Towards the detection of static ATP levels above primary PTPRζ-osteoblastic cells and their knock-out mutants by ATP microbiosensors

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Abstract:
Adenosine-5’-triphosphate (ATP) holds a significant role as omnipresent energy source and as autocrine and paracrine signaling molecule in many cells such as lung cells and bone cells [1]. ATP is considered to be involved in the mechanical stress response of bone cells such as bone-resorbing osteoblast cells or receptor-proteine-tyrosine-phosphatase-zeta (PTPRζ) - osteoblastic cells [2], which are involved in bone formation, bone regeneration and the control of bone volume. ATP release stimulates the proliferation of these P2 - receptor cell types [3,4]. The “deficient” knock-out mutant behaves different in proliferation and differentiation and thus the ATP release above these cells is expected to be altered. A localized detection of ATP at the cellular level is therefore of significant importance. Using amperometric ATP microbiosensors with diameters ranging from 10 - 50 µm enables localized ATP measurements above wild type and knock-out cells.

Key words: ATP, localized determination, biosensor, osteoblastic cells, static conditions.

ATP Microbiosensors
In the present study microelectrodes with diameters of 10 and 50 µm are used as transducer for ATP biosensors. This biosensors base on a competitive assay with glucose converting oxidoreductases (e.g. GOD or PQQ-GDH) and hexokinase (HEX) immobilized at the surface of the microelectrode (5,6). The calibration and characterization of ATP biosensors (a micrograph of such an ATP microbiosensors is shown in figure 1) was performed prior to localized ATP measurements. Calibration was obtained in buffer solution by adding first an aliquot of 300 µl 0.2 M glucose solution and successive aliquots of 1mmol ATP solution; a representative calibration curve is shown in figure 2.

Fig. 1. Micrograph of an immobilized layer at an UME of 10µm diameter
Experimental section

The experimental set-up for the detection of ATP release under static conditions is shown in Fig. 3. Experiments were conducted in a three-electrode setup in combination with a scanning electrochemical microscope for positioning the biosensor close to the cell surface. A dual microelectrode assembly served as working electrodes (WEs), using one electrode for positioning and the second as transducer for the microbiosensor. For positioning of the ATP biosensor, double-electrode assembly was used, where one electrode serves as ATP biosensor and the other remains unmodified for recording current-distance curves using a scanning electrochemical microscopy (SECM) set-up. The current obtained from oxygen reduction at –600 mV vs. Ag/AgCl is recorded in buffered solution (pH = 7.4) while the electrode assembly is approached to the surface (green arrow in figure 4). The approach was stopped at a distance of approx. 10 µm.

References


Then the ATP biosensor is biased at 600 mV and moved towards the cell monolayers.

Fig. 2. Exemplary calibration curve of an ATP biosensor (transducer: 10 µm UME); different aliquots of ATP stock solution (1 mmol) were added to a 0.2 M glucose solution while amperometric measurements at 0.6 V vs. Ag/AgCl reference were performed.

Fig. 3. Set up used for measurements of ATP release from osteoblastic cells under static conditions. A is the stage, adjustable in three axes, B the platinum counter electrode, C is the working electrode, D is the ibidi-dish with the cultivated cells and E is the video microscope.

The cultivation of both cell types were carried out using a double-chamber insert as shown in figure 4. After cell seeding and cultivation, the insert was removed and the whole dish is filled with buffer (ringers solution) for localized ATP detection.

PTPRζ – cells / Knock-out mutants

empty chamber

Fig. 3. Ibidi cultivation dish with insert, one cell type (either PTPζ or K.O-mutants) was seeded in the insert, the other chamber was left empty.