оковокоз INSTRUMENTS high-resolution respirometry O2k-Protocols Demo



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O2k-Fluorometry: HRR and simultaneous determination of mtmembrane potential with safranin

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1. Introduction

The general methodological outline, potential caveats to be observed, and an example of a hypothetical application of combined high-resolution respirometry (HRR) and fluorescence-based determination of mitochondrial transmembrane potential ($\Delta \Psi_{mt}$ or mtMP) are described in the publication by Krumschnabel et al. [1]. Here, a step-by-step description of data acquisition and analysis is described based on the example given in Fig. 9.6 of the publication.

2. The O2k Demo experiment

A substrate-uncoupler-inhibitor titration (SUIT) protocol is applied that examines Complex II-linked respiration of mouse brain first spin homogenate at different coupling states and the concurrent changes of mtMP as assessed by the use of safranin (Saf). Sample preparation as well as the characteristics of the different coupling states elicited by additions in the SUIT protocol are described in detail in [1], [2] and [3].

3. Setting the instrument for data acquisition

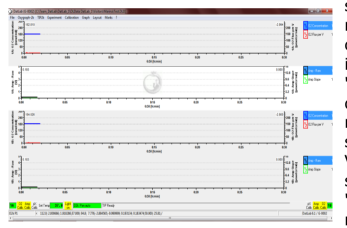
Instrumental setup and preparation with regard to measuring respiration are outlined in [4]. In addition, the O2k-Fluorescence LED2-Module consisting of the Fluorescence-Control Unit and, in the case of using Saf, two Fluorescence Sensors Blue, are connected to the O2k (for details see: [5]) and switched on by pressing the power switch on the front panel of the Fluo Control Unit. Next, DatLab is started and the O2k Control window is opened by pressing F7.

Control of LED-intensity and Amplification: In the Control window the Amperometric tab window is opened and the sensor numbers are defined for documentation. Then, Polarization voltage and Gain for sensor are selected, which for the experiment to be described were 500 mV and 1000 mV, respectively. In general, the ideal settings for polarization

02k Control 0xygen, 02 Amp	erometric, Amp		
02k serial number	G-(
Power-02k			
Chamber	Α	В	
Channel: Amperometric, Amp Gain for sensor	1000 -	1000 -	
Polarization voltage [mV]	500	500	
Amp sensor #	A-0008	B-0117	
		Save	
			Send to Oxygraph-2k
•	Load setup Save s	etup	Cancel

voltage need to be determined in preliminary experiments. Settings are optimal when a signal is obtained which is both large enough to avoid limitation of resolution by digital noise but also well below the range maximum of 10 V, keeping in mind that the maximum signal obtained in the course of an experiment may be larger than the initial one but must also not exceed this value.

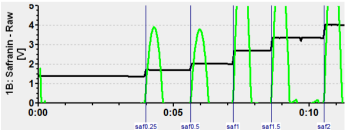
Selecting the appropriate Layout: During data acquisition the experimenter should be able to assess whether the signal obtained is within the desired range and if it stable or has attended a new steady



state following any experimental manipulation. Thus, in the case of using Saf it is appropriate to initially select the lavout 'Safranin signal' which raw displays data related to respirometry according to the standard layout '05 Flux per Volume corrected' and the Saf signal as plots of the raw signal 'Amp Raw' [V] just below each related plot of the oxygen signal.

4. Data acquisition

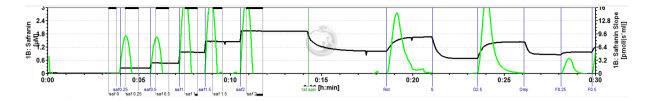
Following these initial adjustments the chambers containing respiration medium MiR06Cr were closed and the chamber lights were switched off.



After recording a baseline, Saf was added from a 200 μ M stock solution in 5 consecutive steps to obtain final Saf concentrations of 0.25, 0.5, 1, 1.5, and 2 μ M in the chamber. Given the

excellent linearity of fluorescence signal over this concentration range (see Fig. 9.1 of [1]), a single addition of 2 μM Saf may be regarded

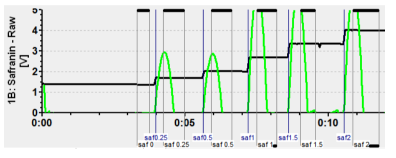
suffcient if a simple conversion of the raw signal to Saf concentration is desired.



Subsequently, mouse brain first spin homogenate (Thom) was injected, resulting in an uptake of the dye and a corresponding decline of Saf fluorescence and this was followed by additions of SUIT chemicals causing changes in respiratory rate and of mtMP as described in [1]. If the experimenter wants to observe Saf fluorescence converted to Saf concentration in real-time, calibration can be done while the experiment is running. Alternatively, calibration can be done during data analysis, both following the same procedure as outlined next.

5. Data analysis

For calibration of the Saf signal the appropriate plot of the raw signal of



chamber A or B are selected and marks are set to data regions before and after stepwise addition of Saf. Clicking into the black bar on the top or bottom of the marks, each mark should be named according to

the final concentration of Saf obtained and the concentration entered in 'Value'. Next, the Calibration window is opened by selecting 'Calibration' and 'Amperometric, Amp' and the marks created are displayed in the

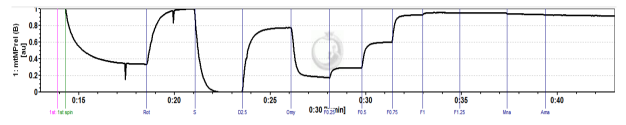
Channel type: 1B: Amp	Active se	nsor #								
Channel label: Safranin		Unit: µM	·							
Signal Slope										
Current calibration			Calibrate							
Calibration source			Select marks	Ente	r conceni	tration			Copy fr	om file
Active file Calib. Sensor #			saf 0		lame	Time	Signal [V]	Slope	Conc.	
			✓ saf 0.25		af 0.25	00:04:36	1.70603		✓ 0.25	5000
Name Signal [V]	Slope	Conc. [µM]	✓ saf 0.5 ✓ saf 1		af 0.5	00:06:16	2.02022			0000
			🖌 saf 1.5	s	af 1	00:07:51	2.69305	-0.17955	✓ 1.00	0000
			✓ saf 2	s	af 1.5	00:09:17	3.36142	-0.25867	✓ 1.50	
				► s	af 2	00:11:19	4.01059	-0.26207	✓ 2.00	0000
										Ŧ
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			Reset to	Intercept [л Г	1.3628	R ² =0.99998	Show graph		
			system default	intercept [, I)					
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								Cance	i li ca	alibrate

center of the window. The marks desired to be included in the calibration are selected by ticking the field next to each mark name and and Saf names concentrations will appear on the right side of the window. DatLab then calculates the parameters of a linear regression analysis

through these data and the values for Sensitivity [V/ μ M] and the Intercept as well as the coefficient of determination, R², are displayed in

the lower part of the window. For a visual inspection of the calibration the data can be viewed by pressing 'Show graph'. Individual calibration marks may now be selected or deselected as required and the final selection is then used for calibration by pressing 'Calibrate'. All data of the selected plot are now converted to Saf concentration and are being displayed as 'Safranin [μ M]'.

Conversion of changes in fluorescence or Saf concentration into mV changes of mtMP may be achieved using the K^+ selective ionophore valinomycin, adding known amounts of KCl solution and applying the Nernst equation, see [6].



Alternatively, the signal may be normalized to detect changes within minimum and maximum fluorescence observed during the experiment, which in the present experiments corresponds to the Saf signal elicited in the CII-linked LEAK state, $S(Rot)_L$, and CII-supported ETS capacity, respectively. Regarding the latter, it should be kept in mind that complete collapse of mtMP may only be observed at an uncoupler concentration that required for stimulation of maximum uncoupled surpassing respiration which may actually already become inhibitory again. Of note, we found that fluorescence may show an apparent drift in the uncoupled state such that the observed maximum signal is not seen in the fully during uncoupled state but Rot-inhibited respiration of tissue homogenate, i.e. in the initial ROX state.

The step-by-step procedure to obtain this normalized plot is as follows:

- * Select plots to display the fluorescence signals as either raw signal or calibrated signal
- * Set marks on the traces corresponding to the minimum and maximum signal observed during the experimental part of the data, NOT the calibration part.
- * Display the marks and keep a note of the minimum and maximum value (or copy and paste marks to Excel)
- * Start the scripting function by pressing [ctrl+Alt+D]
- * Press 'Generate plot', then 'Load macro'; alternatively, if the macro is not available in your program, enter it manually as displayed below; in addition, on the right hand side of the macro window right mouse click on the channel '1B: Safranin - Raw' and select 'A' to name the variable for the macro function. Next, press 'Save macro' if you want to re-use it for subsequent experiment analysis.
- * Select 'mtMPrel(A).DLC' and open it

MiPNet20.13

Edit macro	1A: O2 Raw Signal
	1A: 02 Concentration
Name with Deal (D)	1A: O2 Pressure kPa
Name mtMPrel (B) Unit au	1A: O2 Pressure mmHg
	1A: O2 Slope uncorr.
Editor Debugger Help	1A: O2 Flux per V
Eakor Debugger Help	1A: O2 Flux per unit
	1B: O2 Raw Signal
1 var min, max: Float;	1B: 02 Concentration
2 min:= 2.3048;	1B: O2 Pressure kPa
- min 2.3040,	1B: O2 Pressure mmHg
3 max:=3.6393;	1B: O2 Slope uncorr.
	1B: O2 Flux per V
4	1B: O2 Flux per unit
5	1: Block Temp.
	1: Barom. Pressure
6 FuncResult:= (A-min) / (max-min);	1: Peltier Power
	1A: pX - Raw
	1A: pX
	1A: pX - slope

- * Enter noted values of min and max for Chamber A into the corresponding lines and click 'Update plot'
- * Press 'Generate plot', then 'Load macro'
- * Select 'mtMPrel(B).DLC' and open it
- * Enter noted values of min and max for Chamber B into the corresponding lines and click 'Update plot'
- * Finally, in the 'Select plot' menu tick 'Calc' '1: mtMPrel(A)' and '1: mtMPrel(B)' for displaying the normalized plots of chambers A and B, respectively.

6. References

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http://wiki.oroboros.at/index.php/O2k-Demo experiments#O2k-Demo O2k-MultiSensor

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1B: pX - Raw 1B: pX

1B: pX - slope 1A: Safranin - Raw

1A: Safranin 1A: Safranin - slope

1B: Safranin - Raw 1B: Safranin

1B: Safranin Slope