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Numerical Modeling of Glucose Biosensor With pH-Based Electrochemical Field-Effect Transistor Device — Source link

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Numerical Modeling of Glucose biosensor with pH-based Electrochemical Field Effect Transistor Device

Moussa Ba, Abdou Karim Diallo*, El Hadji Babacar Ly, Jérôme Launay, Pierre Temple-Boyer

Abstract—In this work, we present a new model for glucose detection using an Electrochemical Field Effect Transistor (ElecFET) with a MATLABTM software. ElecFET is a device in which the basic element is a pH-ChemFET (pH-sensitive Chemical Field Effect Transistor) designed with an integrated platinum microelectrode around the sensitive gate dielectric capable to trigger electrochemical reactions. A number of parameters such as polarization time (T_{pol}) and polarization voltage (V_p) have been investigated, as well as the enzymatic unit number per volume (N_{enz}), the Michaelis constant (K_M) and the glucose concentration (C_0). Each of the aforementioned parameters influences the glucose sensor response. Experimental data compared with our numerical model showed a reasonable fit.

Index Terms—Modeling, ChemFET, Microelectrode, ElecFET, Glucose sensor, Biosensor

I. INTRODUCTION

OWADAYS, it is no longer necessary to demonstrate the prominence to detect or monitor the glucose concentration either in the blood or in a daily diet. The detection of glucose can be found in various fields such as, environment, food industry, fuel cells [1], to name a few. However, one of the flagship applications is in clinical field. Detecting or monitoring the level of glucose in the blood is an excellent indicator of diabetes. This chronic and non-communicable disease (NCD) is one of the most important public health challenges in this 21st century. The World Health Organization has registered 450 million cases of diabetes worldwide and this number could reach 700 million in 2045 [2].

From that point of view, several types of biosensors dedicated to glucose detection have been studied. Most of the techniques used for glucose detection are based on optical or electrochemical detections [3, 4]. However, the optical methods are expensive and not easy to use for specific applications. Therefore, the most studied glucose detection devices use electrochemistry and particularly the amperometry technique, while potentiometry is still less explored. Both methods, i.e., amperometry and potentiometry, work in liquid phase.

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Amperometry uses the oxido-reduction properties leading to the development of electrodes and electrochemical cells [5, 6], while potentiometry focuses on the working principle of ion sensitive electrode (ISE) and ChemFET resulting to charge trapping at the solid/liquid interface [7, 8] followed by potential changes. The combination of both techniques is very promising and has been initiated earlier by Sohn et al. [9] and later by Diallo et al. [10]. The device resulting from the combination of the aforementioned techniques has been named ElecFET and comprises a pH-ChemFET associated to an integrated platinum microelectrode around the sensitive gate area. The platinum microelectrode is used to trigger the electrochemical reactions while the pH-ChemFET measures the pH variations around the gate dielectric. ElecFET has been used for the detection of dissolved oxygen [9], hydrogen peroxide (H_2O_2) , lactate and glucose [10-12], opening a promising way for biomolecules produced by reactions related to oxidase-based enzymes. Despite efforts made for understanding the ElecFET working principle, very few models have been studied to predict the sensor response with this device, compared to what is observed in BioFET (Biologically sensitive FET)[13,14] and especially in EnFET (Enzymatic FET)[15-20]. The only models for ElecFET are those proposed by Diallo et al. [21] and Aoun et al. [22] using respectively MATLAB and COMSOL Multiphysics 2D for the detection of hydrogen peroxide and water electrolysis. In this study, we present a model and experimental results of glucose biosensor based on ElecFET device using MATLAB software. We first considered the transformation of glucose into hydrogen peroxide H_2O_2 thanks to the glucose oxidase. Then, H_2O_2 is electrochemically transformed into hydronium ions H_3O^+ through the platinum microelectrode. Finally, the H_3O^+ ions diffuse into the solution leading to local pH changes detected by the pH-ChemFET. Beyond diffusion phenomena of glucose and all the chemical species, the main influential parameters such as enzymatic unit number per volume (N_{enz}) , Michaelis constant (K_M) as well as the polarization (V_p) voltage and polarization time (T_{pol}) on the microelectrode, are studied. Experimental data have been correlated to the theory in order to assess the glucose biosensor response.

II. EXPERIMENTAL SECTION AND MODELING

A. Experimental Section

The realized ElecFET and the enzymatic functionalization have been studied in previous works [10]. Fig. 1a is a

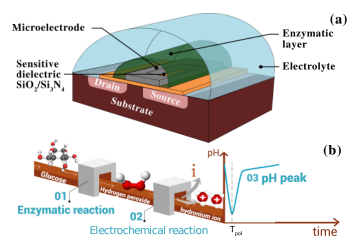


Fig. 1. (a) Schematic representation, (b) glucose detection principle using ElecFET device.

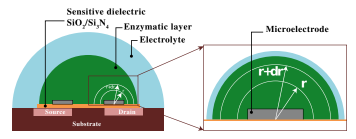


Fig. 2. Description of one-dimensional model in radial coordinate.

schematic representation and Fig. 1b illustrates the glucose detection principle using ElecFET.

B. Modeling

The different reactions during ElecFET operation are briefly described in Fig. 1b. We first considered the enzymatic reaction of glucose into hydrogen peroxide and the diffusion of different biochemical species(step 1). Secondly, by polarizing the platinum microelectrode, electrochemical reaction is triggered and H_2O_2 transformed into hydronium ions H_3O^+ (step 2). The current produced during this electrochemical reaction depends on the V_p and T_{pol} . Finally, the hydronium ions released from this reaction created a local pH change detected by the traditional pH-ChemFET(step 3).

Fig. 2 represents the Cross-section of the ElecFET device describing the one-dimensional model in radial coordinate. To simplify the schema of the description of one-dimensional model in radial coordinate, we only consider one part of the microelectrode in the cross section of fig. 2.

• Modeling of glucose oxidase enzymatic reaction The glucose oxidase (GOD) catalyzes the glucose oxidation according to the following reaction(Reaction 1):

$${\rm glucose} + {\rm O_2} \xrightarrow{\rm glucose~oxidase + H_2O} {\rm gluconic~acid} + {\rm H_2O_2} \; ({\rm Reaction} \; 1)$$

The term of production/consumption (g_{enz}) corresponding to the production of hydrogen peroxide from the glucose substrate is expressed in moles per unit of volume and unit of time and written as:

$$g_{enz} = a_M * N_{enz} \frac{[S]}{[S] + K_M}.$$
 (1)

Where a_M is the maximal enzyme activity, N_{enz} is the enzymatic unit number per volume, [S] is the glucose concentration in solution and K_M is the glucose oxidase Michaelis constant. The term g_{enz} exists only within the enzymatic layer, otherwise it is equal to zero. In the model, this layer thickness is set to $1\,\mu\mathrm{m}$, based on estimates of the enzyme deposition rate.

• Modeling of the electrochemical production of H_3O^+

By applying to the integrated microelectrode a voltage at least greater than the equilibrium potential for H_2O_2 oxidation, the hydrogen peroxide resulting from the enzymatic reaction is oxidized and H_3O^+ ions are released according to the reaction:

$$H_2O_2 + 2H_2O \rightarrow O_2 + 2H_3O^+ + 2e^-$$
 (Reaction 2)

The concentration of the H_3O^+ ions resulting from this reaction is given by the following relation:

$$(\frac{\partial [H_3O^+](r,t)}{\partial t}) = 2g_{elec} = 2[H_2O_2](r,t)\frac{K_{1^+}}{dr}exp(\frac{F(\Delta V_p)}{RT}). \quad (2)$$

Where g_{elec} represents the production/consumption rate according to the associated electrochemical reaction, $\Delta V_p = (V_p - E_{1+})$ with E_{1+} the equilibrium potential for H_2O_2 oxidation, $K_{1+} = 5.10^{-8}$ m/s is the standard rate constant for the H_2O_2 oxidation, F is the Faraday constant (F = 96.485 C/mol), R is the ideal gas constant (R = 8.32 J/(K•mol)), T = 300 K is the absolute temperature.

• Influence of water self-ionization According to water self-ionization :

$$2H_2O \leftrightarrow H_3O^+ + OH^-$$
 (Reaction 3)

The variation of H_3O^+ ions concentration is equal to those OH^- ions. It is noted ϕ and its expression is given by :

$$\phi = \Delta[H_3O^+] = \Delta[OH^-]. \tag{3}$$

$$\phi = \frac{\sqrt{([H_3O^+] - [OH^-])^2 + 4K_w} - [H_3O^+] - [OH^-]}{2}.$$
 (4)

Where K_w is the ionic product of water.

Modeling of diffusion phenomena

The diffusion model is based on the Fick's diffusion equation, assuming a one-dimensional model in spherical coordinates with the radial component r (Fig. 2):

$$\frac{\partial C(r,t)}{\partial t} = D \frac{1}{r^2} \frac{\partial}{\partial r} (r^2 \frac{\partial C(r,t)}{\partial r}) + \epsilon_1 g_{enz}(r,t) + \epsilon_2 g_{elec}(r,t). \quad (5)$$

Where C(r,t) describes the concentration of the species, D the associated diffusion coefficient, $\epsilon_i=\pm 1$ (with i=1 or 2), for a production $\epsilon_i=+1$ while for a consumption $\epsilon_i=-1$, g_{enz} represents the production/consumption according to the associated enzymatic reaction. The term of g_{elec} is obtained only when a polarization voltage is applied on the microelectrode during T_{pol} . As a result, the diffusion equations are:

$$\frac{\partial [glucose](r,t)}{\partial t} = D_{glucose} \frac{1}{r^2} \frac{\partial}{\partial r} (r^2 \frac{\partial [glucose](r,t)}{\partial r}) - g_{enz}(r,t). \tag{6}$$

$$\frac{\partial [H_2O_2](r,t)}{\partial t} = D_{H_2O_2} \frac{1}{r^2} \frac{\partial}{\partial r} (r^2 \frac{\partial [H_2O_2](r,t)}{\partial r}) + g_{enz}(r,t) - g_{elec}(r,t). \tag{7} \label{eq:7}$$

$$\frac{\partial [H_3O^+](r,t)}{\partial t} = D_{H_3O^+} \frac{1}{r^2} \frac{\partial}{\partial r} (r^2 \frac{\partial [H_3O^+](r,t)}{\partial r}) + 2g_{elec}(r,t). \eqno(8)$$

All the diffusion constants have been calculated according to the Einstein's Equation [15; 16]:

$$D = A \frac{kT}{\sqrt[3]{M}}. (9)$$

Where A is a parameter related to the electrolyte, k the Boltzmann's constant, T the absolute temperature and M is the molar mass of the associated molecule.

The initial condition related to mass transport of H_3O^+ ions is given by:

$$[H_3O^+](r,0) = 10^{-pH_0} \Leftrightarrow [OH^-](r,0) = K_w 10^{pH_0}.$$
 (10)

At the upper interface of the electrolyte $(r=r_{max})$ and the lower interface (r=0), the interface is assumed to be impermeable (i.e., there is no diffusion at the upper limits and lower):

$$\left(\frac{\partial C(r,t)}{\partial r}\right)_{r=0} = \left(\frac{\partial C(r,t)}{\partial r}\right)_{r=r_{max}} = 0.$$
 (11)

• Modeling of the pH-ChemFET response

The pH-ChemFET threshold voltage V_T variation also considered as the sensor response was estimated according to the following equation:

$$\delta V_T = S_0 (pH(d,t) - pH_0).$$
 (12)

where S_0 is the pH-ChemFET sensitivity according to the site-binding model and d is the distance between the integrated microelectrode and the pH-ChemFET gate sensitive area.

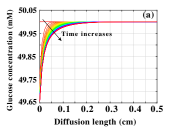
III. RESULTS AND DISCUSSIONS

In previous work, several attempts have been done to demonstrate glucose sensor that used conventional EnFET [23-26], but the major drawbacks of its rapid progress in practical applications is the limitation of the dynamic range of the biosensor response because of the insufficient concentration of dissolved oxygen and the low dissociation constant of gluconic acid [12; 26]. The advantages obtained with this ElecFET results from the platinum microelectrode added to the device. Indeed, this microelectrode is firstly responsible to the H_2O_2 electrolysis producing oxygen which is recycled in glucose oxidation reaction and establishing an extended dynamic range. Secondly, the oxidation reaction of hydrogen peroxide at the platinum microelectrode is fast, maintained over time as wished and produces two hydronium ions compared to gluconic acid dissociation used solely in EnFET. This second point enables to control the electrochemical reaction and therefore the sensor response. The ElecFET model is simulated by MATLAB tool and the parameters and values used for the modeling are summarized in table I.

• Production/consumption of chemical species from enzymatic reaction: diffusion phenomena

TABLE I
PARAMETERS AND VALUES USED IN THIS MODELING

Parameter	Signification	Value
T_{pol}	Polarization time	10 s
V_p	Polarization voltage	1 V
$ \begin{array}{c} T_{pol} \\ V_p \\ E_{1^+} \end{array} $	Equilibrium potential for	0.7 V
	H_2O_2 oxidation	
S_0	Sensitivity	59 mV/pH
pH_0	Initial pH of the solution	7
C_0	Initial concentration of glucose	$5.10^{-2} mol/L$
$[H_2O_2]_0$	Initial concentration of H_2O_2	0 mol/L
N_{enz}	Enzymatic unit number per	$10^4 \ units/cm^3$
	volume	·
K_M	Michealis Constant of glucose	$7.15.10^{-2} \ mol/L$
	oxidase	,



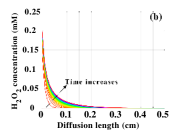


Fig. 3. (a) Glucose concentration and (b) hydrogen peroxide concentration versus diffusion length at different times and keeping constant all the parameters.

Fig. 3a and 3b show respectively the glucose and hydrogen peroxide concentration changes in the enzymatic layer as a function of the diffusion length over time. As we can see in Fig. 3a, close to the sensitive gate, the glucose concentration decreases as the time increases, while beyond this area, i.e., deep into the bulk solution, it remains constant. Opposite behavior is observed with respect to hydrogen peroxide. The concentration of H_2O_2 increases close to the sensitive zone while further in the solution it decreases. These behaviors are explained by the presence of glucose oxidase enzyme which catalyzes the reaction of glucose in the presence of oxygen to hydrogen peroxide. The local production of this H_2O_2 leads to its increase, a consequence of the glucose consumption (concentration decreasing) in the enzymatic layer. The longer the time increases, the more diffusion occurs. It should be noted that beyond the enzymatic layer, the concentration of hydrogen peroxide is very low (or even zero) as shown in Fig. 3b. At the same time, the more the time increases the more diffused glucose into the enzyme layer, thereby leading to its oxidation. Consequently, the glucose concentration decreases near the detection area of the sensor device. This confirms the consumption/production of glucose/ H_2O_2 respectively through the enzymatic reaction.

• Electrochemical reaction and pH detection

The detection of pH by the pH-ChemFET is performed when H_3O^+ ions are produced. The reaction of glucose using their specific glucose oxidase enzyme (Reaction 1) generates H_2O_2 molecules. As mentioned in the previous section, H_2O_2 is converted on the platinum electrode giving oxygen and two H_3O^+ ions (Reaction 2) diffusing into the enzymatic layer and beyond. Fig. 4 illustrates the H_3O^+ concentration changes versus diffusion length. As can be seen, far from the

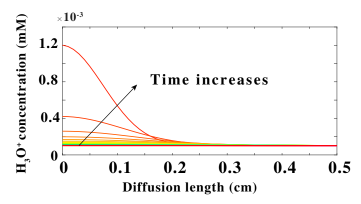


Fig. 4. Hydronium ions concentration versus diffusion length at different times by keeping constant all the parameters.

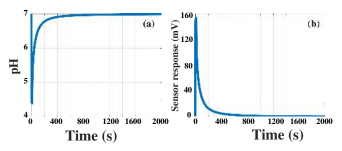


Fig. 5. (a) pH variation and (b)sensor response variation versus time by keeping constant all the parameters.

detection area, i.e., deep into the bulk solution, the H_3O^+ concentration is equal to its initial value $(10^{-7}M)$, whereas close to the Si_3N_4 /solution interface the concentration is higher. Similar principle related to diffusion phenomena mentioned above could explain this change in concentration. When the electrochemical reaction is triggered gradually as time increases, as long as this reaction occurs, the local production of H_3O^+ is maintained and the diffusion phenomena keep going. Therefore, on the platinum microelectrode covered by the enzymatic layer, the concentration of hydronium ions are more and more important and further in the solution it is very low.

By applying a polarization $V_p=1~\rm V$ on the microelectrode for a during $T_{pol}=10~\rm s$ (by keeping C_0 , N_{enz} , K_M , and all the parameters constant), the generation of H_3O^+ ions is triggered. By removing the polarization after T_{pol} , we observe diffusion phenomena with a return to the initial state as observed in fig. 5a. The local pH variations are recorded by the pH-ChemFET and result in a potential change as can be seen in Fig. 5b. The sensor response related to the potential variations, changes from 0 V to 158 mV corresponding respectively to pH=7 and pH=4. This demonstrates that localized impulsional pH variations can be effectively obtained near the integrated microelectrode and detected at the adjacent pH-sensitive surface after glucose oxidation by the glucose oxidase enzyme.

• Influence of main parameters for glucose detection

In this section the main potential parameters influencing the sensor response have been investigated. Such parameters are the glucose concentration, polarization voltage, polarization

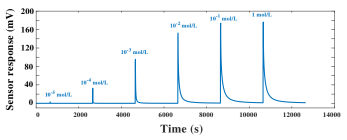


Fig. 6. Sensor response variation over time by successively changing the glucose concentration.

time, enzymatic unit number per volume and the Michaelis constant.

Fig. 6 represents the sensor response over time with different glucose concentrations in the range of $[10^{-5} \text{ mol/L}]$ 1 mol/L]. As observed, the sensor response increases with glucose concentration. It linearly increases from $C_0 = 10^{-4}$ mol/L to $C_0 = 10^{-1}$ mol/L with potential changes of 20 mV and 175 mV respectively, giving thus a linear detection range. Beyond $C_0 = 10^{-1}$ mol/L, we observe a saturation effect that could be the consequence of glucose diffusion phenomena from the solution towards the enzymatic layer. Fig. 7a shows the sensor response versus glucose concentration in logarithmic scale for different polarization voltages. By keeping V_p constant, the sensor response linearly increases before saturating. This saturation behavior starts from 10^{-1} mol/L of glucose regardless of the polarization voltage. From Fig. 7a, we can note that: (i) irst, it is found that the sensor sensitivity decreases for low V_p , and in the same way, the greater the polarization on the platinum microelectrode, the higher the sensitivity. The sensitivity changes from 8 mV/dec to 46 mV/dec for respectively 0.8 V and 0.95 V of V_p . (ii) Then the limit of detection is affected by V_p . As can be seen for Vp = 0.85 V; 0.90 V and 0.95 V, the glucose detection limits are respectively 10^{-3} mol/L, 10^{-4} mol/L and 10^{-5} mol/L. (iii) Finally, the sensor response increases as the polarization increases. This behavior is shown by the figure in insert which represents the sensor response versus V_p . There is clearly an exponential dependence of the sensor response polarization as evidenced by the Butler-Volmer theory.

An increase of polarization time has a very slight influence on the sensitivity. In addition, the limit of detection remains almost unchanged. The only notable variation recorded is the sensor response increase as T_{pol} increases. For $T_{pol} = 0.5$ s and 250 s, the potential variations are respectively 160 mV and 180 mV as illustrated in Fig. 7b. The insert of Fig. 7b represents the sensor response versus polarization time. We note an increase of the sensor response followed by saturation. Polarization on the microelectrode with V_p during T_{pol} maintains the electrochemical reaction of hydrogen peroxide as long as such a polarization keeps going. This enables to keep producing H_3O^+ ions until the phenomena related to chemical species diffusion induce saturation effects on the sensor response. The enzymatic unit number per volume is one of the most important parameters and depends on the enzyme preparation. So, we can also act on this parameter to control the sensor response. Fig. 8a shows the influence

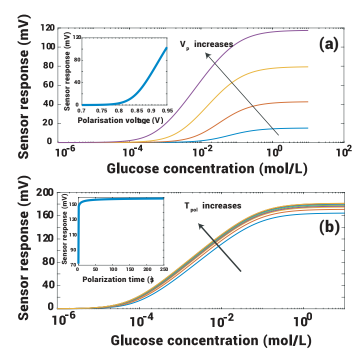


Fig. 7. (a) Sensor response versus glucose concentration in logarithmic scale with different $V_p = 0.85$ V; 0.90 V and 0.95 V (insert we show sensor response versus V_p).(b) Sensor response versus glucose concentration in logarithmic scale with different $T_{pol} = 0.5$ s; 2 s; 5 s; 10 s; 20 s; 50 s; 100 s; 180 s; 200 s and 250 s(insert represents sensor response versus T_{pol}).

of N_{enz} on the sensor response. It is found that this N_{enz} directly affects the biosensor sensitivity, it decreases when the enzymatic layer is weakly concentrated in glucose oxidase. For $N_{enz} \geq 10^3$ units/cm³, there is no sensitivity increase. It is important to mention that the enzymatic unit number is involved in the generation/consumption of species as described by equation (2), so the higher the N_{enz} is, the greater the enzymatic kinetics increases, leading to a sensitivity increase. This increase of N_{enz} promotes the glucose consumption and therefore H_2O_2 creation (H_3O^+ creation). We can also record an increase of sensor response as well as a decrease of the limit of detection when N_{enz} increases. Fig. 8b shows the sensor response versus glucose concentration in logarithmic scale with different Michaelis constant K_M . The Michaelis constant is specific to the enzyme and reflects its affinity with substrate, the larger it is, and the more the substrate is fixed poorly on the enzyme. It assesses the reaction kinetics and the lower the value, the faster the reaction. From Fig. 8b, neither the sensitivity nor the sensor response is affected. However, we can observe a shift of glucose detection limit. When K_M increases the glucose detection limit shifts towards high concentrations. The Michaelis constant help then to determine the concentration ranges related to linear operation of ElecFETbased biosensor, but has no influence on the sensitivity.

• Comparison between theory and experimental data

In this section we compare the experimental data with our model. In order to validate our model different sensor responses have been recorded using the following parameters: $S_0 = 50 \text{ mV/pH}$, $N_{enz} = 150 \text{ units/cm}^3$, $K_M = 100 \text{ mV/pH}$

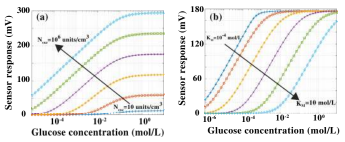


Fig. 8. Sensor response versus glucose concentration in logarithmic scale with (a) different N_{enz} and (b) different K_M .

 10^{-2} mol/L ,pH=7 , $\Delta V_p=0.27$ V.

Fig. 9a and 9b show comparative studies between theory/experiment, and a good similarity is noted. For both fig. 9a and 9b, we can see that the sensor response curve in experimental studies shows a similar shape compared with theoretical curve. Our numerical model shows that the sensor response linearly increases with glucose concentration in solution and saturates for the highest values (Fig. 9a). This saturation phenomenon is related to the ElecFET detection principles as previously mentioned. Indeed, in the presence of glucose, H_2O_2 is created thanks to the glucose oxidase enzyme. By applying a positive voltage on the platinum microelectrode in order to trigger the electrochemical reaction, H_2O_2 is transformed with H_3O^+ ions generation and therefore a local pH decrease. As the glucose concentration increases, the H_3O^+ ions production also increases. Nevertheless, the H_3O^+ ion increase is limited by diffusion and electromigration phenomena, leading to a new chemical equilibrium at the microscale. Fig. 9b shows the reproducibility measurement for T_{pol} =15 min. A reasonable fit between the model and the experimental data is recorded. It may be noted that the sensor responses in the theoretical case for both fig. 9a and 9b show slightly higher variations than in the experimental case. Additionally, the experimental curve in Fig. 9b shows a return to equilibrium slower than that obtained by theory. The discrepancy recorded between simulation and experimental data is probably due to a possible drop of glucose oxidase activation, processing or storage conditions and the effect of buffer properties. Buffer solution is of great importance in biosensor applications. Several works reported on the buffer properties which enhance the response and the stability of the biosensor [22, 27, 28]. In our model the chemical effect of the buffer solution was not considered (and therefore neglected) in order to better understand the ElecFET detection/transduction principles. Despite all these considerations, an acceptable fit has been obtained with experimental results.

IV. CONCLUSION

In this paper we have studied a new model using ElecFET device for glucose detection. The main influential parameters have been investigated. The Michaelis constant helps to determine concentration ranges related to linear operation of the ElecFET-based biosensor, but has a no significant influence on the sensitivity. In the meanwhile, enzymatic unit number increases the sensitivity and the detection range. The polarization time has no effect on the sensitivity and the dynamic

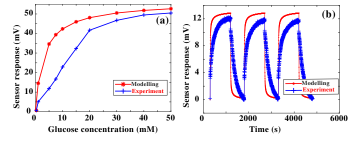


Fig. 9. Comparison between modeling and experimental data: (a) sensor response versus glucose concentration for ΔV_p =0.27 V and T_{pol} =5min.(b) Sensor response over time with successive constant polarization ΔV_p =0.27 V, T_{pol} =15min and $C_0=10^{-3}$ mol/L.

range. Whereas the polarization voltage increases the sensor response, the sensitivity and the dynamic range. Our numerical model compared to experimental data shows a similar shape, giving a significant step on predicting glucose biosensor using ElecFET. However, the model needs to be improved by taking into account the buffer properties of the electrolyte solution. Nevertheless, it enables to better understand the detection principle of biosensor based on ElecFET device.

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