

Partitioning between cytochrome c oxidase and alternative oxidase studied by oxygen kinetics of dark respiration in *Chlamydomonas reinhardtii*: a microalgae model organism

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http://www.mitofit.org/index.php/Di_Marcello_2019_MitoFit_Preprint_Arch



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Editor MitoFit Preprint Archives: Gnaiger E

Introduction

Bioenergetics is the study of how living organisms acquire and transform energy to perform biological work. Energetic coupling between chloroplasts and mitochondria has been described in algae, demonstrating the a good functionality and interaction between both organelles is necessary to maintain metabolic integrity. High-resolution respirometry (HRR) is widely used to assess mitochondrial respiration and other bioenergetics parameters in the biomedical field of mitochondrial research and its clinical applications [1]. In our interdisciplinary study, we adapted the multimodal approach of the Oroboros O2k high-resolution respirometer to investigate algal bioenergetics for biotechnological purposes.

In contrast to mammalian cells, algal mitochondria possess alternative oxidases (AOX), which bypass electron transfer from the Q-junction through Complexes CIII and CIV [2]. Therefore, in algae we can distinguish between AOX-dependent and cytochrome c oxidase-dependent respiration through the Q-AOX and CIII-CIV pathways.

Material and methods

The microalgal model organism *Chlamydomonas reinhardtii* wild-type strain *wt12* was grown at RT in Tris-Acetate-Phosphate (TAP) medium in a 16:8 h light:dark cycle. Oxygen flux, J_{O_2} , was monitored in *wt12* living cells in the exponential growth phase at 25 °C in Oroboros O2k high-resolution respirometers excluding any light in the chambers. Substrate-uncoupler-inhibitor titration (SUIT) protocols were specifically developed to characterise activities of the Q-AOX pathway (SUIT-022 [3]) and CIII-CIV pathway (SUIT-023 O₂ [4]). To quantify the contribution of the Q-AOX pathway to algal dark respiration, we studied the

oxygen kinetics of (1) ROUTINE-respiration in TAP medium, (2) Q-AOX dependent respiration after inhibition of CIV with 1 mM potassium cyanide (KCN), and (3) CIII-CIV dependent respiration after inhibition of AOX with 1 mM salicylhydroxamic acid (SHAM). Oxygen kinetics was obtained from aerobic-anaerobic transitions with high time resolution at a data sampling interval of 0.2 s. p_{50} is the O_2 partial pressure, p_{O_2} , at 50% of maximal respiration, J_{max} [5]. The p_{50} was calculated from hyperbolic fits using the Oroboros O2Kinetics software for automatic O_2 calibration, correction for zero O_2 signal drift, instrumental background O_2 flux and exponential time constant of the polarographic oxygen sensor [6]. A single shifted hyperbolic fit was used to fit J_{O_2} as a function of p_{O_2} in each aerobic-anaerobic transition.

Results and conclusions

p_{50} ranged from 0.06 to 0.08 kPa for ROUTINE-respiration with an excellent fit by a first-order hyperbolic function. This oxygen affinity is comparable to that in small mammalian cells [7]. Upon inhibition of CIV with KCN, J_{O_2} was significantly impaired (Fig. 1A) and p_{50} increased three-fold up to 0.35 kPa. No decline of J_{O_2} and p_{50} was observed relative to ROUTINE-respiration after inhibition of AOX with SHAM (Fig. 1B). In all cases, excellent fits of respiration as a function of oxygen pressure were obtained by a first-order hyperbolic function.

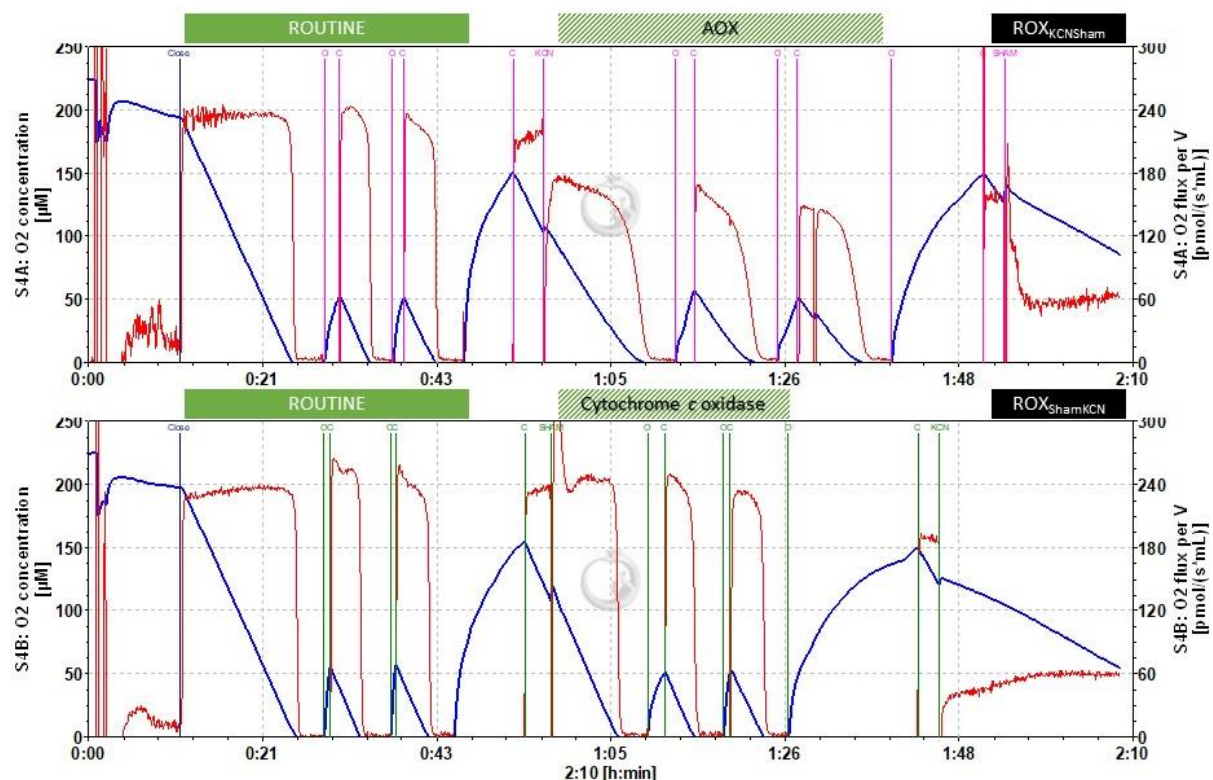


Figure 1. High-resolution respirometry for the study of dark respiration and O_2 kinetics with *C. reinhardtii* wt12. Representative O2k traces showing O_2 concentration and O_2 flux per chamber volume with repeated aerobic-anaerobic transitions (O_2 kinetics) and re-oxygenations. **A:** Protocol SUIT-022: AOX-ce CN+SHAM. **B:** Protocol SUIT-023: AOX-ce SHAM+CN. Note the high technical reproducibility of ROUTINE-respiration in both protocols, and the identical and relatively high residual oxygen consumption, Rox , after titration of both inhibitors in both protocols.

If the potential contribution of the Q-AOX pathway in the ROUTINE-state would be compensated for by increased CIII-CIV pathway flux after addition of SHAM, then the mixed Q-AOX and CIII-CIV pathways would give rise to biphasic hyperbolic oxygen kinetics, with a contribution of the high-affinity CIII-CIV pathway and the low-affinity Q-AOX pathway. Taken together, our results provide evidence against a contribution of AOX to ROUTINE-dark respiration in *wt12* cells under the presently applied culture conditions. Oxygen kinetics provides a fast and simple method for detection of AOX and CIV activities in dark respiration of living microalgal cells, extending our current understanding of the different O₂ affinities of the two pathways in these organisms and their possible effects on the bioenergetics and metabolism of the cells.



Supported by project NextGen-O2k, which has received funding from the European Union's Horizon 2020 research and innovation programme under grant agreement No 859770

References

1. Doerrier C, Garcia-Souza LF, Krumschnabel G, Wohlfarter Y, Mészáros AT, Gnaiger E (2018) High-Resolution FluoRespirometry and OXPHOS protocols for human cells, permeabilized fibers from small biopsies of muscle, and isolated mitochondria. *Methods Mol Biol* 1782:31-70. - www.bioblast.at/index.php/Doerrier_2018_Methods_Mol_Biol
2. Young L, Shiba T, Harada S, Kita K, Albury MS, Moore AL (2013) The alternative oxidases: simple oxidoreductase proteins with complex functions. *Biochem Soc Trans* 41:1305-11.
3. www.bioblast.at/index.php/SUIT-022_O2_ce_D051
4. www.bioblast.at/index.php/SUIT-023_O2_ce_D053
5. Gnaiger E (2001) Bioenergetics at low oxygen: dependence of respiration and phosphorylation on oxygen and adenosine diphosphate supply. *Respir Physiol* 128:277-97. - www.bioblast.at/index.php/Gnaiger_2001_Respir_Physiol
6. Meszaros AT, Haider M, Di Marcello M, Gnaiger E (2018) High-resolution mitochondrial oxygen kinetics as diagnostic tool in Complex IV impairments. Abstract Mitochondrial Medicine 2018 Hinxton UK. - www.bioblast.at/index.php/Meszaros_2018_Mt_Med_Hinxton
7. Scandurra FM, Gnaiger E (2010) Cell respiration under hypoxia: facts and artefacts in mitochondrial oxygen kinetics. *Adv Exp Med Biol* 662:7-25. - www.bioblast.at/index.php/Scandurra_2010_Adv_Exp_Med_Biol