Tailoring the Electrochemical and Morphological Properties of Electropolymerized and Dropcast rGO:PSS-PEDOT:PSS Transducers for Ion-Selective Sensors †

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Abstract: Fabrication of ion-selective sensors for continuous measurement in fluids depends on understanding the electrochemical and morphological properties of transducers. Electropolymerized nanomaterials essentially offer stable transducers that can reduce measurement drifts. This study aims to elucidate the electrochemical and morphological characteristics of electropolymerized reduced graphene oxide stabilized in polystyrenesulfonate and poly(3,4-ethylenedioxythiophene): polystyrenesulfonate composites on screen-printed carbon electrodes (rGO:PSS-PEDOT:PSS/SPCEs) using scanning electron microscopy (SEM) and cyclic voltammetry (CV) in 0.1 M ferricyanide (Fe(CN)64/3−). We fabricated the rGO:PSS-PEDOT:PSS/SPCEs by two different techniques: electropolymerization deposition (EPD) and drop-casting (DC). Results revealed smaller peak-to-peak potential separation (ΔEp) of 360 mV for EPD rGO:PSS-PEDOT:PSS/SPCEs, compared to 510 mV for the DC rGO:PSS-PEDOT:PSS/SPCEs. A smaller ΔEp indicates higher reversibility and faster electron-transfer rate at the electrode-analyte interface. SEM results showed EPD rGO:PSS-PEDOT:PSS/SPCEs have the roughest surface among electrodes; homogeneous globular structures with diameter range of 1.4–5.3 µm covered the electrode surface. In terms of electrode integrity in fluids, cracks can be seen on the surface of DC PEDOT:PSS/SPCEs after undergoing CV in 0.1 M Fe(CN)64/3−, whereas rGO:PSS-PEDOT:PSS/SPCEs for both deposition methods maintained their integrity. Globular structures of rGO:PSS-PEDOT:PSS using EPD methods remained after undergoing CV. The results suggest that EPD serves as a potential method to fabricate a stable transducer for ion-selective sensing. This study aims to elucidate performance of nanocomposites via EPD methods, to develop stable ion-selective sensors for physiological and environmental applications.

Keywords: electropolymerization; drop-cast; reduced graphene oxide; PEDOT:PSS; rGO:PSS; all-solid-state ion-selective sensors

1. Introduction

Ion-selective sensors have been an attractive choice of sensing technology for clinical, food, and environmental applications owing to sensitive, low-cost, and straightforward fabrication processes, while offering in addition the ability to monitor analytes in fluids [1–6]. Driven by advancements in nanotechnology, modern ion-selective sensors have evolved from conventional liquid-filled ion-selective electrodes (ISEs) into all-solid-state ISEs (AS-ISEs) that are easier to miniaturize and integrate into sensing platforms [7,8]. However, prolonged use of AS-ISEs in fluids can affect the
integrity of sensors and result in measurement drifts. Continuous real-time monitoring is required for various applications encompassing water-quality monitoring [9–11], in-vivo measurement for physiological processes [12–15], and nutrient monitoring in food [16]. Yet, conventional ISEs with inner filling solution are sensitive to evaporation and osmotic pressure, resulting in liquid volume change and hence frequent calibration [17]. Furthermore, handling of solutions requires microfluidics and automation. In this regard, there is a need to develop a robust and stable sensor that can function continuously in fluids with minimal measurement drift and less need for calibration during use. Such minimal drift with high potential stability of AS-ISEs can be achieved using solid-contact materials [18,19].

Previous work reported several techniques to deposit solid-contact materials, such as drop-casting (DC) [20,21], spin coating [22,23], inkjet printing [24–26], and electropolymerization deposition (EPD) [27,28]. DC is the simplest, consisting of two easy steps: casting the materials onto the WEs followed by drying the electrodes. However, transducer materials deposited by drop-casting are relatively thick, lack homogeneity along the WEs, and possess weak adhesion to electrode surfaces. As a result, electrodes modified via drop-casting are not mechanically robust; measurements results can vary between electrodes, so the technique is unsuitable for scaling-up purposes. EPD, however, is a promising method to fabricate robust and stable AS-ISEs owing to stronger molecular interaction between the polymers and the electrode surface, homogeneous thin-film deposition whose thickness can be controlled by changing the number of potential sweep cycles, and the capability to coat nanosize WEs [29]. Electropolymerization of 3,4-ethylene dioxythiophene (EDOT) in poly(sodium-4-styrenesulfonate) (NaPSS) and lithium perchlorate solution (LiClO₄) occurs when electrodes oxidize the EDOT monomers, making the resulting cations combine with themselves, thus producing higher oligomers of electropolymerized PEDOT:PSS [30]. The electropolymerized PEDOT:PSS has been tested for stability of the charge/discharge cycle using cyclic voltammetry (CV) and for its adhesion to electrode surfaces in flowing aqueous media [31].

Besides electropolymerized PEDOT:PSS, several studies have employed electropolymerization of graphene-based nanomaterials and PEDOT as a composite. PEDOT was electropolymerized from an aqueous solution with graphene oxide (GO) as the only charge-compensating counterion. During electropolymerization, GO can be incorporated into the positively charged PEDOT matrix because of the polar hydroxyl and epoxy groups, thus making the GO surface negatively charged. At present, the electropolymerization of rGO-PEDOT:PSS composite has been done with two-step processes, either by doing electropolymerization on the rGO surface [32], or by reducing GO to rGO electrochemically on top of the electropolymerized PEDOT layer [33].

In this study, we employed a one-step electropolymerization of EDOT in a complex electrolyte solution with an aqueous dispersion of reduced graphene oxide. The electropolymerized reduced graphene oxide stabilized in polystyrenesulfonate and poly(3,4-ethylenedioxythiophene): polystyrenesulfonate composite on screen-printed carbon electrodes (rGO:PSS-PEDOT:PSS/SPCEs) was then electrochemically characterized by cyclic voltammetry measurement. We used reduced graphene oxide stabilized in polystyrenesulfonate (rGO:PSS) to avoid rGO stacking while promoting hydrophilicity of the nanomaterial, allowing rGO:PSS to be dissolved in aqueous solution. In addition to electrochemical characterization, we also performed surface-morphology characterization to investigate the effect of fluids on electrode-surface morphology after CV measurement. A schematic of the experimental setup is illustrated in Figure 1. This study essentially offers a new platform to develop robust and stable AS-ISEs, which are particularly useful for monitoring ion concentrations continuously and in real-time for physiological and environmental applications.
Figure 1. All-solid-state ion-selective sensors. (a) Schematic of all-solid-state ion-selective sensors for continuous monitoring in fluids. A: sample fluid, B: screen-printed carbon electrode (SPCE), C: connector between electrode and pocketSTAT™, D: pocketSTAT™, E: IVIUM software. Fabrication steps of rGO:PSS-PEDOT:PSS/SPCEs by (b) drop-casting (DC) and (c) electropolymerization deposition (EPD) techniques. B1: expanded view of SPCEs showing the working electrode (WE).

2. Materials and Methods

2.1. Materials and Instrument

Screen-printed carbon electrodes (SPCEs, RRPE1001C) with diameter Φ = 2 mm were purchased from Pine Instrument Company, Grove City, PA, USA. A PocketSTAT™ was purchased from IVIUM Technologies, Eindhoven, the Netherlands. Reduced graphene oxide stabilized with poly(sodium 4-styrenesulfonate) (rGO:PSS, 900197), poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT:PSS, 483095), LiClO₄, EDOT, and NaPSS were purchased from Sigma-Aldrich, St. Louis, MO, USA. 98% concentrated sulfuric acid (H₂SO₄, 1407-80), and potassium ferricyanide (K₃Fe(CN)₆, 6969-00) were purchased from R&M Chemicals, Selangor, Malaysia. Analytical-grade chloride salts of potassium and sodium were procured from Sigma-Aldrich, St. Louis, MO, USA. All chemicals of analytical grade were used as received without further purifications. Deionized water (DI) was used throughout the experiment.
2.2. Fabrication of Drop-Cast and Electropolymerized rGO:PSS-PEDOT:PSS/SPCEs

SPCEs were activated via cyclic voltammetry for 3 cycles in 0.1 N H₂SO₄ at potentials of 2.5 V to -2.5 V and scan rate of 100 mV/s. The activated SPCEs were rinsed in DI water and left to dry. To prepare rGO:PSS-PEDOT:PSS/SPCEs, two techniques were used in modifying the working electrode: drop-casting (DC) and electropolymerization deposition (EPD). For DC, rGO:PSS was mixed with PEDOT:PSS to prepare a rGO:PSS-PEDOT:PSS composite. To fabricate rGO:PSS-PEDOT:PSS/SPCEs, 3 µl rGO:PSS-PEDOT:PSS was drop-cast onto the WE of an SPCE. For EPD, the NaPSS, EDOT, and rGO:PSS were dissolved in 0.1 M LiClO₄. The galvanostatic technique by applying constant current was used to electropolymerize EDOT and rGO:PSS to rGO:PSS-PEDOT:PSS. Both DC and EPD rGO:PSS-PEDOT:PSS/SPCEs were left to dry for 24 h under a laminar flow hood (Esco Micro, ADC—4C1, Shah Alam, Selangor, Malaysia).

2.3. Cyclic Voltammetry of rGO:PSS-PEDOT:PSS/SPCEs

Cyclic voltammetry (CV) was performed for unmodified SPCEs, DC rGO:PSS-PEDOT:PSS/SPCEs, and EPD rGO:PSS-PEDOT:PSS/SPCEs in 0.1 M potassium ferricyanide, K₃Fe(CN)₆, over a potential range from -0.5 V to 1.0 V at scan rates of 75, 100, 150, 200, and 250 mV/s. Electrochemical parameters such as peak current (Iₚ), effective surface area (Aₑ), and separation of peak-to-peak potential (ΔEₚ) were analyzed. The Iₚ obtained from the measurement can be used to calculate the Aₑ of electrodes using the Randles-Sevcik equation:

\[ I_p = (2.69 \times 10^5)n^{3/2}D^{1/2}C_Aν^{1/2} \]  
\[ A_e = \frac{k}{(2.69 \times 10^5)n^2D^2C} \]

2.4. Scanning Electron Microscopy (SEM) of rGO:PSS-PEDOT:PSS/SPCEs

Surface morphologies of SPCEs, DC rGO:PSS-PEDOT:PSS/SPCEs, and EPD rGO:PSS-PEDOT:PSS/SPCEs were observed using a scanning electron microscope (JSM-IT100, JEOL, JAPAN). The electrodes were coated with gold/palladium particles using the Quorum SC7620 Sputter Coater prior to imaging. SEM images were obtained with a power setting of 8.0 kV and magnification of 300×. The average diameter of rGO:PSS-PEDOT:PSS globules was determined using Image J software for n = 15.

3. Results

3.1. Cyclic Voltammetry of rGO:PSS-PEDOT:PSS/SPCEs

CV analysis was performed for bare SPCEs, EPD rGO:PSS-PEDOT:PSS/SPCEs, and DC rGO:PSS-PEDOT:PSS/SPCEs in 0.1 M Fe(CN)₆⁴⁻ to investigate the changes in Iₚ, Aₑ, and ΔEₚ. We determined the Iₚ of EPD and DC rGO:PSS-PEDOT:PSS/SPCEs compared to bare SPCEs by analyzing the difference in anodic peak current (ΔI) between the bare and modified electrodes at a scan rate of 100 mV/s. There was a noticeable increase in the Iₚ of EPD and DC rGO:PSS-PEDOT:PSS/SPCEs compared to that of bare SPCEs (Iₚ, Bare = 0.247 mA, Iₚ, EPD = 0.415 mA, and Iₚ, DC = 1.045 mA; ΔI, EPD-Bare = 0.168 mA, and ΔI, DC-Bare = 0.798 mA). The increase in Iₚ suggests that rGO:PSS-PEDOT:PSS can facilitate accessibility of the Fe(CN)₆⁴⁻ to the electrode surface by improving interfacial electron transfer, thus enhancing the Iₚ signals. Incorporation of PEDOT:PSS to the rGO increased the electron transport owing to π-π interactions between PEDOT:PSS and rGO. The availability of π-bonding aromatic rings increases and contributes to a larger number of π-π overlaps, leading to more electron delocalization around the aromatic ring and thus improving the electrical conductivity of the modified electrodes [34–36]. Note that rGO:PSS is rGO layers stabilized in polystyrenesulfonate,
which inhibits rGO stacking while promoting hydrophilicity of the nanomaterial, thus allowing rGO:PSS to be dissolved in aqueous solution [37].

In addition, Aₑ was also determined by applying the Randles-Sevcik equation. The Aₑ for bare SPCEs, EPD rGO:PSS-PEDOT:PSS/SPCEs, and DC rGO:PSS-PEDOT:PSS/SPCEs were 3.1 mm², 5.7 mm², and 6.7 mm², respectively, showing a ~2-fold increase for modified SPCEs compared to bare SPCEs. The increase in Aₑ of the EPD and DC rGO:PSS-PEDOT:PSS/SPCEs in comparison to the bare SPCEs could be attributed to the increase in surface-to-volume ratio of the nanomaterials, as well as to increase in current density and charge transferability [21].

Further analyses of CV graphs compared the effect of deposition method on separation of peak-to-peak potentials (ΔEₚ = Eₚa - Eₚc), where lower ΔEₚ implies an enhanced electrocatalytic ability of Fe(CN)₆⁴⁻⁻ at the electrode-analyte interface. We analyzed the ΔEₚ of EPD rGO:PSS-PEDOT:PSS/SPCEs in comparison to DC rGO:PSS-PEDOT:PSS/SPCEs. The EPD rGO:PSS-PEDOT:PSS/SPCEs showed a lower ΔEₚ of 360 mV, 1.4 times lower than those of DC rGO:PSS-PEDOT:PSS/SPCEs (ΔEₚ = 510 mV). The lower ΔEₚ suggests higher reversibility and faster electron transfer at the electrode-analyte interface; thus, the diffusion of analyte to the electrode surface is a dominant contributor to the peak potential [38]. The structure of PEDOT:PSS and rGO:PSS composites formed from the EPD process could potentially assist the electron transfer between electrode surface and analyte, making the nanocomposite suitable for ion-selective sensing.

Figure 2. (a) Cyclic voltammetry (CV) measurements of bare SPCEs, EPD rGO:PSS-PEDOT:PSS/SPCEs, and DC rGO:PSS-PEDOT:PSS/SPCEs at a scan rate of 100 mV/s in 0.1 M Fe(CN)₆⁴⁻⁻. ΔIₚ is shown. CV measurements of (b) SPCEs, (c) DC rGO:PSS-PEDOT:PSS/SPCEs, and (d) EPD rGO:PSS-PEDOT:PSS/SPCEs at scan rates of 75, 100, 150, 200, 250 mV/s in 0.1 M Fe(CN)₆⁴⁻⁻. ΔEₚ is shown.
3.2. Surface Morphology of rGO:PSS-PEDOT:PSS/SPCEs

The surface morphology of SPCEs modified with EPD and DC of rGO:PSS-PEDOT:PSS was characterized under SEM. SEM images were taken after all electrodes were run through CV. The morphology could provide an insight on the effect of fluids on the nanocomposite after undergoing CV, as CV is a consumptive process that can affect morphology of nanomaterial transducers. Figure 3 shows the surface morphology of bare SPCEs, PEDOT:PSS/SPCEs, DC rGO:PSS-PEDOT:PSS/SPCEs, and EPD rGO:PSS-PEDOT:PSS/SPCEs. Bare SPCEs display a rough and flaky-like morphology (Figure 3a), PEDOT:PSS/SPCEs appear smooth with some bumps, and DC rGO:PSS-PEDOT:PSS/SPCEs resemble PEDOT:PSS/SPCEs (Figure 3b) but with the appearance of tiny grains on the electrode surface (Figure 3c). EPD rGO:PSS-PEDOT:PSS/SPCEs have the roughest surface and homogeneous globular structures can be seen to cover the electrode surface (Figure 3d). Detailed analysis using Image J software revealed that the globule diameters are in the range of 1.4–5.3 µm. In terms of electrode integrity in fluids, in this case after electrodes had undergone CV in 0.1 M Fe(CN)₆⁴⁻/₃⁻, cracks can be seen on DC PEDOT:PSS/SPCE surfaces (Figure 3b), whereas rGO:PSS-PEDOT:PSS/SPCEs for both deposition methods of DC (Figure 3c) and EPD (Figure 3d) maintained their integrity. The results suggest that PEDOT:PSS alone can become brittle with prolonged measurement and storage in fluids. In PEDOT:PSS layers, PEDOT is surrounded by a PSS network that attracts a considerable amount of water, allowing transport and penetration of ions into the PEDOT layers. Ion transport, and subsequently electrochemical activity, become possible with such hydrated pathways. This, however, brings a challenge to maintain layer integrity since the layers are exposed to aqueous environment such as biological fluids for in-vivo applications and flowing water for environment applications [39]. In this regard, incorporation of rGO:PSS into PEDOT:PSS as a composite can improve the mechanical strength of rGO:PSS-PEDOT:PSS when electrodes undergo CV measurements. The results are consistent with previous studies that found that PEDOT:PSS/graphene nanocomposite [40] and PEDOT:PSS/carboxylic group functionalized graphene composite [41] are stable in aqueous media owing to the properties of graphene derivatives, which improve flexibility, enhance adhesion, and increase mechanical strength of the composite.

In addition, globular structures of rGO:PSS-PEDOT:PSS using EPD methods remained after undergoing CV, indicating a robust and stable transducer in fluids. Besides morphological properties, the CV of EPD rGO:PSS-PEDOT:PSS have shown smaller ∆Eₚ, which implies higher reversibility and faster electron transfer at the electrode-analyte interface. Tailoring the morphological and electrochemical properties of the EPD rGO:PSS-PEDOT:PSS/SPCEs, we can conclude that rGO:PSS-PEDOT:PSS via EPD can be a potential technique to fabricate a robust transducer for ion-selective sensing in biological and environmental fluids. Similar results were observed in previous work, where PEDOT:PSS drop-casting techniques produced smoother images when the electrodes were soaked in PBS; however, globular structures remained for electropolymerized PEDOT:PSS [31].
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Figure 3. Surface morphology of (a) bare SPCEs, (b) PEDOT:PSS/SPCEs, (c) DC rGO:PSS-PEDOT:PSS/SPCEs, and (d) EPD rGO:PSS-PEDOT:PSS/SPCEs. Bare SPCEs display a rough and flaky-like morphology, PEDOT:PSS/SPCEs appear smooth with some bumps, and DC rGO:PSS-PEDOT:PSS/SPCEs resemble PEDOT:PSS/SPCEs but with the appearance of tiny grains on the electrode surface. The roughest surface is EPD rGO:PSS-PEDOT:PSS/SPCEs where a homogeneous globular structure can be seen to cover the electrode surface. Detailed analysis using Image J software revealed that the globule diameters are in the range of 1.4–5.3 µm. In terms of electrode integrity in fluids, in this case after electrodes underwent CV in 0.1 M Fe(CN)$_6^{3-/4^-}$, cracks can be seen on DC PEDOT:PSS/SPCE surfaces, whereas rGO:PSS-PEDOT:PSS/SPCEs for both deposition methods of DC and EPD maintained their integrity. The results suggest that PEDOT:PSS alone can become brittle with prolonged measurement and storage in fluid. Incorporation of rGO:PSS into PEDOT:PSS as a composite improves the mechanical strength of rGO:PSS-PEDOT:PSS when electrodes undergo CV measurements.

4. Conclusions

We compared CV and surface morphology of rGO:PSS-PEDOT:PSS/SPCEs by two different techniques: electropolymerization deposition (EPD) and drop-casting (DC). Higher peak current ($I_P$) was observed for nanocomposite-modified SPCEs with $I_{P,EPD} = 0.415$ mA and $I_{P,DC} = 1.045$ mA, compared to bare SPCEs with $I_{P, Bare} = 0.247$ mA. The results indicate that the nanocomposite improves the conductivity of the working electrodes (WEs). Larger effective surface area ($A_e$) was also observed for nanocomposite-modified SPCEs with $A_{e,EPD} = 5.7$ mm$^2$ and $A_{e,DC} = 6.7$ mm$^2$, showing a ~2-fold increase compared to bare SPCEs with $A_{e, Bare} = 3.1$ mm$^2$. In addition, we analyzed the effect of deposition method on the electrochemical properties of the composite. Electropolymerized rGO:PSS-PEDOT:PSS/SPCEs showed a smaller peak-to-peak potential separation ($\Delta E_p$) of 360 mV compared to the drop-cast rGO:PSS-PEDOT:PSS/SPCEs (510 mV). The smaller $\Delta E_p$ of electropolymerized rGO:PSS-PEDOT:PSS suggests higher reversibility and faster electron transfer at the electrode-analyte interface. SEM results showed that bare SPCEs display a rough and flaky-like morphology,
PEDOT:PSS/SPCEs appear smooth with some bumps, and DC rGO:PSS-PEDOT:PSS/SPCEs resemble PEDOT:PSS/SPCEs but with appearance of tiny grains on the electrode surface. The roughest surface is EPD rGO:PSS-PEDOT:PSS/SPCEs, where a homogeneous globular structure can be seen to cover the electrode surface. Detailed analysis using Image J software revealed that the globule diameters are in the range of 1.4–5.3 µm. In terms of electrode integrity in fluids, in this case after electrodes underwent CV in 0.1 M Fe(CN)₆⁴⁻/₃⁻, cracks can be seen on DC PEDOT:PSS/SPCE surfaces, whereas rGO:PSS-PEDOT:PSS/SPCEs for both deposition methods of DC and EPD maintained their integrity. The results suggest that PEDOT:PSS alone can become brittle with prolonged measurement and storage in fluids. Incorporation of rGO:PSS into PEDOT:PSS as a composite improves the mechanical strength of rGO:PSS-PEDOT:PSS when electrodes undergo CV measurements. Globular structures of rGO:PSS-PEDOT:PSS using EPD methods remained after undergoing CV. The results suggest that EPD serves as a potential method to fabricate a stable transducer for ion-selective sensing. Future work includes investigating the effect of ion-selective membranes on EPD rGO:PSS-PEDOT:PSS/SPCEs, and testing the fabricated sensors for continuous monitoring in fluids. This study essentially offers a scalable platform to develop robust and stable ion-selective sensors, which are particularly useful for monitoring ion concentrations continuously in real-time for physiological and environmental applications.

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