Research Paper

The interplay between mitochondrial reactive oxygen species formation and the coenzyme Q reduction level

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ABSTRACT

Our aim was to elucidate the relationship between the rate of mitochondrial reactive oxygen species (mROS) formation and the reduction level of the mitochondrial coenzyme Q (mQ) pool under various levels of engagement of the mQ-reducing pathway (succinate dehydrogenase, complex II) and mQH₂-oxidizing pathways (the cytochrome pathway and alternative oxidase pathway, (AOX)) in mitochondria isolated from the amoeba Acanthamoeba castellanii. The mQ pool was shifted to a more reduced state by inhibition of mQH₂-oxidizing pathways (complex III and complex IV of the cytochrome pathway, and AOX) and the oxidative phosphorylation system. The mQ reduction level was lowered by decreasing the electron supply from succinate dehydrogenase and by stimulating the activity of the cytochrome or AOX pathways. The results indicate a direct dependence of mROS formation on the reduction level of the mQ pool for both mQH₂-oxidizing pathways. A higher mQ reduction level leads to a higher mROS formation. For the cytochrome pathway, mROS generation depends nonlinearly upon the mQ reduction level, with a stronger dependency observed at values higher than the mQ reduction level of the phosphorylating state (~35%). AOX becomes more engaged at higher mQ pool reduction levels (above 40%), when mROS production via the cytochrome pathway increases. We propose that the mQ pool reduction level (endogenous mQ redox state) could be a useful endogenous reporter that allows indirect assessment of overall mROS production in mitochondria.

1. Introduction

Mitochondrial coenzyme Q (mQ, ubiquinone) is an essential electron carrier that plays a central role in the mitochondrial electron transport respiratory chain [1,2]. Mitochondrial Q shuttles electrons between dehydrogenases and the oxidizing pathway(s) of the mitochondrial respiratory chain and is also involved in the formation of superoxide from semiubiquinone radicals by the respiratory chain, which can lead to mitochondrial oxidative damage [3–5]. It is widely accepted that mitochondrial reactive oxygen species (mROS) production depends on the level of reduction of mitochondrial electron carriers, especially on mQ. Surprisingly, there are only few data relating mQH₂/mQ ratios [6] or mQ reduction level [7] with mROS generation.

The amoeba Acanthamoeba castellanii is a nonphotosynthesizing free-living protozoan belonging to the group that diverged from the animal/fungal line after the split from plants [8]. As an opportunistic pathogen that can cause serious diseases in humans, A. castellanii is an evolutionarily and medically important amoebozoan. This species presents features, including mitochondrial physiology, that are common to plants, fungi and animals. In addition to several dehydrogenases, the plant-type respiratory chain of A. castellanii mitochondria contains two ubiquinol (mQH₂)-oxidizing pathways, namely, the classical antimycin A- and cyanide-sensitive cytochrome pathway and the alternative benzohydroxamate- and propyl gallate-sensitive ubiquinol oxidase (AOX) pathway [9–11]. The mQ pool plays a central role in the respiratory chain; respiratory substrate-oxidizing dehydrogenases reduce mQ to mQH₂, and the two oxidizing pathways convert mQH₂ to mQ. Electron transfer via the AOX pathway does not result in proton pumping and is therefore not coupled to the mitochondrial production of ATP. The study of mitochondrial respiration of succinate (complex II substrate) in A. castellanii allows investigation of the kinetics of two mQH₂-oxidizing pathways; one proton electrochemical gradient

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The pathway was titrated with increasing concentrations of malonate (an $\Delta\mu$H$^+$-independent pathway, the AOX pathway). Because mROS production depends on $\Delta\mu$H$^+$ [4], investigation of *A. castellanii* mitochondria enables the determination of the relationship between mROS formation and the mQ reduction level at different mitochondrial membrane potential (m$\Delta\Psi$) values depending on the engagement of the two mQH$_2$-oxidizing pathways.

The aim of our work was to elucidate the relationship between mROS formation and the reduction level of the mQ pool under a variety of mitochondrial respiration conditions, i.e., at varying degrees of engagement of mQ-reducing the pathway (succinate dehydrogenase, complex II) and mQH$_2$-oxidizing pathways (the cytochrome pathway and AOX) in isolated *A. castellanii* mitochondria. The mQ reduction level was increased by decreasing electron flow out of the mQ pool via inhibition of the mQH$_2$-oxidizing pathways (complex III, complex IV, or AOX) or inhibition of the oxidative phosphorylation (OXPHOS) system (ATP synthase or ATP/ADP antiporter). The mQ pool was shifted to a more oxidized state by decreasing the electron supply from complex II via inhibition of the mQ-reducing pathway (substrate dehydrogenase) or by stimulation of the activities of the mQH$_2$-oxidizing pathways under uncoupling conditions (the cytochrome pathway) or under GMP activation (AOX). We measured the mQ reduction level under given mitochondrial oxygen consumption and mitochondrial membrane potential (m$\Delta\Psi$) conditions in terms of H$_2$O$_2$ formation.

2. Materials and methods

2.1. Acanthamoeba castellanii cell culture and isolation of mitochondria

Trophozoites of the *A. castellanii* strain Neff (ATCC®30010TM) were cultured as described previously [10]. Cells from 72-h cultures were inoculated (time 0) to a final density of approximately 2.5 ± 0.4 × 10$^5$ cells × ml$^{-1}$. After approximately 40 h of exponential growth with a generation time (cell doubling time) of 8 h, the amoeba cultures reached the intermediate growth phase and then the stationary phase, the latter preceding transformation into cysts within a few hours. In this study, trophozoites of *A. castellanii* were harvested 48 h after inoculation, in the intermediate phase (6.8 ± 0.5 × 10$^6$ cells × ml$^{-1}$). Mitochondria were isolated in an isolation medium containing 0.25 M sucrose, 10 mM Tris/HCl (pH 7.4), 0.5 mM EGTA, and 0.2% bovine serum albumin (BSA) and then purified on a self-generating Percoll gradient (28%) for 45 min at 40,000 × g [9]. Purified mitochondria were washed in isolation medium without BSA and EGTA. Protein concentrations of the isolated mitochondria were determined using the biuret method.

2.2. General measurement conditions

All measurements were performed in a standard incubation medium (28 °C) containing: 120 mM KCl, 20 mM Tris/HCl (pH 7.4), 3 mM KH$_2$PO$_4$, 8 mM MgCl$_2$, 1 mM EGTA, and 0.2% BSA with continuous stirring. Mitochondria (0.33 mg of protein/ml) were incubated in 0.5 ml of the standard incubation medium (see above) with 5 mM succinate as an oxidizable substrate of complex II (in the presence of rotenone (2 µM) to block electron input from complex I). Respiratory rate, m$\Delta\Psi$, mQ reduction levels, and mROS formation were measured under (i) resting (nonphosphorylating, State 4) conditions, i.e., in the absence of exogenous ADP, and (ii) phosphorylating (State 3) conditions, i.e., in the presence of 1–2 mM ADP. The mQ-reducing pathway was titrated with increasing concentrations of malonate (an inhibitor of succinate dehydrogenase, complex II). The activity of a major mQH$_2$-oxidizing pathway (the cytochrome pathway) was varied with increasing concentrations of (i) antimycin A or cyanide (inhibitors of complex III and complex IV of the cytochrome pathway, respectively), (ii) oligomycin or carboxyatractysolide (inhibitors of ATP synthase and ATP/ADP antiporter, respectively), or (iii) uncoupler (carbonyl cyanide p-trifluoromethoxyphenylhydrazone, FCCP) in the presence of AOX inhibitor. Parameters of the cytochrome pathway were measured in the presence of benzohydroxamate (1 mM) (respiratory rate, m$\Delta\Psi$, and mQ reduction level) and propyl gallate (3 µM) (H$_2$O$_2$ formation). The activity of the alternative mQH$_2$-oxidizing pathway (AOX) was varied with increasing concentrations of GMP (an allosteric activator of AOX, [12,13]) in the presence of 0.65 mM cyanide or 90 nM antimycin A (to exclude the activity of the cytochrome pathway). The relationships between the studied respiratory chain parameters (respiratory rate, m$\Delta\Psi$, mQ reduction levels, and H$_2$O$_2$ formation) and increasing concentrations of the respiratory chain and OXPHOS system modulators are shown in the Supplementary materials (Figs. S1–S6).

2.3. Mitochondrial oxygen consumption and m$\Delta\Psi$ measurements

Oxygen uptake was measured polarographically with a Clark-type oxygen electrode (Rank Brothers, Cambridge, UK) in 3.0 ml of incubation medium (28 °C) with 1 mg of mitochondrial protein. Only high-quality *A. castellanii* mitochondrial preparations, i.e., those with ADP/O values of ~1.40 (with succinate as a respiratory substrate) and respiratory control ratios of ~3.5, were used in all the experiments. For OXPHOS control, phosphorylating respiration (State 3) was measured after an ADP pre-pulse (50 µM) using 150 µM ADP as the main pulse. The total amount of oxygen consumed during phosphorylating respiration was used to calculate the ADP/O ratio. The m$\Delta\Psi$ measurements allowed fine control of the duration of phosphorylating respiration. Values of O$_2$ uptake are given in nmol O$_2$ × min$^{-1}$ × mg protein$^{-1}$.

The m$\Delta\Psi$ was measured simultaneously with oxygen uptake using a tetraphenylphosphonium (TPP$^+$)-specific electrode as described previously [14–16]. The TPP$^+$-specific electrode was calibrated with three sequential additions (0.8, 0.8, and 1.6 µM) of TPP$^+$. After each run, 0.5 µM FCCP was added to release TPP$^+$ for baseline correction. For calculation of the m$\Delta\Psi$ value, the matrix volume of the amoeba mitochondria was assumed to be 2.0 µl × mg$^{-1}$ protein. The calculation assumes that the TPP$^+$ distribution between the mitochondria and medium followed the Nernst equation. The values of ΔΨ are given in mV.

2.4. Assay of H$_2$O$_2$ production by isolated mitochondria

The mitochondrial H$_2$O$_2$ production rate was measured by the Amplex Red-horseradish peroxidase method (Invitrogen) [17]. Horseradish peroxidase (0.1 U × ml$^{-1}$) catalyzes the H$_2$O$_2$-dependent oxidation of nonfluorescent Amplex Red (5 µM) to fluorescent resorufin red. Fluorescence was kinetically followed for 10 min at 545 nm (excitation), 590 nm (emission), and gain 150 using an Infinite M200 PRO Tecan multimode reader with 24-well plates. Mitochondria (0.17 mg of mitochondrial protein) were incubated in 0.5 ml of the standard incubation medium (see above) with 5 mM succinate as an oxidizable substrate in the presence of rotenone (2 µM). Because, at high concentrations, AOX inhibitors, benzohydroxamate and propyl gallate, are known to inhibit horseradish peroxidase activity and scavenge free radicals [18–21], H$_2$O$_2$ formation related to the cytochrome pathway activity was measured in the presence of a low concentration of propyl gallate (1.5 µM), which completely inhibits AOX activity. Because cyanide inhibits horseradish peroxidase activity [22], the H$_2$O$_2$ formation rate related to AOX activity was measured in the presence of 90 nM antimycin A to inhibit cytochrome pathway activity. Therefore, titration of H$_2$O$_2$ formation related to cytochrome pathway activity was performed only in the presence of increasing concentrations of antimycin A not cyanide. Reactions were monitored with constant stirring at 28 °C and calibrated with known amounts of H$_2$O$_2$ in the absence or presence of 1.5 µM propyl gallate. H$_2$O$_2$ production rates were determined from slopes calculated from readings obtained from several repeated 10-min measurements. Values of H$_2$O$_2$ production are given in...
pmol H$_2$O$_2$ × min$^{-1}$ × mg protein$^{-1}$.

2.5. Determination of the mQ reduction level

The reduction level of mQ (mQH$_2$/mQtot), i.e., the ratio of reduced mQ (ubiquinol, mQH$_2$) to the total endogenous pool of mQ in the inner mitochondrial membrane under steady-state respiration was determined by extraction followed by HPLC detection [23]. Mitochondria (1–2 mg of mitochondrial protein) in the steady state were chemically quenched with 4 ml of 0.65 M HClO$_4$ in methanol (0°C), and mQ was subsequently extracted with 3 ml of petroleum ether. Detection of the oxidized and reduced forms of mQ (at 275 nm and 290 nm, respectively) was performed by HPLC with a reverse-phase Lichrosorb 10 RP 18 column (4.6 mm × 250 mm). A completely oxidized extract was obtained during incubation in the absence of a reducing substrate using an evaporation/ventilation step. A completely reduced extract was obtained upon anaerobiosis and in the presence of respiratory substrate (5 mM succinate), 1 mM cyanide and 1 mM benzohydroxamate. As previously determined, the endogenous Q in A. castellanii mitochondria is Q$_0$ [12]. Commercial Q$_0$ (Sigma) was used for peak calibration. The reduction level of mQ is expressed as the percentage of total mQ (mQH$_2$/mQtot).

3. Results

Cytochrome pathway activity was measured in the presence of AOX inhibitor (1 mM benzohydroxamate or 1.5 μM propyl gallate). The activity of the alternative mQH$_2$-oxidizing pathway (AOX) was measured in the presence of 0.65 mM cyanide or 90 nM antimycin A to exclude cytochrome pathway activity.

3.1. The relationship between mROS formation and the reduction level of the mQ pool under various degrees of engagement of the cytochrome pathway

At first, the mQ pool was progressively shifted to a more oxidized state by stimulating the activity of the mQH$_2$-oxidizing pathway (the cytochrome pathway) by uncoupling with increasing concentrations of FCCP (up to 500 nM) (Fig. 1, S1). Fig. 1 and B show changes in respiratory rate (progressively increasing from ~55 to ~195 pmol O$_2$ × min$^{-1}$ × mg protein$^{-1}$) and H$_2$O$_2$ formation rate (progressively decreasing from ~100 to ~390 pmol H$_2$O$_2$ × min$^{-1}$ × mg protein$^{-1}$) when the mΨ of nonphosphorylating mitochondria was gradually decreased (from ~189 to ~160 mV) by the uncoupler. The uncoupler-induced changes in the respiratory and H$_2$O$_2$ formation rates were accompanied by a gradual decrease in the mQ reduction level (from ~69 to ~28%) (Fig. 1C–E). A linear dependence of mΔΨ on the mQ reduction level for the cytochrome pathway under the uncoupling conditions was observed (Fig. 1E).

Next, the mQ reduction level of the cytochrome pathway under phosphorylating conditions was gradually increased by inhibition of the OXPHOS system with increasing concentrations of carboxyatractyloside (up to 325 nM) or oligomycin (120 nM) (Fig. S2). Fig. 2 shows the relationships between the respiratory rate and H$_2$O$_2$ formation versus mΔΨ (Fig. 2A, B) and mQ reduction level (Fig. 2C, D) when the cytochrome pathway activity of phosphorylating mitochondria (in the presence of 1 mM ADP) was gradually varied with the inhibitors of ADP/ATP antiporter and ATP synthase. During carboxyatractyloside/oligomycin titration, an increase in mΨ from ~160 mV (uninhibited phosphorylating state) up to ~192 mV (fully inhibited phosphorylating state) was accompanied by an increase in mQ reduction from ~30% (uninhibited phosphorylating state) to ~72% (nonphosphorylating state in the presence of OXPHOS inhibitors). A linear dependence of mΔΨ on the mQ reduction level for the cytochrome pathway under phosphorylating conditions (during the carboxyatractyloside/oligomycin-induced State 3 to State 4 transition) is presented in Fig. 1E. Moreover, a progressive decrease in the respiratory rate from ~195 to ~50 pmol O$_2$ × min$^{-1}$ × mg protein$^{-1}$ and a progressive increase in H$_2$O$_2$ formation from 110 to ~440 pmol H$_2$O$_2$ × min$^{-1}$ × mg protein$^{-1}$ accompanied the carboxyatractyloside/oligomycin-induced State 3 to State 4 transition (Fig. 2A–D).

Rates of H$_2$O$_2$ formation were elevated up to 4 times in the resting State 4 (in the absence of or presence of OXPHOS inhibitors) compared to the rates in the phosphorylating State 3 or uncoupled state (Figs. 1, 2), because higher (up to ~30 mV) mΔΨ and higher (up to 40%) mQ reduction levels were observed.

Furthermore, the mQ reduction level of the cytochrome pathway was gradually increased by decreasing the electron flow out of the mQ pool via inhibition of complex III by increasing the concentrations of antimycin A (up to 60 nM) and complex IV by increasing the concentrations of cyanide (up to 0.4 mM) (Fig. S3). Measurements were performed under phosphorylating (State 3) and nonphosphorylating (State 4) conditions. Because cyanide inhibits peroxidase activity, measurements of H$_2$O$_2$ formation were performed only during antimycin A titrations. Fig. 3 shows the relationships between respiratory rate and H$_2$O$_2$ formation versus mΔΨ (Fig. 3A, B) and versus mQ reduction level (Fig. 3C, D) when the cytochrome pathway activity was gradually inhibited. The relationships obtained during the antimycin A and cyanide titrations were similar. A much steeper linear dependence of mΔΨ on the mQ reduction level was obtained for the cytochrome pathway under nonphosphorylating conditions than under phosphorylating conditions (Fig. 3E). During antimycin A/cyanide titration under nonphosphorylating conditions, a progressive decrease in mΔΨ from ~189 mV (uninhibited State 4) up to ~136 mV (fully inhibited State 4) was accompanied by a relatively moderate increase in mQ reduction level from ~69% (uninhibited State 3) to ~81% (fully inhibited State 3) (Fig. 1E). This 12% increase in the mQ reduction level was accompanied by an ~1.2-fold increase in the H$_2$O$_2$ formation rate (Fig. 3D). Under phosphorylating conditions, a progressive decrease in mΔΨ from ~160 mV (uninhibited State 3) up to ~141 mV (fully inhibited State 3) was accompanied by a progressive increase in mQ reduction from ~30% (uninhibited State 3) to ~77% (fully inhibited State 3) (Fig. 1E). This 47% increase in the mQ reduction level accompanying the inhibition of the cytochrome pathway in State 3 was accompanied by an ~4.2-fold increase in the H$_2$O$_2$ formation rate (Fig. 3D).

The results indicate a direct dependence of mROS formation on the reduction level of the mQ pool (and mΔΨ) under various levels of engagement of the cytochrome pathway. The higher mQ reduction level (and the larger mΔΨ) the bigger mROS production.

3.2. The relationship between mROS formation and the reduction level of the mQ pool under varying activity of complex II when the cytochrome pathway is engaged

The dependence of the rate of electron transfer through the mQ reducing enzyme (complex II) on the reduction level of the mQ pool (and mΔΨ) when the cytochrome pathway is active was studied by malonate titration under phosphorylating and nonphosphorylating conditions. Malonate titration enabled lowering of the mQ reduction level below 30% (reached with FCCP titration, Fig. 1C–E), leading to an ~10% reduction level for both respiratory states (Fig. 4C–E). At levels below the mQ pool reduction level (and mΔΨ) of uninhibited State 3, i.e., below 30% of the mQ reduction level (and below ~160 mV), the kinetic relationships between H$_2$O$_2$ formation versus mΔΨ (Fig. 4B) and versus mQ reduction level (Fig. 4D) overlapped for both respiratory states. Under both phosphorylating and nonphosphorylating conditions, H$_2$O$_2$ formation depends nonlinearly upon the mQ pool reduction level (and mΔΨ). The threshold values for the mQ pool reduction level and mΔΨ are slightly greater than the values for the uninhibited State 3, i.e., an ~35% reduction level of mQ and ~165 mV of mΔΨ. At values greater than these thresholds, a sharply increased dependence of...
H₂O₂ formation on the mQ reduction level (and mΔΨ) was observed (Fig. 4B, D); therefore, a small increase in the mQ reduction level (and mΔΨ) leads to a high rate of H₂O₂ production by mitochondria.

3.3. The relationship between mROS formation and the reduction level of the mQ pool under varying activity of AOX

When the cytochrome pathway was inactive (in the presence of high concentrations of cyanide or antimycin A), the activity of the unstimulated AOX pathway led to low levels of oxygen uptake (~ 20 mol O₂ × min⁻¹ × mg protein⁻¹) accompanied by very high mQ reduction levels (~ 77%) and H₂O₂ formation (~ 230 pmol H₂O₂ × min⁻¹ × mg protein⁻¹) (Fig. 5). The AOX-mediated mROS formation is independent of mΔΨ (and ΔμH⁺), since no proton pumping occurs in the respiratory pathway when AOX works with complex II. Increasing the concentration of GMP progressively stimulated AOX activity, leading to up to ~ 4.5 times higher respiratory rates, up to ~ 1.3 times lower H₂O₂ formation, and a considerable decrease in mQ reduction levels (up to ~ 59%). Fig. 5 shows the approximately linear relationships between respiratory rate (Fig. 5A) and H₂O₂ formation (5B) versus mQ reduction level (ranging from 59% to 77%) when the AOX activity was gradually stimulated by GMP.

3.4. The relationship between mROS formation and the reduction level of the mQ pool under varying activity of complex II when AOX is engaged

The dependence of the rate of electron transfer through the mQ-reducing enzyme (complex II) on the reduction level of the mQ pool when AOX is active was studied by malonate titration in the absence or presence of GMP (Fig. 6). A progressive decrease in respiratory rate and
H$_2$O$_2$ formation was observed with lowering of the mQ pool reduction level from ~77 to ~52% and from ~59 to ~42% for unstimulated and GMP-stimulated succinate-sustained AOX, respectively. The kinetic relationships between H$_2$O$_2$ formation versus mQ reduction level (Fig. 6B) overlapped for both conditions and were approximately linear in the range of the mQ reduction level of AOX during succinate oxidation. These results indicate that AOX becomes engaged at only high reduction levels of the mQ pool (greater than 40%).

3.5. Inactive mQ pool

In A. castellanii mitochondria, a completely oxidized mQ pool (0% mQ reduction level) was obtained after incubation of mitochondria in the absence of a reducing substrate. Upon anaerobiosis and in the presence of a respiratory substrate (5 mM succinate), 1 mM cyanide, and 1 mM benzoxyatractylside, a completely reduced mQ pool with ~85% reduction was obtained. This result indicates that in A. castellanii mitochondria isolated from the intermediate phase of growth, ~15% of the mQ pool is unreducible.

4. Discussion

The mQ reduction shift that occurs when mitochondria switch from nonphosphorylating conditions (State 4, a more reduced state) to phosphorylating conditions (State 3, a more oxidized state) is of utmost physiological significance. In vivo, mitochondria can shift rapidly between these conditions inducing a change in the mQ reduction level and thereby in mROS formation. Our study combined quantitatively important mitochondrial parameters: mΔΨ, oxygen uptake, mROS formation, and mQ pool reduction level, with a special focus on the
relationship between mROS formation and mQ reduction level. In *A. castellanii* mitochondria, during nonphosphorylating respiration, high mQ reduction levels were accompanied by increased mΔΨ values and mROS formation. In contrast, during phosphorylating respiration, the relative oxidation of the mQ pool was accompanied by decreased mΔΨ values and therefore by decreased mROS formation. In isolated mitochondria, the relationship between mROS formation and the mQ pool reduction level can be elucidated by varying the mQ pool reduction level (mQH2/mQtot) by using agents that cause both stimulation and inhibition of respiratory chain electron transport.

In *A. castellanii* mitochondria, the respiratory activity of the cytochrome pathway is much higher than that of AOX. The cytochrome pathway becomes engaged at much lower mQ pool reduction levels (from ~10%) (Fig. 4C–E) than the AOX pathway (from ~40%) (Fig. 6).

Comparing the relationships between mROS formation and the mQ reduction level of the mQ pool for both mQH2-oxidizing pathways (Fig. 7C), it is clear that at higher mQ reduction levels, the cytochrome pathway produces much more (even twice as much) mROS compared to AOX. At ~40% of mQ reduction level, the mROS formation is similar for both mQH2-oxidizing pathways (~130 pmol H2O2 × mg protein−1).

The relationship between mROS formation and the mQ pool reduction level is much steeper for the cytochrome pathway than for AOX; for the cytochrome pathway, H2O2 formation depends strongly upon the mQ pool reduction level in the 40–80% range (Fig. 7C). Above the threshold value of the mQ pool reduction level, i.e., slightly above the value for the uninhibited State 3 (~35%), which corresponds to ~165 mV mΔΨ (Fig. 7A, B), a small increase in the mQ reduction level gives rise to a high rate of H2O2 production by mitochondria, with oxidation...
occurring via the cytochrome pathway. It has been shown previously that in nonphosphorylating rat heart mitochondria, mROS generation strongly but nonlinearly depends upon $m\Delta\Psi$, increasing at $m\Delta\Psi$ greater than that of State 3. Our results confirm this observation, however indicating the contribution of the $mQ$ reduction level to mROS formation.

To date, at least 11 sites that produce superoxide anion and/or $H_2O_2$ have been identified in mammalian mitochondria. Under our experimental conditions, during succinate oxidation, the flavin site of mitochondrial complex II (site II$F$ of complex II) produced mROS when AOX was active, while site II$F$ of complex II and the Qo site of mitochondrial complex III (site IIIQo) participated in mROS production when the cytochrome pathway was active. The highest $H_2O_2$ formation was observed in the presence of antimycin A (Fig. 7A, C), a Q site inhibitor of complex III. It has been shown that contributions of specific sites of the mitochondrial respiratory chain to the production of ROS in mitochondria depend very strongly on the substrates being oxidized. Our results indicate that the production of ROS in mitochondria depends not only on the engagement of mROS formation sites but also on the engagement of QH$_2$-oxidizing pathways.

There are only few studies relating $mQ$ reduction level with mROS generation, which is likely due to difficulties in the measurement of the $mQ$ reduction level. In different cell lines, it has been shown how the mitochondrial electron transport chain is optimized to better oxidize different fuels using the reducing status of $mQ$ (mQH$_2$/Q ratio) as a metabolic sensor and mROS generated by complex I by reverse electron transport as an executor. In rat heart mitochondria, mROS production by reverse electron transport at complex I is steeply dependent on $m\Delta\Psi$ and on $mQ$ reduction level, indicating the sensitivity of superoxide anion production by this respiratory chain site to the two

Fig. 4. The relationships between respiratory rate (A) and $H_2O_2$ formation (B) versus $m\Delta\Psi$ and the relationships between respiratory rate (C), $H_2O_2$ formation (D), and $m\Delta\Psi$ (E) versus $mQ$ reduction level when the cytochrome pathway is engaged. Complex II (CII) activity was varied with increasing concentrations of malonate (up to 4 mM). Measurements were performed under phosphorylating (State 3) and nonphosphorylating (State 4) conditions, in the presence of 1 mM benzhydroxamate or 1.5 $\mu$M propyl gallate (H$_2$O$_2$ formation). Arrows indicate the starting point and direction of titrations.
physiological variables [7]. Marphy’s group has shown that in the presence of antimycin A, the production of superoxide at the Qo site is specifically linked to local accumulation of reduced cytochrome b566 when the Qi site is inhibited by antimycin A, and this effect is independent of mΔΨ [7]. Our results indicate that mROS production is a direct function of the mQ pool reduction level and is independent of mΔΨ when the Qi site is inhibited by antimycin A and when reverse electron transport is excluded by rotenone during succinate oxidation.

Complex I is believed to be a major site of mROS production in the mitochondrial electron transport chain, either from FAD- or NAD-driven electron flux [3]. It is also generally assumed that the generation of mROS when succinate is used as the substrate depends mostly on reverse electron transfer through complex I. Our study shows that during succinate oxidation, the production of mROS may be very high even when complex I is inhibited by rotenone. Further studies are needed to elucidate whether reverse electron transfer through complex I is the major source of mROS in A. castellanii mitochondria.

The present study indicates that in A. castellanii mitochondria, AOX becomes more engaged at higher mQ pool reduction levels, when mROS production via the cytochrome pathway increases (Fig. 7C). Activation of cyanide- and antimycin A-resistant AOX-mediated respiration by GMP significantly decreased mQ reduction levels and H2O2 formation (Figs. 5, 4), confirming the antioxidant role of the oxidase. It has been previously shown that in A. castellanii mitochondria, AOX may play a role not only in the energetic status of the cell by decreasing the yield of OXPHOS [12] but also in preventing the generation of mROS [27], which are maintained at a constant level throughout the growth cycle of amoeba batch culture [28]. In the present study, we found an ~15% unreducible mQ pool in mitochondria isolated from A. castellanii cells from an intermediate phase of growth. A question arises as to whether the size of the unreducible mQ pool in the inner mitochondrial membrane influences mQ reduction level-dependent mROS formation. Further studies are needed to answer this question. Our previous study indicates that, compared to A. castellanii cells from the intermediate phase of growth, decreased reducible mQ levels in intensively dividing cells (i.e., at the exponential phase of growth) accompanied by increased mQ reduction levels could lead to increased mROS formation and thereby to increased AOX activity (which depends on the mQ reduction level) [11,28].

Mitochondrial ROS production has been described to be a direct function of mΔΨ [4,29]. Fig. 7A is a summative description of the relationships between H2O2 formation versus mQ reduction level when AOX activity was varied with increasing concentrations of GMP (up to 1 mM). Measurements were performed in the presence of 0.65 mM cyanide or 90 nM antimycin A. Arrows indicate a starting point and direction of titrations.

Fig. 5. The relationships between respiratory rate (A) and H2O2 formation (B) versus mQ reduction level when AOX activity was varied with increasing concentrations of GMP (up to 1 mM). Measurements were performed in the presence of 0.65 mM cyanide or 90 nM antimycin A. Arrows indicate a starting point and direction of titrations.

Fig. 6. The relationships between respiratory rate (A) and H2O2 formation (B) versus mQ reduction level when AOX is engaged in the absence or presence of 1 mM GMP. Complex II (CII) activity was varied with increasing concentrations of malonate (up to 4 mM). Measurements were performed in the presence of 0.65 mM cyanide or 90 nM antimycin A. Arrows indicate the starting point and direction of titrations.
observations and the fact that AOX-mediated mROS formation is mΨ independent, we postulate that mROS production is a direct function of the mQ pool reduction level rather than of mΔΨ. The dependence of H2O2 production on the mQ reduction level is clearly evidenced in this study for both QH2-oxidizing pathways (Fig. 7C). Therefore, the mQ pool reduction level (endogenous mQ redox state) could be a useful endogenous reporter that allows indirect assessment of overall mROS production in mitochondria. Most of the mitochondrial sites of superoxide anion/H2O2 formation are related to the reduction level of the mQ pool that connects the dehydrogenase and oxidase sides of the electron transport chain. For example, the rates of mitochondrial superoxide anion/H2O2 production from site IF of complex I and site Qo of complex III can be assessed indirectly by measuring endogenous reporters such as the mitochondrial NAD(P)H redox state and the cytochrome b566, respectively [25,26]. Measurement of the endogenous mQ pool reduction level could provide insight into overall mROS production related to mitochondrial respiratory chain complexes. The present study analyses the dependence of H2O2 production on the mQ pool reduction level in mitochondria, by using A. castellanii mitochondria as an example, under conditions in which concentrations of agents that cause both stimulation and inhibition of respiratory chain electron transport (substrate, inhibitors and activators) were varied. In future studies, the measurements of the mQ pool reduction level under physiological conditions in the absence of inhibitors of electron transport may be helpful for the assessment of overall intrinsic production of mROS in mitochondria and the physiological role of these species as signaling molecules and in pathologies.

5. Conclusions

We elucidated for the first time the relationship between the rate of mROS formation and the reduction level of the endogenous mQ pool varied by agents that cause both stimulation and inhibition of the mitochondrial respiratory electron transport chain. Our results indicate that mROS production is a direct function of the mQ pool reduction level rather than of mΔΨ. The production of ROS in mitochondria depends not only on the engagement of mROS formation sites but also on the engagement of QH2-oxidizing pathways. The reduction level of the mQ pool could be a useful endogenous reporter that allows indirect assessment of overall mROS production in mitochondria.

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Conflict of interest/declarations of interest

The authors declare no conflict of interest.

Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.redox.2018.07.018.

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