Supporting Information: Graphene BioFET sensors for SARS-CoV-2 detection: A multiscale approach

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SUPPORTING NOTE 1: SELF-CONSISTENT GRAPHENE-BASED BIOFET SENSOR SIMULATION

The numerical description of the Graphene BioFET is based on the solution of the Poisson equation in combination with the continuity equation for electrons and holes along the semiconductor channel. A self-consistent scheme is implemented to solve this set of equations in a two-dimensional cross-section of the structure that allows us to evaluate the charge distribution in their different regions.

In the graphene channel the continuity equation is solved in the diffusive regime assuming a common Fermi level ($E_F$) for both electrons and holes\textsuperscript{1–3}. $E_F$ jointly with the DoS are used to evaluate the electron and hole density along the graphene channel and then the source-drain current according to the expression:

$$I_{DS} = q (n\mu_n + p\mu_p) \nabla V_{E_F}$$

where $q$ is the elementary charge, $V_{E_F}$ is the Fermi level potential ($E_F = -qV_{E_F}$), $n$ ($p$) is the electron (hole) density and $\mu_n$ ($\mu_p$) the electron (hole) mobility. The latter parameter includes longitudinal-electric field ($|E_x|$) dependencies and velocity saturation ($v_{sat}$) effects\textsuperscript{4}:

$$\mu = \frac{\mu_0}{\left[1 + \left(|E_x| \frac{\mu_0}{v_{sat}}\right)^\beta\right]^{1/\beta}}$$

For the electrolyte region we solve the modified Poisson-Boltzmann equation\textsuperscript{5} that includes steric effects (a limitation in the concentration due to the finite size of ions) in the distribution of the $i$-th ion:

$$c_i = c_{i,0} e^{qz_i(V-V_{ref})/(k_BT)} e^{-V_{PMF,i}}$$

where, $c_{i,0}$ is the bulk concentration, $c_{\text{max},i}$ the maximum allowed concentration, $z_i$ the ion valence and $V_{PMF,i}$ the Potentials of Mean Force profile. The latter is an additional element that includes the interactions at the electrolyte-solid interface. The value of these profiles are extracted from references\textsuperscript{6,7} where Molecular Dynamics simulations were carried out to evaluate the interactions for different types of ions. As a result, they provide a set of distance-to-surface dependent profiles. These came along with a variable water permittivity $\varepsilon_w$ profile associated to a position dependent water density. In order to prevent unrealistic hydrogen and hydroxyl ion concentrations, PMF profiles for these ions are considered in accordance with the aforementioned water density profiles. We define exponential PMF profiles that result in a negligible concentration below the position of the Gibbs dividing surface reported by\textsuperscript{6,7}.

In addition to the potential dependent model for the distribution of ions, we also include the reactions involving the components of the Phosphate Buffer (PBS) following\textsuperscript{8}. This is only considered for the elements that constitute

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the PBS and is included as an additive component. The reactions involved in the pH regulation are the following ones:

\[
H_3PO_4 \overset{pK'_{a_1,T}}{\rightleftharpoons} H^+ + H_2PO_4^- \tag{4}
\]

\[
H_2PO_4^- \overset{pK'_{a_2,T}}{\rightleftharpoons} H^+ + HPO_2^{2-} \tag{5}
\]

\[
HPO_2^{2-} \overset{pK'_{a_3,T}}{\rightleftharpoons} H^+ + PO_3^{3-} \tag{6}
\]

where \( pK'_{a_i,T} \) are the temperature-dependent reaction constants. These values depend on the ionic strength \( I \) that is evaluated locally (in each point of the grid). So that, a local value of the reaction constants is obtained using the expressions:

\[
\begin{align*}
\left\{ \begin{array}{l}
pK'_{a_i,T} = pK_{a_i,T} + (2z_{a_i} - 1) \left[ \frac{A\sqrt{T}}{1 + \sqrt{T}} - 0.1I \right] \\
pK_{a_i,T} = pK_{a_i} + \frac{dpK_{a_i}}{dT}(T - 298.15K)
\end{array} \right.
\tag{7}
\end{align*}
\]

where \( A \) is a temperature-dependent constant, \( z_{a_i} \), the charge of the conjugate acid species and \( pK_{a_i} \) is the reaction constant at \( T = 298.15K \). The concentration of \( H_2PO_4^- \), \( HPO_2^{2-} \) and \( PO_3^{3-} \) can be calculated from the reactions described by expressions (4) to (6):

\[
\begin{align*}
[H_3PO_4] &= [H_3PO_4] 10^{pH-pK'_{a_1,T}} \tag{8} \\
[HPO_2^{2-}] &= [H_2PO_4^-] 10^{pH-pK'_{a_2,T}} \tag{9} \\
[PO_3^{3-}] &= [HPO_2^{2-}] 10^{pH-pK'_{a_3,T}} \tag{10}
\end{align*}
\]

The concentration \([H_3PO_4]\) is the only value missed in this calculus, however, the sum of all the components of these reactions (\([H_3PO_4]\), \([H_2PO_4^-]\), \([HPO_2^{2-}]\) and \([PO_3^{3-}]\)) has to be equal to the concentration of PBS.
### Table I. Values used in the evaluation of $pK_{a,i,T}$, which are extracted from\(^8\).

| Parameter       | Value  | Units |
|-----------------|--------|-------|
| $pK_{a_1}$ ($25^\circ$C) | 2.15   |       |
| $pK_{a_2}$ ($25^\circ$C) | 7.21   |       |
| $pK_{a_3}$ ($25^\circ$C) | 12.33  |       |
| $dpK_{a_1}/dT$ | 0.0044 | $K^{-1}$ |
| $dpK_{a_2}/dT$ | -0.0028 | $K^{-1}$ |
| $dpK_{a_3}/dT$ | -0.028  | $K^{-1}$ |
| $z_{a_1}$       | 0      |       |
| $z_{a_2}$       | -1     |       |
| $z_{a_3}$       | -2     |       |
| $A$ ($25^\circ$C) | 0.5114 |       |

\[ [H_3PO_4] + [H_2PO_4^-] + [HPO_2^-] + [PO_3^-] = [PBS] \] (11)

Therefore, the concentration of $H_3PO_4$ can be calculated as

\[ [H_3PO_4] = \frac{[PBS]}{1 + 10^{pH-pK'_{a_1,T}} \left( 1 + 10^{pH-pK'_{a_2,T}} \left( 1 + 10^{pH-pK'_{a_3,T}} \right) \right)} \] (12)

As we considered a PBS based on NaH$_2$PO$_4$ there is also a contribution to the sodium ion concentration:

\[ [N_a^+] = -z_{H_2PO_4^-} [H_2PO_4^-] - z_{HPO_2^-} [HPO_2^-] - z_{PO_3^-} [PO_3^-] \] (13)

All the parameters required for the calculation of the reaction constants $pK'_{a,i,T}$ are extracted from\(^8\) and summarized in Table I. The electrolyte considered is a 1×PBS the composition of which is $[\text{NaCl}] = 140\text{mM}$, $[\text{KCl}] = 2.7\text{mM}$ and $[\text{NaH}_2\text{PO}_4] = 10\text{mM}$.

In addition to the charge associated to ions in the solution, we need to include the charge provided by the receptor molecules. We described the protein structures in atomic detail and assigned a partial charge to each atom using the PDB2PQR protocol\(^9\). This protocol includes protonation of the protein structures in order to optimize the hydrogen-bond network based on an empirical pKa estimation for titratable residues. The charges were assigned using the default PARSE force field parameters\(^10,11\). This task is performed in a three-dimensional (3D) description, so we need to translate this information to a 2D description in order to integrate it into the device level simulation. Figure S.3 schematically describes the procedure. We start with the 3D molecular charge profile and project it into a plane that is defined by the desired orientation for the molecule. This plane is discretized in agreement with the grid that will be later used in the device simulation so to achieve a straightforward insertion of the molecular charge.

**Supporting Note 2: Minority Carrier Concentration Under Receptors**

Figure S.4 depicts the minority carrier concentration corresponding to two gate biases ($V_{FG}$), electrons in the p-branch (-0.5V) and holes in the n-branch (0.5V). The behavior of these profiles mimics that of the main carriers, analyzed in the main text. Beginning with electrons (Figure S.4a), they show a tightening of the region where their concentration decreases and, as a consequence, a more steep profile. In the case of holes (Figure S.4b), they depict a quite large increase (around ×3) when compared with the profile obtained for the p-branch (around ×1.5), although the magnitude of the charge in this latter scenario is much greater.
FIG. S.3. Projection of the 3D molecule charge distribution to obtain the 2D profiles used in the device simulation. Vector \( \hat{n} \) sets the desired orientation for the projection plane that is later discretized to define the 2D charge profiles of the molecules. The extracted profiles are used in the device simulations by replicating them along the longitudinal axis according to the defined positions.

\[ V_{FG} = -0.5 \text{V} \]

FIG. S.4. Superimposed minority carrier concentrations, that is, electrons in the p-branch (left) and holes in the n-branch (right), under each receptor. We consider a region of length \( l_0 \) under each receptor and \( \alpha = 0.6 \) (6 receptors are activated) to observe the changes in the carrier concentrations when the receptor is activated. The profiles behave in the same way as those for the majority carriers does, electron concentration drops under the molecule while holes concentration increases. The latter is the one with the most remarkable behavior as hole concentration in the n-branch changes in a larger extent than in the p-branch.

**SUPPORTING NOTE 3: MOLECULE - ELECTROLYTE MODEL VALIDATION**

Aiming at the validation of our numerical approach, we have considered the work by Lud *et al.*\(^{12}\) which addresses the detection of peptides and proteins by a thin-film resistor. More specifically, we have focused on the results given for the detection of aspartic acid. The fabricated structure is quite large (80 \( \mu \text{m} \times 80 \mu \text{m} \)), but, in contrast to most of references, their measurements are based on the surface potential which enables the analysis in a reduced structure (as long as the aspect ratio is kept) making it suitable to be reproduced and thus employed to validate the calculations at the molecule model level. To this end, the experimental structure considered for the simulations is depicted in Figure S.5a.

Figure S.5a shows the simulated structure, reproducing the experimental realization: a 30 nm-thick Si layer sandwiched in between two SiO\(_2\) layers, a 20 nm-thick substrate and a 2 nm-thick cover. The latter is coated by the 1.5 nm-thick lipid layer that hosts the NTA headgroups that act as receptors and has a -1\( \text{q} \) charge.

We first computed the profile of the surface potential as a function of the number of aspartic acid units considered in the histidine-aspartic acid complex attached to the receptors, and compared it with the experimental results provided by Lud *et al.* This profile is obtained for two Debye lengths \( \kappa^{-1} \) of the electrolyte, assuming 0.1xPBS solution with different KCl concentrations. For the modelling of the complex, histidine tag is considered electrically neutral, while each aspartic acid unit has a net charge of -1\( \text{q} \). Their sizes are set according to the data provided in\(^{12,13}\) for these two Debye lengths. The results depicted in Figure S.5b \((\kappa^{-1} = 1.1 \text{ nm})\) and Figure S.5c \((\kappa^{-1} = 0.7 \text{ nm})\), demonstrate that numerical simulations reproduce to a very good agreement the behaviour of the experimental data provided by\(^{12}\). The trend of the surface potential as a function of the number of aspartic...
acids units is accurately replicated by the simulations for the two Debye lengths (ion concentrations) considered.

The validation is extended to fit the values provided in the same work for the Green Fluorescent Protein (GFP). The same structure as in Figure S.5a but changing the molecule parameters to fit those for the GFP molecule: a 4 nm-height and 3 nm-width block with a net charge of -8\( q \). The data provided for this case corresponds to the change in the surface potential as a function of the protein concentration in the sample. This requires an additional step to translate the protein concentration \( c_{\text{GFP}} \) into number of activated receptor, which is the value considered in the simulations. This is done using the association constant extracted from the Langmuir isotherm fitting carried out in the paper:

\[
N_{\text{act}} = N_{\text{rec}} \frac{K_d c_{\text{GFP}}}{1 + K_d c_{\text{GFP}}} \Rightarrow c_{\text{GFP}} = \frac{1}{K_d} \frac{1}{\frac{N_{\text{act}}}{N_{\text{rec}}} - 1}
\]

(14)

where \( N_{\text{rec}} \) is the total number of receptor sites and \( N_{\text{act}} \) the number of activated sites. These two expressions make possible to translate between \( N_{\text{act}} \) and \( c_{\text{GFP}} \) using the \( K_d \) provided (6.5\( \cdot 10^5 \) M\(^{-1} \)) and the number of receptor sites in the simulation (\( N_{\text{rec}} = 10 \)). The result of these simulations are depicted in Figure S.6.

It is worth to highlight that this validation, at the molecule-charge and sensor-surface potential levels, is the most relevant to demonstrate the capability of the computational approach proposed here, i.e. the possibility to treat, in a multiscale fashion, the details about the molecular properties into a complete device level study. This validation is, indeed, complementary to those others of the electrolyte ions interaction\(^{14} \) and the semiconductor\(^{15} \) that we already accomplished in less sophisticated versions of the implemented tool and with other purposes.
FIG. S.6. Change of the surface potential as a function of the protein concentration in the sample. The structure considered is the same as the one depicted in Figure S.5a but using the Green Fluorescent Protein (GFP) instead of aspartic acid, which depicts a higher net charge (-8q) and size (4 nm×3 nm).

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