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DEVELOPMENT OF pH-BASED ELECFET BIOSENSORS FOR LACTATE ION DETECTION

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Abstract

In this work, we report on ElecFET (Electrochemical Field Effect Transistor) devices potentially of interest for the detection of different molecules in solution. ElecFET are electrochemical microsensors in liquid phase, based on two elements: (i) a pH-sensitive chemical field effect transistor (pH-ChemFET) and (ii) a metallic microelectrode deposited around the sensitive gate. The coexistence of these two elements combines (i) potentiometric and (ii) amperometric detection effects at the microscale. Design, fabrication and experimental validation of ElecFETs based on silicon and polymer micro-technologies, are reported. We first demonstrate the detection of hydrogen peroxide (H₂O₂) in solution, showing a sensitivity of 5 mV/mM in the [10-100mM] concentration range. The ElecFET concept is then extended to the detection of glucose and lactate in the [1 – 30 mM] and [1 – 6 mM] concentration range respectively. The sensitivities are between 2-6 mV/mM and 8-20 mV/mM respectively.

Keywords: electrochemical sensor, ChemFET, microelectrode, hydrogen peroxide, glucose, lactate, potentiometric detection
1. Introduction

The monitoring of L-lactate biosensors is gaining interest for clinical use [1-3] and food quality [4] assessment. Lactate ion analysis in saliva can be used as a diagnostic tool for cystic fibrosis. It can also be used for monitoring lactate concentrations in athletes (i.e. 400m runners) during and after a training session, in order to estimate their capability to quickly metabolize lactic acid, and thus evaluate effectiveness of the training program. Furthermore, the food quality monitoring requires suitable analytical methods for process and quality control, methods that will be rapid, reliable, specific and cost-effective.

Lactate oxidase (LOD) is widely used for lactate detection because of its simple reaction; LOD catalyzes the lactate oxidation to pyruvate and hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}). In most of the publications, lactate detection is based on the amperometric technique, which employs microelectrodes for the monitoring oxido-reduction reactions at a conductive solid/liquid interface [5-8]. This technique faces a severe technological issue. As the microelectrodes size becomes significantly lower than the diffusion length, spherical (rather than linear) diffusion phenomena occur at the microelectrode surface. As an effect, the related amperometric response is no longer time-dependent. Another technique used to measure the electrochemical potential at an insulative solid/liquid interface is potentiometry. The potentiometry technique is based on chemical field effect transistors (ChemFET) [9-11]. ChemFETs attracted attention in the development of biosensors, and were extended to enzyme field-effect transistors (EnFET). EnFETs are based on pH-dependent biocatalytic reactions leading to charge modification at the gate surface, according to the substrate concentration. Several EnFETs based on a variety of enzymes [9] such as glucose-oxidase (GOD) [12-14], horseradish peroxidase [15] urease [16], lactate dehydrogenase [17, 18], have been reported so far. For example, Xu et al. have reported a ChemFET-based lactate biosensor based on the LOD functionalization on the sensitive gate [19]. The authors immobilized lactate oxidase and MnO\textsubscript{2} nanoparticles in poly(dimethyldiallylammonium chloride) (PDDA) films via
layer-by-layer self-assembly to construct (PDDA/MnO\textsubscript{2}/PDDA/LOD)\textsubscript{n} multilayer films. MnO\textsubscript{2} powder acts as a catalyst to stimulate the disproportionation of H\textsubscript{2}O\textsubscript{2} to O\textsubscript{2} and H\textsubscript{2}O. MnO\textsubscript{2} nanoparticles react with hydrogen peroxide to form Mn\textsuperscript{2+} and O\textsubscript{2}, with two hydrogen ions H\textsubscript{3}O\textsuperscript{+} consumption.

The combination of amperometric and potentiometric techniques is a promising method for lactate detection. The concept involves the functional integration of a microelectrode with a ChemFET on a single chip [20-22]. Such a device is indicated as an electrochemical field effect transistor (ElecFET). By combining these two transduction principles at the microscale, the ElecFET technology provides new electrochemical detection potentialities. First, it enables the adaptation of the pH-ChemFET-based potentiometric principles to the detection of electro-active chemical species and, as a consequence, their extension to new enzymatic family such as oxidases (production of hydrogen peroxide H\textsubscript{2}O\textsubscript{2}). Second, it allows electrochemical measurement in liquid phase without any current monitoring at very low level (nano- or pico-amperes), whose control is still a challenge in term of drift and signal to noise ratio. Finally, it is compatible with the silicon-based technology and can be still improved thanks to further technological integration.

In this paper, we report on design, fabrication and study of ElecFET microdevices, which combine potentiometric and amperometric detection/transduction principles at the microscale, thanks to the common integration of a metallic microelectrode with a pH-sensitive ChemFET. First we apply this concept to the detection of H\textsubscript{2}O\textsubscript{2} by studying the influences of the main polarization parameters, i.e. voltage (V\textsubscript{P}) and duration (t\textsubscript{P}) onto the platinum microelectrode. Then, we focus on the detection of glucose by functionalizing the gate surface of the transistor with an enzymatic layer containing glucose oxidase, and finally extended to the lactate detection, with a lactate oxidase enzyme immobilization.

2. Experimental
N-channel, SiO$_2$/Si$_3$N$_4$-gate, pH-sensitive chemical field effect transistors (pH-ChemFET) were fabricated on (100)-oriented N-type (500 Ω.cm) silicon wafers using a standard P-well technology. Such technology was studied in previous work, evidencing threshold voltage $V_t$ around 0.7 V and low leakage current ($I_{off} < 1$ μA) [23]. Then, a tantalum/platinum (thickness: 50nm / 200nm) layer was deposited by physical vapor deposition and patterned by lift-off process in order to integrate the microelectrode all around the pH-ChemFET sensitive gate (mean distance: 100 μm). Finally, a SU-8 passivation was performed at the wafer level, leaving the pH-ChemFET sensitive surface and the Ta/Pt microelectrode active area uncovered. All the technological processes (figure 1a) were developed in cleanroom (class 10000 and 100). The ElecFET microdevices were manufactured on 3.5 x 3.5 mm$^2$ chips. These chips were stuck on specifically coated printed circuit using an epoxy insulating glue. After wire bonding, packaging was finally performed at the system level using a silicone glop-top in order to adapt the final sensor to the detection in liquid phase (figure 1b).

ElecFET detection/transduction principles were first studied for the hydrogen peroxide H$_2$O$_2$ detection in phosphate buffer (PBS, pH = 7.2) solutions (~ 20 mL) containing different stabilized concentrations [H$_2$O$_2$] ranging from 10 to 100 mM. Then, enzymatic functionalization process and the ElecFET enzymatic detection properties were tested for the glucose detection using glucose oxidase because of their properties in terms of activity, stability and cost. At last, when the whole protocol was optimized, the lactate detection was investigated using lactate oxidase. The enzymatic membranes (GOD: G6125-10kU and LOD: L0638-50U purchased from SIGMA) were prepared by the cross-linking of enzymes with bovine serum albumin (BSA) in glutaraldehyde solution (5%). A mixture of 50 μL of enzyme solution (0.5 mg of protein) with 0.5 mg of BSA was prepared in 100 mM phosphate buffer pH 7.2 with 5 μL of glycerol added into the mixture. Glycerol was used as a plasticizer to avoid the formation of cracks in the enzyme membrane during storage and to result in a better homogeneity of the membrane and its adhesion on the surface of the microsensor. 0.3 μL of
the enzymatic solution were deposited on the sensitive area using a micropipette then, 0.6 µL of glutaraldehyde (5%) were dropped onto the enzyme layer and left to dry for 60 minutes in air. The microsensor was then stored at 4°C overnight.

All experiments were performed in ambient atmosphere at room temperature according to a specific electrical scheme (figure 1a). The different electrical operations in solution were performed with a commercial reference calomel electrode in order to bias the electrolyte to the mass \( V_{\text{GRE}} = V_{\text{electrolyte}} = 0 \). A suitable electrical polarization (potential \( V_p \), time \( t_p \)) was applied to the Ta/Pt microelectrode using a Keithley 2400 voltage source in order to trigger specific oxido-reduction phenomena. The pH-ChemFET characterization was performed in saturation mode while using constant drain-source voltage \( V_{DS} \) and drain-source current \( I_{DS} \) (typically \( V_{DS} = 1 \) V and \( I_{DS} = 0.1 \) mA). A ChemFET-meter measurement interface was used to measure the pH-ChemFET gate-source voltage \( V_{GS} \), also called output voltage \( V_{out} \), enabling the local pH monitoring in the microelectrode surroundings depending on the polarization conditions.

3. Results and discussion

3.1. Hydrogen peroxide detection

The ElecFET concept was initially used for the detection of hydrogen peroxide \( H_2O_2 \) oxidation in water-based solutions:

\[
- H_2O_2 \text{ oxidation} (V_p > E_{1+}); \quad H_2O_2 + 2H_2O \rightarrow O_2 + 2H_3O^+ + 2e^- \quad (1)
\]

where \( E_{1+} \) is the equilibrium potential of the \( O_2/H_2O_2 \) redox couple.

In the presence of hydrogen peroxide \( H_2O_2 \), a positive bias on the platinum microelectrode produces hydronium \( H_3O^+ \) ions. Therefore the local pH decreases and finally the pH-ChemFET-
meter output voltage $V_{out}$ increases. Since the $E_{1+}$ is $\sim-0.7$ V [24], a similar polarization voltage $V_p$ was applied on the platinum microelectrode at different polarization times ($t_p$) while maintaining a constant $[\text{H}_2\text{O}_2]$ concentration at 50 mM in the analyzed buffer solution (pH = 7.2). Positive shifts of $V_{out}$ (i.e. local pH decreases) are clearly evidenced with the 50 mmol/L H$_2$O$_2$ concentration in solution, in agreement with equation 1 (figure 2a). $V_{max}$ (maximum $V_{out}$) increases with $t_p$, and the voltage variation $\Delta V_{out}$ reaches $\sim150$ mV ($\Delta \text{pH} \approx 3$ according to the pH-ChemFET sensitivity around 50 mV/pH) for the highest polarization time ($t_p = 60s$).

The influence of the polarization voltage $V_p$ was also studied (figure 2b). Starting from $V_p = 0.5$ V (lower than the H$_2$O$_2$/O$_2$ equilibrium potential $E_{1+}$ [20]) impulsional voltage variations (i.e. impulsional pH variations) are probed in the buffered solution: $V_{max}$ increases with $V_p$. However, in the H$_2$O$_2$ electrolysis case, a constant level ($\Delta V_{out} \approx 225$ mV or $\Delta \text{pH} \approx 4.5$ according to the pH-ChemFET sensitivity) appears in the [1.1 – 1.3 V] polarization voltage range. This level can be easily explained. According to the Butler-Volmer theory applied to the H$_2$O$_2$ oxidation, the anodic current depends on the $[\text{H}_2\text{O}_2]$ concentration in solution [24]:

$$E_{1+} = \text{H}_2\text{O}_2/\text{O}_2$$

where $E_{1+}$ is the equilibrium potential of the O$_2$/H$_2$O$_2$ redox couple, F is the Faraday constant, S is the microelectrode active surface, $k_{1+}$ and $\alpha_{1+}$ are the standard rate constant and the anodic transfer coefficient of the O$_2$/H$_2$O$_2$ redox couple respectively, R is the ideal gas constant and T is the temperature.

Since hydrogen peroxide is consumed by the electrochemical reaction (equation 2), the $[\text{H}_2\text{O}_2]$ concentration at the microelectrode surface decreases with time until equilibrium is reached between the H$_2$O$_2$ oxidation (equation 2) and the H$_2$O$_2$-related diffusion phenomena. This self-limited equilibrium is finally responsible for the constant level phenomenon. Finally, for $V_p > 1.3$V, H$_2$O oxidation starts to occur at the microelectrode and interferes with the H$_2$O$_2$ oxidation (equation 2), leading to a further increase of $V_{max}$. 

\[
- \text{Faradaic current and \text{H}_2\text{O}_2} \\
+ \text{Chemical current and \text{O}_2} \\
\text{H}_2\text{O}_2 + 2\text{H}^+ + 2e^- \rightarrow \text{H}_2\text{O}_2 + 2\text{H}_2\text{O}
\]
The ElecFET responses obtained with and without hydrogen peroxide were studied and compared (figure 2b inset). Therefore ElecFET devices can be used for the potentiometric detection of hydrogen peroxide $H_2O_2$ in solution. The polarization voltage $V_P$ must range between 0.5 V and 1.3 V to prevent any water electrolysis interferences. The increase of $V_P$ is responsible for a sensitivity increase and for a detection range decrease. A compromise was found by choosing the following parameters for our detection experiments: $V_P = 0.7$ V and $t_P = 30$ s. Different buffered solutions (pH = 7.2) with various hydrogen peroxide $H_2O_2$ concentrations were studied using ElecFET microdevices (figure 3). Results demonstrate that the amplitude of the impulsional voltage variations increases with the $[H_2O_2]$ concentration in solution, and saturates for the highest values. This saturation phenomenon is related to the previously discussed ElecFET detection principles. When the $[H_2O_2]$ concentration increases, the $H_3O^+$ ions production also increases (equations 1 and 2). Nevertheless, the $H_3O^+$ ion increase is limited by diffusion and electromigration phenomena and by the buffer properties, leading to a new chemical equilibrium at the microscale. Hydrogen peroxide $H_2O_2$ potentiometric detection was finally demonstrated in the [10 - 100 mM] range with a global sensitivity of $\sim$2 mV/mM (figure 3 inset).

3.2. Glucose and lactate detection

To adapt ElecFET microdevices to bio-electrochemical detection, we used an enzymatic functionalization based on glucose oxidase (GOD) and lactate oxidase (LOD). LOD was preferred to lactate dehydrogenase (LDH) since it does not require the NADH/NAD$^+$ cofactor in solution to be effective. The biosensor using LDH has some drawbacks, including the fact that they are unstable, use an additional intermediate reaction and the electrochemical oxidation of NADH occur, generally, at high over potentials [25-27]. GOD and LOD catalyze the conversion of glucose and lactate to gluconic acid and pyruvic acid respectively. The catalysis produces also hydrogen peroxide, which can be oxidized at the microelectrode surface, according to the following reactions:
Glucose + O₂ ----> Gluconic acid + H₂O₂ \hspace{1cm} (3)
L-lactate + O₂ ----> pyruvic acid + H₂O₂ \hspace{1cm} (4)

Since the dissociation constants of acids produced by the reactions are very low, a simple pH-ChemFET would not significantly detect any changes in term of pH. Thus we exploited the electrochemical technique to achieve the target. The produced hydrogen peroxide was electrochemically oxidized as shown in Equation 1. Their associated reaction kinetics is rapid and is sustained over time through the polarization time. The production of hydronium ions H₃O⁺ remains higher and the pH may be measured by pH-ChemFET-metry at the microscale.

Figures 4a and 4b show the output voltage variations $\Delta V_{out}$ versus time for different lactate and glucose concentrations respectively and for different concentrations of phosphate buffer solution ($V_P = 0.7$ V and $t_P = 5$ min). When the lactate (respectively glucose) concentration increases, the $\Delta V_{out}$ value also increases. In fact, the lactate (respectively glucose) reaction in the presence of LOD (respectively GOD) produce hydrogen peroxide H₂O₂. For a potential $V_P = 0.7$ V, the electrochemical molecule was oxidized on the platinum microelectrode, leading to hydronium ions H₃O⁺ release and therefore to a local pH change measured by the pH-ChemFET. As soon as the polarization on the platinum microelectrode was interrupted, the ElecFET microsensor response tends to turn to back equilibrium because of diffusion phenomenon and solution buffer effect.

Plotting the maximal $\Delta V_{out}$ value versus lactate and glucose concentrations, typical detection curves were obtained for different buffered concentrations, i.e. 6 and 25 mM for the lactate detection, 25 and 100 mM for the glucose detection (figures 5a and 5b). It appears that the sensitivity of the sensor depends on the buffer concentration, which is a classical result in pH-ChemFET-metry. However, results enable to determine detection sensitivities between 8 and 20 mV/mM for the lactate detection between 2 and 6 mV/mM for the glucose one, in concentration
range between [1-6 mM] and [1-30 mM] respectively. For lactate detection, ElecFET microsensors give higher sensitivity and lower detection range sensitivity than classical ChemFET-based ones (reported value around 10 mV/mM in the [1-5 mM] concentration range) [19]. On the contrary, for glucose detection ElecFET are characterized by lower sensitivity on similar detection range [14,21]. Such results discrepancy is related to differences between the LOD-based and GOD-based enzymatic layers. Indeed, experiments were performed according to enzymes initial conditioning and properties, leading to different enzymatic activities and finally to different detection sensitivities and different detection range according to different buffer effects. Detection limit was finally estimated around 0.05 mM for both lactate and glucose detection. Such result is worse than that obtained for conventional amperometry technique [28-30]. Nevertheless, the ElecFET detection properties can be improved by optimizing the enzymatic layer properties, the microdevice geometry and integration level, as well as the polarization conditions.

Finally, the evolution of ElecFET response was studied as a function of polarization time $t_P$ for lactate and glucose detection. Figure 6a was obtained for 1 mM of lactate in different phosphate buffers concentrations, i.e. 6 mM and 25 mM, with a polarization voltage $V_P = 0.7V$. The variations of $\Delta V$ were 20mV and 135 mV respectively. Figure 6b shows the glucose ElecFET response under the same conditions for two different phosphate buffer concentrations, i.e. 25 mM and 100 mM. In both cases, the sensor response increases linearly with the polarization time $t_P$ and reaches a saturation value. The sensor response was even more pronounced when the buffer concentration decreased. When the polarization time was continuously increased, reactions 1, 3, and 4 increased the concentration of hydronium ions $\text{H}_3\text{O}^+$ and consequently the decrease of the local pH. This behaviour was kept until equilibrium was established between the $\text{H}_3\text{O}^+$ ions production, the diffusion phenomena and the buffer effect.

4. Conclusion
To summarize, we studied a promising detection concept named ElecFET microsensor, which uses a pH ChemFET with a platinum microelectrode and combines potentiometric and amperometric techniques. This new concept was first applied to the detection of H$_2$O$_2$. The polarization duration $t_P$ and voltage $V_P$ parameters were studied and the limit of the microelectrode polarization was determined to avoid any interference with the water oxidation reaction. A sensitivity of 5 mV/mM was obtained in a concentration range of hydrogen peroxide ranging from 10 to 100 mM. The detection of H$_2$O$_2$ opens new opportunities for the H$_2$O$_2$–based enzymatic detection of biomolecules. This concept was finally applied to the lactate and glucose detection by functionalizing the gate surface with specific enzymes (lactate oxidase and glucose oxidase). Depending on the phosphate buffer concentration, lactate sensitivities between 8 and 20 mV/mM were evidenced in the [1 - 6 mM] concentration range.

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**References**


Biographies

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Laurent Mazenq was born on May 30, 1982. He received his University Institute of Technology’s Degree from the University Paul Sabatier de Toulouse (France) in 2002. Then, he joined the Laboratoire d'Analyse et d’Architecture des Systèmes of the French Centre National de la Recherche Scientifique (LAAS-CNRS) as an engineer in micro/nanotechnologies. He is more especially working on the development of optical microlithography processes for the technological realisation of micro/nanodevices.

Jérôme Launay was born on March 11, 1975. He received the degree in electronic engineering from the Institut National des Sciences Appliquées de Toulouse" (France) in 1998. He joined the Laboratoire d'Architecture et d'Analyse des Systèmes from the French "Centre National de la Recherche Scientifique" (LAAS-CNRS) in 1998 and received the PhD degree from the Institut National des Sciences Appliquées de Toulouse (France) in 2001. In 2002, he became lecturer at the Université Paul Sabatier de Toulouse (France). His research activities include the development of chemical microsensors for the detection in liquid phase.
William Sant was born on May 18, 1969. He received his Master’s Degree in electronics in 1996 and his Diplome d'Etudes Approfondies in Microelectronics from the Université Paul Sabatier de Toulouse (France) in 1999. He joined the Laboratoire d'Architecture et d'Analyse des Systèmes of the French Centre National de la Recherche Scientifique (LAAS-CNRS) in 2000 and received the PhD degree from the Université Paul Sabatier de Toulouse (France) in 2004. Since then, he has joined the HEMODIA society (France) as an R&D engineer and has been working on the development of ChemFET microsensors for medical applications.

Pierre Temple-Boyer was born on October 25, 1966. He received his Engineer Degree in electronic engineering from the Ecole Supérieure d'Electricité (Paris – France) in 1990 and his Master’s Degree in microelectronics from the Université Paul Sabatier de Toulouse (France) in 1992. He joined the Laboratoire d'Architecture et d'Analyse des Systèmes of the French Centre National de la Recherche Scientifique (LAAS-CNRS) in 1992 and received the PhD degree from the Institut National des Sciences Appliquées de Toulouse (France) in 1995. Since then, as a senior CNRS researcher, he has been working on the development of microtechnologies for the realisation of integrated microsystems.
Figures Caption

Figure 1: Cross-section (a) and details (b) of the ElecFET device (chip size: 3.5 x 3.5 mm²).

Figure 2: Detection of impulsional pH variations due to the H₂O₂ electrolysis ([H₂O₂] = 50 mM) in 100 mM phosphate buffer solutions, influence of the polarization time tₚ (a) and influence of the polarization voltage Vₚ with and without hydrogen peroxide (b and inset).

Figure 3: Detection of impulsional pH variations related to the H₂O₂ electrolysis in phosphate buffer solutions: influence of the H₂O₂ concentration and (inset) ElecFET response for the H₂O₂ detection.

Figure 4: ElecFET microsensor response versus time (Vₚ = 0.7 V) for the lactate (a) and glucose (b) detection in phosphate buffer solutions.

Figure 5: ElecFET lactate (a) and glucose (b) detection for different phosphate buffer concentrations.

Figure 6: ElecFET responses ΔVₜₐₐₜ versus polarization time tₚ (Vₚ = 0.7 V) for lactate (a) and glucose (b) in different buffer concentrations.
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Figure 2: Detection of impulsive pH variations due to the $\text{H}_2\text{O}_2$ electrolysis ($[\text{H}_2\text{O}_2] = 50 \text{ mM}$) in $100\text{mM}$ phosphate buffer solutions, influence of the polarization time $t_P$ (a) and influence of the polarization voltage $V_P$ with and without hydrogen peroxide (b and inset).
Figure 3: Detection of impulsional pH variations related to the H$_2$O$_2$ electrolysis in phosphate buffer solutions: influence of the H$_2$O$_2$ concentration and (inset) ElecFET response for the H$_2$O$_2$ detection
Figure 4: ElecFET microsensor response versus time ($V_p = 0.7$ V) for the lactate (a) and glucose (b) detection in phosphate buffer solutions.
Figure 5: ElecFET lactate (a) and glucose (b) detection for different phosphate buffer concentrations.
Figure 6: ElecFET responses $\Delta V_{out}$ versus polarization time $t_P$ ($V_P = 0.7V$) for lactate (a) and glucose (b) in different buffer concentrations.