochrome c transfers electrons further to CIV. The ubiquinol generated at the Q₁ site is recycled by binding to the Qₒ site of CIII (Hunte et al 2003; Trumpower 1990; Trumpower, Gennis 1994).

Kröger and Klingenberg analyzed the kinetic control of the Q-redox state in submitochondrial particles. An ETS-reactive Q-pool Q₉ is distinguished from an inactive pool mtCoQ₉a. At steady state the redox state of Q₉ is proportional to respiratory rate and Q₉ has been considered to be homogenous, characterized as Q-pool behavior (Kröger, Klingenberg 1966, 1973a, 1973b; Ernster et al 1969; Hackenbrock et al 1986). However, according to Gutman (1985) there is inhomogeneity of the Q₉-pool with different redox states of Q at various reduction sites (Cottingham, Moore 1983). A third Q-pool is suggested as an exo-mtCoQ-pool, reduced by NADH added to intact pigeon heart mitochondria and located on the outer face of the mtIM (Jørgensen et al 1985). Considering that lateral diffusion of Q is high in the lipid bilayer and not rate-limiting for electron transfer, the inhomogeneity can be explained by SC₁III₉IVₙ supercomplex formation (NADH oxidation through the CI-Q-III branch) in contrast to homogenous ‘Q-pool behavior’ between CI (and other dehydrogenases) and CIII (succinate oxidation; Bianchi et al 2004; Estronell et al 1992; Rauchova et al 1997; Stoner et al 1984; Enriquex, Lenaz 2014). According to the solid-state model (Rich 1984), Q-intermediates are transferred in currently considered supercomplexes Qₛ₉ by substrate channeling preventing equilibration with the free Q-pool Q₉free. The Q₉free-pool is a reservoir for slow binding to SC₁III₉IVₙ and uncoupling proteins (Bianchi et al 2003; Echtay et al 2000; Lenaz, Genova 2009). The solid-state and liquid-state (random-collision) models describe the extremes of a dynamic molecular organization, with intermediary behavior possible depending on factors such as kinetic or physical limitations (Rich 1984) leading to the plasticity model (Enriquex, Lenaz 2014).

Q-extraction is an established method for measurement of the redox state not only of the ETS-reactive Q-pool but also of the inactive mtCoQ-pool, involving extraction and determination of the total concentration of reduced quinol and oxidized quinone by high-performance liquid chromatography (Reed, Ragan 1987; Takada et al 1984; Zannoni, Moore 1990; Van den Bergen et al 1994) or electrochemical detection (Tang, Miles 2012). It has the advantage that (1) the concentrations can be determined, and (2) applications are possible in the presence of inhibitors which interfere with the Q-electrode system. In the present study, we describe the Q-Module of the NextGen-02k for real-time and simultaneous measurement of O₂ consumption and the redox state of the Q-mimetic CoQ2 based on Rich (1988) and Moore et al (1988). The advantages of this technique are (1)
real-time monitoring of continuous changes in the Q-redox state, and (2) sensitivity to the redox state of the ETS-reactive Q-pool without interference by inactive mtCoQ (Van den Bergen et al 1994).

2. Materials and methods

2.1. Reagents

ADP, adenosine 5’-diphosphate potassium salt (Calbiochem 117105) stock 500 mM with 300 mM MgCl₂·6 H₂O: 501.3 mg ADP dissolved in 1.2 mL H₂O; pH neutralized with 5 M KOH; 122.0 mg MgCl₂·6 H₂O added, adjusted to pH 7.#1

Ama, antimycin A (Sigma Aldrich A8674) stock 5 mM: 5.4 mg dissolved in 2 mL EtOH.#2

CCCP, carbonyl cyanide 3-chlorophenylhydrazone carbonate (Sigma Aldrich C2759) stock 1 mM: 1.02 mg CCCP dissolved in 5 mL EtOH#2

CoQ2 (Sigma Aldrich C8081) stock 10 mM: commercial vial (2 mg CoQ2) dissolved in 628 µL EtOH. Stock 1 mM: 50 µL of 10 mM CoQ2 stock diluted with 450 µL EtOH.#2

M, malate (Sigma Aldrich M1000) stock 400 mM: 268.2 mg dissolved in 3 mL H₂O, pH adjusted to 7.0 with 5 M KOH, volume adjusted to 5 mL.#1

P, pyruvate (Sigma Aldrich P2256) stock 2 M: 44 mg dissolved in 180 µL H₂O every day fresh.

Rot, rotenone (Sigma Aldrich R8875) stock 1 mM: 0.39 mg dissolved in 1 mL EtOH.#2

S, succinate (Sigma Aldrich S2378) stock 1 M: 1.351 g dissolved in 3 mL H₂O, pH adjusted to 7.0 with 1 M HCl, final volume adjusted to 5 mL.#1

Suspension buffer: 10.25 g mannitol (225 mM), 6.42 g sucrose (75 mM) and 0.095 g EGTA (1 mM) dissolved in 250 mL H₂O. pH adjusted to 7.4 with KOH or HCl if needed.#1

Isolation buffer A: 62.5 mg fatty acid free BSA (2.5 g/L) dissolved in 25 mL suspension buffer fresh every day.

Isolation buffer B: 5 mg subtilisin dissolved in 10 mL isolation buffer A, fresh every day.

Isolation buffer C: 0.25 g fatty acid free BSA (2.5 g/L) dissolved in 500 mL isolation buffer D every day freshly.

Isolation buffer D: 27.4 g sucrose (320 mM), 0.303 g Tris-Cl (10 mM), 0.093 g K-EDTA (1 mM) dissolved in 250 mL H₂O, pH adjusted to 7.4 with KOH or HCl if needed.#1

MiR05, mitochondrial respiration medium (Gnaiger et al 2000): 0.5 mM EGTA, 3 mM MgCl₂·6H₂O, 60 mM lactobionic acid, 20 mM taurine, 10 mM KH₂PO₄, 20 mM 2-[4-(2-hydroxyethyl)piperazin-1-yl]ethanesulfonic acid HEPES, 110 mM sucrose, 1 g/L BSA; pH 7.1; prepared using MiR05-Kit (Oroboros Instruments, Austria 60101-01) and BSA.#1

#1 Aliquots stored at -20 °C in plastic vials; #2 aliquots stored at -20 °C in dark glass vials.

Further reagents were obtained from Sigma Aldrich (cat. N°): MES, 2-(N-morpholino)ethanesulfonic acid hydrate (M8250); ATP, adenosine 5’-triphosphate disodium salt hydrate (A2383); BSA, fatty acid-free bovine serum albumin (A6003); CaCO₃ (C4830); D-sucrose (S7903); dithiothreitol (D0632); EGTA, ethylene glycol tetraacetic acid (E4378); imidazole (S6750); KCl (60130); KH₂PO₄ (P5655); KOH (P1767); mannitol (M4125); phosphocreatine disodium salt (P7936); subtilisin, protease from
*Bacillus licheniformis* Type VIII, lyophilized powders, 7-15 mg/unit (P5380). Bartelt, Austria: EtOH, ethanol 99.9 %, (CL0005055000); Scharlab: MgCl₂·6H₂O magnesium chloride hexahydrate (MA0036); Evoqua Water Technologies GmbH: H₂O, deionized ultra-pure water (Ultra Clear™ TP UV UF TM).

2.2. **Animals**

C57BL/6N wild-type young adult mice (male and female) were housed in clear plastic cages (maximum five mice per cage) in the animal facility of the Medical University of Innsbruck. Mice were kept in a controlled environment (23 ± 3 °C, 12/12 h light/dark cycle) and fed *ad libitum* with free access to water. After cervical dislocation, heart and brain were removed and immediately placed in ice-cold BIOPS. All procedures involving animals were conducted in accordance with the Austrian Animal Experimentation Act in compliance with the European convention for the protection of vertebrate animals used for experimental and other scientific purposes (Tierversuchsgesetz 2012; Directive 2010/63/EU; BMWFM-66.011/0128-WF/V/3b/2016). According to the 3Rs principle the number of animals was minimized.

2.3. **Isolation of mitochondria**

A glass/Teflon Potter-Elvehjem homogenizer (WiseStir HS-30E, Wisd laboratory instruments) and centrifuge (Rotina 380R, Andreas Hettich GmbH & Co. KG, Tuttingen, Germany) were used. All procedures were carried out in an ice bath or at 4 °C.

Mouse heart mitochondria were isolated following Fontana-Ayoub et al (2015). Briefly, wet mass of the whole heart was determined, washed with ice-cold BIOPS and minced with scissors in ice-cold BIOPS (1 mL). The tissue was transferred into a pre-cooled glass/Teflon potter and homogenized at ~1000 rpm (five strokes) in 2 mL isolation buffer B. The homogenate was transferred to a 50-mL Falcon tube containing 3 mL isolation buffer B and centrifuged at 800 g for 10 min. Using a new 50-mL Falcon tube, the supernatant was centrifuged at 10 000 g for 10 min. The supernatant was discarded, the pellet was resuspended in isolation buffer A (final volume 2 mL), and centrifuged at 10 000 g for 10 min. The supernatant was discarded, and the mitochondrial pellet was finally resuspended in 200 µL suspension buffer.

Mouse brain mitochondria were isolated following Sumbalova et al (2016). Briefly, wet mass was determined, and the tissue was cut into small particles with sharp scissors in isolation buffer C. The medium was discarded, the tissue suspended in isolation buffer C (0.1 g tissue/1 mL), transferred to a pre-cooled glass/Teflon potter, and homogenized at 1000 rpm (five strokes). The homogenate was transferred to a 50-mL Falcon tube (0.5 g tissue/20 mL homogenate) and centrifuged at 1000 g for 10 min. The pellet was discarded, and the supernatant was centrifuged at 6200 g for 10 min. The supernatant was removed, the pellet resuspended in isolation buffer D (0.5 g tissue/10 mL), and recentrifuged at 6200 g for 10 min. The supernatant was discarded, and the mitochondrial pellet was finally suspended in 500 µL isolation buffer D.

The mitochondrial suspension was gently mixed with a 200-µL pipette (five up-down cycles), and 10 µL or 20 µL of heart or brain mitochondrial suspension was injected
with a 50-µL Hamilton syringe into the O2k-chamber through the titration capillary of the stopper, respectively.

2.4. Mitochondrial protein content

Mitochondrial protein was determined based on Lowry et al (1951) using the DC™ Protein Assay (Bio-Rad, Hercules, CA, US) following the manufacturer instructions. The absorbance was measured at 620 nm in a Tecan Infinite™ F200 spectrophotometer (Tecan, Männedorf, Switzerland). 0.025 mg/mL isolated heart mitochondria and 0.09 mg/mL isolated brain mitochondria were applied in the experiments.

2.5. High-resolution respirometry

O₂ flux and the CoQ2-redox states were measured in the NextGen-O2k (Oroboros Instruments, Austria), recorded, and analyzed using DatLab 7.4 (Oroboros Instruments, Austria). The O2k monitors the O₂ signal of polarographic oxygen sensors (POS) over time and plots O₂ consumption of a biological sample continuously. The two chambers of the O2k were calibrated at experimental volumes of 2 mL. Instrumental quality control was performed routinely as a standard operating procedure of HRR: (1) daily oxygen sensor test including air calibration in MiR05 at experimental temperature of 37 °C (stability ±0.002 °C), and stability of flux better than ±1 pmol·s⁻¹·mL⁻¹, and (2) monthly instrumental O₂ background test including zero calibration of the POS (Doerrier et al 2018; Gnaiger 2001; 2008). The medium was continuously stirred with a polyether ketone (PEEK)-coated magnetic stirrer bar at 750 rpm for optimum signal stability of the POS and mixing of the suspended sample and dissolved substances.

Volume-specific oxygen flux $J_{O2}$ [pmol·s⁻¹·mL⁻¹] was calculated real-time as the negative time derivative of the O₂ concentration by DatLab 7.4. The O₂ flux was corrected for (1) the instrumental O₂ background flux $J_{O2}^{bg}$, (2) dilution of the sample by titrations, and (3) residual oxygen consumption $J_{VOX}$ measured in the presence of isolated mitochondria without any respiratory fuel substrates and ADP or after inhibition of the electron transfer system.

2.6. Q-Module

The Q-Module of the NextGen-O2k provides the basis for continuous monitoring of the redox state of an added Q-mimetic in isolated mitochondria and chloroplasts (Figure 1). According to the original description (Rich 1988), a three-electrode system and a mobile short-chain Q-mimetic (CoQ1 or CoQ2) are required to indirectly detect the redox state of the ETS-reactive Q-pool trapped in the mitochondrial inner membrane mtIM. CoQ2 reacts both with the biochemical sites of the ETS and the measuring electrode. Q-mimetics do not react directly with the long isoprenyl chain CoQ in the ETS, rather they are reduced by e.g. CI and CII, oxidized by CIII at the Q₀ site and reduced at the Q₁ site. If the redox state of the Q-mimetic is in equilibrium with the redox state of ETS-reactive CoQ, the redox state of the Q-mimetic reflects the redox state of ETS-reactive mtCoQ. In the present study, a low concentration of CoQ2 (1 µM) was used, as described by Moore et al (1988; 1991).
The three-electrode system consists of a glassy carbon working electrode (GCE) set at a fixed potential relative to the silver/silver chloride (Ag/AgCl) reference electrode (Rich 1988). The potential set on the GCE is chosen to be sufficient to either oxidize the reduced or reduce the oxidized CoQ2. A platinum (Pt) counter electrode completes the electronic circuit. If the GCE is set to a potential oxidizing CoQ2, then CoQ2 reduced by the ETS undergoes oxidation on the GCE surface, resulting in a current between the GCE and Pt electrode. In this case the concentration of the reduced CoQ2 is proportional to the current measured between GCE and Pt electrodes: the current increases in proportion to the concentration of reduced CoQ2. Current $I$ [A] is converted into a voltage $U$ (electric potential difference [V]) and amplified: $U=I\cdot R$ ($R$: resistance). Conversely, if the GCE is set to the CoQ2 reduction potential, the oxidized CoQ2 undergoes reduction on the GCE surface and current flows into the opposite direction. In the present study, the GCE was set to the oxidation peak potential $E_{p1}$ when measuring the CoQ2 redox state. The GCE and Pt electrodes are built-in fixed parts of the Q-stopper, whereas the reference electrode can be inserted through a separate inlet (Figure 1).

**Figure 1.** Q-Module with Q-sensor and stopper. The glassy carbon electrode (black) and platinum electrode (shiny silver) are built-in as fixed parts of the PEEK stopper. (a) Top view showing the central titration capillary and the inlet for the reference electrode (not inserted). (b) Bottom view with conical center guiding gas bubbles to the capillary, double Viton O-rings. (c) Top view with reference electrode. (d) Q-sensor with reference electrode. (e) Q-stopper inserted into the chamber of the NextGen-O2k prototype (front view).

### 2.7. Cyclic voltammetry

Cyclic voltammetry CV was controlled and recorded by DatLab 8.0 (Oroboros Instruments, Austria) before and after experiments. O2k-chambers were washed three times for 5 min with H$_2$O. In the meantime, the Q-sensor (Figure 1b) was polished with wetted 0.3 µm aluminum oxide powder on a polishing cloth, performing a figure eight motion in a vertical position 10-15 times, then polished with 0.05 µm aluminum powder in the same way. Afterwards, the Q-sensor and the separate reference electrode were rinsed with H$_2$O and wiped with a soft tissue (Komlodi et al 2021).
After adding 2.3 mL MiR05 into the O2k-chamber, the Q-stopper was inserted with the mounted reference electrode into the O2k-chamber and excess medium was siphoned off the stopper receptacle. The background CV was determined without rotation of the stirrer. Then 30 µM CoQ2 (6 µL of 10 mM stock) was titrated into the chamber and the stirrer switched on to mix the CoQ2 solution. The stirrers were switched off and CV was started to determine the oxidation and reduction peak potentials of the Q-mimetic. Finally, the O2k-chamber, stopper, and reference electrode were washed with H2O, 99.9% EtOH, and H2O again.

3. Results

3.1. Instrumental oxygen background test

The Q-stopper is equipped with the titration capillary, a large capillary for inserting the reference electrode, and the fixed GC- and Pt-electrodes (Figure 1). The design was optimized for minimum O2 diffusion through the stopper, comparable with the specifications of HRR using the standard O2k-stopper with a single injection capillary (Gnaiger 2001).

![Figure 2. Instrumental O2 background flux as a function of O2 concentration: NextGen-O2k with Q-stoppers and O2k with regular O2k-stoppers. (a) Superimposed traces in two 2-mL chambers with Q-stoppers. \( j_o \) to \( j_4 \) refer to background O2 flux monitored at sequentially lowered O2 concentrations. Excess dithionite (100 µL) added to deplete O2 for zero calibration of the POS. Experiment 2019-08-28_PQ1-01. (b) 12 O2k-chambers using different Q-stoppers. (c) 20 O2k-chambers with regular O2k-stoppers. Average ± SD for the intercept \( a^o \) and slope \( b^o \) of linear regressions for individual chambers. Lines show linear regressions calculated through all data points.](image-url)
Correction for instrumental background O$_2$ flux is a standard procedure in HRR (Gnaiger 2001). The instrumental background O$_2$ flux is due to the O$_2$ consumption of the POS and O$_2$ diffusion into and out of the aqueous medium in the O2k-chamber, part of which may occur through diffusion leaks in the stopper. The instrumental background O$_2$ flux $f_O^o_{2}$ was measured in the absence of biological sample in the closed chamber in the range of experimental O$_2$ levels at four different O$_2$ concentrations: near air saturation $\sim$170 µM, $\sim$90 µM, $\sim$45 µM, $\sim$20 µM (Figure 2a). O$_2$ levels were reduced by dithionite titrations using the TIP2k (Titration-Injection microPump) and maintained for 20 min. O$_2$ flux was a linear function of O$_2$ concentration. The intercept ($a^o$: flux at zero O$_2$ concentration) and slope ($b^o$) were calculated from linear regressions for each individual chamber. $a^o$ was -2.6 ± 0.7 pmol·s$^{-1}$·mL$^{-1}$ using the Q-stopper, not significantly different from the intercept measured with the regular O2k-stoppers (-2.3 ± 0.4 pmol·s$^{-1}$·mL$^{-1}$; Figure 2b and c).

3.2.  Cyclic voltammetry

Cyclic voltammetry CV is applied in studies of the redox chemistry of CoQ (Gulaboski et al 2016). We applied CV at experimental conditions as quality control to (1) determine the oxidation- and reduction-peak potentials of CoQ2 under specific experimental conditions, (2) check the quality of the Q-sensor, and (3) test the interference of chemicals used in the HRR assay with the Q-sensor. In CV, the electrical potential between the GCE and the Ag/AgCl electrode is varied over time in cycles, while the current is recorded between the GC- and Pt-electrodes. The current is plotted as a function of the applied electrical potential in the cyclic voltammogram, where the characteristic peaks refer to the maximum rate of CoQ2 oxidation (oxidation peak potential, $E_{p1}$) and the maximum rate of reduction (reduction peak potential, $E_{p2}$; Figure 3). These values are set automatically by DatLab 8.0 and used to poise the GCE (Rich 1982; Rich 2004).

Figure 3 shows $E_{p1}$ and $E_{p2}$ determined after careful polishing of the GCE and Pt electrode using different Q-sensors in various chambers of the NextGen-O2k. The $E_{p1}$ of CoQ2 was 31.8 ± 6.5 mV, and $E_{p2}$ was -269.9 ± 11.1 mV using freshly polished electrodes. Reproducibility of the CV measurements was high using different Q-sensors in various NextGen-O2k prototypes. In MiR05, 30 µM CoQ2 was optimal for CV. Lower CoQ2 concentrations did not result in detectable peaks at gain 1 V/µA, whereas the limit of detection was reached at higher than $\sim$90 µM CoQ2. The lowest concentration of CoQ2 should be applied which gives well-defined $E_{p1}$ and $E_{p2}$. Stirring of the solution is avoided during CV to minimize convection. Upon stirring in the presence of quinone (oxidized), only a peak related to quinone reduction is visible, whereas the peak of quinol oxidation cannot be observed, because the quinol is stirred off from the surface of GCE. At the start of CV an initial potential of +30 mV was used, which is close to the oxidation peak potential $E_{p1}$ of CoQ2. To avoid coating of the GCE, the initial potential must be close to the peak potential (Graham 2018). A polarization window was chosen of -500 mV to +500 mV versus Ag/AgCl. The narrowest possible range of potentials should be applied during CV scanning. Excessively high and low potentials might lead to chemical modification or coating of the GCE (Graham 2018). Coating of the GCE can inhibit electron transfer on the electrode surface.
Figure 3. Cyclic voltammogram in the absence and presence of 30 µM CoQ2, non-stirred MiR05, 37 °C (NextGen-O2k).

Initial potential difference: +30 mV, polarization window from -500 mV to +500 mV, scanning speed 100 mV/s, gain 1. (a) Red plot: background CV without CoQ2; blue plot: CV in the presence of 30 µM CoQ2; Experiment: 2019-06-14_PQ2.

(b) Oxidation peak potential $E_{p1}$, and (c) reduction peak potential $E_{p2}$ of CoQ2 with six different Q-sensors. $E_{p1}$ and $E_{p2}$ [mV] are shown as average ± SD; n=51.

We found an optimum of five cycles from -500 mV to +500 mV and back to -500 mV in standard CV applications. Although well-defined peaks for $E_{p1}$ and $E_{p2}$ were observed after one cycle, it is recommended to run more cycles to check whether additional peaks are detected or the shape of $E_{p1}$ and $E_{p2}$ changes over the cycles owing to side-reactions. 100 mV/s was applied as scanning speed, which should allow for diffusion as the controlling process of exchange of CoQ2 between the surface of GCE and the medium. If the scanning speed is very slow, CoQ2 might be transported to and from the electrode surface by convection rather than diffusion. At too fast scanning speeds, the double layer charging current increases due to the rearrangement of solution molecules at the surface of the GCE. This results in high baseline currents that obscure the diffusion-controlled cyclic voltammogram (Graham 2018).

CV serves additionally as an essential quality control to evaluate the function of the Q-sensor, for minimizing directional long-term drift and non-directional short-term noise of the signal. No peaks should be observed in the background CV of complete incubation media without CoQ2, while the peaks in the presence of 30 µM CoQ2 should be well-defined and sufficiently sharp (Figure 3). If the peaks are not sharp enough and not well-defined, or additional peaks are observed (with and without CoQ2), the GCE and Pt-electrode are polished with aluminum powder, the Q-sensor and O2k-chamber are washed with H2O, 70 % ethanol, 99.9 % EtOH and H2O, the glass barrel of the reference electrode is re-filled with 3 M KCl solution, and the quality of the porous vycor frit of the glass barrel of the reference electrode should be evaluated (Komlodi et al 2021).
Figure 4. Coupling-control in the succinate-pathway S; SUIT-006. Simultaneous measurement of O$_2$ flux and redox state of the ETS-reactive Q-pool in mitochondria isolated from mouse heart. Simultaneous with SUIT-031 (Figure 5). (a) Blue plot: O$_2$ concentration [µM]; red plot: O$_2$ flux per volume [pmol·s$^{-1}$·mL$^{-1}$]. (b) Non-calibrated (raw) Q-signal $U_{raw}$ [V] and reduced Q-fraction $x_{Qred}$. Ren and Rox, residual oxygen consumption with endogenous substrates and after inhibition by rotenone (Rot) when fully oxidized CoQ2 ($x_{Qred} = 0$) was calibrated ($\times$). Further titrations: 1S, S-linked LEAK respiration $L$; 2D (ADP), S-linked OXPHOS capacity $P$; 3U (uncoupler CCCP; 1 µM), S-linked ET capacity $E$. Fully reduced CoQ2 ($x_{Qred} = 1$) calibrated under anoxia (+). The effect of antimycin A (Ama) on the Q-signal was not due to the carrier EtOH. (c) Coupling/pathway control diagram.

3.3. Substrate-uncoupler-inhibitor titration protocols

SUIT protocols are used to study respiratory control in different pathway- and coupling-control states in a single experimental assay. With a coupling-control protocol (SUIT-006 Q mt D071) and a coupling-pathway control protocol (SUIT-031 Q mt D072) we investigated O$_2$ flux and the redox state of the ETS-reactive Q-pool simultaneously (Figure 4 and 5). Harmonized SUIT protocols are applied with common cross-linked respiratory states, which can be considered as replicate measurements and therefore, allow harmonization of data obtained in different SUIT protocols. $S$(Rot)$_P$ and $S$(Rot)$_E$ are harmonized respiratory states in SUIT-006 and SUIT-031 without or with pyruvate &
malate, respectively. In chemical background tests, titrations in the absence of mitochondria did not exert any effect on the Q-signal in both SUIT protocols.

**The steps of coupling-control protocol SUIT-006 (Figure 4):**

After addition of isolated mitochondria in the absence of fuel substrates and ADP, residual oxygen consumption with endogenous substrates Ren is due to oxidation of endogenous substrates that remained after mitochondrial isolation and residual O₂ consumption Rox not related to the ETS.

1. Addition of CoQ2 (1 µM): residual endogenous respiration Ren.

2. Rotenone (Rot; 0.5 µM) and succinate (S; 10 mM): Rox after inhibition of respiration of endogenous substrates by Rot. Additionally, Rot prevents inhibition of succinate (S)-linked respiration caused by oxaloacetate accumulation (Gnaiger 2020). In the absence of rotenone, oxaloacetate is formed from malate in the reaction catalyzed by malate dehydrogenase MDH. Rot inhibits CI and oxidation of NADH, which results in an increase of the NADH/NAD⁺ ratio and consequently to feed-back inhibition of MDH, thus blocking formation of oxaloacetate. Succinate is formed in the tricarboxylic acid cycle and is the substrate of CII. It is oxidized to fumarate and supports electron transfer through CII to Q. Succinate with rotenone supports S-linked LEAK respiration and leads to reduction of CoQ2, reflected in an increase of the Q-signal.

3. ADP (D; 2.5 mM) was added at kinetically saturating concentration to stimulate S-OXPHOS capacity. A decrease of the Q-signal indicates partial oxidation of CoQ2.

4. Uncoupler CCCP was titrated (U; 0.5 µM/step) after stimulation by ADP as a measure of electron transfer capacity E. Neither O₂ flux nor the Q-redox state changed after CCCP titrations, showing that OXPHOS capacity was not limited by the phosphorylation system.

5. Anoxia was reached after the mitochondria consumed the O₂ in the O2k-chambers, the ETS and CoQ2 become reduced. Anoxia was used for calibration of fully reduced Q (Section 3.4). Antimycin A (Ama; 2.5 µM) is a Q-site inhibitor of CIII and was added to check its effect on the fully reduced Q-pool under anoxia. The Q-signal in the presence of Ama did not show dependence on the O₂ concentration (data not shown).

**The steps of coupling-pathway control protocol SUIT-031 (Figure 5):**

1. Addition of CoQ2 (1 µM): residual endogenous respiration Ren.

2. Pyruvate & malate (PM; 5 mM P and 2 mM M) were added in immediate sequence to induce NADH-linked LEAK respiration. Pyruvate is converted to acetyl-CoA in the reaction catalyzed by pyruvate dehydrogenase. Malate serves as a co-substrate and after entering the mitochondria it is oxidized to oxaloacetate catalyzed by MDH. In both reactions NADH, the substrate of CI, is formed. Oxidation of NADH leads to reduction of the Q-pool through CI (Q-signal increased). PM caused only partial reduction of CoQ2 in the LEAK state compared to S(Rot) in SUIT-006.

3. ADP (D; 2.5 mM) was added at kinetically saturating concentration to initiate N-OXPHOS capacity. A decrease of the Q-signal indicated partial oxidation of CoQ2.
4. Succinate (S; 10 mM) was added to induce NS-convergent electron transfer. Succinate further increased the O₂ flux and reduced CoQ2 in the OXPHOS state when added in the presence of PM, showing an additive effect in the Q-junction.

5. Rotenone (Rot; 0.5 µM) blocked N-linked respiration and led to oxidation of CoQ2 via CI inhibition leading to S-OXPHOS. The two protocols are harmonized at state S(Rot). E.

6. Uncoupler CCCP (U; 0.5 µM/step) was titrated (1 µM in total) to initiate S-ET capacity which is a common respiratory state to SUIT-006, S(Rot). E. Neither O₂ flux nor the Q-redox state changed in mouse cardiac mitochondria showing that the S-OXPHOS capacity was not limited by the phosphorylation system.
7. Anoxia corresponds to the state where CoQ2 is fully reduced (Section 3.4). Antimycin A (Ama; 2.5 µM).

3.4. Calibration: redox states of ETS-reactive Q

The redox state of ETS-reactive Q (Q_{ra} \equiv Q) is expressed as the mole fraction (Cohen et al. 2008) \( x_{Q_{red}} \) of reduced Q in each steady state of a SUIT protocol. To calculate \( x_{Q_{red}} \), the raw CoQ2 signal \( U_{raw} \) is calibrated against (1) the fully oxidized CoQ2 signal \( U_{ox} \) and (2) the fully reduced CoQ2 signal \( U_{red} \). \( x_{Q_{red}} \) is calculated as a fraction of fully reduced CoQ2 (Table 1). The sum of the oxidized and reduced fractions of Q equals 1, \( x_{Q_{red}} + x_{Q_{ox}} = 1 \). In this formalism the intermediate redox state of semiquinone is not taken into account.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition [Unit]</th>
<th>Quinol fraction ( x_{Q_{red}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( U_{ox} )</td>
<td>fully oxidized raw Q-signal [V]</td>
<td>( x_{Q_{red}} = 0 )</td>
</tr>
<tr>
<td>( U_{red} )</td>
<td>fully reduced raw Q-signal [V]</td>
<td>( x_{Q_{red}} = 1 )</td>
</tr>
<tr>
<td>( U_{raw} )</td>
<td>raw (non-calibrated) Q-signal [V]</td>
<td>( x_{Q_{ox}} = 1 ) calibrated fully oxidized Q_{ra} fraction, quinone ( x_{Q_{red}} = (U_{ox}-U_{ox})/(U_{ox}-U_{ox}) = 0 )</td>
</tr>
<tr>
<td>( x_{Q_{red}} = 1 )</td>
<td></td>
<td>( x_{Q_{red}} = (U_{red}-U_{ox})/(U_{red}-U_{ox}) = 1 )</td>
</tr>
<tr>
<td>( x_{Q_{red}} )</td>
<td>reduced Q_{ra} fraction</td>
<td>( x_{Q_{red}} = (U_{raw}-U_{ox})/(U_{red}-U_{ox}) )</td>
</tr>
</tbody>
</table>

\( U_{ox} \) is measured in the presence of CoQ2 and isolated mitochondria, and absence of reduced fuel substrates. Isolated mitochondria may contain endogenous substrates which can slightly reduce Q_{ra} in the calibration state for \( U_{ox} \). Then the CI inhibitor rotenone acting upstream of the Q-junction may inhibit respiration of endogenous substrates Ren further to Rox and decrease a small fraction of reduced Q in the REN state to the fully oxidized signal \( U_{ox} \) in the ROX state. The \( x_{Q_{red},REN}^* \) calibrated against \( U_{ox} \) after rotenone addition in SUIT-006 (Figure 4) is used for correction of \( x_{Q_{red}} \) in SUIT-031 measured simultaneously in parallel O2k-chambers (Figure 5).

\[
x_{Q_{red}} = x_{Q_{red}}^* + x_{Q_{red},REN}^* (1 - x_{Q_{red}}^*)
\]

Eq. 1

Rotenone exerted no or a minor effect on mitochondrial respiration and the Q-signal when added after REN. The median \( x_{Q_{red},REN}^* \) was 0.04 (range 0.03-0.05) in brain mitochondria while respiration declined by 100 pmol·s⁻¹·mg⁻¹ from Ren to Rox. The median \( x_{Q_{red},REN}^* \) was 0.00 (range -0.01--0.01) in cardiac mitochondria without inhibitory effect of rotenone (\( Ren = Rox \); Figure 4).

The easiest and most accurate calibration of \( U_{red} \) at fully reduced Q_{ra} is performed under anoxia after mitochondria reduced the O₂ concentration practically to zero in the O2k-chamber with minimum O₂-backdiffusion (Figure 2). To do so, a mt-protein concentration >0.05 mg/mL is used which consumes relatively fast the O₂ in the closed chamber (Figure 4 and 5). Alternatively, the O₂ concentration can be decreased by
nitrogen or hydrogen gas injection into the gas phase of an opened O2k-chamber, intermittently moving the stopper slightly upwards to form a gas phase above the stirred aqueous phase, and then allowing respiration to establish anoxia in the closed chamber.

### 3.5. Optimization of CoQ2 concentration

The lowest practicable CoQ2 concentration is applied to avoid side reactions on the ETS caused by the mimetic. Our study confirmed that 1 µM CoQ2 was sufficient to detect the redox states of ETS-reactive Q without any influence on respiration (Moore et al 1991). The use of 0.5-1.5 µM CoQ2 did not influence the Q-redox state measured at constant concentration of mouse heart mitochondria (Figure 6b). It is recommended to test the effect of CoQ2 on each type of mitochondria under experimental conditions.

### 3.6. Technical reproducibility

Figures 6a and b show $x_{Q_{\text{red}}}$ of technical replicates performed in parallel using different Q-sensors. In Figure 6a, the coupling-control protocol SUIT-006 (S-pathway) was applied in mouse brain mitochondria (representative trace: Figure 4). In Figure 6b, both SUIT protocols (S- and N-pathways; representative traces: Figure 4 and 5) were used with mouse heart mitochondria. The results indicate a high reproducibility in every pathway- and coupling-control state. A range of CoQ2 concentrations from 0.5 µM to 1.5 µM, keeping the same concentration of sample for every experimental chamber, did not impact the redox state of the ETS-reactive Q-pool (Figure 6).

### 3.7. Oxygen flux, reduced molar Q-fraction, and additivity

In the S-linked LEAK state, CoQ2 was highly reduced in brain ($x_{Q_{\text{red}}} = 0.80$; range 0.78 to 0.82) and heart ($x_{Q_{\text{red}}} = 0.92; 0.89$ and 0.95; Figure 6a and b). In contrast, $x_{Q_{\text{red}}}$ was lower (less reduced) in the N-linked LEAK state ($x_{Q_{\text{red}}} = 0.24; 0.22$ to 0.28; Figure 6b).

ADP-induced stimulation of respiration increases the load on the ETS. Under these conditions of increasing the load from the LEAK to OXPHOS state, CoQ2 became more oxidized in brain ($x_{Q_{\text{red}}} = 0.16; 0.13$ to 0.19) and heart ($x_{Q_{\text{red}}} = 0.62; 0.58$ and 0.66; Figure 6c). In heart, $x_{Q_{\text{red}}}$ was higher in the S-linked ($x_{Q_{\text{red}}}$=0.56; range 0.51 to 0.61) than in the N-linked ($x_{Q_{\text{red}}}$=0.18; range 0.13 to 0.23) OXPHOS state (Figure 6b and d). This is consistent with the high ET-capacity of the S- compared to the N-pathway (Figure 5).

The higher electron supply capacity of the S-branch drives Q into a more reduced state. Uncoupler titrations (ET state) after stimulation by a kinetically saturating ADP concentration (OXPHOS state) did not affect respiration and Q-redox state in brain and heart, indicating that OXPHOS capacity is not limited by the capacity of the phosphorylation system in these mitochondria (Gnaiger 2020). Whereas coupling control decreased $x_{Q_{\text{red}}}$ (more oxidized) by increasing the load (higher flux; Figure 6c), pathway control increases $x_{Q_{\text{red}}}$ (more reduced) by increasing the drive of electron input into the Q-junction (higher flux; Figure 6d).

OXPHOS capacity in heart mitochondria was low in the N-pathway (CI-linked) $J_N = 780$ (range 720 to 840) pmol·s⁻¹·mg⁻¹, higher in the S-pathway (CII-linked) $J_S = 1397$
(range 1317 to 1477) pmol·s⁻¹·mg⁻¹ and showed an additive effect in the combined NS-pathway \( J_{NS} = 1837 \) (range 1706 to 1968) pmol·s⁻¹·mg⁻¹. The NS-linked \( O_2 \) flux \( (J_{NS}) \), however, was lower than \( J_s + J_e \) demonstrating incomplete additivity. \( S \) was the dominant \( \alpha \)-pathway with \( J_s > J_e \). Then flux control ratios are \( \alpha = J_s/J_{NS} \) and \( \beta = J_e/J_{NS} \). Additivity \( A_{\alpha\beta} \) is defined as \((1 - \alpha)/\beta\) (Gnaiger 2020). Complete additivity \((A_{\alpha\beta} = 1)\) is obtained when the linear sum of the component N- and S-pathway ET capacities \((J_n + J_s)\) equals the ET capacity of the convergent NS-pathway with the NS-substrate combination \((J_{NS})\). In heart, \( A_{\alpha\beta} \) was 0.57 (range 0.54 to 0.60) indicating incomplete NS-additivity of \( O_2 \) flux.

![Image](https://www.mitofit.org/)

**Figure 6. Reduced Q-fraction \( x_{Q_{\text{red}}} \) and \( O_2 \) flux in mouse brain and heart mitochondria using six different Q-sensors.** (a) Brain; SUIT-006; 1 µM CoQ2. Fully oxidized Q \( (x_{Q_{\text{red}}}=0) \) calibrated in the presence of rotenone (\( \times \)). Respiratory states: Residual endogenous respiration REN; residual oxygen consumption after addition of rotenone ROX; S-linked LEAK respiration \( S_L \); S-linked OXPHOS capacity \( S_{ET} \); S-linked ET capacity \( S_E \). Calibration of fully reduced Q \( (x_{Q_{\text{red}}}=1) \) under anoxia (\( + \)). Experiments: 2020-04-23_PN1-02; 2020-04-23_PN2-03; 2020-04-23_PQ2-02. (b) Heart; SUIT-006 (filled symbols) and SUIT-031 (open symbols); CoQ2 concentrations as indicated. REN* from SUIT-006 used to correct fully oxidized Q \( (x_{Q_{\text{red}}}=0) \) in SUIT-031 (\( \times \)). N-linked LEAK respiration \( N_L \); N-linked OXPHOS capacity \( N_{ET} \); NS-linked OXPHOS capacity \( NS_{ET} \); S-linked OXPHOS capacity after inhibition by rotenone \( S_{ET} \); S-linked ET capacity \( S_E \). Calibration of fully reduced Q \( (x_{Q_{\text{red}}}=1) \) under anoxia (\( + \)). (c) Coupling control: effect of increased load — from LEAK- to OXPHOS- and ET-states — on \( x_{Q_{\text{red}}} \) as a function of \( O_2 \) flux per protein mass \( J_{O_2} \) [pmol·s⁻¹·mg⁻¹] at constant S-pathway state (from panels a and b). (d) Pathway control: effect of increased drive — with electron input into the Q-junction by separate or combined convergent pathways — on \( x_{Q_{\text{red}}} \) as a function of \( J_{O_2} \) [pmol·s⁻¹·mg⁻¹] at constant OXPHOS-coupling state (from panel b). Inverted regression analysis was performed (Gnaiger 2021).
Q-redox was directly proportional to the OXPHOS capacity under pathway control — increasing from 0.18 (range 0.13 to 0.23) for N, 0.56 (range: 0.51 and 0.61) for S, to 0.63 (range 0.6 and 0.66) for NS — resulting in a linear dependence of $Q_{\text{red}}$ on respiratory rate. Proportionality — i.e. linearity and zero intercept — between reduction of CoQ2 and ET capacities (N- and S-pathway) indicates that CoQ2 equilibrates equally with the CI-Q-CIII and CII-Q-CIII branches of the electron transfer system and thus reflects the redox state of the ETS-reactive Q-pool (Table 2).

**Table 2. Q-pools and redox-reactivity with the Q-mimetic CoQ2**

<table>
<thead>
<tr>
<th>Q-pool</th>
<th>Symbol</th>
<th>Definition (CoQ2: reactive or non-reactive)</th>
</tr>
</thead>
<tbody>
<tr>
<td>coenzyme Q</td>
<td>CoQ</td>
<td>total CoQ in the cell, in mt-preparations, or added experimentally</td>
</tr>
<tr>
<td>mitochondrial CoQ</td>
<td>mtCoQ</td>
<td>total CoQ in mitochondria; $mtCoQ = Q_{\text{mt}} + mtCoQ_{\text{ia}}$</td>
</tr>
<tr>
<td>inactive mtCoQ</td>
<td>mtCoQ$_{\text{ia}}$</td>
<td>insensitive to changes in mitochondrial respiratory states; partially oxidized under anoxia and partially reduced in aerobic mt-preparations incubated without fuel substrates; (CoQ2: non-reactive)</td>
</tr>
<tr>
<td>ETS-reactive Q</td>
<td>$Q_{\text{ra}} \equiv Q$</td>
<td>fully reduced under anoxia, fully oxidized in aerobic mt-preparations incubated without external and internal fuel substrates and/or addition of inhibitors upstream of Q: $Q = Q_{\text{free}} + Q_{\text{SC}}$; (CoQ2: reactive or possibly partially reactive)</td>
</tr>
<tr>
<td>free Q</td>
<td>$Q_{\text{free}}$</td>
<td>free pool of ETS-reactive Q according to the fluid-state model (random-collision model); (CoQ2: reactive)</td>
</tr>
<tr>
<td>supercomplexed Q</td>
<td>$Q_{\text{SC}}$</td>
<td>pool of $Q_{\text{ra}}$ bound to supercomplexes according to the solid-state model, not equilibrated with $Q_{\text{free}}$ (CoQ2: reactive or possibly partially reactive)</td>
</tr>
</tbody>
</table>

4. **Discussion**

Evidence within the literature suggests that a large fraction of CI is organized as a supercomplex in conjunction with CIII and CIV (respirosome) contributing to the supercomplex-bound Q-pool $Q_{\text{SC}}$, whereas CII is not organized in a supercomplex but reacts with the free Q-pool $Q_{\text{free}}$ according to the fluid-state model (Enriquez, Lenaz 2014). Even $Q_{\text{free}}$ may not be homogenous, with central localization of ubiquinone in the lipid bilayer and more extensive interaction of ubiquinol with the peripheral phospholipid acyl chains (Ausili et al 2008). A direct link can be made between supercomplex channeling, interaction with the free Q-pool, and additivity of respiratory NS-pathway capacity. Complete channeling through the supercomplex SCI$_{\text{II}}$III$_{\text{n}}$IV$_{\text{n}}$ predicts complete additivity ($A_{\alpha\beta} = 1$; Gnaiger 2020). Without interaction between the redox components in the channel and the free redox intermediates, there is no interaction between the N- and S-pathways which implies complete additivity. Importantly, the apparent excess capacity of CIV over the combined NS-ET capacity is high in mouse heart mitochondria (Lemieux et al 2017). Therefore, incomplete NS-additivity was not due to limitation of ET capacity downstream of Q.
In the present work we optimized the simultaneous measurement of the redox state of the ETS-reactive Q-pool and respiration in isolated mitochondria using the amperometric three-electrode sensor.

4.1. Advantages

**Real-time and continuous detection of redox state:** Monitoring the redox state of the ETS-reactive Q-pool in real-time is one of the main advantages of the electrochemical Q-sensor in contrast to the Q-extraction method. The Q-Module of the NextGen-O2k yields a significantly higher signal-to-noise ratio and lower drift compared to the original Q-electrode system (Moore et al 1991).

Zero to 50 % of the total mtCoQ-pool is not ETS-reactive (Urban, Klingenberg 1969; Kröger, Klingenberg 1973b; Jørgensen et al 1985; Van den Bergen et al 1994). The ETS-inactive mtCoQ-pool does not interact with the ETS nor CoQ2. Thus, redox changes reported by the electrochemical Q-sensor are related to respiratory activity (Table 2).

**Simultaneous measurement of Q-redox state and O2 flux:** The Q-Module of the NextGen-O2k allows for simultaneous measurement of redox state of the ETS-reactive Q-pool and O2 flux in a closed chamber. Multiple titrations can be carried out via the titration capillary of the specifically designed stopper, which closes the O2k-chamber.

**Controlled O2 concentrations and high resolution:** Owing to the near air-tight experimental O2k-chamber, the O2 concentration can be increased or decreased (between 0 and 1000 µM the POS gives a linear response), which allows measurement not only at air saturation, but also in hypoxic and hyperoxic ranges. Minimizing the O2 diffusion is essential to obtain anoxic conditions for calibration of fully reduced Q. Using the original Q-electrode system (Rich 1988; Moore et al 1988; Dry et al 1989) resolution of the oxygen sensor was limited and O2 diffusion into the closed chamber posed a problem. Therefore, high mitochondrial concentrations were required.

4.2. Limitations

**Q-pool compartmentalization:** CoQ2 does not interact directly with free mtCoQ. However, provided that both CoQ2 and Qr have the same relative rates of reaction with the quinone-reactive complexes, CI, CII and CIII, then both CoQ2 and ETS-reactive Q of the mtCoQ-pool will follow the same redox transitions. However, Q-compartmentalization may occur between a free ETS-reactive Q-pool in the lipid phase of the mtIM behaving according to the liquid-state or random-collision model and a bound ETS-reactive Q-pool tightly associated with respiratory supercomplexes. Such possible compartmentalization of Q needs to be considered in the interpretation of the amperometric signal of the Q-Module. This is particularly important if dissociation of supercomplexes is under control of the $pmF$ (Enriquez, Lenaz 2014). Then equilibration of CoQ2 with compartmentalized ETS-reactive Q relates to different pool sizes in the LEAK state at high $pmF$ and the OXPHOS- and ET-states at lower and very low $pmF$, respectively (Figure 6). Pool size of the free mobile Q should not matter. In this context it is interesting to note that uncoupler titrations inducing the transition from S(Rot)$^+$ to S(Rot)$^-$ did not affect the Q-redox state nor O2 flux in the presence or absence of pyruvate & malate (Figures 4 and 5). Inaccessible
ETS-reactive Q-sites — and therefore inaccessible CoQ2 non-reactive sites — in e.g. supercomplexes, however, would not be reported by the monitored CoQ2 redox state. Importantly, the reduced Q-fraction \( x_{\text{Qred}} \) increased linearly and proportionally with respiratory flux from N-pathway control involving the supercomplex \( \text{SCl}_{3}, \text{III}_{4}\text{V}_{n} \) to S-pathway control (Figure 6d), suggesting that CoQ2 is similarly reactive with the CI-Q-CIII and CII-Q-CIII branches irrespective of heterogeneity of the ETS-reactive Q-pool (Table 2).

**mtCoQ concentration:** Electrochemical determination of mtCoQ concentration is not possible (however, see Petrova et al 2014). If total mtCoQ is of interest (ETS-inactive and ETS-reactive), Q-extraction is the method of choice (Zannoni, Moore 1990; Van den Bergen et al 1994). CoQ2 may interact with non-mitochondrial redox-reactive CoQ-pools which may interfere with the Q-signal in crude isolated mitochondria.

**Chemical interference:** Some inhibitors and chemicals applied in HRR interfere and may even damage the Q-sensor. Dithionite, cytochrome c (Osakai et al 2019), ascorbate, TMPD (tetramethyl-p-phenylenediamine dihydrochloride), CIV inhibitors (i.e. potassium cyanide and azide) and cyclohexylammonium salts of some chemicals (e.g. glycerol-3-phosphate) interfered with the Q-signal. The alternative ubiquinol oxidase inhibitor benzohydroxamate and NADH cannot be applied with the Q-electrode (Van den Bergen et al 1994). Thus, it is advisable to perform a chemical background test in the absence of biological sample, and CV in the absence and presence of 30 \( \mu \text{M} \) CoQ2. If the shape of the CV has changed or additional “peaks” in the current in CV are detectable, a non-interfering alternative working electrode potential can be selected, changing between an oxidizing potential (to remove electrochemical interference from an oxidant) or reducing potential (to remove electrochemical interference from a reductant).

**4.3. Conclusions**

Electrochemical monitoring of the redox state of the ETS-reactive Q-pool adds a further dimension to coupling- and pathway-control analysis of isolated mitochondria. The Q-Module of the NextGen-O2k enables real-time monitoring of redox changes of the ETS-reactive Q-pool without interference by the inactive mtCoQ-pool. The reduced Q-fraction declines with respiratory rate stimulated by coupling control from LEAK to OXPHOS and ET states, but increases with respiration controlled by N-, S-, and convergent NS-pathway capacities. This powerful approach expands studies in mitochondrial physiology providing a greater insight into the role and regulation of mitochondrial function in health and disease.

**Acknowledgements**

The hardware and electronics of the NextGen-O2k Q-Module was developed in collaboration with Phillip Gradl and his team (WGT-Elektronik GmbH & Co KG). Lukas Gradl and Markus Haider (H-Tech) were our partners in software development (DatLab 7.4 and DatLab 8.0 CV-Module, respectively). We thank Marco Di Marcello for excellent technical support in chemical and buffer preparation, mitochondrial isolation, and maintenance of the NextGen-O2k, and Chris Donnelly for stimulating discussions. This work was supported by project NextGen-O2k which has received
funding from the European Union’s Horizon 2020 research and innovation programme under grant agreement N° 859770.

Abbreviations

- $A_{\alpha\beta}$: additivity; $A_{\alpha\beta} = (1-\alpha)/\beta$
- $\alpha$: dominant FCR; $\alpha = J_S/J_{NS}$
- Ama: antimycin A
- $\beta$: sub-dominant FCR; $\beta = J_N/J_{NS}$
- CCCP: carbonyl cyanide m-chlorophenyl hydrazone
- CI to CIV: Complex I to Complex IV
- CoQ: coenzyme Q; mtCoQ: mitochondrial coenzyme Q
- CV: cyclic voltammetry
- $E$: electron transfer capacity, ET capacity
- $E_{p1}, E_{p2}$: oxidation peak potential, reduction peak potential [mV]
- ETS: electron transfer system
- FCR: flux control ratio
- GCE: glassy carbon electrode
- HRR: high-resolution respirometry
- imt: isolated mitochondria
- J: pathway O$_2$ flux
- L: LEAK respiration
- M: malate
- N: NADH-linked pathway
- P: pyruvate
- $P$: capacity of oxidative phosphorylation, OXPHOS capacity
- POS: polarographic oxygen sensor
- Q, $Q_{ra}$: mitochondrial ETS-reactive coenzyme Q
- Ren: residual endogenous respiration; REN: state of Ren
- Rot: rotenone
- Rox: residual oxygen consumption; ROX: state of Rox
- S: succinate
- SUIT: substrate-uncoupler-inhibitor titration
- U: uncoupler

References


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