


 Cite this: *Sens. Diagn.*, 2024, 3, 1935

## Challenges in aptamer-based sensor development using carbon nanotube networks†

 Laura Ferrer Pascual, <sup>a</sup> Eero Gustafsson, <sup>a</sup> Juha Siitonen, <sup>b</sup> Vasuki Durairaj <sup>c</sup> and Tomi Laurila <sup>\*ab</sup>

Electrochemical aptamer-based (EAB) sensors represent a promising biosensing platform, leveraging the selectivity of aptamers and the advantages of electrochemical methods. These sensors offer high sensitivity, rapid response, low limits of detection, cost-effectiveness, and miniaturization potential. While gold electrodes have been predominantly used in EAB sensors, alternatives such as carbon nanotubes (CNTs) are gaining attention. CNTs offer advantages like large surface area and conductivity but pose challenges due to their reactivity and 3D network structure. In this study, we explore the development of EAB sensors using single-wall carbon nanotube (SWCNT) networks, emphasizing on the challenges and electroanalytical insights. Three key electrochemical parameters are proposed for assessing EAB sensor performance: (i) variations in peak current, (ii) shifts in peak position, and (iii) the restoration of the background current. Focusing solely on peak current changes can be misleading, as factors like aptamer surface depletion can influence it. Additionally, both partial and integrated currents should be monitored in square wave voltammetry (SWV) analysis, considering both ON and OFF behaviours across frequencies. This comprehensive approach provides a preliminary assessment of successful binding and surface passivation in EAB sensors when combined with surface analytical techniques such as surface plasmon resonance (SPR) measurements.

 Received 9th July 2024,  
 Accepted 26th September 2024

DOI: 10.1039/d4sd00250d

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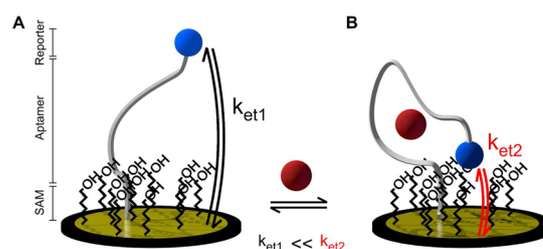
### 1. Introduction

Electrochemical aptamer-based (EAB) sensors offer significant promise in the field of biosensing. EAB sensors combine the high specificity of aptamers with the high sensitivity, rapid response, low limits of detection, low cost, and miniaturization of electrochemical methods.<sup>1,2</sup> In addition, EAB sensors are promising for multi-drug screening because their aptamer sequence can be customised for different target molecules and their signal arises from the binding of the target instead of its chemical reactivity.<sup>3</sup>

Aptamers are short nucleic acid strands with high selectivity towards their target arising from systematic evolution. The aptamer refolds when binding to the target, changing its three-dimensional conformation. The folding of the strand changes the distance between a redox reporter attached to the aptamer and the electrode surface, producing

a change in the electron transfer kinetics (Fig. 1). The signal is classified as “signal on” or “signal off” depending on whether the binding-induced conformational changes moves the redox label closer to or further away from the electrode surface, respectively.<sup>4</sup>

As of date, most electrochemical aptamer-based studies are focused on the use of gold electrodes as sensing material due to the widely known gold-alkanethiol chemistry.<sup>6,7</sup> Despite their potential, gold-based systems face a limitation in their operational lifespan due to the inherent instability of the weak thiol-bond. This instability hinders the continuous application of voltage during measurements, as it can lead to the desorption of the aptamer-SAM monolayer.<sup>8</sup> In addition, gold



**Fig. 1** Working principle of an ideal electrochemical aptamer-based sensor in A) the absence and B) presence of target.<sup>5</sup>

<sup>a</sup> Department of Electrical Engineering and Automation, School of Electrical Engineering, Aalto University, PO Box 13500, 00076 Aalto, Finland.

E-mail: [tomi.laurila@aalto.fi](mailto:tomi.laurila@aalto.fi)

<sup>b</sup> Department of Chemistry and Materials Science, School of Chemical Engineering, Aalto University, PO Box 16200, 00076 Aalto, Finland

<sup>c</sup> Sustainable Products and Materials, VTT Technical Research Centre of Finland, P. O. Box 1000, FI-02044 Espoo, Finland

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4sd00250d>



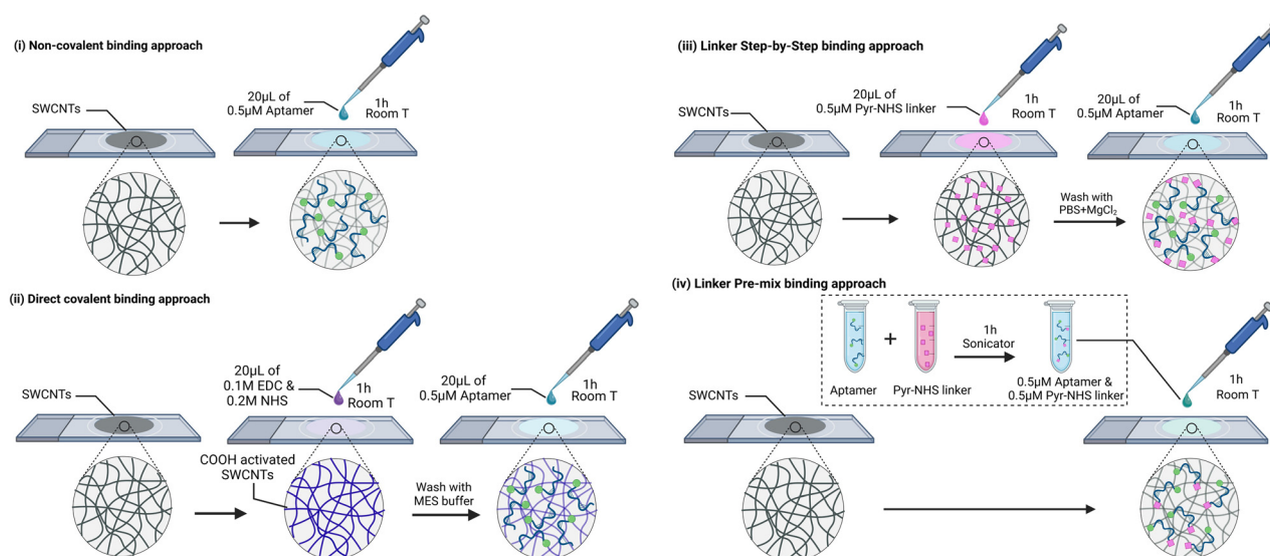
presents a high cost compared to other available sensing materials such as carbon nanotubes (CNTs). There have been a few studies investigating alternative materials such as glassy carbon,<sup>1,9,10</sup> silicon,<sup>11</sup> or graphene<sup>12</sup> for EAB sensor development. Carbon nanotubes are often considered promising electrode materials for electrochemical sensing due to their large surface area, high conductivity, good sensitivity, wide potential windows, and fast response times.<sup>13</sup> While the elevated reactive surface area of CNTs is typically advantageous, it also results in increased non-specific interactions, which are unfavourable for aptamer binding. Moreover, the 3D structure of CNTs promotes volume interactions over surface interactions, hindering the uniform orientation of aptamers during immobilization. Lastly, the typically advantageous flexibility in the functionalisation of carbon surfaces proves disadvantageous for EAB sensors, as it leads to surfaces with large heterogeneity.<sup>14,15</sup>

Recently, there has been increasing scrutiny regarding the practical functionality of electrochemical aptamer sensors, even when deployed on traditional gold (Au) surfaces. In particular, the oversimplified depiction of their operation, as illustrated in Fig. 1, has faced criticism.<sup>16</sup> As mentioned earlier, substituting carbon-based materials for Au in these sensors amplifies the contribution of non-specific interactions, thereby adding complexity to the underlying challenges. In this context, we present the findings of our recent investigations aimed at developing electrochemical aptamer-based sensors integrated with single-wall carbon nanotube (SWCNT) networks.

Our system under investigation involves SWCNT network electrodes on a glass support, onto which aptamers specific to vancomycin have been immobilized using four distinct strategies. The first strategy is centred on the direct

immobilization of aptamers through non-specific interactions with the SWCNTs. The second strategy involves the covalent binding of the amine terminal group of the aptamers to the carboxylic groups of SWCNTs *via* EDC/NHS coupling. The last strategy employs a linker molecule (1-pyrenebutyric acid *N*-hydroxysuccinimide ester, Pyr-NHS) with two different protocols. In one protocol, termed the 'layer-by-layer (LbL)' approach, the linker is first deposited, followed by the drop-casting of the aptamer. The alternative protocol involves pre-mixing the aptamers and the linker molecule before drop-casting the solution onto the SWCNTs. In summary, these four protocols are denoted as (i) direct non-covalent, (ii) direct covalent, (iii) linker LbL, and (iv) linker pre-mixed, respectively (Fig. 2). Detailed descriptions of these immobilization protocols, along with the materials and methods used, are provided in the experimental section. Vancomycin (VA) is selected as the target molecule because it is classified as one of the critical antibiotics for human medicine by the World Health Organization (WHO). It serves as the last resort in treating resistant MRSA and holds the top position on the US Center for Disease Control's list of antibiotics associated with high costs in treating antibiotic-resistant infections.<sup>17</sup> Our goal is to shed light on the challenges that arise when transitioning from flat Au electrodes to the three-dimensional world of SWCNT networks. Furthermore, we aim to provide readers with valuable electroanalytical insights into the system, aiding in the assessment of successful aptamer binding to the electrode. We argue that to evaluate the electroanalytical performance of an EAB sensor, it is crucial to consider three key parameters:

- (i) Variations in the measured peak current,
- (ii) Shifts in the peak position, and
- (iii) The restoration of the original background current and peak position after the target measurement.



**Fig. 2** Aptamer binding approaches of this work: (i) non-covalent immobilization through aptamer adsorption to SWCNTs, (ii) direct-covalent immobilization through amide bond formation, (iii) indirect immobilization through linker LbL method, and (iv) indirect immobilization through linker pre-mix method.



At present, the assessment of EAB sensors relies exclusively on changes in peak current—a criterion that may be prone to misinterpretation, given that alterations in peak current can stem from factors such as the gradual depletion of aptamers from the electrode surface. Furthermore, when employing the commonly used Square-Wave Voltammetry (SWV) technique, it is crucial to monitor the three key parameters defined above using both partial and integrated currents. Focusing solely on the latter can obscure vital information regarding peak position and peak current. Additionally, it is imperative to examine both the ON and OFF behaviours as a function of frequency. This comprehensive analysis is essential to confidently affirm the successful binding and surface passivation of the sensor. Alongside electrochemical investigations, physicochemical characterizations are essential to substantiate our arguments. However, it is crucial to recognize the limitations of commonly used techniques like Infrared spectroscopy, Surface Plasmon Resonance (SPR), and atomic force microscopy, especially when applied to physically and chemically complex 3D electrode structures, as demonstrated in this manuscript. In this study, SPR measurements are conducted to comprehend adsorption interactions at the SWCNT networks and elucidate the efficacy of our different aptamer binding protocols. Nonetheless, deriving quantitative conclusions about the efficacy of aptamer binding to the electrode surface or their target response requires consideration of the three aforementioned electrochemical response parameters, as discussed in the following manuscript.

To clarify the primary focus of our investigation, it is important to delineate the scope of this study. This manuscript focuses on the fundamental challenges encountered in the development of aptamer-based sensors, emphasizing conditions relevant to physiological environments, with all measurements conducted in phosphate-buffered saline at pH 7.4. The inherent selectivity of aptamers, combined with the passivation of the sensor's surface with the linker molecules, is anticipated to reduce interference from electrochemically active inner-sphere probes (ISR). While studies on pH variations and interferents are beyond the scope of this work, these will be explored in a forthcoming study focused on sensor performance optimization.

## 2. Experimental

Sulfo-NHS (*N*-hydroxysulfosuccinimide), EDC (*N*-(3-dimethylamino-propyl)-*N*'-ethylcarbodiimide hydrochloride), 0.1 M MES buffer (2-(*N*-morpholino)ethanesulfonic acid, 0.9% NaCl, pH 4), and vancomycin hydrochloride were purchased from Thermo Fisher. Magnesium chloride (MgCl<sub>2</sub>), 1× phosphate buffered saline (PBS), Atto MB2 methylene blue derivative, Pyr-NHS linker (1-pyrenebutyric acid *N*-hydroxysuccinimide ester) was purchased from Sigma Aldrich. Potassium chloride (KCl) was purchased from Merck Suprapur. The modified aptamer was purchased from Integrated DNA Technologies. The aptamer sequence: 5'-CG AGG GTA CCG CAA TAG TAC TTA TTT GCC TAT TGT GGG TCG G-3' with modification on the 5' with an

amino group and the redox reporter methylene blue (ATTO-MB2) on the 3' end. The cartoon model (made by M-Fold) is shown in the ESI† (Fig. S11).

### 2.1 Electrode fabrication

The single-wall carbon nanotubes (SWCNTs) were provided by Canatu Oy. Briefly, the SWCNT networks were fabricated in a laminar flow reactor by chemical vapor deposition and were collected using a membrane filter. The SWCNTs fabrication protocol is described in detail in.<sup>18</sup> The electrode fabrication consisted of press-transferring the SWCNTs onto a glass substrate where the network was then densified by adding 99.7% ethanol and letting it evaporate in air. Electrical contact was made by adding conductive silver paste and letting it dry for a minimum of 30 min. Then a conductive copper tape was used to contact the silver paste to the copper slide. PTFE-tape was used to insulate the electrode from other regions except a 3 mm hole.

### 2.2 Aptamer binding methods

Four different EAB sensors were fabricated based on the aptamer binding method (Fig. 2): (i) in the non-covalent binding approach, the aptamer was attached to the CNT surface due to the adsorption of the DNA strand and methylene blue. To do so, 20 μL of 0.5 μM aptamer solution in PBS + 2 mM MgCl<sub>2</sub> was drop-casted onto the electrodes for one hour at room temperature and then cleaned with PBS + 2 mM MgCl<sub>2</sub>. (ii) In the direct covalent binding approach, 20 μL of a solution of 0.1 M EDC and 0.2 M NHS in MES buffer was drop-casted to the electrode for 1 h at room temperature. Then, the electrode was cleaned with MES buffer and immediately 20 μL of 0.5 μM aptamer solution in PBS + 2 mM MgCl<sub>2</sub> was drop-casted for another hour at room temperature. Finally, the electrode was cleaned with PBS + 2 mM MgCl<sub>2</sub>. The EDC/NHS concentration and ratio was optimised based on the largest peak shift upon VA binding (Table S11†). For the linker approach, two different protocols were used: a so-called “layer-by-layer (LbL)” functionalization and a “pre-mix” functionalization. (iii) The “LbL” functionalization consisted in first drop-casting 20 μL of 0.5 μM Pyr-NHS linker to the electrode for 1 h at room temperature. Then, the electrode was cleaned with PBS + 2 mM MgCl<sub>2</sub> and 20 μL of 0.5 μM aptamer solution in PBS + 2 mM MgCl<sub>2</sub> was drop-casted for 1 h at room temperature. Finally, the electrode was cleaned with PBS + 2 mM MgCl<sub>2</sub>. (iv) The “pre-mix” functionalization consisted of bath sonication for 1 h a solution containing 0.5 μM of aptamer and 0.5 μM of linker. The temperature of the bath increased to 44 °C. Then, 20 μL of the solution was drop-casted to the electrodes for 1 h and then cleaned with PBS + 2 mM MgCl<sub>2</sub>.

### 2.3 Washing methods

After the electrode functionalization, the electrodes underwent washing using two different protocols to remove most of the loosely attached aptamers. One method involved immersing the electrodes in a 500 μM vancomycin bath solution in PBS with 2



mM MgCl<sub>2</sub> for 30 minutes. The second method consisted of repetitive washings in a 6 M GuHCl solution: 40 μL of GuHCl solution was drop-casted onto each electrode for 15 minutes, followed by immersion in PBS with 2 mM MgCl<sub>2</sub>. This procedure was repeated two additional times. Following the washing protocols, the electrodes were placed in PBS with 2 mM MgCl<sub>2</sub> before the SWV measurements.

## 2.4 Surface plasmon resonance

The SWCNT networks were press-transferred onto pre-cleaned (UV-ozone treated) Au-coated SPR sensors (BioNavis Ltd.) and densified with 99.7% ethanol, followed by air drying. The SPR measurements were carried out using a multi parametric surface plasmon resonance instrument (MP-SPR model Navi 200, BioNavis Ltd.). All CNT modified Au sensors were first stabilised in the measurement buffer for 2 hours following which the adsorption studies were carried out from different test solutions at 10 μl min<sup>-1</sup> flow rate. Following each adsorption step, the sensors were rinsed with the measurement buffer solution at 10 μl min<sup>-1</sup> flow rate for 25 min.

## 2.5 Electrochemical techniques

Electrochemical investigations were carried out using both conventional Cyclic Voltammetry (CV) and Square-Wave Voltammetry (SWV). Experiments were conducted with a Gamry Reference 600 + potentiostat. A three-electrode cell was used with an Ag/AgCl reference electrode (Radiometer Analytical) and a Pt wire (Goodfellow) counter electrode. The solutions were purged with N<sub>2</sub> for 30 min before the experiments. All the measurements were carried out at room temperature in a Faraday cage.

## 2.6 Measurement protocols

All measurements were performed using four different sensor types, with three nominally identical electrodes per type ( $N = 3$ ). For each sensor type, the electrodes measured the same solution, representing technical repeats. This setup allowed us to assess the precision and consistency of the biosensor system across replicates.

**2.6.1 Frequency study and effect of various linker chemistry protocol.** SWV was used with frequencies ranging from 10 to 2000 Hz with a 50 mV pulse size. Before the measurement, the electrodes were washed with 6 M GuHCl. Then, the electrodes were first measured in background (BG) which consisted of a solution of PBS + 2 mM MgCl<sub>2</sub>. Then, the electrodes were measured with saturating target concentration (500 μM VA buffer solution) and after the measurement they were placed in a beaker containing PBS + 2 mM MgCl<sub>2</sub> solution. Finally, the electrodes were measured again in PBS + 2 mM MgCl<sub>2</sub> (background after, BGA).

**2.6.2 Stability study protocol.** SWV was used from 10 to 2000 Hz with a 50 mV pulse size. Different washing methods were used: no cleaning, VA washing, and GuHCl washing (see Washing methods section). Then, the electrodes were first

measured in background (PBS + 2 mM MgCl<sub>2</sub>) and consequently in saturating target concentration (500 μM VA buffer solution) and finally in background solution again. This procedure was repeated two more times. After each vancomycin measurement, the electrodes were placed in a beaker containing PBS + 2 mM MgCl<sub>2</sub> solution.

**2.6.3 Vancomycin concentration series protocol.** SWV was used from 120 to 180 Hz with a 50 mV pulse size. The electrodes were first measured in background (BG) which was a solution of PBS + 2 mM MgCl<sub>2</sub>. Then, specific volumes of a 500 μM VA in PBS + 2 mM MgCl<sub>2</sub> stock solution were added to the cell to achieve the different VA concentrations: 1, 10, 25, 50, 100 and 250 μM. In between the different VA concentration measurements, the electrodes were placed in a beaker containing PBS + 2 mM MgCl<sub>2</sub> solution. Finally, the electrodes were measured again in PBS + 2 mM MgCl<sub>2</sub> (background after, BGA).

**2.6.4 Passivation protocol.** The electrodes were first measured with CV in background which was a 1 M KCl aqueous solution. Then, the electrodes were measured with CV in a 500 μM ATTO-MB2 solution and after the measurement they were placed in a beaker containing 1 M KCl solution. Finally, the electrodes were measured with CV again in 1 M KCl for 80 cycles.

## 2.7 Data analysis

The partial currents in the SWVs have been potential corrected by the addition of 50 mV to forward currents and subtraction of 50 mV to reverse currents. The “current decrease” and “signal change” are calculated from eqn (1):

$$\text{Signal change (\%)} = \frac{(BG - VA)}{BG} \cdot 100 \quad (1)$$

Signal loss is calculated from eqn (2):

$$\text{Signal loss (\%)} = \frac{(BG - BG_{\text{after}})}{BG} \cdot 100 \quad (2)$$

Where BG and BG<sub>after</sub> are the forward peak current in PBS + 2 mM MgCl<sub>2</sub> before and after measuring vancomycin, respectively. VA is the forward peak current when measuring vancomycin.

Limit of detection (LOD) is calculated with eqn (3):

$$\text{LOD} = \frac{3.3 \cdot \text{SD}_{BG}}{\text{slope}} \quad (3)$$

Where the standard deviation (SD<sub>BG</sub>) is the mean of the five last stabilization SWV currents of three electrodes for each sensor type and the slope is calculated from the calibration curves in Fig. 8.

# 3. Results

## 3.1 Surface plasmon resonance

Au coated SPR sensors were first pre-cleaned through UV-ozone treatment, followed by rinsing with MilliQ water and air drying. SWCNT network was then press-transferred on top of the Au sensor surface and densified by adding a drop of



99.7% ethanol and letting it evaporate in air. Note that this sample preparation was done solely for the purpose of SPR measurements. SPR sensograms obtained using a 670 nm laser wavelength for the different adsorption studies carried out in this work are shown in Fig. 2. The angle at which the SPR effect occurs for a particular wavelength is highly sensitive to the refractive index (measured in refractive index unit – RIU) of the medium that is in contact with the metal surface of the SPR sensor.<sup>19</sup> Adsorption events occurring at the sensor surface thus result in a change in the SPR angle, corresponding to the refractive index change due to the adsorbed layer. The thickness of the adsorbed layer can be determined using the eqn (4):

$$d = \frac{l_d \Delta \text{angle}}{2x(n_a - n_0)} \quad (4)$$

where  $\Delta \text{angle}$  is the change in the SPR angle,  $l_d$  is a characteristic evanescent electromagnetic field decay length (estimated as  $0.37 \times \text{laser wavelength}$ ),  $x$  is a sensitivity factor for the sensor obtained after calibration of the SPR instrument (here  $109.94^\circ \text{ RIU}^{-1}$ ),  $n_0$  is the refractive index of the bulk solution ( $1.334 \text{ RIU}$  for water),  $n_a$  is the refractive index of the adsorbed substance. The mass ( $\Delta m$ ) of the adsorbed layer per unit area can be further estimated from the thickness  $d$  and packing density  $\rho$  of the adsorbed material, using the eqn (5):

$$\Delta m = d \cdot \rho \quad (5)$$

The refractive index  $n_a$  and density  $\rho$  values used for the different materials in this study are tabulated in Table 1, along with the estimated thickness and mass of the adsorbed materials corresponding to the SPR sensograms in Fig. 3. The objectives of this study are to observe the trends in binding of aptamers to the carbon surface by various binding protocols, and their response to vancomycin. Reference measurements indicate that the vancomycin molecules themselves strongly adsorb onto the SWCNT network. Further, all other studied

molecules also show strong adsorption onto the SWCNT network and, in multiple injection studies, the first injection step shows the highest adsorption indicating that the strong non-specific interactions occurring at the pristine carbon surface are indeed the most dominant effect as discussed earlier. Interestingly, the direct covalent activation protocol indicates a strong irreversible adsorption, likely from the EDC by-products.<sup>20</sup> Consequently, the amount of aptamer adsorbed is estimated to be the lowest ( $1.5 \text{ mg m}^{-2}$ ) in this protocol. The linker LbL protocol also indicates a similar, relatively low amount of aptamer ( $2.7 \text{ mg m}^{-2}$ ) adsorption, whereas with the direct non-covalent approach, a considerably higher amount of aptamer ( $18.5 \text{ mg m}^{-2}$ ) is adsorbed onto the carbon surface. Due to the complexity of adsorption mechanisms from the equimolar linker + aptamer pre-mixed solution, it is difficult to estimate quantitatively the amount of adsorbed aptamers alone in this system. Using a simplified numerical average approximation of the linker and aptamer refractive index and density values, the mass of adsorbed linker + aptamer layer is estimated to be  $11.4 \text{ mg m}^{-2}$ .

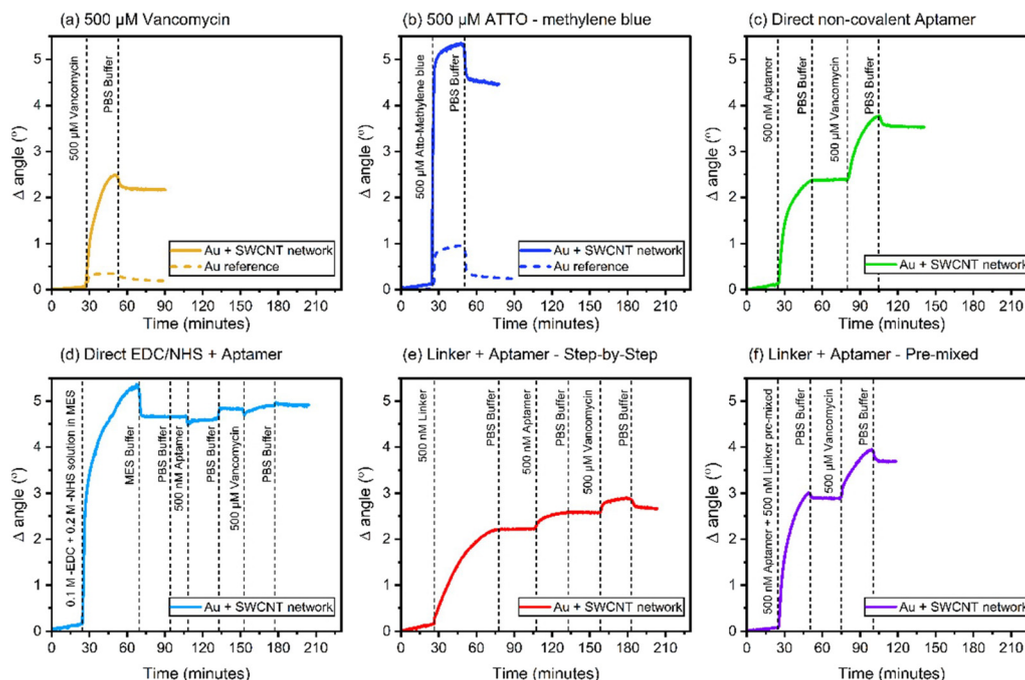
According to this SPR results, the direct covalent and linker LbL systems exhibit similar behaviour in terms of their aptamer adsorption characteristics, hinting at the presence of ordered aptamers on their surfaces. It is crucial to note that these observed adsorption patterns sharply contrast with the other two system types. Both the direct covalent and linker LbL systems display relatively modest levels of both aptamer and vancomycin adsorption, when compared to direct non-covalent and linker pre-mixed approaches which demonstrate approximately ten times greater adsorption of both aptamers and vancomycin. These results suggest that the direct covalent and the linker LbL systems yield more controlled aptamer responses to vancomycin, possibly indicating a degree of ordering in the arrangement of aptamers on their surfaces. Conversely, the alternative systems result in less organized and more random responses. Further, the SPR results suggest that there is some degree of coupling of the aptamers to the SWCNT network especially in the case of direct covalent linking.

**Table 1** Thickness and mass of different adsorbed materials based on SPR study, along with the refractive index and density approximations used for each material

Aptamer binding	Material adsorbed	Refractive index (RI)	Density ( $\text{g cm}^{-3}$ )	Thickness (nm)	Mass ( $\text{mg m}^{-2}$ )
—	Vancomycin <sup>21</sup>	1.735	1.7	6.0	10.3
—	ATTO-MB2 (ref. 22)	1.676	1.2	7.3	8.7
Direct non-covalent	Aptamer <sup>23–25</sup>	1.520	1.3	14.2	18.5
	Vancomycin	1.735	1.7	3.2	5.4
Direct covalent	EDC/NHS <sup>26</sup>	1.528	1.6	45.7	36.5
	Aptamer	1.520	1.3	1.2	1.5
Linker LbL	Vancomycin	1.735	1.7	0.3	0.5
	Linker <sup>27</sup>	1.852	1.2	4.8	5.7
	Aptamer	1.676	1.2	2.0	2.7
Linker pre-mixed	Vancomycin	1.520	1.3	0.2	0.4
	Linker + aptamer <sup>a</sup>	1.686 <sup>a</sup>	1.3 <sup>a</sup>	9.2	11.4
	Vancomycin	1.735	1.7	2.2	3.8

<sup>a</sup> Average values of aptamer and linker refractive index and density have been used here to estimate the adsorption parameters from this equimolar mixture.





**Fig. 3** SPR sensograms showing the adsorption of (a) 500  $\mu\text{M}$  vancomycin, (b) 500  $\mu\text{M}$  ATTO-MB2, (c) 500 nM aptamer via direct non-covalent binding, (d) direct covalent activation followed by 500 nM aptamer, (e) 500 nM linker followed by 500 nM aptamer (LbL approach) and (f) 500 nM linker + 500 nM aptamer – pre-mixed solution. The responses of adsorbed aptamers to buffer rinse and 500  $\mu\text{M}$  vancomycin injection were also followed for each aptamer linking protocol (c–f).

However, due to the limited number of carboxylic groups on the SWCNT network materials (as documented in ref. 28 and 29), it is likely that most of the applied EDC/NHS reacts further, leading to the formation of by-products,<sup>30</sup> which then partially foul the surface (Fig. 3d).

### 3.2 Electrochemical characterizations

In this section, we present a comprehensive analysis of the electrochemical performance of our designed electrochemical aptamer-based sensors. Our investigation focuses on four different aptamer binding methods to evaluate their efficacy and stability. Additionally, we examine the optimal cleaning methods for removing non-specifically bound aptamers, comparing no cleaning, guanidinium hydrochloride (GuHCl) cleaning, and vancomycin cleaning.

The electrochemical characterizations were performed using square wave voltammetry (SWV). In SWV, the current is measured at the end of each potential pulse. This technique allows us to study the current's behaviour based on the relationship between the rate of the electron transfer reaction and the frequency of the square wave pulse. Specifically, the current shows a quasireversible maximum at a critical frequency, which is related to the surface standard rate constant ( $k_{\text{sur}}$ ).<sup>31</sup> By studying the current change at different SWV frequencies, we can approximate the rates of the aptamer's bound and unbound states to a target. This facilitates the identification of regions exhibiting signal ON and OFF behaviours in EAB sensors.<sup>32,33</sup> In addition, the

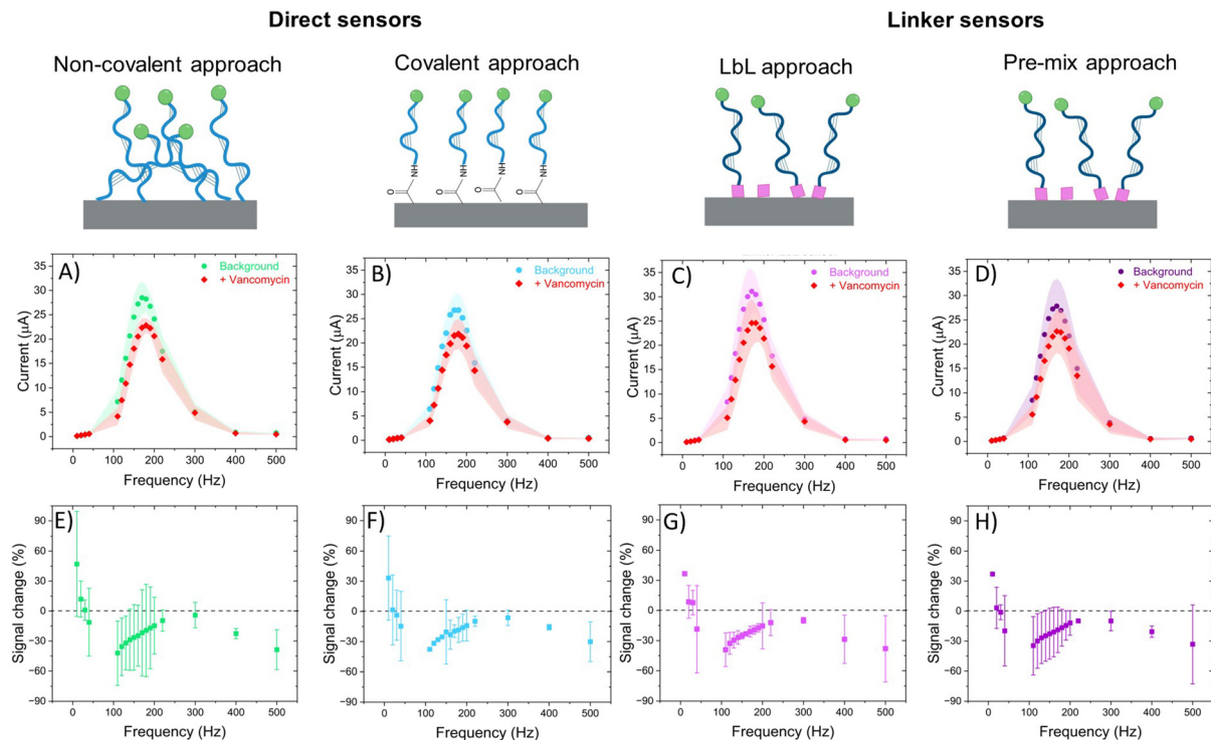
electrochemical reversibility and thus the peak location of a surface bound redox probe in SWV measurements is mainly controlled by the dimensionless kinetic parameter  $\omega = k_{\text{sur}}/f$ , where  $f$  is the measurement frequency.<sup>34–36</sup> When a target is measured with an EAB electrode at a specific frequency, all changes in  $\omega$  can be taken to arise from the changes in  $k_{\text{sur}}$ . If  $k_{\text{sur}}$  decreases and thus  $\omega$  decreases, there will be an anodic shift in the peak position for the oxidation process, while opposite is true in the case of increase in  $k_{\text{sur}}$ . Therefore, with SWV we can also evaluate the nature of the signal changes with EAB sensors.

Through our analysis, we investigate the frequency response of the sensors and evaluate the impact of different linker chemistries on their signal output. We also examine the sensors' stability and repeatability, which are crucial for reliable long-term measurements. Furthermore, we assess the sensors' response to varying concentrations of VA to establish their sensitivity and detection limits. Lastly, we explore strategies for passivating the sensor surface to minimize non-specific binding of aptamers and methylene blue, thereby enhancing the specificity and accuracy of the sensors.

Each aspect of this investigation provides detailed insights into the mechanisms underlying sensor performance and offers guidelines for optimizing EAB sensor design.

**3.2.1 Frequency study and effect of various linker chemistry.** To assess the distinct responses of the four EAB sensor types, we conducted SWV measurements across a broad frequency range (10–2000 Hz) in both the absence and presence of saturating vancomycin concentration in PBS +



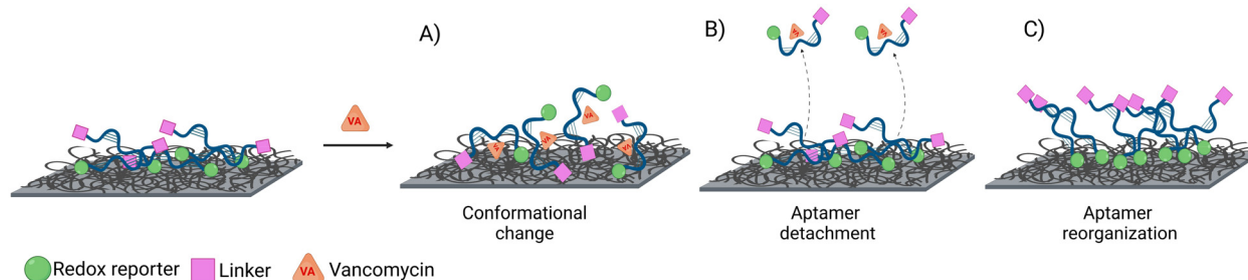


**Fig. 4** Frequency behaviour study of EAB sensors with different aptamer immobilization methods. Direct immobilization was performed with two approaches: non-covalent (A and E) and covalent binding (B and F). The indirect immobilization by the use of linker was performed by two approaches: layer-by-layer (C and G) and premix (D and H) methods. The frequency behaviour was studied with current vs. frequency plots (A–D) in absence (background) and presence (+vancomycin) of target as well as with signal change vs. frequency plots (E–H) ( $N = 3$ ).

$\text{MgCl}_2$  solution. This study aimed to approximate the surface rate constant, identify the optimal interrogation frequency for maximal signal change, and evaluate signal ON/OFF behaviours.

All sensors exhibited a maximum current at a frequency of 170 Hz, irrespective of the presence of the target (Fig. 4A–D), corresponding to a surface rate constant of approximately  $170 \text{ s}^{-1}$  for both bound and unbound states. Despite the similar rate constant for both binding states, the signal change plot reveals frequency-dependent behaviour. Specifically, a signal-ON region is observed from 10 to 30 Hz, while a signal-OFF region is evident from 40 to 2000 Hz. However, this frequency dependent behaviour cannot be

directly associated with conformational changes since unbound and bound states present the same  $k_{\text{sur}}$ . The observed signal-OFF behaviour could be attributed to both a detachment of non-covalently bound aptamers from the electrode surface and the conformational change of the aptamers upon target binding (Fig. 5A and B). However, the former will not produce a consistent peak shift. In our case, all sensor systems exhibited an anodic shift of the forward (oxidation) peak position upon VA addition (Fig. S14<sup>†</sup>), indicating that the reaction becomes kinetically less feasible. In addition, the magnitude of peak position changes depended on the applied SWV frequency. At lower frequencies (40 Hz), the sensors showed larger peak shifts



**Fig. 5** Scheme of the three possible scenarios producing a signal change upon target addition: A) conformational change of the aptamer, B) detachment of the non-covalently bound aptamers, and/or C) reorganization of the non-covalently attached aptamers.



and higher peak position restoration than at medium frequencies (160 Hz).

Our hypothesis for the observed signal-ON behaviour is a possible reorganization of non-covalently bound aptamers upon vancomycin binding (Fig. 5C). This binding event displaces some aptamers from the electrode surface, creating space for others to adsorb to the carbon nanotubes *via* their methylene blue (MB) end. The adsorption of MB induces two effects: an anodic peak shift and a slight increase in background peak current. The shift results from the higher overpotential required to drive the reaction due to the adsorption of the reduced form of MB (the reactant) on the SWCNT surface. However, this peak shift will be of much smaller magnitude than that produced by a conformational change. Simultaneously, the small current increase is attributed to the higher concentration of redox reporters on the electrode surface within the effective electron transfer distance. The SPR studies (Fig. 3) corroborate the pronounced adsorption tendency of the redox probe on the CNT networks, providing further support for our observations. Therefore, it is vital to also monitor the restoration of the background current to its original peak position (prior to vancomycin binding), considering that a comparable anodic shift may result from two vastly distinct physical mechanisms. In the context of conformational changes, a return is anticipated since the aptamer will get reoriented upon vancomycin removal; yet such a return is not observed in the case of the strong adsorption of MB. Among all sensor types, the linker LbL sensor presented the most samples with both anodic peak shift and peak position restoration at 160 Hz.

The signal loss was defined as the change in background peak current (in PBS + MgCl<sub>2</sub>) before and after vancomycin addition. This signal loss was similar across all four sensor types (Table S12<sup>†</sup>) and is likely primarily due to the detachment of aptamers from the electrode surface. Additionally, the similar

signal loss observed in the direct non-covalent sensor compared to the other sensors again suggests strong adsorption of the aptamers to the CNT network, an observation supported by the SPR results.

**3.2.2 Stability and repeatability of the sensors.** As observed in the frequency study, some non-specifically bound aptamers could be randomly oriented, contributing to background noise signals and detaching during sensor use. Due to their weak interaction with the electrode surface, it is likely that these aptamers are located at the outer layers of the electrode surface, which increases their contribution to redox reactions occurring at low frequencies, whereas strongly bound aptamers participate in faster reactions due to their proximity to the electrode surface. Therefore, to reduce background noise signals arising from non-specifically bound aptamers, we evaluated the efficiency of the cleaning method and assessed the stability of the sensors.

First, we studied the effect of 6 M GuHCl solution on non-covalent aptamer sensors and a control group (no cleaning). We used SWV at both low (40 Hz) and high (2000 Hz) frequencies during repetitive measurements of saturating concentrations of VA, monitoring signal loss after each VA measurement. The guanidinium ion (Gu<sup>+</sup>) possesses flat hydrophobic faces that can participate in hydrophobic interactions and directional H-bonding *via* its three NH<sub>2</sub> groups. Therefore, Gu<sup>+</sup> has been shown to denature protein secondary structure by competing for H-bonds and is additionally capable of stacking with nonpolar side chains, particularly those containing planar aromatic groups.<sup>37,38</sup> In a similar fashion, aptamers can readily bind to guanidinium ions, potentially releasing any bound species as they do so.<sup>39,40</sup> Our result showed that GuHCl cleaning reduced the signal loss at low frequencies compared to non-cleaned electrodes, suggesting removal by Gu<sup>+</sup> ions of non-specifically bound aptamers from the outer layers of the

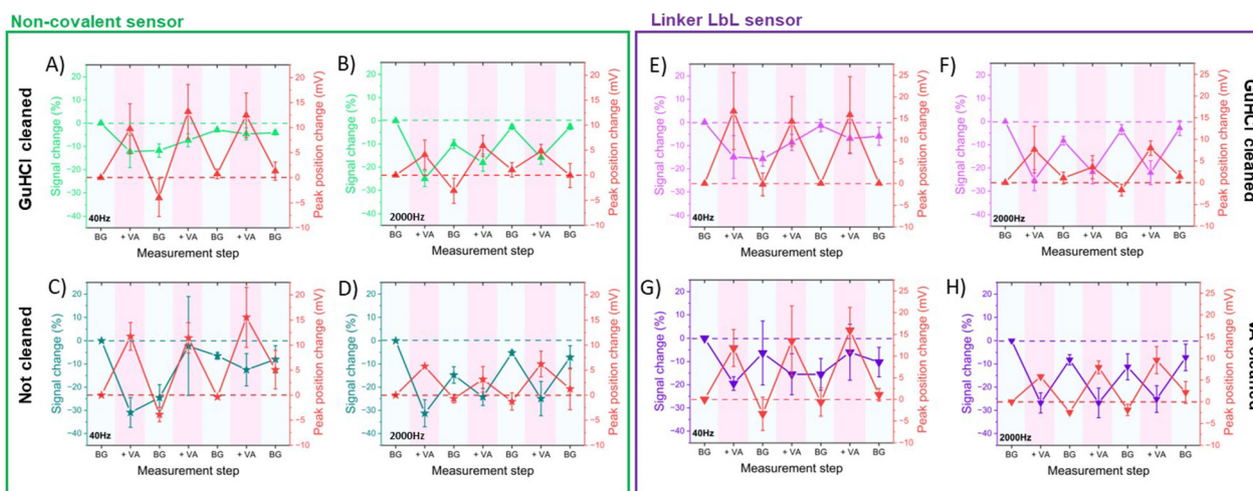
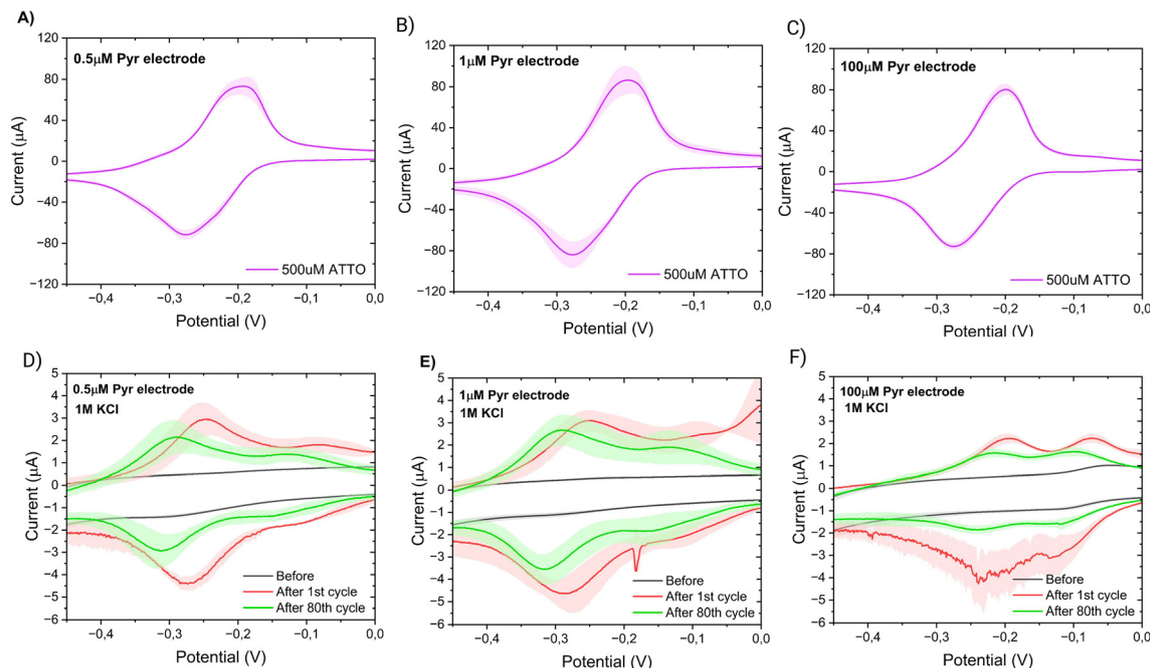


Fig. 6 Stability upon repetitive measurements of vancomycin at 40 and 2000 Hz of non-covalent sensors previously cleaned with GuHCl (A and B) and without a previous cleaning (C and D). Additionally, the stability upon repetitive measurements of vancomycin at 40 and 2000 Hz of linker LbL sensors previously cleaned with GuHCl (E and F) and with VA cleaning (G and H) is presented. The stability was assessed in terms of signal and peak position changes ( $N = 3$ ).





**Fig. 7** Passivation against methylene blue of a SWCNTs electrode with different linker concentrations. CV response of passivated electrodes to 500  $\mu\text{M}$  of methylene blue (ATTO) (A–C) ( $N = 3$ ). CV response of passivated electrodes in background (KCl) before and after the exposure of the electrodes to methylene blue (D–F) ( $N = 3$ ).

electrode surface (Fig. 6A and B). Additionally, the non-cleaned electrodes exhibited higher standard deviations and total signal loss at low frequencies (Table S13<sup>†</sup>). Conversely, at high frequencies, the signal loss was similar regardless of the cleaning method, indicating that aptamers close to the electrode surface were strongly bound and unaffected by  $\text{Gu}^+$  ions.

Observing significant signal loss after the initial VA measurement in non-covalent non-cleaned sensors (Fig. 6C and D), we hypothesized that VA may indirectly detach non-specifically bound aptamers from the electrode surface. This detachment could occur as the aptamers preferentially bind to vancomycin rather than to the electrode surface. To test this hypothesis, we evaluated the effectiveness of  $\text{GuHCl}$  and VA cleaning methods in linker LbL sensors (Fig. 6E–H). Both methods exhibited similar signal loss, albeit with slightly higher variability in the VA cleaning method (Table S13<sup>†</sup>). This suggests that both proposed cleaning methods effectively removed unbound aptamers and prepared the aptamers to be in their unbound state.

In terms of sensor stability during repetitive measurements of saturating concentrations of VA, both direct non-covalent and linker LbL sensors exhibited changes in signal and peak position upon exposure to VA at high frequencies, which were nearly restored during background measurements following VA measurement. Therefore, the sensors demonstrated good stability and repeatability when measuring saturating concentrations of VA at high frequencies. On the other hand, the signal change at low frequencies upon VA or BG measurement was inconsistent, likely due to the presence of

more randomly oriented and loosely bound aptamer at the outer layers of the electrode surface.

**3.2.3 Passivation against non-specific aptamers and MB with linker.** Due to reported strong adsorption of the redox reporter methylene blue to carbon nanotubes<sup>41–43</sup> and the present SPR results, we investigated the potential of the linker to prevent MB adsorption on the CNT network. Initially, we examined the adsorption of a methylene blue derivative (ATTO-MB2) on a plain electrode (SWCNTs on a glass substrate). Following exposure to MB and subsequent measurement in KCl, the plot displayed a peak at  $-0.3$  V associated with MB. Notably, this peak exhibited only a 31% decrease after 80 cycles (Fig. S17<sup>†</sup>). Next, we tested SWCNT electrodes coated with various linker concentrations. However, the CVs revealed no significant decrease in peak current when measured in 1 M KCl for 80 cycles (Fig. 7). In addition, the passivation was similar regardless of the linker concentration, suggesting that a full coverage of the electrode surface is already obtained with “low” concentration of linker ( $0.5$   $\mu\text{M}$ ). When compared to a plain electrode, the passivation achieved with the linker showed no discernible difference in peak current decrease. This observation indicates an unsuccessful passivation against MB adsorption, regardless of the linker concentration employed (Table 2). However, even though the linker fails to passivate the surface against non-specific interactions, it seems to help orient the aptamers based on the SPR and electrochemical results obtained with the linker LbL sensors.

**3.2.4 Concentration response.** Next, the four EAB sensors were tested *via* SWV at 160 Hz for the detection of vancomycin across a concentration range from 1 to 250  $\mu\text{M}$  (Fig. S18<sup>†</sup>). The



**Table 2** Percentage decrease in peak current (−0.25 V, CV) between the 1st and 80th cycles in 1 M KCl, post ATTO-MB2 measurement, for plain and passivated electrode ( $N = 3$ )

Plain electrode	Passivated electrode		
	0.5 $\mu\text{M}$ linker	1 $\mu\text{M}$ linker	100 $\mu\text{M}$ linker
31%	$33 \pm 7\%$	$8 \pm 14\%$	$45 \pm 10\%$

sensors displayed similar responses to vancomycin at low concentrations (1–25  $\mu\text{M}$ ). However, due to the progressive signal loss from aptamer detachment, only the 100 and 250  $\mu\text{M}$  VA concentrations can be considered reliably detected by all four sensor types (*i.e.*, when signal change > signal loss in Table 3). As expected, the signal loss was greatest for the direct non-covalent sensors, while the direct covalent sensors exhibited the lowest loss. Notably, the direct non-covalent sensors also showed significant variability, especially at high VA concentrations.

Regarding peak shifts upon VA addition, considerable variability in sensor performance was observed within the same sample type. Some sensors exhibited an anodic shift, while others showed a cathodic shift upon VA addition (Table S14<sup>†</sup>). This supports the hypothesis that there is a mix of differently attached aptamers on the surface exhibiting a variety of local reaction constants. Additionally, none of these sensors showed restoration of the peak position when tested in PBS +  $\text{MgCl}_2$  after the VA concentration series measurements.

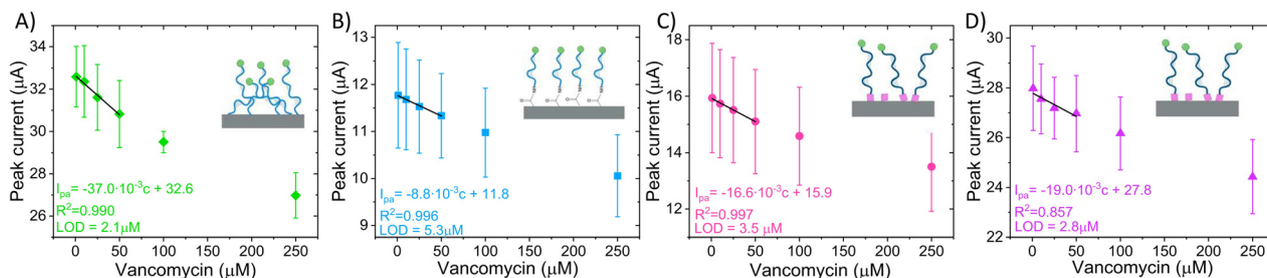
Based on the aforementioned results, a calibration curve could be obtained to detect different concentrations of vancomycin in the physiological range (1–50  $\mu\text{M}$ ) (Fig. 8) for all the sensor types. The calibration curves revealed that direct non-covalent sensors

exhibit the highest sensitivity and LOD (Table 4), which might misleadingly suggest enhanced performance compared to the other sensors. However, this sensor fails to meet our predefined criteria for proper operation. Furthermore, the signal generated by this sensor does not originate from a target-induced conformational change. One can apply the same reasoning to the other EAB sensor found in the literature (Table 4), where although it presents a current change with the addition of VA, there is no data on peak shift, thus it cannot be confirmed that the sensor signal arises from a conformational change of the aptamer. Consequently, when employing these commonly used calibration curves, it is crucial to carefully consider the underlying physical signal generation mechanism.

To conclude, for a proper sensor operation, it is imperative to address the two central challenges: (i) efficient aptamer binding and (ii) effective surface passivation. While the former affects the stability of the sensor the latter causes a substantial amount of background noise originating from the randomly oriented non-specifically bound aptamers. One potential approach to solve these challenges could involve transitioning from a 3D to a 2D carbonaceous material, allowing for better control over the orientation of aptamers and simultaneously reducing the adsorption volume. In

**Table 3** Signal change to different concentrations of vancomycin (1–250  $\mu\text{M}$ ) and total signal loss of the sensors ( $N = 3$ )

[VA] ( $\mu\text{M}$ )	Signal change (%)			
	Direct non-covalent	Direct covalent	Linker LbL	Linker pre-mixed
1	$0.5 \pm 0.3$	$0.4 \pm 0.5$	$0.3 \pm 1.0$	$2.0 \pm 2.0$
10	$1.2 \pm 1.6$	$1.1 \pm 0.0$	$1.5 \pm 1.0$	$3.5 \pm 1.0$
25	$3.5 \pm 0.8$	$2.3 \pm 0.6$	$2.9 \pm 1.2$	$4.7 \pm 1.1$
50	$5.9 \pm 0.7$	$3.9 \pm 1.4$	$5.5 \pm 1.4$	$5.5 \pm 1.1$
100	$9.8 \pm 2.7$	$7.0 \pm 0.6$	$8.7 \pm 1.4$	$8.3 \pm 0.8$
250	$17.4 \pm 5.7$	$14.8 \pm 0.8$	$15.5 \pm 0.6$	$14.4 \pm 1.1$
Signal loss (%)	$11.3 \pm 6.8$	$6.9 \pm 0.5$	$8.5 \pm 0.2$	$9.9 \pm 1.1$

**Fig. 8** Calibration curves and linear fit from 1–50  $\mu\text{M}$  VA for A) direct noncovalent, B) direct covalent, C) linker LbL, and D) linker pre-mixed electrodes ( $I_{\text{pa}}$  = forward partial peak current,  $c$  = vancomycin concentration, LOD = limit of detection,  $N = 3$ ).

**Table 4** LOD and sensitivity values of vancomycin sensors found in the literature

Material	LOD ( $\mu\text{M}$ )	Sensitivity ( $\mu\text{A } \mu\text{M}^{-1}$ )	Ref.
rGO-modified glassy carbon	0.2	0.810	44
Au-SAM EAB	NA	0.006 <sup>a</sup>	45
Hybrid graphene-Au nanocomposite	0.3	0.768	46
Direct noncovalent EAB ( $N = 3$ )	2.1	0.037	This work
Direct covalent EAB ( $N = 3$ )	5.3	0.009	
Linker premix EAB ( $N = 3$ )	2.8	0.019	
Linker LbL EAB ( $N = 3$ )	3.5	0.016	

<sup>a</sup> Approximated from Fig. 1C (1–10  $\mu\text{M}$ ) from ref. 44.

addition, we are currently working on a bio-derived linker which is displaying extremely promising results with respect to surface passivation, together with the known ability for it to be used for direct linking of biomolecules using EDC/NHS chemistry.

## 4. Conclusions

While the literature contains numerous aptamer-based electrochemical sensors, their operational depiction is oversimplified, preventing their optimization, and understanding of the phenomena taking place on the electrode surface. Moreover, it is paramount to understand how the aptamers' conformational changes and factors such as redox reporter adsorption are shown in the SWV signal, as this understanding is pivotal for advancing the field. Here we have proposed three electrochemical parameters to determine if a conformational change of the aptamer occurs in an EAB sensor: peak shift, current amplitude change, and the restoration of the original background current and peak position after the target measurement. These parameters, supported by the SPR measurements, provide us with insight about the plausible interfacial structure of the EAB sensors in different cases. Hereby we have monitored the behaviour of four types of SWCNTs EAB sensors using different aptamer binding methods: direct (covalent and non-covalent) and indirect using a linker molecule. At this stage, the sensors could only be used as single-use sensors if an appropriate calibration curve is performed. In the future, it will be necessary to optimize binding and passivation, in order to use these EAB sensors in continuous and accurate measurements.

## Data availability

The authors confirm that the data supporting the findings of this study are available within the article and its ESI.† Raw data that supports the findings of this study are available from the corresponding author upon request.

## Author contributions

LFP: investigation, methodology, writing – original draft, writing – review & editing, formal analysis, visualization. EG: investigation & methodology. JS: conceptualization, writing – original draft, writing – review & editing. VD: investigation,

methodology, writing – original draft, writing – review & editing, formal analysis, visualization. TL: conceptualization, methodology, writing – original draft, writing – review & editing, resources, supervision.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge Canatu Oy for providing the SWCNT networks and Elli Leppänen for her help during the investigation process.

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