Lab-on-a-Chip System to Monitor the Oxygen Consumption of Mammalian Cells

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Abstract:
We present a Lab-on-a-Chip system to monitor the uptake of oxygen by mammalian cells during their metabolism. The oxygen consumption indicates the state of the cells and their response to the physical or chemical stimuli and is thus of interest for biological analyses. Although oxygen measurements are topic of investigations for a long time already, such a measurement system is still complex with a high number of external components which hamper their usage for biological researchers. Our system consists of a microfluidic chip with an oxygen sensitive phosphorescent film out of dye PtTFPP in a matrix of polystyrene. All external components of the optical system and the temperature control are low-cost components that are integrated in the assembly. Therefore, the entire system has a small footprint, an easy usage and a high degree of automation. With this system, the oxygen uptake of HaCaT cells is reproducibly measured as 10, 20 and 22 amol/(cell ∙ s) for temperatures of 34°C, 37°C and 40°C.

Key words: Lab-on-a-Chip, oxygen sensing, oxygen consumption rate, PtTFPP, phosphorescence

Introduction
One of the most important characteristics of living mammalian cells is the metabolism where glucose or pyruvate is burned under the uptake of oxygen. The metabolism is influenced by physical or chemical stimuli so that the state of the cells can be monitored by measuring continuously the oxygen concentration in the culture medium.

The oxygen consumption rate (OCR) is in the range of 5 – 50 amol/(cell ∙ s) [1] which can only be measured reliably with high cell densities in small, closed chambers that are common in microfluidics. An attractive sensing principle are phosphorescent films where the intensity of the emitted light correlates with the oxygen concentration [2]. The film can be integrated into the chamber [3-5]. Apart from the sensing chip, such a system requires external components for the excitation light source, an optical read-out system, a control unit and a temperature control because the metabolism of cells as well as the phosphorescent film are very temperature sensitive. These components are expensive and bulky such as a microscope or SLR camera or incubators [3,4] which hamper the application in biological labs.

In this contribution, we present a Lab-on-a-Chip (LoC) system that is based on a microfluidic chip with oxygen sensitive luminescent elements and with an integrated heater and temperature sensor. The read-out and the control is realized with a Raspberry Pi while low-cost LEDs are used to excite the film which results in a compact assembly with a small footprint and without any bulky or expensive external equipment. With this system, the OCR of human keratinocyte cells is determined for different cell concentrations and temperatures.

Design of the System
The LoC system to measure the concentration of the dissolved oxygen consists of a microfluidic chip out of glass and silicon with an oxygen sensitive film at 5 sensing spots out of PtTFPP which is embedded in polystyrene as described earlier [3]. Additionally, two electrical conductors are patterned on the backside to heat the chip and to measure the temperature based on the resistance change (see fig. 1.) [5].

The chip is clamped into a 3D-printed holder for the microfluidic and electrical connection as shown in fig. 2 [6]. A Raspberry Pi camera is mounted above the chip to measure the phosphorescent light of the film that is excited...
with a LED with the wavelength of 395 nm. Additionally, the system consists of a Raspberry Pi as control unit and a small electrical circuit to control the LED and the temperature.

As shown in fig. 4, the slope of the oxygen concentration correlates with the number of cells. Furthermore, higher temperatures result in higher OCR. In particular, the rates of 10, 20 and 22 amol/(cell-s) were determined for temperatures of 34°C, 37°C and 40°C.

Experiments and Results

The microfluidic chip is calibrated according to the Stern-Volmer-equation with an error of only 0.6 % (air) in the range between 0 % (air) and 20 % (air). HaCaT-cells with DMEM-medium (cell density $12.9 \pm 1.2 \times 10^8$ cell/L) are filled into the microfluidic chip, which is heated to $37\pm0.5^\circ$C. The oxygen concentration decreases immediately and within $3.04 \pm 0.06$ h the entire oxygen is consumed (see fig. 3). In comparison, the oxygen concentration in pure medium decreased by less than 1 % (air) in this time.

Conclusions

The presented system is based on a microfluidic chip and small and low-cost optical and electronical components to measure the oxygen consumption of mammalian cells. Because of its simplicity, compactness and ease of use, this lab-on-a-chip system is a major progress to make oxygen sensing systems available for biological researchers.

References


