O2k-Protocols SOP: O2k Quality Control 1

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Updates: http://wiki.oroboros.at/index.php/MiPNet06.03 POS-Calibration-SOP



O2k Quality Control 1: Polarographic oxygen sensors and accuracy of calibration – DatLab 8

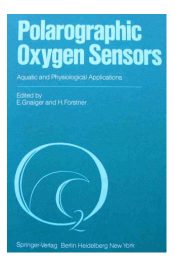
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Summary: High-resolution respirometry (HRR) critically depends on maintenance (MiPNet19.18B) and accurate calibration of the polarographic oxygen sensors (OroboPOS, POS). Calibration errors > 10 % as commonly encountered in the literature cannot be accepted in HRR. Standard operating procedures (**O2k-SOP**) are described: (1) Cleaning and preparation for use of O2k-chambers (MiPNet19.03); (2) Quality control for evaluation of proper POS function (SOP: O2 sensor test, MiPNet06.03); and (3) Accurate POS calibration (MitoPedia: O2-Calibration - DatLab, MiPNet06.03). This is Part 1 of O2k Quality Control, a component of the *Oroboros Quality Management*.

1. Oxygen concentration and partial pressure

A POS responds to partial pressure of oxygen, p_{0_2} . Expressing the oxygen signal in terms of p_{0_2} has advantages. However, respiration is expressed in molar units related to biochemical stoichiometries. Conversion of partial pressure of oxygen, p_{0_2} , to oxygen concentration, c_{0_2} , is based on accurate barometric pressure (measured electronically) and oxygen solubilities in experimental media.

The oxygen solubility of mitochondrial respiration media MiR05, and MiR06 (MiR05 with added catalase) relative to pure water (oxygen solubility factor, $F_{\rm M}$) is 0.92, accurately determined for MiR05 at 37 °C and 30 °C. At air saturation, standard barometric pressure (100 kPa), and 37 °C, $p_{\rm O_2}$ is 19.63 kPa, and $c_{\rm O_2}$ is 190.7 μ M in MiR05 or MiR06.

2. Polarographic oxygen sensor

Each O2k-chamber is equipped with an $\underline{\text{OroboPOS}}$ which is developed for optimum function of the O2k. The signal is linear in the large p_{0_2} range from pure oxygen (100 kPa) to 20 kPa (air saturation) and down to 0 kPa. Thus, the OroboPOS is superior to optical sensors.

The OroboPOS requires minimal service interventions, operates at a high sensitivity and stability for periods of > 3 months without needing to change the POS membrane.

Oxygen diffuses from the sample to the cathode surface through (1) an unstirred layer of the sample at the outer POS membrane surface, (2) the POS membrane and (3) the POS electrolyte layer. To minimize the unstirred layer of the sample, a high and constant stirring of the sample medium is required. At the cathode, the partial pressure of oxygen is effectively reduced to zero. Under steady-state conditions, the oxygen flow to the cathode depends on the external oxygen pressure, and the electrochemical reduction of oxygen yields an oxygen-dependent consumption of oxygen by the POS. This gives rise to an electric current which is converted into a voltage.

In short, the POS produces its electrical signal by consuming the oxygen that diffuses across the oxygen-permeable membrane to the cathode. The cathode and anode reactions are,

$$O_2 + 2 H_2O + 4 e^- \rightarrow 4 OH^-$$

4 Ag $\rightarrow 4 Ag^+ + 4 e^-$

At air saturation, the signal of the POS is c. 2 μ A. From the stoichiometry (above) and Faraday constant (2.591 pmol $O_2 \cdot s^{-1} \cdot \mu A^{-1}$), oxygen consumption by the POS at air saturation in a 2 cm³ chamber is theoretically 2.6 pmol·s⁻¹·cm⁻³, in direct agreement with experimental observations (MiPNet14.06).

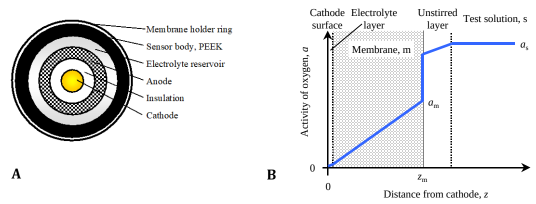
2.1. Cathode

The OroboPOS has a gold cathode, which is generally superior to platinum, since it has a broader plateau in the polarographic measuring circuit (Gnaiger et al., 1983). The sensitivity of polarographic oxygen sensors is a function of cathode area. Long-term stability increases with a high electrolyte volume and a high ratio of anode to cathode area.

The signal to noise ratio increases and the relative signal drift at zero oxygen decreases with cathode area. Therefore, the OroboPOS has a relatively large cathode area (2 mm diameter), yielding a high sensitivity owing to a stable zero current. Signal noise decreases with decreasing oxygen to less than \pm 0.002 kPa (recorded near zero oxygen over 100 data points and 0.2 s intervals) which is of particular advantage for measurements at physiological intracellular oxygen levels.

2.2. Anode

The silver-silver chloride anode has a large area compared to the cathode. The anode may become dark grey-black requiring periodical cleaning with ammonia.



The polarographic oxygen sensor (A) consists of a gold cathode and a silver-silver chloride anode, connected by KCl electrolyte covered with an oxygen-permeable membrane. Oxygen diffusion profile (B) at the polarographic oxygen sensor under steady-state conditions in a stirred test solution.

2.3. Electrolyte

KCl solution (1 mol·dm⁻³; 74.56 g potassium chloride per liter deionized water). Dissolve 1.49 g KCl in deionized water to yield a total volume of 20 mL. A high quality of deionized or distilled H_2O is critically important. Before filling the electrolyte into the receptacle of the POS, warm it to c. 40 °C, particularly after storage at 4 °C, to avoid formation of gas bubbles in the electrolyte reservoir of the POS.

An alkaline electrolyte with KOH did not improve stability of the signal, had no positive effect on the long-term behavior of the time constant and is less convenient for handling. For these reasons, we do not use a KOH electrolyte.

For a H_2S insensitive mode of operation at high sulfide concentrations, a special electrolyte is freshly prepared: Equilibrate distilled water with nitrogen gas. Dissolve 100 g K_2S -9 H_2O in 1 liter deionized water, stirring for a long time. Filter the black precipitate and store in the dark, never longer than 6 weeks. The polarizing voltage must be changed from 800 mV to 100 mV.

2.4. Membrane

At a given oxygen concentration in the test solution, the signal of a POS depends on the properties of the membrane, increasing with diffusion coefficient and oxygen solubility (the product of which is the permeability coefficient), and decreasing with membrane thickness. While a high signal is desirable in terms of a high electronic signal to noise ratio, and a low membrane thickness and high diffusion coefficient increase the time resolution, these advantages are offset by a high background oxygen



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consumption in the respirometer chamber, an increased sensitivity to the stirring of the sample, and a shortened lifetime of the anode and electrolyte. Therefore, the choice of the membrane requires optimization according to specific requirements. OroboPOSmembranes (FEP, 25 μ m thickness) are used for HRR. Application of a new membrane is simplified by the OroboPOS-Service Kit.

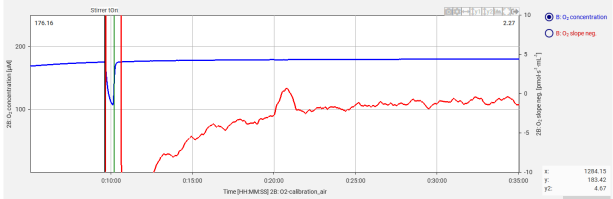
3. Calibration and quality control (02k-SOP)

3.1. Preparation

- 1. Switch on the O2k, connect it to DatLab. Edit the O2k channel settings in DatLab. Clean the O2k-chambers using the DL-protocol "O2k-cleaning BeforeUse" (MiPNet19.03).
- 2. Increase the temperature of the stock of experimental medium to slightly above experimental temperature. Add 2.1-2.5 mL medium to each 2.0-mL 02k-chamber and 0.54 mL to each 0.5-mL 02k-chamber. This helps avoid the formation of gas bubbles and minimizes disturbance of the 02k temperature.
- 3. With the stirrer on (typically 750 rpm = 12.5 Hz), insert the stopper fully; check that no air bubbles are contained in the volume-calibrated chamber.
- 4. Siphon off excess medium from the top of the stopper.
- 5. Lift the stopper to the stopper spacer position.
- 6. Start a new file in DatLab and select the DL-protocol "O2-calibration air" or "O2-calibration air and zero".

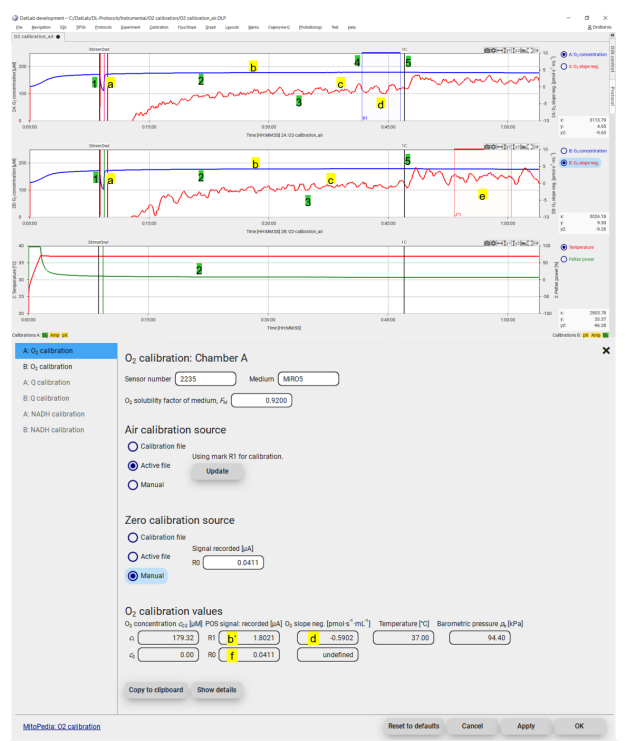
3.2. The O₂ sensor test

1. Even before final stabilization of O_2 slope neg., perform a stirrer test [F9], switching both stirrers automatically off and on. The default period is 30 s, for experiments at 37 °C. At lower experimental temperature, this period should be prolonged (60 s at 25 °C).



Stirrer test for quality control of the POS (standard 30 s) displayed with Graph Layout

"02-Calibration - Background" (MiR05; 37 °C; data recording interval: 2 s; slope smoothing: 80 s; 30 min time scale). 2021-09-09 P2-01.DLD



POS Quality control using the DatLab protocol (DL-Protocol) O2_calibration_air.DLP: Plot of the O₂ sensor test (above; File 2017-02-06 P2-01.DLD; time scale is 1:05 h:min) and oxygen calibration window (below).

Quality control **a**: Upon automatic re-start of the stirrer (On), the increase of the oxygen signal should be rapid and monoexponential.

2. About 20 min are required for approximate air equilibration after temperature equilibration of the incubation medium, visualized as stabilization of the Peltier power.

Quality control **b**: The raw signal (blue trace) should be between 1 and 3 μ A when at sea level up to 1000 m altitude (p_b 90 - 101 kPa,) and temperature is 25 - 37°C.

3. Within 40 min, the oxygen signal should be stable with O₂ slope neg. (uncorrected) close to zero.

Quality control \mathbf{c} : Signal noise should be low, reflected in the noise of the O_2 slope neg. (red trace) within \pm 2 (\pm 4 is acceptable) pmol·s⁻¹·mL⁻¹ for both 2.0-mL and 0.5-mL chambers at a data recording interval of 2 s and slope smoothing 80 s selected for calculation of the slope.

- **4**. Set a mark on the oxygen signal (R1 the mark is automatically named R1 when using the DL-protocol) and click on Calibration to open the DatLab O_2 calibration window. Three options are available for applying O_2 calibration values:
- (1) clicking on "calibration file" loads the values from a calibration file; (2) clicking on 'Active file', uses the values from mark R1 of the current file; (3) clicking on 'Manual', allows for the values to be entered manually. If "Active file" is selected, the R1 will be shown in \mathbf{b}' . Add the O₂ solubility factor of the medium, F_{M} (O₂ solubility factor).

Quality control **d**: The slope uncorrected should be \pm 1 pmol·s⁻¹·mL⁻¹ for 2.0-mL and 0.5-mL chambers if averaged across the section of the experiment marked as R1 for air calibration (**d**). The recorded POS signal should be close to the previous calibration under identical experimental conditions. Click on Apply.

5. Close the chamber and if required, perform a zero oxygen calibration (section 4), or continue with a complete instrumental O₂ background test (MiPNet14.06).

Quality control **e**: After closing the chamber, select plot Y2 and set mark J°1. The O_2 slope (neg.) should be 3 ± 1 pmol·s⁻¹·mL⁻¹ for the 2.0-mL chamber (MiPNet14.06) and 10.0 ± 4 pmol·s⁻¹·mL⁻¹ for the 0.5-mL chamber. Flux values higher than 4.0 or 14.0 pmol·s⁻¹·mL⁻¹ for the 2.0- or 0.5-mL chamber, respectively, may indicate a biological contamination.

Quality control f: The zero signal at mark R0 obtained at zero calibration (section 4) should be < 2 % of R1 (stable at < 5 % is acceptable).

4. Zero oxygen calibration

Zero calibration can be done by one of the following methods:

4.1. Zero calibration with instrumental O_2 background test: TIP2k

02k-SOP: » MiPNet14.06 Instrumental 02 Background

4.2. Zero calibration: manual titration of dithionite (02k-SOP)

- 1. Prepare "zero solution": Dissolve two spatula tips or 20 mg of dithionite (sodium hydrosulfite, Na-dithionite, Na₂S₂O₄; O2-Zero Powder in the OroboPOS-Service Kit) in 0.5 mL deionized water. Mix in a small vial with minimum gas space. Use fresh dithionite. Dithionite is oxidized during prolonged storage and needs to be replaced.
- 2. Inject 20 μ L zero solution into the closed O2k-chamber using a 50μ L microsyringe.

- 3. Oxygen depletion is very rapid, and zero oxygen is reached within a few minutes. However, a few more minutes may be required until a stable signal is obtained, R_0 [μ A].
- 4. Inject another 10 μ L zero solution. Repeat as long as the signal responds by a further decline. Siphon off excess medium from the stopper.
- 5. On the final titration when no further decline is seen, the zero signal stabilizes quickly ($< \pm 0.2$ or $< \pm 0.8$ pmol·s⁻¹·mL⁻¹ for the 2.0-mL or 0.5-mL chamber respectively).
- 6. Set a mark over the stable "zero" signal (R_0), to complete the two-point oxygen calibration [F5]. Select Mark R1 and Mark R0 for R_1 and R_0 in the O_2 calibration window.

4.3. Zero calibration: mitochondrial respiration

Due to the high oxygen affinity of isolated mitochondria, intact cells and tissue homogenate, residual traces of oxygen are insignificant after respiratory oxygen depletion. Therefore, you can use your experimental sample for such zero-oxygen calibration. Alternatively, prepare a stock of baker's yeast, with 200 mg dry yeast in 2 mL physiological salt solution. Stir heavily to obtain a homogenous suspension of yeast cells and add 50 μL yeast suspension into the 2-mL chamber through the cannula of the stopper, using a microsyringe.

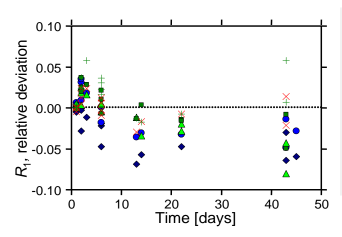
More details: »Gnaiger et al (1995), »Gnaiger (2001).

5. O₂-Calibration list: quality control



Oroboros FileFinder: Click on the icon "O2k-Manual". Go to 'O2k-Qualtiy control and SOPs' and move to the right to open the Excel file "O2-calibration.xlsx". Save a copy of this Excel template

and paste the calibration parameters into new lines sequentially for chamber (A) and (B), thus generating a data base for quality control of instrumental calibration.



Stability of the signals of six OroboPOS at air calibration, R1, over a period of > 1 month at constant temperature (25 °C). Membranes were not exchanged, and the sensors were left mounted to the O2k-chambers, which were filled with 70 % ethanol during storage, and with mitochondrial respiration medium during calibrations (from Gnaiger 2008).

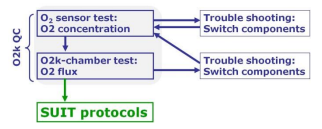
Trends over time can thus be evaluated, and possible irregularities of sensor performance are quickly recognized for intervention by sensor service.

» MiPNet19.18B POS-service

6. O₂ sensor test and zero oxygen calibration: when?

The O₂ sensor test (section 3) should be performed:

- Every day after switching on the O2k.
- During troubleshooting procedures, when switching components between the two chambers, a quick sensor test is performed after each step (stirrer test, raw sensor signal).



After application of a new membrane and POS Service. In some cases, the signal
of the OroboPOS improves (higher signal stability, less noise, shorter response
time), when leaving the O2k switched on overnight (O2k-chambers filled with
deionized H₂O at 25° C with stopper in closed position and the illumination
switched off).

Zero oxygen calibration should be performed:

- Every few weeks.
- When working at low oxygen levels it is recommended to measure R0 before and after each measurement using the experimental sample for zero calibration (Section 4.3).

 O_2 sensor test and zero calibration are also performed at the beginning and end of the O2k-chamber test (instrumental O_2 background test).

7. References

Gnaiger E (2001) Bioenergetics at low oxygen: dependence of respiration and phosphorylation on oxygen and adenosine diphosphate supply. Respir Physiol 128:277-97. »Bioblast link

Gnaiger E (2008) Polarographic oxygen sensors, the oxygraph and High-Resolution FluoRespirometry to assess mitochondrial function. In: Mitochondrial Dysfunction in Drug-Induced Toxicity (Dykens JA, Will Y, eds) John Wiley:327-52. » <u>Bioblast link</u>

Gnaiger E, Forstner H, eds (1983) Polarographic Oxygen Sensors. Aquatic and Physiological Applications. Springer, Berlin, Heidelberg, New York:370 pp. »Bioblast link

Gnaiger E, Steinlechner-Maran R, Méndez G, Eberl T, Margreiter R (1995) Control of mitochondrial and cellular respiration by oxygen. J Bioenerg Biomembr 27:583-96. »Bioblast link

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Supplement A: Calibration of time constant for signal correction

Correction for the time response by using an accurate time constant is essential for high-resolution analysis of kinetic studies, such as ADP pulse titrations and oxygen kinetics involving rapid transitions to anoxia (Gnaiger 2001).

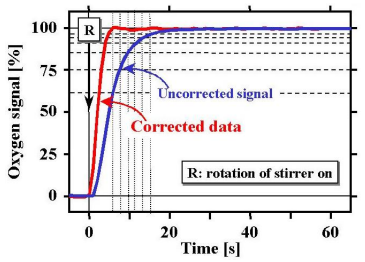


Figure 1. Sensors respond with a time delay to rapid changes of oxygen (uncorrected signal). A step change is simply achieved by switching the stirrer off at air saturation, allowing for a local depletion of oxygen at the cathode, followed by switching the stirrer on. The oxygen signal is expressed in % of the total step change. Is the oxygen sensor sufficiently fast for kinetic studies? DatLab yields the answer, gives the exponential time constant (3 s in the present example) and displays the time-corrected data (modified after Gnaiger 2001).

The signal of polarographic oxygen sensors responds with a time delay to rapid changes in the partial pressure of oxygen in the medium (Fig. 1). This convolution of the signal is due to the separation of the oxygen sensor from the experimental medium by a membrane and an electrolyte layer. Consequently, the signal at the cathode responds to a change in oxygen only after oxygen diffusion has taken place through the membrane to the cathode. The time response to changes of p_{02} depends mainly on the thickness of the sensor membrane (z_m) , the oxvgen permeability membrane, temperature, and the unstirred boundary layer of the experimental solution.

The response time of the oxygen sensor is characterized by an exponential time constant, τ . Knowledge of τ is crucial both for

quality control of the POS and for the time correction of O2k recordings in high-resolution respirometry, particularly in kinetic studies. A fast response of the sensor is indicative of a high quality of sensor maintenance. Prolonged use or storage of the sensor without anode cleaning may increase the response time of the sensor. Such a sensor may be used only if the signal is stable and a high time resolution is not required.

 τ can be experimentally determined by pulse-titration of anoxic into air-saturated medium or by a stirrer test, i.e. turning the stirrer off and on (Fig. 2). Both methods yield identical results. The response is fitted to an exponential function which yields the value of τ [s].

au depends on experimental temperature, with a Q_{10} of c. 0.69 (Fig. 2). As expected for a diffusion-controlled process, the time constant au strongly depends on the experimental temperature. A semilogarithmic plot of time constant au vs. temperature results in a straight line (Fig. 2), indicating a 31 % decrease in au for a 10 °C increase in temperature.

Stirring speed influences τ theoretically only when (1) mixing is slow of the injected (anoxic) solution with the (air-saturated) oxygraph medium (i.e., if the time constant of the mixing process is in the same range or higher than the time constant of the oxygen sensor), or when (2) unstirred layers play a significant role in oxygen diffusion limitation to the cathode. τ is virtually constant between 100 and 700 rpm in anoxic injection

experiments, indicating that complete mixing is achieved within a few seconds. A 5 % increase of τ between 700 and 100 rpm is consistent with the corresponding 5 % decrease of the oxygen signal recorded in air-saturated water. This points to more pronounced unstirred layer effects at lower stirring speeds and, at the same time, excludes a significant contribution of the mixing process to τ . Similarly, an increase in viscosity associated with the addition of 10 % dextran to the experimental medium does not significantly affect the time constant.

More details: »Gnaiger (2001)

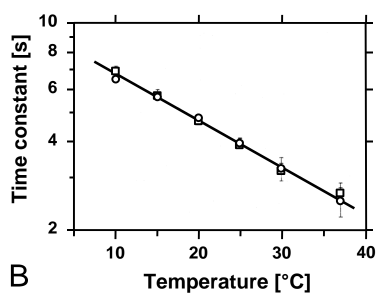


Figure 2. Effect of temperature on the time constant τ . The temperature was varied between 10 and 37 °C, and the time of both constants sensors (chamber A and B in the same Oxygraph) were determined by the titration method. Stirring speed 300 rpm; chamber volume 2 cm³; titration volume 200-250 mm³. Each value represents the mean ± SD of 5-6 measurements (from Gnaiger 2001).

Supplement B: O₂ calibration window in DatLab

In the calibration window click on "view details" to see the oxygen calibration parameters and equations calculated by DatLab.

Concentration: Parameters are displayed for conversion of the raw signal to concentration.

Calibration factor for concentration, F_c [μ M· μ A-¹]: This is the multiplication factor, F_c , calculated to convert the recorded current (corrected for the zero signal) into oxygen concentration (Supplement C - Eq. 2).

Calibration offset, a_c [μ A]: This is the POS zero signal at zero oxygen concentration, which is subtracted from the current before multiplication with the calibration factor (Supplement C - Eq. 3).

Pressure: Parameters are displayed for conversion of the POS signal current to partial pressure of oxygen. These are the fundamental parameters for evaluation of signal stability over periods of several months since the POS responds to partial pressure of oxygen in the medium rather than concentration.

 p_1 [kPa]: p_{0_2} at air saturation, $p_{0_2}^*$, a function of temperature and barometric pressure.

 p_0 [kPa]: Usually p_{O_2} at zero oxygen concentration, or any other p_{O_2} at the second calibration point, p_0 .

 I_1 [µA]: POS signal as a current, at air saturation.

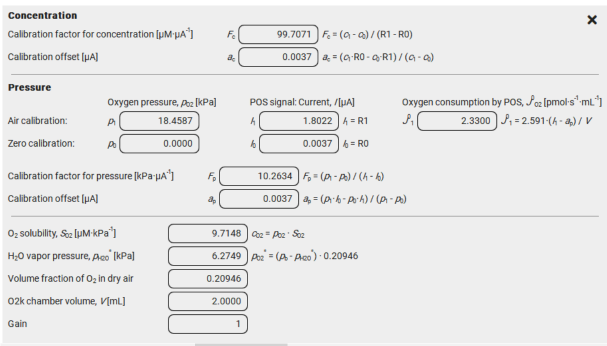


Figure 3. Upon clicking 'show details' in Calibration window oxygen calibration parameters are displayed as calculated by DatLab.

 I_0 [µA]: POS signal as a current, at zero oxygen concentration, or any other p_{0_2} at the second calibration point.

Oxygen consumption by the POS, $J^{\circ}_{0_2}$ [pmol·s⁻¹·mL⁻¹]: Theoretical oxygen consumption of the oxygen sensor at air saturation under experimental conditions (Supplement C - Eq. 9).

Calibration factor for pressure, F_p [kPa· μ A⁻¹]: This is the multiplication factor, F_p , calculated to convert the current of the POS (corrected for the zero current) into partial pressure of oxygen (Supplement C - Eq. 5).

Calibration offset, a_p [µA]: This is the POS zero current, at zero partial pressure of oxygen, which is subtracted from the current before multiplication with the calibration factor (Supplement C - Eq. 4).

O₂ **solubility**, S_{O_2} [μ M·kPa⁻¹]: a function of temperature and oxygen solubility factor of the medium (Supplement C - Eq. 7).

 H_2O vapor pressure, $p_{H_2O}^*$ [kPa]: a function of temperature, is subtracted from the barometric pressure, p_b . (Supplement C – Eq. 9).

Volume fraction of O₂ in dry air: 0.20946, when multiplied with the pressure ($p_b - p_{H_2O}^*$), it yields the partial pressure of oxygen.

O2k chamber volume, V [mL]: The effective aqueous volume of the closed O2k-chamber.

Gain, *G*: The gain is used for current to voltage conversion. Gain is fixed to 1.

Supplement C: Equations for oxygen calibration

C1. Oxygen concentration and recorded signal

The recorded oxygen signal, R_t , at experimental time t, is calibrated in terms of oxygen concentration at time t, $c_{O_2}(t)$,

$$c_{\Omega_2}(t) = (R_t - a_c) \cdot F_c \tag{1}$$

where F_c is the calibration factor based on concentration,

$$F_c = \frac{c_1 - c_0}{R_1 - R_0} \tag{2}$$

and a_c is the POS signal at zero oxygen concentration,

$$a_c = \frac{c_1 \cdot R_0 - c_0 \cdot R_1}{c_1 - c_0} \tag{3}$$

 $c_1 = c_{02}^*$ is the oxygen concentration at equilibrium with air (air saturation). Typically, R_1 and R_0 are the calibration recordings at air saturation and zero oxygen (if $c_0 = 0$ μ M, then $a_c = R_0$).

C2. Oxygen pressure and POS current

In the more general case, the oxygen sensor responds to partial pressure of oxygen, and a linear oxygen calibration can be performed at any two calibration pressures of oxygen, p_1 and p_0 . The corresponding oxygen signals in terms of current [μ A] are I_1 and I_0 . A sensor current of 1 μ A yields a raw signal of 1 V at a gain setting of 1 V/ μ A.

The zero current or offset, a [μ A], is

$$a = \frac{p_1 \cdot I_0 - p_0 \cdot I_1}{p_1 - p_0} \tag{4}$$

If the calibration point p_0 is chosen at zero oxygen concentration, then $a = I_0$. The corresponding calibration factor, related to partial pressure of oxygen and current, is F_p [kPa/ μ A],

$$F_{p} = \frac{p_{1} - p_{0}}{I_{1} - I_{0}} \tag{5}$$

After calibration, comparable to Eq. (1), the partial pressure of oxygen, $p_{0_2}(t)$, can be calculated from the POS signal current, I_t , at any time t,

$$p_{O_2}(t) = (I_t - a) \cdot F_p \tag{6}$$

C3. Oxygen concentration and partial pressure of oxygen

The partial pressure of oxygen is related to the oxygen concentration, $c_{0_2}(t)$ [μ M=nmol·mL⁻¹], by the oxygen solubility, S_{0_2} [μ M·kPA⁻¹], which is calculated by DatLab based on experimental temperature and the oxygen solubility factor of the experimental medium, FM.

$$c_{0_2}(t) = p_{0_2}(t) \cdot S_{0_2} \tag{7}$$

C4. Oxygen signal and background oxygen consumption

The oxygen-related POS current, $(I_t - a)$ [μ A] (Eq. 6), results from the steady-state oxygen diffusion from the medium across the membrane and oxygen consumption at the cathode of the POS. Based on the stoichiometry of 4 electrons per molecule O_2 reduced at the cathode and the Faraday constant (96,485 C·mol⁻¹), oxygen consumption is expected at 2.591 pmol O_2 ·s⁻¹· μ A⁻¹. The oxygen consumption by the POS, per volume of the O2k chamber, V [mL], is $J^{\circ}_{O_2}$ [pmol·s⁻¹·mL⁻¹], calculated as

$$J^{\circ}_{0_2} = 2.591 \cdot (I_t - a_p) / V \tag{8}$$

When the O2k-chamber is closed after equilibration at air saturation, the measured instrumental background oxygen consumption, $J^{\circ}_{O_2}$, can be compared with this theoretical value. At a volume of 2 mL, therefore, the expected instrumental O₂ background at air saturation is 2.6 pmol O₂·s-1·mL-1 (Eq. 8; MiPNet14.06).

Supplement D: O₂ solubility and concentration at air saturation

D1. Partial pressure of oxygen and concentration

It is practical to calculate the saturation concentration for pure water, which then is corrected by the solubility factor of the medium, F_M , to account for the reduced O_2 solubility in salt media. Due to the salting-out effect, F_M must be <1.0 in salt media used for respiratory studies of mitochondria, cells, and tissues.

 $F_{\rm M}$ is typically near 0.9 for O2k media (0.92 for MiR06 and MiR05). Several oxygen solubilities reported in the literature must be criticized on the basis of physicochemical considerations.

Water in equilibrium with air contains an oxygen concentration proportional to the oxygen solubility and the partial oxygen pressure of air. In the gas-liquid boundary, air is saturated with water vapor at the partial pressure of $p_{\rm H_2O}^*$. The water vapor pressure is subtracted from the total barometric pressure, $p_{\rm b}$, to obtain the partial pressure of dry air, $p_{\rm b}$ - $p_{\rm H_2O}^*$. The volume fraction of dry air is constant at $\Phi_{\rm O_2}$ = 0.20946. Therefore, the partial pressure of oxygen at air saturation is, for any temperature and barometric pressure,

$$p_{02}^* = (p_b - p_{H20}^*) \cdot 0.20946$$
 (9)

The saturation O_2 concentration depends on the O_2 solubility, S_{O_2} [μ M·kPa⁻¹],

$$c_{0_2}^* = p_{0_2}^* \cdot S_{0_2} \tag{10}$$

Oxygen solubility is a function of temperature and oxygen solubility factor of the medium. In other words, oxygen solubility, S_{0_2} , is defined as the ratio of partial pressure of oxygen and concentration,

$$S_{0_2} = c_{0_2}^*/p_{0_2}^* \tag{11}$$

D2. Temperature effect on saturation O2 concentration

 $p_{\mathrm{H_2O}}^*$ (Eq. 13) is the saturation water vapor pressure at experimental temperature. $p_{\mathrm{H_2O}}^*$ is a function of absolute temperature, T [K], obtained from the experimental temperature, θ , recorded in units °C,

$$T = \theta + 273.15^* \tag{12}$$

The saturation water vapor pressure [kPa] is (Table 1),

$$p_{\text{H}_2\text{O}}^* = \exp[(-216961 \cdot T^{-1} - 3840.7) \cdot T^{-1} + 16.4754]$$
 (13)

Until recently, the atm-standard pressure has been used: 1 atm = 760 mmHg = 101.325 kPa. For pure water in equilibrium with air at this atm-standard pressure, the 'unit standard concentration' of oxygen, C^* , is calculated by the polynomial expression,

$$C^* = \exp \{ [(-8.621949 \cdot 10^{11} \cdot T^{-1} + 1.243800 \cdot 10^{10}) \cdot T^{-1} - 6.642308 \cdot 10^7] \cdot T^{-1} + 1.575701 \cdot 10^5 \} \cdot T^{-1} - 135.90202 \}$$
(14)

Table 1. Saturation water vapor pressure, $p_{\rm H_2O}^*$, partial pressure of oxygen, $p_{\rm O_2}^*$, and oxygen concentration, $c_{\rm O_2}^*$, at air saturation and standard barometric pressure, $p_{\rm b}^\circ = 100$ kPa, in pure water as a function of temperature. $S_{\rm O_2}$ is the oxygen solubility, independent of choice of standard pressure. f° is the multiplication factor to convert partial O₂ pressures and concentrations given at atm-standard pressure (1 atm = 101.325 kPa) to the IUPAC standard pressure of 100 kPa (compare Eq. 15), $f^\circ = (100-p_{\rm H_2O}^*)/(101.325-p_{\rm H_2O}^*)$

θ	T	$p_{\mathrm{H}_2\mathrm{O}^*}$	$p_{0_2^*}$	$c_{0_2^*}$	f°	S_{O_2}
°C	K	kPa	kPa	μmol∙dm ⁻³		µmol∙dm-³∙kPa-1
40	313.15	7.38	19.40	197.6	0.9859	10.18
37	310.15	6.27	19.63	207.3	0.9861	10.56
35	308.15	5.62	19.77	214.2	0.9862	10.83
30	303.15	4.24	20.06	233.0	0.9864	11.62
25	298.15	3.17	20.28	254.8	0.9865	12.56
20	293.15	2.34	20.46	280.4	0.9866	13.70
15	288.15	1.70	20.59	310.9	0.9867	15.10
10	283.15	1.23	20.69	348.1	0.9868	16.83
5	278.15	0.87	20.76	393.9	0.9868	18.97
4	277.15	0.81	20.78	404.3	0.9868	19.46

D3. Barometric pressure and saturation O2 concentration

The unit standard concentration and the oxygen concentration at air saturation (Table 1) and actual barometric pressure are related by (compare f° in Table 1),

$$c_{0_{2}}^{*} = C^{*} \cdot p_{0_{2}}^{*} / [(101.325 - p_{H_{2}0}^{*}) \cdot 0.20946] \cdot F_{M}$$

$$= C^{*} \cdot (p_{b} - p_{H_{2}0}^{*}) / (101.325 - p_{H_{2}0}^{*}) \cdot F_{M}$$
(15)

D4. The barometric altitude relation (BAR)

The partial pressure of oxygen declines with altitude. Hypoxia causes a limitation of maximal aerobic capacity. The VO_{2max} of acclimatized persons declines at high altitude by c. 11 % per 1,000 m, whereas the partial pressure of oxygen declines by 12 % to 14 % per 1,000 m up to 6,000 m, and by 15 % to 17 % per 1,000 m between 6,000 and 9,000 m. The quadratic model atmosphere equation, MAE, was introduced by John B. West to describe the dependence of average barometric pressure and altitude with high accuracy. An exponential function is the basis of the ICAO Standard Atmosphere, which can be fitted to realistic reference data comparable to the MAE. This leads to the barometric altitude relation, BAR, which expresses the relationship between barometric pressure, p_b , and altitude, h [m], with an even superior fit (Tab. 2):

$$p_{\rm b} = p_{\rm b}^{\,\circ} \cdot \left(1 - \frac{0.00616 \cdot h}{288.15}\right)^{5.256} \tag{16}$$

The standard pressure at average sea level, p_b° , is inserted with 101.325 kPa or 760 mmHg to calculate barometric pressure in the respective unit. Compared to the ICAO, only the temperature gradient of -6.5 °C/km (ICAO) was replaced by the parameter -0.00616 °C/m (BAR) which was obtained by a mathematical fit to the reference data in the range of 0 to 9,000 m. 288.15 K is the air temperature of 15 °C at sea level. Deviations between

MAE und BAR are less than \pm 0.06 kPa (0.4 mmHg) in the range of 0 to 9 km altitude. In this context the relevance of mitochondrial oxygen kinetics is discussed briefly. The p_{50} of mitochondrial respiration is 0.01 to 0.1 kPa (0.08 to 0.8 mmHg; this is the partial pressure of oxygen at which mitochondrial respiration drops to 50 % of maximum values). These generally very low p_{50} values are important for our understanding of some apparently paradoxical mechanisms of muscular acclimatization and adaptation to hypoxia at extreme altitude (Gnaiger 2013).

Table 2. Barometric pressure, p_b , and partial pressure of oxygen, p_{0_2} , in dry air and respiratory air saturated by water vapor as a function of altitude, h. The decline of respiratory air p_{0_2} is expressed relative to sea level or per 1,000 m change of altitude (from Gnaiger 2013). \boldsymbol{a} : Innsbruck, A (95.0 kPa; Jul-Aug 2013); \boldsymbol{b} : Schröcken, Körbersee, AT (83.6 kPa; Oct 2013); \boldsymbol{c} : Monte Rosa, IT (58.4 kPa; Aug-Sep 2004); \boldsymbol{d} : Mt Chacaltaya (54.2 kPa; Aug 2012); \boldsymbol{e} : Everest Base Camp (52.7 kPa; Mar 2013); \boldsymbol{f} : Mt Everest (12, 13). Numbers in parentheses are measurements of p_b during respirometric studies with the Oroboros O2k.

h	$p_{\rm b}$	$p_{\rm b}$	Dry air $p_{O_2,da}$	Respiratory air p_{0_2}		Change rel.	Rel. change
[m]	[kPa]	[mmHg]	[kPa]	[kPa]	[mmHg]	to sea level	$p_{0_2}/1.000 \text{ m}$
0	101.3	760	21.2	19.9	149		
1,000	90.4	678	18.9	17.6	132	-0.11	-0.12
2,000	80.5	604	16.9	15.6	117	-0.22	-0.13
3,000	71.5	536	15.0	13.7	103	-0.31	-0.13
4,000	63.3	475	13.3	12.0	90	-0.40	-0.13
5,000	55.9	420	11.7	10.4	78	-0.48	-0.14
6,000	49.2	369	10.3	9.0	68	-0.55	-0.14
7,000	43.2	324	9.1	7.7	58	-0.61	-0.15
8,000	37.8	284	7.9	6.6	50	-0.67	-0.16
9,000	33.0	247	6.9	5.6	42	-0.72	-0.17
575a	94.9	712	19.9	18.6	139	-0.07	
1,675b	83.7	627	17.5	16.2	122	-0.19	
4,559c	59.1	443	12.4	11.1	83	-0.44	
5,240 ^d	54.3	407	11.4	10.1	75	-0.50	
5,364e	53.4	401	11.2	9.9	74	-0.50	
8,848 ^f	33.7	252	7.1	5.7	43	-0.71	

D5. O2 solubility factor in salt solutions

The salting out effect is responsible for the reduced oxygen solubility in aqueous solutions compared to pure water (Fig. 4). Detailed equations are available for calculating the oxygen solubility of sea water at different salinities (Forstner and Gnaiger 1983). Physiological solutions commonly used in Oxygraph studies (Rasmussen, Rasmussen 2003; Reynafarje, Costa, Lehninger 1985) are compared with pure water and 20 ‰ sea water in Fig. 4. The corresponding polynomial equations are summarized in Tab. 3 for calculating the oxygen saturation concentration in equilibrium with air at various temperatures and standard pressure (Tab. 4).

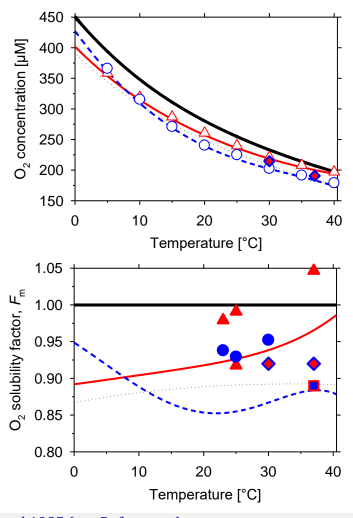


Figure 4. Oxygen concentration at saturation and standard barometric pressure (100 kPa; top) and oxygen solubility factor (bottom) in MiR05 (diamonds), KCl medium (open triangles, full line; 150 mmol·dm⁻³ KCl) and sucrose medium (open circles, dashed line: 250 mmol·dm⁻³ sucrose; data for both media from Reynafarje et al 1985), compared to pure water (upper full line) and 20 ‰ sea water (lower dotted line). For the parameters of the polynomials see Table 2. The solubility factor for serum is shown by the full square (bottom). Literature data (bottom) on KCl (closed triangles) and sucrose media (closed circles) show (i) the wide scatter of solubility data, (ii) the erroneous use of values even higher than solubility established for pure water, and (iii) a trend to higher values, particularly in sucrose medium, compared to Reynafarje

et al 1985 (see References).

Characteristic temperatures are commonly used in experimental studies. Under these conditions it is convenient to use oxygen solubility factors for the medium, $F_{\rm M}$ (Fig. 4). This factor is independent of barometric pressure, but $F_{\rm M}$ changes with temperature (compare Fig. 4). The solubility factors are compiled in Tab. 5 for different salinities of sea water (Forstner and Gnaiger 1983) and two typical media used with isolated mitochondria (Reynafarje, Costa, Lehninger 1985). The latter values have been criticized on methodological grounds by Rasmussen and Rasmussen (2003), and the complex temperature dependence of $F_{\rm M}$ compared to sea water is doubtful from a thermodynamic perspective Fig. 4). The oxygen solubility factor of MiR05 (MiR06) is 0.92, at 30 °C and 37 °C (Rasmussen, Rasmussen 2003), corresponding to an oxygen concentration in equilibrium with air under standard conditions ($c_{0_2}^*$) of 214.4 and 190.7 μ M, respectively. The oxygen solubility of serum is 9.4 nmol O_2 ·cm⁻³·kPa⁻¹ at 37 °C (Baumgärtl and Lübbers 1983). In comparison to the oxygen solubility in pure water (10.56 nmol O_2 ·cm⁻³·kPa⁻¹ at 37 °C; Tab. 1), this corresponds to a solubility factor for serum of $F_{\rm M}$ = 0.89 (Fig. 4) and $c_{0_2}^*$ of 184.5 μ M.

Table 3. Parameters of the polynomial fits of oxygen saturation concentration in equilibrium with air at p_0 ° = 100 kPa, for sea water (0 ‰ and 20 ‰) and typical Oxygraph incubation media, in the range of θ from 5 to 40 °C. Instead of the theoretically based plot of $\ln(S_{0_2})$ versus T^{-1} , the fits were performed on the untransformed data, with temperature, θ , in units of °C ($r^2 \ge 0.999$ in all cases). The equation in nested form is,

 $c_{0_2^*} = \{[(b_4 \cdot \theta + b_3) \cdot \theta + b_2] \cdot \theta + b_1\} \cdot \theta + a$

Medium	A	B_1	b_2	b_3	b_4
0 ‰	450.5946	-12.60381	0.2712233	-0.003808	$2.379 \cdot 10^{-5}$
20 ‰	390.8769	-10.2165	0.2051415	-0.002746	$1.621 \cdot 10^{-5}$
KCl	401.9152	-10.70002	0.2291496	-0.003283	$2.492 \cdot 10^{-5}$
Sucrose	427.411	-14.4983	0.2762108	-0.0003628	-3.606·10 ⁻⁵

Table 4. Oxygen solubility, S_{02} [µM.kPa⁻¹], for seawater at various salinities (10 ‰, 20 ‰, 30 ‰ and 36 ‰), and for two typical Oxygraph media (concentrations given in mmol·dm⁻³); "Sucrose": 250 sucrose, 5 KCl, 3 K-Hepes, pH 7.05; "KCl": 150 KCl, 3 K-Hepes, pH 7.05.

θ		S _{O2} for se	S_{O_2} for exp. Medium			
°C	10 ‰	20 ‰	30 ‰	36 ‰	Sucrose	KCl
40	9.62	9.08	8.58	8.29	8.96	10.01
37	9.98	9.43	8.90	8.61	9.33	10.19
35	10.24	9.67	9.14	8.83	9.54	10.36
30	10.98	10.37	9.80	9.47	10.07	10.90
25	11.86	11.20	10.57	10.21	10.74	11.64
20	12.92	12.19	11.49	11.09	11.70	12.58
15	14.21	13.38	12.59	12.14	13.07	13.75
10	15.79	14.82	13.91	13.39	14.95	15.22
5	17.75	16.60	15.53	14.92	17.42	17.04
4	18.19	17.00	15.89	15.26	17.99	17.45

Table 5. Oxygen solubility factor of the medium, F_M , for seawater at various salinities (10 ‰, 20 ‰, 30 ‰ and 36 ‰), and for two typical O2k media (concentrations given in mmol·dm⁻³); "Sucrose": 250 sucrose, 5 KCl, 3 K-Hepes, pH 7.05; "KCl": 150 KCl, 3 K-Hepes, pH 7.05.

θ		Fм for se	F _M for exp. Medium			
°C	10 ‰	20 ‰	30 ‰	36 ‰	Sucrose	KCl
40	0.945	0.892	0.842	0.814	0.880	0.983
37	0.945	0.893	0.843	0.815	0.884	0.966
35	0.945	0.893	0.844	0.815	0.881	0.956
30	0.945	0.893	0.843	0.815	0.867	0.938
25	0.944	0.892	0.842	0.813	0.855	0.926
20	0.943	0.889	0.838	0.809	0.853	0.918
15	0.941	0.886	0.833	0.804	0.865	0.911
10	0.939	0.881	0.827	0.796	0.889	0.904
5	0.936	0.875	0.819	0.786	0.918	0.898
4	0.935	0.881	0.817	0.784	0.925	0.897

References

Baumgärtl H, Lübbers DW (1983) Microaxial needle sensor for polarographic measurement of local O₂ pressure in the cellular range of living tissue. Its construction and properties. In: Polarographic Oxygen Sensors, (Gnaiger E, Forstner H, eds) Springer, Berlin, Heidelberg, New York:37-65. »Bioblast link

Beavis AD, Lehninger AL (1986) The upper and lower limits of the mechanistic stoichiometry of mitochondrial oxidative phosphorylation. Stoichiometry of oxidative phosphorylation. Eur J Biochem 158:315-22.

Brand MD, Harper ME, Taylor HC (1993) Control of the effective P/O ratio of oxidative phosphorylation in liver mitochondria and hepatocytes. Biochem J 291: 739-48.

Chapell JB (1964) The oxidation of citrate, isocitrate and cis-aconitate by isolated mitochondria. Biochem J 90:225-37.

Forstner H, Gnaiger E (1983) Calculation of equilibrium oxygen concentration. In: Polarographic Oxygen Sensors. Aquatic and Physiological Applications. (Gnaiger E, Forstner H, eds) Springer, Berlin, Heidelberg, New York:321-33. »Bioblast link

Gnaiger E (2001) Bioenergetics at low oxygen: dependence of respiration and phosphorylation on oxygen and adenosine diphosphate supply. Respir Physiol 128:277-97. »Bioblast link

Gnaiger E (2013) Die barometrische Höhenrelation (BAR): Form und Funktion. Jahrbuch 2013 Österr Ges Alpin- Höhenmed:51-63.

Gnaiger E, Steinlechner-Maran R, Méndez G, Eberl T, Margreiter R (1995) Control of mitochondrial and cellular respiration by oxygen. J Bioenerg Biomembr 27:583-96. »Bioblast link

Hinkle PC, Yu ML The phosphorus/oxygen ratio of mitochondrial oxidative phosphorylation (1979) J Biol Chem 254:2450-5.

Lemasters JJ The ATP-to-Oxygen Stoichiometries of Oxidative Phosphorylation by Rat Liver Mitochondria (1984) J Biol Chem 259:13123-30. »Bioblast link

Rasmussen HN, Rasmussen UF (2003) Oxygen solubilities of media used in electrochemical respiration measurements. Analyt Biochem 319:105-13.

Rettich TR, Battiono R, Wilhelm E (2000) Solubility of gases in liquids. 22. High-precision determination of Henry's law constants of oxygen in liquid water from T=274 K to T=328 K. J Chem Thermodyn 32:1145-56.

Reynafarje B, Costa LE, Lehninger AL (1985) O_2 solubility in aqueous media determined by a kinetic method. Analyt Biochem 145: 406-18

Stoner CD (1987) Determination of the P/2e- stoichiometries at the individual coupling sites in mitochondrial oxidative phosphorylation. Evidence for maximum values of 1.0, 0.5, and 1.0 at sites 1, 2, and 3. J Biol Chem 262:10445-53.

Wilhelm E, Battino R, Wilcock RJ (1977) Low-pressure solubility of gases in liquid water. Chem Rev 77:219-62.