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# Deriving a novel methodology for nano-BioFETs and analysing the effect of high-k oxides on the amino-acids sensing application



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ABSTRACT

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#### 1. Introduction

Since 1952, when Shockley invented Field-Effect Transistors (FETs), there have been undergoing many significant modifications to cater to the needs of sensing applications [1]. In 1970, Bergveld developed an ion-sensitive Field-Effect Transistor (ISFET) that is used for ion detection in a chemical environment [2]. With further development of technology and its application for biosensing, novel functionalized ISFET devices have been fabricated for protein and DNA detection [3 4]. Various simulation methodologies and fabrication techniques have been carried out to further expand the knowledge of the impact of sensor surface functionalization on the device performance [5 6].

In this paper, we have adopted a design-of-experiment approach based on an analytical model. The analytical model is implemented in a MATLAB simulation environment and it effectively combines the Gouy-Chapman-Stern model with the Site-binding model, which allows us to simulate various amino acids (AAs) immobilized on different high-k dielectrics as gate oxides [7 8]. Three AAs with different side chains are chosen to show the effect of high-k dielectric functionalization on the device sensitivity. The chosen AAs are Arginine (R), Aspartic Acid (D) and Proline (P). The high-k oxides used are titanium oxide (TiO<sub>2</sub>), hafnium oxide (HfO<sub>2</sub>) and silicon dioxide (SiO<sub>2</sub>). The Gouy-ChapmanStern layer model combines the Helmholtz and the diffuse double layer of the Gouy-Chapman model to explain the development of the double-layer capacitance on the sensor's surface [9]. The Site-binding model is used to describe the surface charge developed at the oxide/ electrolyte interface [8]. This paper presents a new methodology that combines these two theories and our new method is applied to AAs immobilized on the sensor surface. Moreover, our simulation work shows that each AA has a unique signature (fingerprints) that can be used to distinguish and identify each AA [10].

In this paper, we present simulation results of a BioFET using methodology based on the Gouy-Chapman-Stern

and the Site-binding models. The derived simulation approach is used to sense different amino acids, such as Arginine (R), Aspartic Acid (D) and Proline (P), functionalized with the help of a linker connected to the gate-

oxide. The performance of the BioFETs is optimized while analyzing the effect of high-k dielectrics as the gate

oxide. In general, the channel oxides are responsible for tuning the parameters such as sensitivity, surface po-

tential and intrinsic buffer capacity. The variation of total surface capacitance, the second derivative of drain

current and surface potential are used to uniquely identify the signatures of different amino acids. The proposed

method can be helpful in defining an efficient method for label-free protein sequencing.

Fig. 1 shows the schematic diagram of the BioFET implemented in the simulations. As an example, D is immobilized over the linker with a carboxylic terminal which eliminates the affinity corresponding to the C- terminal. Similarly, the chosen AAs can be immobilized by either carboxylic or amine terminal leaving the AA with lower affinity sites. The surface potential ( $\Psi_0$ ) is calculated assuming the point charges that are separated from the oxide surface by the length and the effective permittivity of the AAs. Our simulation approach allows us to calculate  $\Psi_0$ , the sensitivity factor ( $\alpha$ ), the intrinsic buffer capacity ( $\beta$ ) and the zeta potential ( $\Psi_{\epsilon}$ ) for different oxides [7]. From this, we can also calculate the depletion width ( $W_D$ ), the drain current ( $I_{SD}$ ) and the 2nd order derivative of  $I_{SD}$  for each immobilized AAs (R, D and P). More importantly, all of these characteristics can be used as unique fingerprints for

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**Fig. 1.** Schematic diagram of the BioFET sensor with immobilised amino acids over the oxide. AA denotes the amino acid with carboxyl sidechains (example-Aspartic Acid). At the very bottom, the light blue colour is the Silicon nanowire channel. Immediately on top of it, as seen in purple colour, is the oxide that can be TiO<sub>2</sub>, HfO<sub>2</sub> or SiO<sub>2</sub>. The Silanol groups (SiOH) are represented directly above that as will be the case when SiO<sub>2</sub> is an oxide. The top of a silanol group is covered with light green colour which is the linker also known as a polyethylene glycol (PEG) layer. PEG is covalently bonded to amino acid on one side and to silanol on the other. The AA has the possibility to be protonated/ deprotonated or to have active sites (for the receptors to attach).

the identification of AAs and polypeptides.

#### 2. Methodology

For each gate oxide and immobilized AAs, we calculated first the surface charge density ( $\sigma_i$ ) (with i = 1 to 5 depending on the AA radicals) using the site-binding model for the three C or N terminals immobilized AAs that we calculated in this work: C-Imm. D [ $\sigma_1$ ], C-Imm. R [ $\sigma_2$ ], C-Imm. P [ $\sigma_3$ ], N-Imm. D [ $\sigma_4$ ], N-Imm. R [ $\sigma_1$ ] and N-Imm. P [ $\sigma_5$ ] where

$$\sigma_1 = qN_S \left(\frac{cH_s^2 - K_a K_b}{K_a K_b + K_b cH_s + cH_s^2}\right)$$
(1)

$$\sigma_2 = qN_S \left(\frac{cH_s^2 + cH_SK_a}{K_aK_b + K_bcH_S + cH_S^2}\right)$$
(2)

$$\sigma_3 = q N_S \left(\frac{c H_S}{c H_S + K_a}\right) \tag{3}$$

$$\sigma_4 = qN_S \left(\frac{-cH_SK_a - K_aK_b}{K_aK_b + K_bcH_S + cH_S^2}\right)$$
(4)

$$\sigma_5 = q N_S \left( \frac{-K_a}{cH_S + K_a} \right) \tag{5}$$

and,  $cH_s = cH_Bexp \frac{-\Psi_0}{2V_T}$ ,  $cH_B = 10^{-pH_B}$ ,  $V_T = \frac{K_BT}{q}$ ,  $cH_s$  &  $cH_B$  represents the surface and bulk proton concentration respectively,  $pH_B$  is the bulk pH,  $N_S$  is the total surface states,  $K_a$  &  $K_b$  are the dissociation constants of the corresponding reactive sites of each AA and  $V_T$  is the thermal voltage.  $\sigma_i$  is equated and calculated iteratively with the  $\sigma_{DL}$  (surface charge density of Gouy-Chapman stern layer model) until the error < tolerance (10<sup>-50</sup>). Bi-section method is adopted to find the point-of-zero charge (pH<sub>pzc</sub>) [isoelectric point] and zeta potential ( $\Psi_{\xi}$ ) for a particular surface charge density optential drop across the stern layer ( $\Psi_{Stern}$ ) and is given by the following relation:

$$\Psi_0 = \Psi_{\text{Stern}} + \Psi_{\xi} = \frac{Q_0 \sinh(\Psi_{\xi}/V_T)}{C_{stern}} + \Psi_{\xi}$$
(6)

where  $C_{Stern}$  is the capacitance of the stern layer.  $C_{Stern}$  is considered to be 0.8F/m<sup>2</sup> [11]. Using  $\Psi_0$ , sensitivity factor ( $\alpha$ ), intrinsic buffer capacity ( $\beta$ ) and 2nd order derivative of surface potential  $\left(\frac{\partial^2 \Psi_0}{\partial D H^2}\right)$  can be

calculated as shown below [12].

$$\alpha = \frac{1}{1 + 2.303 \frac{k_B T C_{diff}}{q^2 \beta}}$$
(7)

$$\frac{\delta \delta \sigma_{DL}}{\delta p H_S} = -q\beta \tag{8}$$

$$\frac{\partial^2 \psi_0}{\partial p H^2} = -2.3 \frac{kT.\partial \alpha}{q.\partial p H_B}$$
(9)

A high-aspect-ratio junctionless p-type FinFET structure is used for current simulations as it gives a larger surface area for the target molecules to come in contact with the receptors [13]. The depletion width ( $W_D$ ) and drain current ( $I_{SD}$ ) calculations are done as mentioned in ref. [14,15]. This occurs while solving the continuity equation across the electrolyte-oxide–semiconductor interfaces, using  $\Psi_0$  as the variable parameter.

# 3. Results and discussions

Fig. 2(a) shows the variation of zeta potential ( $\Psi_{\xi}$ ) with respect to pH<sub>B</sub>. The point-of-zero-charge (pH<sub>pzc</sub>) for SiO<sub>2</sub>, HfO<sub>2</sub> and TiO<sub>2</sub> can be seen as the pH at which the  $\Psi_{\xi}$  crosses the zero value. Table 1 shows the summary of *pH<sub>pzc</sub>*.  $\Psi_{\xi}$  is the potential between the stern potential and the shear plane which is most negative for the SiO<sub>2</sub> due to its higher affinity for losing protons and  $\Psi_{\xi}$  is most positive for HfO<sub>2</sub> because of its higher pK value of protonated sites. For SiO<sub>2</sub>, the pH<sub>pzc</sub> is in agreement with the experimental value. In the case of HfO<sub>2</sub> and TiO<sub>2</sub>, the pH<sub>pzc</sub> simulation value differs from the data available in the literature. This may be attributed to the ways in which oxides are grown and impurities present on the surface of oxides [18].

As shown in Fig. 2(b),  $\alpha$  is minimum at pH<sub>pzc</sub> for all three oxides. For HfO<sub>2</sub>, pH<sub>pzc</sub> is around 7, which is higher than SiO<sub>2</sub> and TiO<sub>2</sub>, since the change in  $\Psi_{\xi}$  with respect to  $pH_B$  is higher for enhanced sensitivity. The increment in  $\alpha$  is observed for all gate oxides before and after  $pH_{pzc}$  as this is due to the increase in  $\Psi_{\xi}$  away from the pH<sub>pzc</sub>.  $\beta$  has a similar trend with respect to pH<sub>B</sub> as it denotes the change in surface charge density  $(\sigma_0)$  with respect to surface pH (pH<sub>s</sub>) as shown in Fig. 2(c). Fig. 3(a) shows the  $\Psi_0$  variation with respect to pH for all three gate oxides. This follows the  $\Psi_{\xi}$  with the addition of  $\Psi_{Stern}$  as per equation 6. Non-linearity of  $\Psi_0$  for TiO<sub>2</sub> is due to the large difference between the affinity constants [16]. Fig. 3(b) is the 2nd order derivative of  $\Psi_0$  with respect to pH<sub>B</sub> that shows the transition of reactive sites over the oxide from protonation to deprotonation clarifying the  $pH_{pzc}$  value. Fig. 3(c) shows the total capacitance  $(C_T)$  that consists of C<sub>stern</sub>, double-layer capacitance (C<sub>DL</sub>) and intrinsic AA capacitance (C<sub>AA</sub>), where all of them are connected in series with each other as shown below.

$$\frac{1}{C_T} = \frac{1}{C_{stern}} + \frac{1}{C_{DL}} + \frac{1}{C_{AA}}$$
(10)

TiO<sub>2</sub> shows better resolution for ion-sensing with higher total capacitance ( $C_T$ ) as compared to the other two oxides, SiO<sub>2</sub> and HfO<sub>2</sub>. As the last step, we consider SiO<sub>2</sub> and TiO<sub>2</sub> as an oxide surface for the immobilization of the three different AAs (R, D and P). Two variants of each AA are considered - C-Imm. and N-Imm. (carboxylic or amine terminal is immobilized respectively). The full surface coverage (no active silanol active groups on the surface) by the AA is considered for simulation. Fig. 4(a) and 5(a) show the depletion width (W<sub>D</sub>) variation with respect to pH<sub>B</sub> for different AAs immobilized over the SiO<sub>2</sub> and TiO<sub>2</sub> as gate oxide respectively. Considering the FET to be completely depleted at the lowest pH, a shift in potential towards negative decreases the depletion width. With the same depletion width at the lowest pH, the higher permittivity of TiO<sub>2</sub> amplified the effect of  $\Psi_0$  to further decrease the depletion width as compared to the SiO<sub>2</sub>. Fig. 4(b) and 5(b) represent the drain current (I<sub>SD</sub>) variation with respect to the pH<sub>B</sub> for SiO<sub>2</sub> and



**Fig. 2.** (a) Zeta potential  $(\Psi_{\xi})$ , (b) sensitivity factor ( $\alpha$ ) and (c) intrinsic buffer capacity ( $\beta$ ) variation in relation to pH for three different gate oxide materials.

#### Table 1

Values of  $pH_{pzc(sim)}$  for the three different oxides obtained from simulation results vs points of zero charge taken from literature for the three oxides ( $pH_{pzc}$  (<sub>lit</sub>)).

Gate Oxide	pH <sub>pzc(sim)</sub>	pH <sub>pzc(lit)</sub>
SiO <sub>2</sub>	2	2 [7]
HfO <sub>2</sub>	7.65	7.4 [16]
TiO <sub>2</sub>	6.33	6.7 [17]

TiO<sub>2</sub> respectively. As the depletion width increases in a pMOS, the I<sub>SD</sub> decreases and vice versa. From both graphs, it can be observed that the TiO<sub>2</sub> helped in distinguishing the difference in current to identify the AAs. This can be attributed to the dissociation constants and the oxide capacitance. Figs. 4(c) and 5(c) show the BioFET behavior with immobilized AAs in terms of  $\frac{\partial^2 I_{SD}}{\partial Dt^2}$ . Indeed, each AA shows very unique behavior

of the isoelectric and inflection points which can be used as unique fingerprints to identify each AA. C-Imm. R and P exhibit positive charges on the oxide surface across the pH range due to the presence of only amine sites. The positive charge acquired by C-Imm. R is more than C-Imm. P due to an extra amine sidechain. C-Imm. D is more balanced due to the presence of C-terminal sidechain and N-terminal with an isoelectric point at pH = 6.625. N-Imm. D and P result in negative charges over the surface across the pH range due to the presence of only carboxylic sites with an extra sidechain in N-Imm. D. whereas, N-Imm. R is balanced with an isoelectric point at pH = 7.325 similar to the C-Imm. D.

Figs. 4(c) and 5(c) represent the 2nd order derivative of  $I_{SD} \left(\frac{\partial^2 I_{SD}}{\partial p H^2}\right)$  for SiO<sub>2</sub> and TiO<sub>2</sub> respectively in presence of AA which varies depending on the affinities of reactive sites. Different site-binding relations are used to calculate the  $\Psi_0$ . The affinity of the remaining amine/ carboxyl sites and sidechains is responsible for the distinct fingerprints of the AAs in the



Fig. 3. (a) Surface potential ( $\Psi_0$ ), (b) 2nd order derivative of  $\Psi_0\left(\frac{\partial^2\Psi_0}{\partial bH^2}\right)$  and (c) total capacitance ( $C_T$ ) vs pH. Graphs shown three different gate oxide materials.



**Fig. 4.** (a) Depletion width ( $W_D$ ) (b) drain current ( $I_{SD}$ ) and (c) 2nd order derivative of  $I_{SD} \left( \frac{\partial^2 I_{SD}}{\partial p H^2} \right)$  vs pH. Graphs shown are for Arginine (R), Aspartic Acid (D) & Proline (P) amino acids considering SiO<sub>2</sub> as a gate oxide with carboxyl group immobilised (solid line: C-Imm) and amine group immobilised (dash line: N-Imm).



**Fig. 5.** (a) Depletion width ( $W_D$ ) (b) drain current ( $I_{SD}$ ) and (c) 2nd order derivative of  $I_{SD} \left( \frac{\partial^2 I_{SD}}{\partial p H^2} \right)$  vs pH. Graphs shown are for Arginine (R), Aspartic Acid (D) & Proline (P) amino acids considering TiO<sub>2</sub> as a gate oxide with carboxyl group immobilised (solid line: C-Imm) and amine group immobilised (dash line: N-Imm).

form of I<sub>SD</sub> and  $\frac{\partial^2 I_{SD}}{\partial p H^2}$  which follows the  $\Psi_0$  of the corresponding AAs. TiO<sub>2</sub> helped in generating more distinct fingerprints in terms of inflection points while maintaining the same zero-crossover points as compared to the SiO<sub>2</sub>.

#### 4. Conclusion

This paper presents a comprehensive analytical model that allows us to uniquely identify the signatures of different amino acids. The main parameters that are used to distinguish between the AA are the 2nd order derivative of drain current  $I_{SD}\left(\frac{\partial^2 I_{SD}}{\partial p H^2}\right)$  and the surface potential  $\Psi_0\left(\frac{\partial^2 \Psi_0}{\partial p H^2}\right)$ . Hence, our analytical model indeed proves the possibility of using FET-based sensors to model proteomics. More importantly, this model can provide additional information while calibrating and benchmarking the simulated results with the experimental data.

# **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

# Data availability

Data will be made available on request.

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