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# Electrochemical paraquat sensor based on lead oxide nanoparticles†

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1,1-Dimethyl-4,4-bipyridinium dichloride known as paraquat is a popular well-known herbicide that is widely used in agriculture around the world. However, paraquat is a highly toxic chemical causing damage to vital organs including the respiratory system, liver, heart, and kidneys and death. Therefore, detection of paraquat is still necessary to protect life and the environment. In this work, an electrochemical sensor based on lead oxide nanoparticles (PbO-NPs) modified on a screen-printed silver working electrode (SPE) has been fabricated for paraquat detection at room temperature. The PbO-NPs have been synthesized by using a sparking method *via* two Pb metal wires. The electrochemical paraquat sensors have been prepared by a simple drop-casting of PbO-NPs solution on the surface of the SPE. The PbO-NPs/SPE sensor exhibits a linear response in the range from 1 mM to 5 mM with good reproducibility and high sensitivity (204.85  $\mu\text{A mM}^{-1} \text{cm}^{-2}$ ) for paraquat detection at room temperature. The PbO-NPs/SPE sensor shows high selectivity to paraquat over other popular herbicides such as glyphosate, glufosinate-ammonium and butachlor-propanil. The application of the PbO-NPs/SPE sensor is also demonstrated *via* the monitoring of paraquat contamination in juice and milk.

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## 1. Introduction

Paraquat (1,1-dimethyl-4,4-bipyridinium dichloride) is one of the well-known herbicides that have been widely used in the world for more than 50 years.<sup>1</sup> Over 130 countries worldwide have applied paraquat in agriculture to inhibit grass and weed growth due to its quick-acting, non-selective contact with green plant tissue and it being easily dissolved in water at room temperature.<sup>2</sup> However, paraquat is highly toxic and has long-term affected the environment such as through spreading and being residual in the air, soil, water, plants and animals. The mechanism of toxicity of paraquat occurs *via* the single electron redox cycling in biological systems. The redox cycling of paraquat can generate activated oxygen such as superoxide anion, superoxide radical and hydrogen peroxide which are relevant to the development of toxicity.<sup>3,4</sup> Although, many countries have banned paraquat, it slowly degrades in the environment with a half-life of more than 10 years.<sup>5,6</sup> Moreover, paraquat causes serious damage to human health and is related to Parkinson's disease.<sup>7,8</sup>

Conventional methods for paraquat detection are based on spectroscopic techniques such as gas chromatography-mass spectrometry (GC-MS)<sup>9</sup> high performance liquid chromatography (HPLC)<sup>10</sup> surface-enhanced Raman scattering (SERS) spectroscopy<sup>11</sup> surface-assisted laser desorption/ionization time-of-flight mass spectrometry (SALDI-TOF MS)<sup>12</sup> and so on. Although, these methods offer very high accuracy to identify quantity of paraquat in samples, they are very expensive, complicated procedures, long time-consuming and unavailable under on-site field analysis. Alternative methods based on bio/electrochemical sensors have received much attention in recent years.<sup>13–15</sup> For examples, Sant'Anna *et al.*<sup>16</sup> fabricated an electrochemical sensor based on nanocomposite of activated biochar and reduced graphene oxide to detect paraquat in food samples such as coconut water, wastewater, honey, lettuce and lemon. Kavazoi *et al.*<sup>17</sup> synthesized tetrasulfonated phthalocyanine for paraquat sensing. Ali *et al.*<sup>18</sup> developed a colorimetric sensor based on imidacloprid stabilized silver nanoparticle that showed high sensitive and low limit of detection toward paraquat. Deroco *et al.*<sup>19</sup> demonstrated a paper-based silver electrode prepared by the inkjet-printing method in the determination of paraquat. Souza *et al.*<sup>20</sup> used beeswax incorporating carbon black nanoparticles to determine paraquat in *Apis mellifera* honey. Jiang *et al.*<sup>21</sup> showed the excellent photo-electrochemical performance for paraquat detection based on ITO/Au/Cu<sub>2</sub>O electrode. Although, several materials have successfully used for paraquat sensing, new materials with low-

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cost and simple preparation methods are still need to enable the electrochemical paraquat sensor into commercialization.

Lead oxide has been served for electrode coating due to its superior reaction sites and intrinsic chemical properties.<sup>22</sup> There are many oxide forms including PbO, Pb<sub>2</sub>O<sub>3</sub>, Pb<sub>3</sub>O<sub>4</sub>, and PbO<sub>2</sub> which can be applied in  $\gamma$ -ray shielding,<sup>23</sup> photoconductor,<sup>24</sup> memristor application,<sup>25</sup> radiation attenuation application,<sup>26</sup> energy-storage application,<sup>27</sup> and lithium ion battery.<sup>28</sup> The lead oxide nanoparticles (PbO-NPs) have been synthesized by several methods such as sol-gel, chemical precipitation, chemical bath deposition, spray paralysis, hydrothermal, and laser ablation.<sup>29</sup> However, most methods require high-temperatures operation, high-vacuum systems, high energy, multistep processes, long operation times leading to expensive and harmful to the environment. A new sparking method<sup>30</sup> prepared nanoparticles with high porosity under atmospheric pressure with inexpensive, simple and rapid may be considered as an alternative and effective method for PbO-NPs synthesis.

In this work, the PbO-NPs as the sensing material for paraquat detection were synthesized by a sparking method at room temperature. In addition, we have developed the electrochemical sensor based on the PbO-NPs modified screen-printed silver electrode (SPE). The determination of paraquat was carried out using cyclic voltammetry (CV) technique under optimal conditions.

## 2. Experimental details

### 2.1 Synthesis of lead oxide nanoparticles

The PbO-NPs were synthesized by sparking method as shown in Fig. 1 with the four pairs of pure lead wires (purity 99.0%, diameter = 0.5 mm) used as sparking tips. The four pairs sparking tips were placed horizontally about 2 mm above the center of the glass substrate. The spacing of sparking wires were set at 1 mm. The sparking method was controlled by a constant voltage of 7.60 V and current of 3.65 A. Firstly, the glass substrate was cleaned by acetone, distilled water and dried. Next, the lead wires were sparked into 1 ml of deionized water

on glass substrate for 45 min at room temperature. The PbO-NPs suspension was ultra-sonicated for 20 min and kept at a room temperature for further use.

### 2.2 Fabrication of the PbO-NPs modified SPE sensor

The screen-printed electrodes (SPEs) were purchased from Quasense. It consists of three main parts as follows: a silver (Ag) working electrode, a carbon (C) counter electrode and a silver/silver chloride (Ag/AgCl) reference electrode. Fig. 2 shows the procedure for preparing the PbO-NPs modified SPE sensor. The 5  $\mu$ L PbO-NPs suspension was dropped cast on the surface of SPE (7.065 mm<sup>2</sup> area) for three times. The modified SPE was then dried in air at room temperature and ready for using in paraquat detection. It should be noted that a standard carbon working electrode can be also used for coating PbO-NPs to detect paraquat. However, the silver working electrode gave a higher current over carbon working electrode at the same concentration of paraquat. Thus, silver working electrode has been selected for this study.

### 2.3 Characterization of PbO-NPs

The morphology of PbO-NPs was characterized by a high-magnification and high-resolution transmission electron microscopy (TEM and HR-TEM, JEOL model: JEM-3100 (HR)) and field-emission scanning electron microscopy (FE-SEM: SU5000, Hitachi). The size distribution of nanoparticles was measured from TEM images using Image J. The crystal phase and crystallinity of PbO-NPs were evaluated by X-ray diffraction (XRD, model: Bruker D8 Advance) with Cu K $\alpha$  radiation (Bruker,  $\lambda = 1.5406 \text{ \AA}$ ,  $2\theta = 20\text{--}80^\circ$ ). The element composition and chemical state in PbO-NPs were analyzed by X-ray photoelectron spectrometry (XPS, model: Axis Ultra DLD, Kratos analytical, Manchester UK).

### 2.4 Electrochemical measurement

The sensing performance of the PbO-NPs SPEs sensor was investigated by cyclic voltammetry (CV) with a Sensit Smart potentiostat (EmStat Pico module) from PalmSens BV and it was

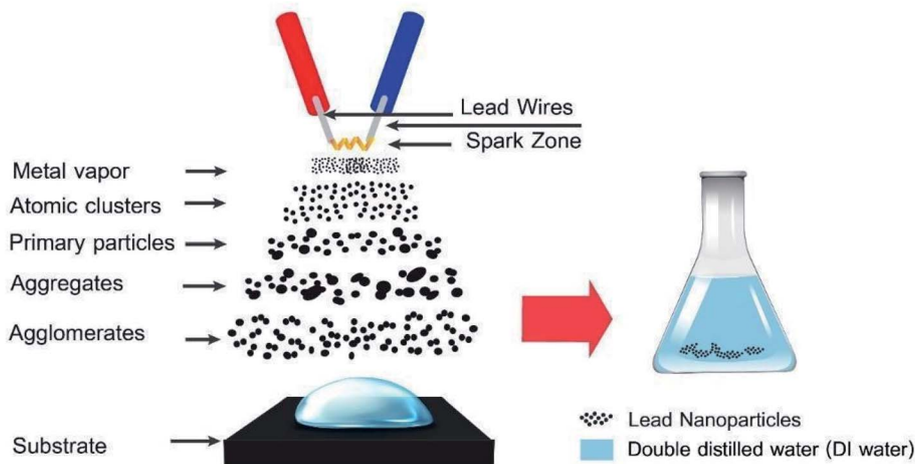


Fig. 1 Schematic illustration of the PbO-NPs synthesis by using sparking method.



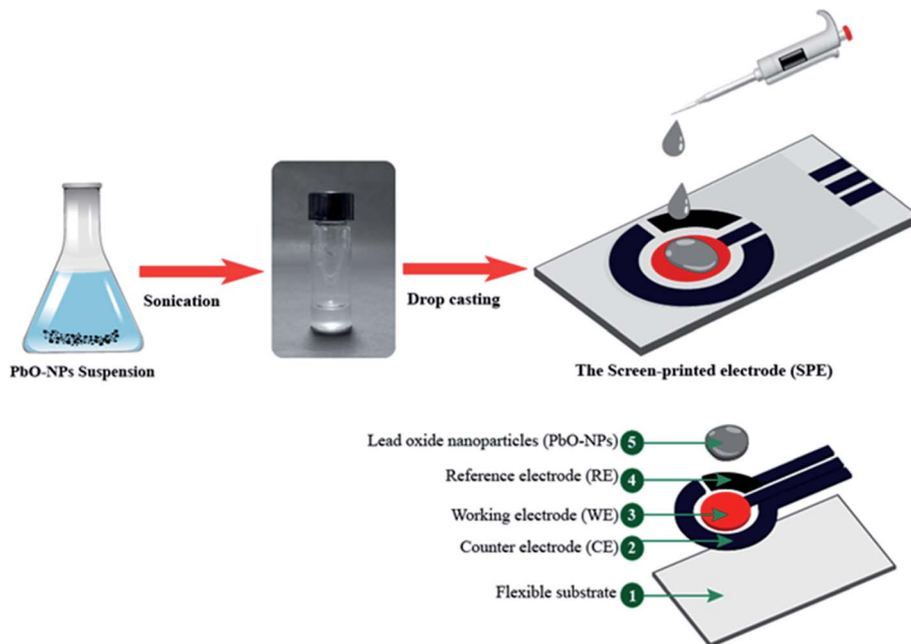


Fig. 2 Schematic illustration of PbO-NPs SPE sensor fabrication.

controlled by a PSTrace software. The cyclic voltammetric characterizations were performed in the potential range between 0 V to  $-1.4$  V and scan rate of  $100 \text{ mV s}^{-1}$ . The PbO-NPs SPEs sensor was equilibrated in 0.1 M Tris solution (electrolytic solution) by performing cyclic voltammetry until the current and potential signal constant. All electrochemical measurements were repeated for three times (four independent fabricated PbO-NPs SPEs sensors). The volume of paraquat and other herbicides for any tests was  $100 \mu\text{L}$ .

## 3. Results and discussion

### 3.1 Characterization of PbO-NPs

The sparking PbO-NPs sensing film exhibits the dense structure and nearly spherical shape with various sizes ranging from 10 to 80 nm as shown in Fig. 3a. The average particle size is  $22.95 \pm 5.46$  nm. The d-spacing of the lattice fringes is  $\sim 0.298$  nm closing to the (101) diffraction plane of the PbO<sup>31</sup> as displayed in Fig. 3b. The XRD peaks reveal three strong peaks at  $2\theta = 31.15^\circ$ ,  $35.83^\circ$  and  $51.96^\circ$  corresponding to the (101), (002) and (220) planes of PbO (JCPDS card, no. 01-085-1288), respectively, as shown in Fig. 3b. It can confirm a pure phase of synthesized orthorhombic lead oxide ( $\beta$ -PbO) *via* simple sparking method. It's well known that  $\alpha$ -PbO transforms to  $\beta$ -PbO at approximately  $490^\circ\text{C}$  (ref. 32) resulting to more stable  $\beta$ -PbO at high temperature. During sparking process, high energy was applied to Pb atoms to form PbO nanoparticles at high temperature. Thus,  $\beta$ -PbO is more contribution than  $\alpha$ -PbO phase. The average crystallite sizes of the PbO crystallites can be determined using Debye–Scherrer formula as given in eqn (1). The average crystallite size is found to be 30.78 nm that is smaller than that of the PbO nanoparticle samples prepared by other methods.<sup>33,34</sup> This small size may provide high surface area to interact with paraquat molecules.

$$D = \frac{k\lambda}{\beta \cos \theta} \quad (1)$$

where  $D$  is the average crystallite size,  $k$  is the constant ( $\sim 0.9$ ),  $\lambda$  is the wavelength of Cu K $\alpha$  line ( $1.5405 \text{ \AA}$ ),  $\beta$  is full width at half maximum (FWHM),  $\theta$  is the Bragg's diffraction angle.

The chemical states and nature bonding of the synthesized PbO-NPs were evaluated by XPS as shown in Fig. 3c and d. The core level of Pb displays the doublet in tetravalent oxidation state of lead due to the spin orbit splitting into  $4f_{5/2}$  and  $4f_{7/2}$ . The de-convoluted Pb 4f shows the two peaks binding energies at 143.5 eV and 138.5 eV as shown in Fig. 3c which correspond to the core levels of Pb<sup>2+</sup> and Pb, respectively. The higher peak intensity of Pb–O bonds presented the more content of the PbO formation with an energy separation of 4.9 eV.<sup>35–39</sup> In Fig. 3d, the de-convoluted O 1s state shows the signal peak that can be fitted from five peaks by Gaussian–Lorentzian curves. The de-convoluted peaks can be divided into five types of oxygen species with the binding energies at 529.9 eV, 530.9 eV, 531.9 eV, 532.9 eV and 534.2 eV. The peak located at 529.9 eV corresponds to oxygen atoms (O<sup>2-</sup>) in the lead oxide lattice (O–Pb bond). The lattice O<sup>2-</sup> peak located at 530.9 eV relates to the OH<sup>-</sup> group from the PbO–OH. The peak located at 531.9 eV corresponds to adsorption or chemisorption of atomic oxygen (O<sub>ads</sub>) which also related to the presence of water molecules. The peak at 532.9 eV attributes to the OH<sub>(ads)</sub> onto the surface of Pb–O.<sup>40</sup> Based on XPS results, it can confirm purity and formation PbO-NPs. However, OH functional groups can be found at the surfaces of PbO-NPs which may help to increase the interactions for paraquat molecules adsorption.

The surface morphology of Ag working electrode after coating of PbO-NPs was characterized by using FE-SEM and EDS analysis as displayed in Fig. 4. Due to small size of PbO-NPs, the



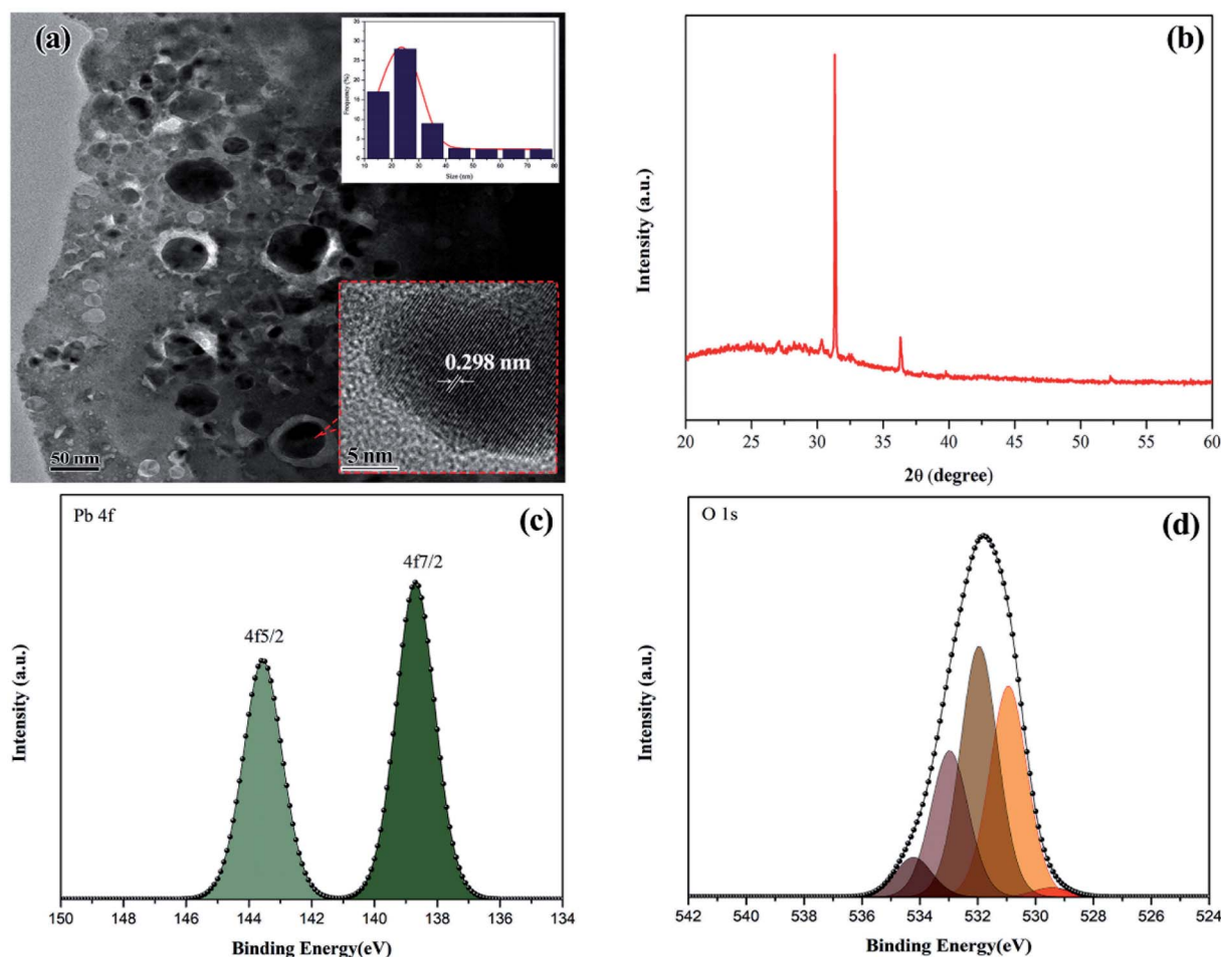


Fig. 3 (a) TEM image of the synthesized PbO-NPs by using sparking method. The insets show their size distribution and HR-TEM image of the lattice fringes. (b) XRD pattern and XPS spectra of high-resolution (c) Pb 4f and (d) O 1s.

PbO-NPs are well distributed on the Ag working electrode. Some agglomeration of PbO-NPs cluster can be also found on the Ag working electrode because of room-temperature evaporation of PbO-NPs solution. Based on EDS spectrum, it can confirm high quality and quantity of PbO-NPs modified on the Ag working electrode by simply drop-casting method. No observed color change on the working electrode was found after coating of PbO-NPs as shown in Fig. S1 in ESI.† Strong attractive forces between PbO-NPs and Ag working electrode without using any binder or adhesive membrane may be contributed from chemisorption that usually found in Ag sensing material.

### 3.2 Electrochemical behaviors of the PbO-NPs sensor

The electrochemical behaviors of PbO-NPs sensor for paraquat detection were investigated *via* cycle voltammetry. The electrochemical response of the PbO-NPs sensor in the presence of paraquat shows the strong four-peak response (two cathodic peaks and two anodic peaks) as displayed in Fig. 5a. The peak potential separation ( $\Delta E = E_{pc} - E_{pa}$ ) were 0.130 V ( $P_1 - P_4$ ) and 0.289 V ( $P_2 - P_3$ ), respectively, for potential scan rate of 100 mV s<sup>-1</sup>. It indicates a quasi-reversible system of redox reaction process. The two anodic reaction equations can be followed as

$PQ \rightarrow PQ^+ + e^-$  and  $PQ^+ \rightarrow PQ^{2+} + e^-$ . The cathodic peak potential at  $-0.85$  V ( $P_1$ ) is the first reduction process involved one electron in the reaction resulting in the formation of the radical cation  $PQ^{*+}$  ( $PQ^{2+} + 1e^- \rightarrow PQ^{*+}$ ). The second cathodic peak potential at  $-1.25$  V ( $P_2$ ) is the second reduction process in which  $PQ^{*+}$  is further reduced *via* one electron process to form neutral molecules ( $PQ^0$ ). The influences of the pH values on  $P_1$  and  $P_2$  are displayed in Fig. 5b and c, respectively. The peak  $P_1$  current shows a maximum value at pH 7 and decreases with rising pH. The positive shift in reduction peak can be observed with decreasing of pH suggesting in reaction with proton-transfer process in the reduction of PQ. At low or high pH values, they decrease in the peak current due to the reaction of disproportionation and degree of protonation<sup>41,42</sup> that reduce amount of oxide on the PbO surface resulting to a decrease in peak current corresponding to previous works.<sup>43,44</sup> In case of  $P_2$ , the peak currents are quite stable in range of pH 7–10 due to the most contribution of first reduction process. By considering the response currents and behavior of the sensor, the pH 7.0 has been selected as the optimum pH value for the supporting electrolyte. The first reduction peak has been chosen to



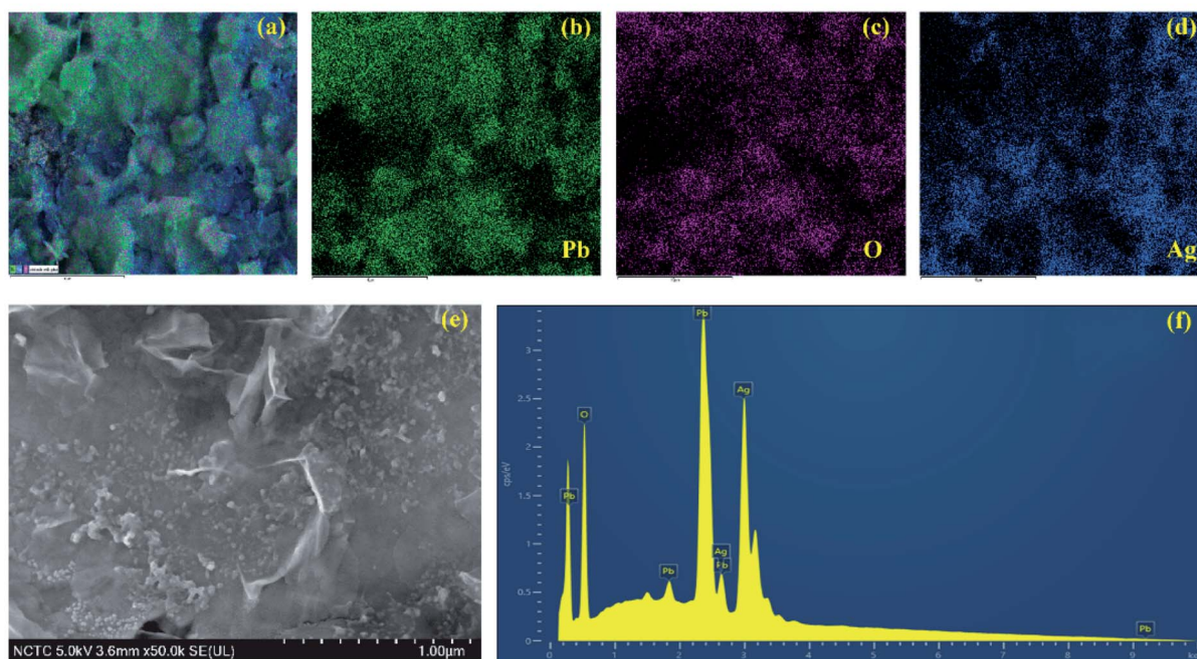


Fig. 4 (a)–(d) EDS mapping, (e) FE-SEM image and (f) EDX spectrum of Ag working electrode after coating of PbO-NPs.

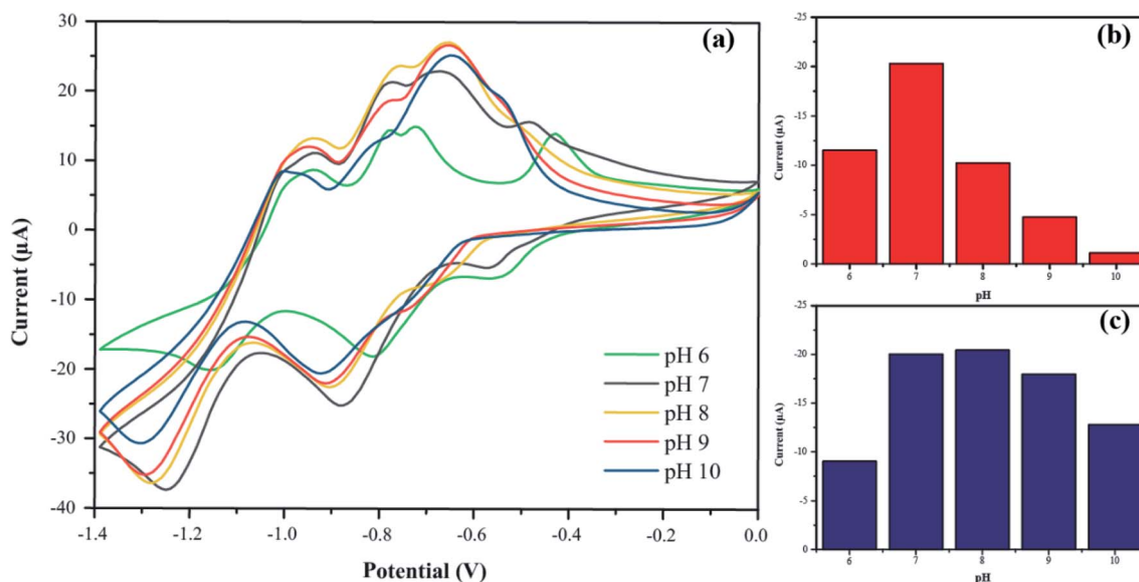


Fig. 5 (a) Cyclic voltammograms responses of PbO-NPs sensor at the different pH values in the presence of 1 mM paraquat. Their current responses of (b)  $P_1$  at  $-0.8$  V and (c)  $P_2$  at  $-1.1$  V.

determine the optimal conditions and calibration curve for paraquat determination.

To understand the dynamic performance of paraquat onto the PbO-NPs sensor, the influence of scan rate was studied from  $100 \text{ mV s}^{-1}$  to  $500 \text{ mV s}^{-1}$  in the Tris buffer (0.1 M, pH 7) containing paraquat (1 mM) as shown in Fig. 6a. It can be observed that peak current increases with increasing scan rates according to typical diffusion-controlled reaction. Meanwhile,

the peak potentials shift to more negative potentials with increasing scan rate due to over reduction by the PbO-NPs. The peak currents are plotted as a function of square root of scan rate as shown in Fig. 6b. A good linear relationship can be obtained in the range of  $100\text{--}500 \text{ mV s}^{-1}$ . This indicates that the process is fully controlled by the diffusion of the electroactive species on the Pb-O NPs.<sup>45</sup> The results of the peaks current can be explained by the Randles-Sevcik's equation as given in eqn

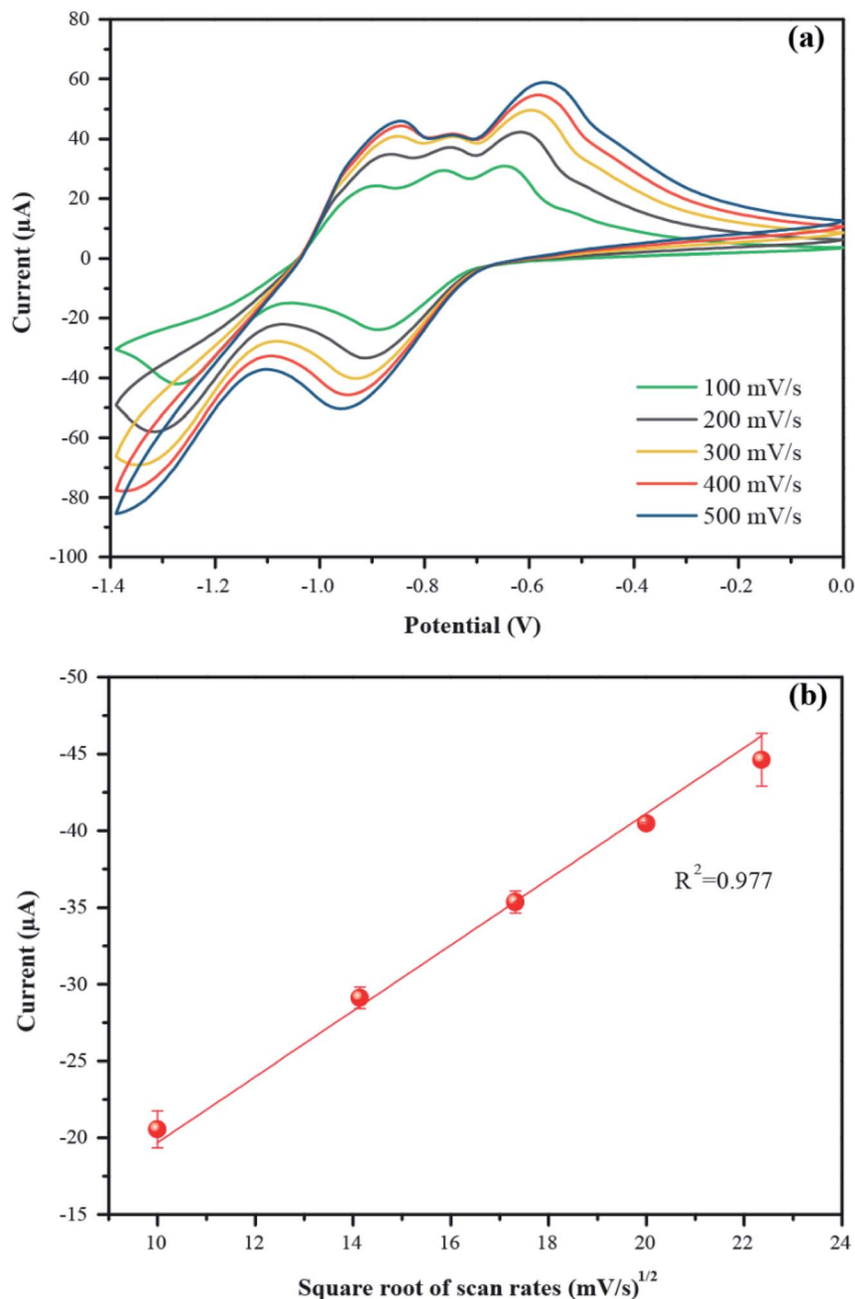


Fig. 6 (a) Cyclic voltammograms of PbO-NPs sensor at the scan rates in the presence of 1 mM paraquat and (b) their cathodic peak currents versus scan rates.

(2). The linear regression equation was  $I_p(\mu\text{A}) = -2.886 v^{1/2} (m v^{1/2} s^{-1/2}) + 0.119$  for  $P_1$  ( $R^2 = 0.998$ ). In

addition, the value of  $A_{\text{eff}}$  is calculated from the slope of fitted line based on eqn (3) as  $6.53 \times 10^{-3} \text{ cm}^2$  for  $P_1$ .

$$I_p = 2.69 \times 10^5 n^3 A_{\text{eff}} D^{1/2} C_0 v^{1/2} \quad (2)$$

where  $I_p$  = the cathodic peak current,  $n$  = the number of electrons transferred in the redox event,  $A$  = the effective surface area of the electrode ( $\text{cm}^2$ ),  $D$  = diffusion coefficient of tris hydroxymethyl

aminomethane ( $\text{cm}^2 \text{ s}^{-1}$ ),  $C_0$  = the bulk concentration of the analyte ( $\text{mol cm}^{-3}$ ),  $v$  = the scan rate ( $\text{mV s}^{-1}$ ).

$$A_{\text{eff}} = \left( \frac{m}{2.69 \times 10^5 n^3 C_0 D^{1/2}} \right) \quad (3)$$

For the paraquat in 0.1 M Tris buffer,  $m$  is to slope,  $n = 1$ ,  $C_0 = 1 \times 10^{-6} \text{ mol cm}^{-3}$  and  $D = 2.7 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$ .

The optimum conditions (pH 7,  $100 \text{ mV s}^{-1}$ ) were selected for studying the electrochemical performance of PbO-NPs



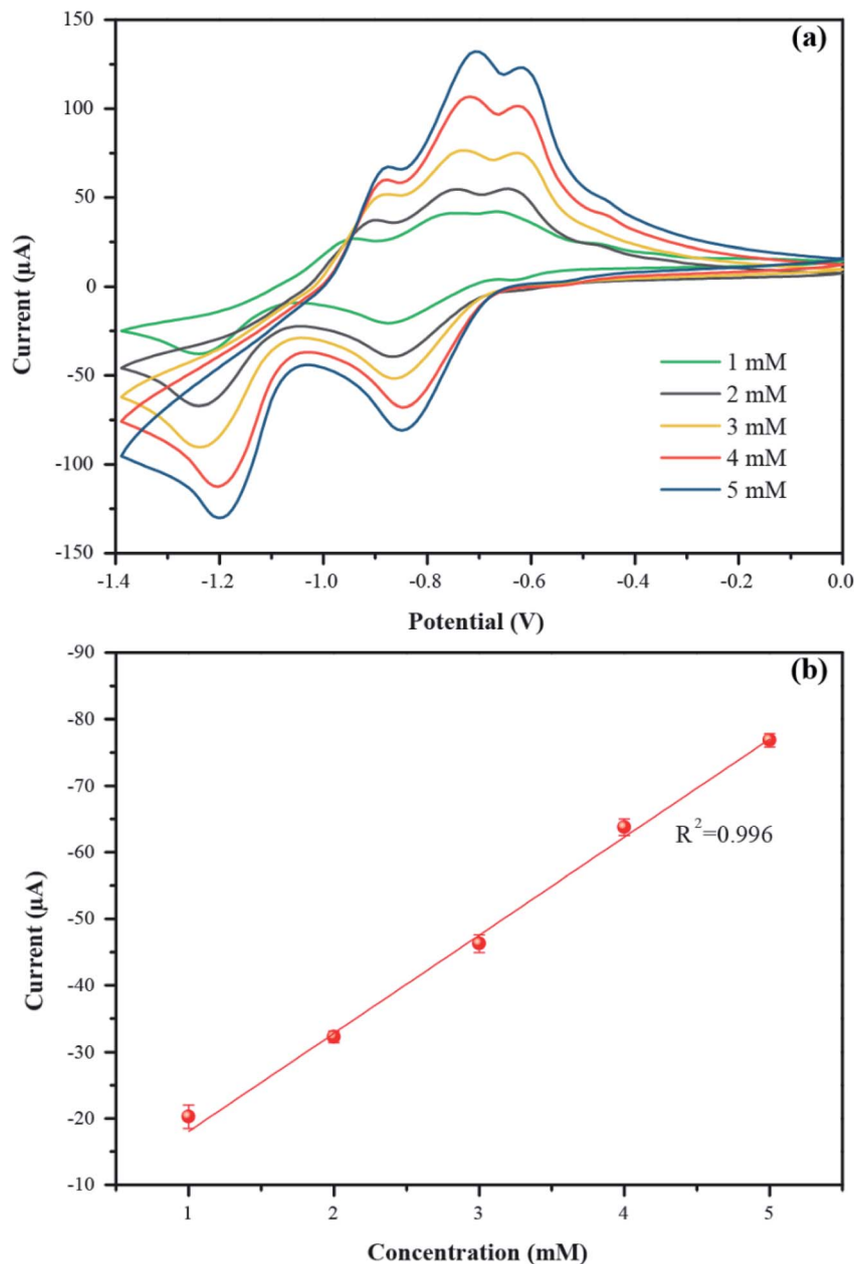


Fig. 7 (a) Cyclic voltammety responses of PbO-NPs sensor in the different concentrations of paraquat (0.1 M Tris buffer, pH 7, scan rate  $100 \text{ mV s}^{-1}$ ) and (b) calibration plot between cathodic peak currents and paraquat concentrations.

sensor in determination of paraquat at different concentrations ranging from 1 mM to 5 mM as shown in Fig. 7a. The cathodic peak currents increase linearly as increasing of paraquat concentrations. The linear regression equation can be well fitted as  $I_{P1} (\mu\text{A}) = 14.74C (\text{mM}) + 3.32$  with correlation coefficients  $R^2$  of 0.996 as shown in Fig. 7b. The sensitivity is calculated to be  $208.63 \mu\text{A mM}^{-1} \text{ cm}^{-2}$  based on eqn (4).<sup>46,47</sup> The limit of detection (LOD) is 1.1 mM as follows eqn (5). The PbO-NPs SPE sensor reveals high sensitivity and low LOD. High sensitivity of PbO-NPs SPE sensor to paraquat may be attributed from good electrostatic interactions. In fact, the paraquat molecules are quaternary ammonium compounds that can be ionized to di-positively charged leads to enhancement of electrostatic

interactions between negatively charged PbO-NPs/SPE and positively charged paraquat ions.<sup>48</sup>

$$\text{sensitivity} = \frac{m}{A} \quad (4)$$

where  $m$  = slope of calibration plot ( $\mu\text{A mM}^{-1}$ ),  $A$  = area of the working surface electrode ( $\text{cm}^2$ ).

$$\text{the limit of detection (LOD)} = \frac{3.3\text{SD}_{\text{blank}}}{m} \quad (5)$$

where SD = standard deviation of peak current of blank ( $5.12 \mu\text{A}$ ) for 30 cycles,  $m$  = slope of the calibration curve ( $14.74 \mu\text{A mM}^{-1}$ ).



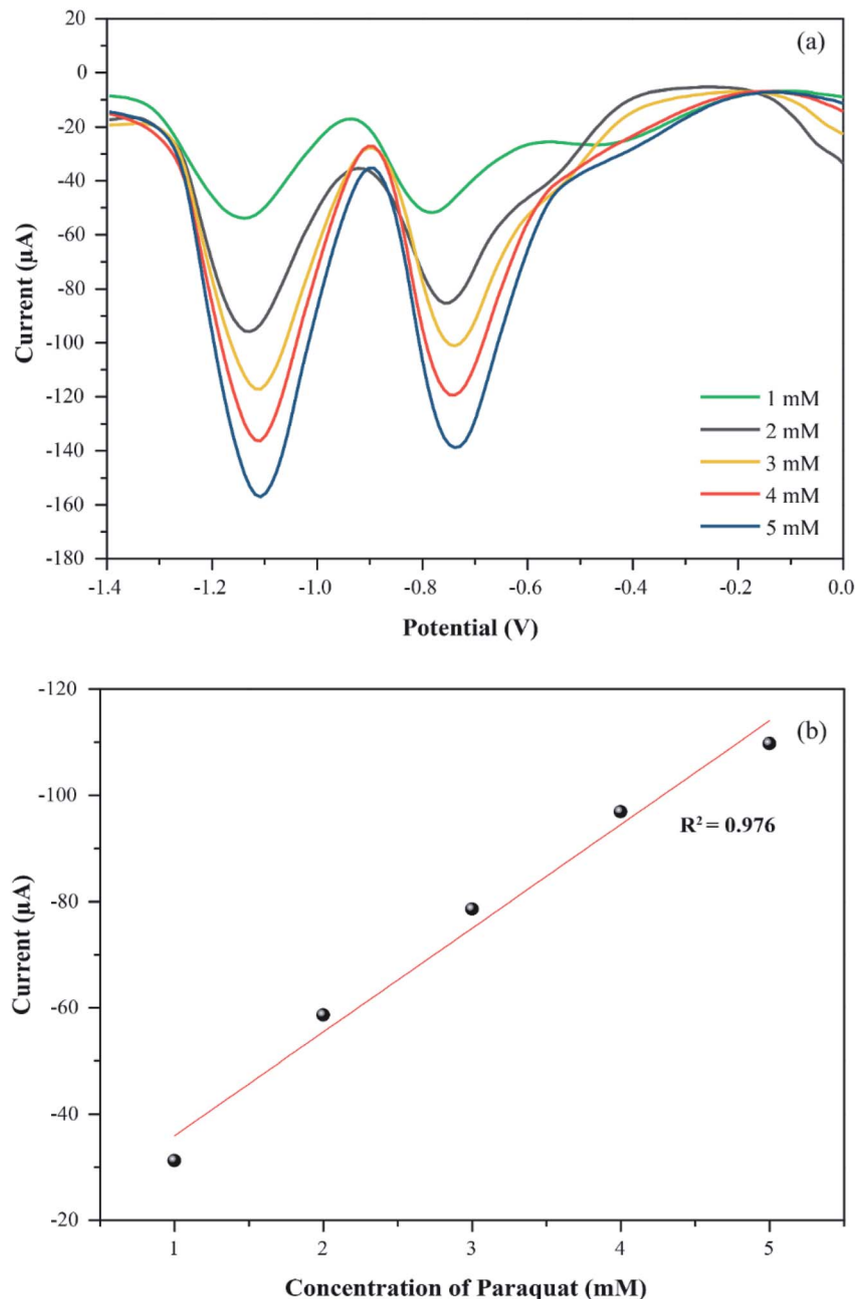


Fig. 8 (a) Differential pulse voltammetry responses of PbO-NPs sensor in the different concentrations of paraquat (0.1 M Tris buffer, pH 7) and (b) calibration plot between cathodic peak currents and paraquat concentrations.

To verify the accuracy of CVs technique for quantitative analysis of paraquat, the differential pulse voltammetry (DPV) was used to compare the electrochemical performance of the PbO-NPs SPE sensors to various paraquat concentrations as shown in Fig. 8. The DPV results exhibit the similar behaviors on cathodic peak currents and voltage according to CVs measurements. The DPV current peaks increase linearly with increasing paraquat concentration ( $I_{P1} (\mu A) = 19.53C (\text{mM}) + 16.41$ ,  $R^2 = 0.976$ ). The sensitivity based on DPV is  $276.63 \mu A \text{ mM}^{-1} \text{ cm}^{-2}$  that is higher than CVs technique because DPV is more sensitive than CV. Therefore, the CVs are good enough

and suitable technique for the quantitative detection of paraquat in this work.

To evaluate the reproducibility and repeatability, the four independent PbO-NPs sensors were fabricated and tested with 10 mM paraquat under the optimal electrochemical condition. The CVs of PbO-NPs sensors were carried out for 30 cycles as shown in the Fig. 9a. It can be seen that the cathodic peaks current increases a bit with increasing the number of scans due to the charge accumulation on sensing film. No any large shift in voltages and current were found. The average standard deviation (SD) values of 4 times measurements were



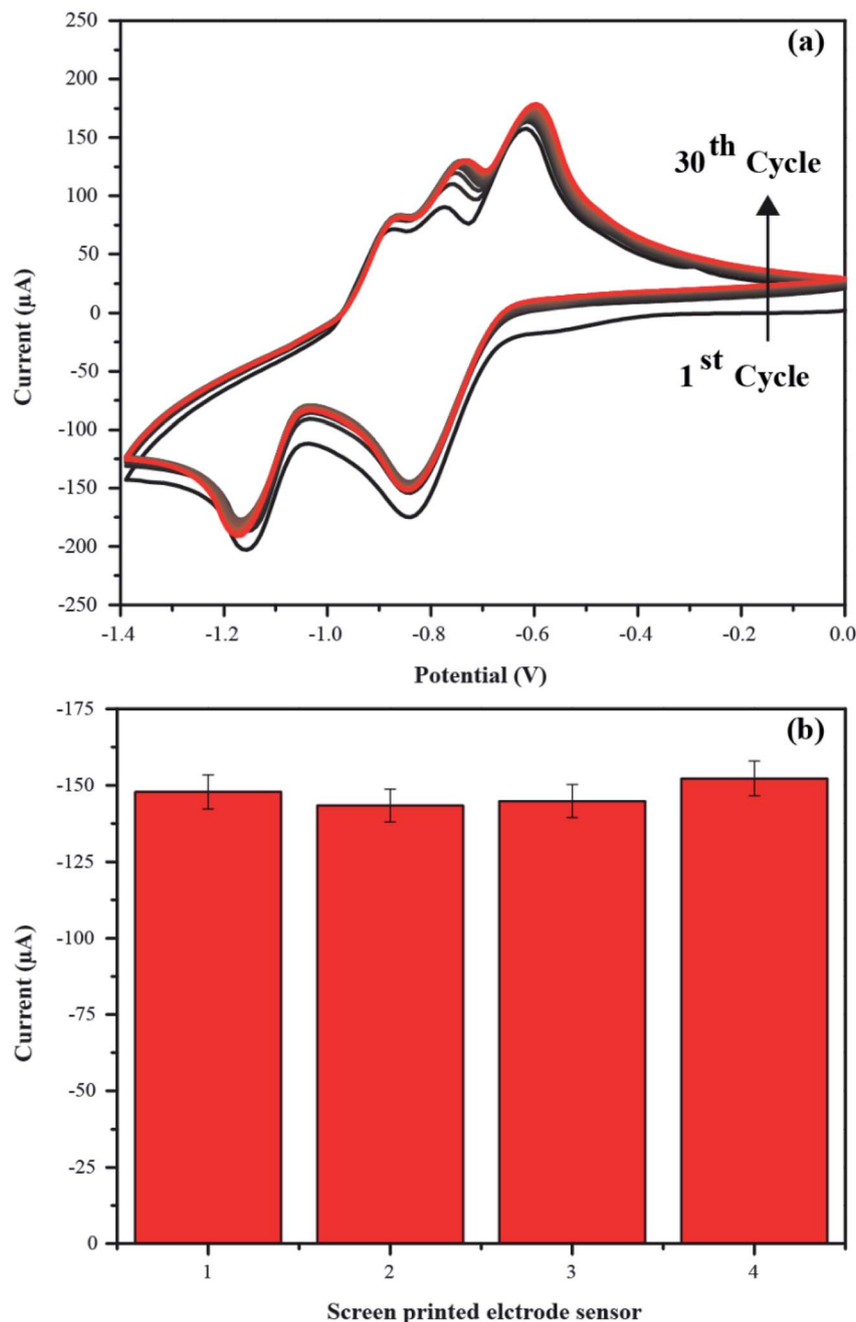


Fig. 9 (a) Cyclic voltammety responses of PbO-NPs sensor in presence of 10 mM paraquat as a function of the number of cycles and (b) their cathodic peak currents of four independent fabricated PbO-NPs sensors.

approximately 4.56% for peak 1 as shown in the Fig. 9b. The CVs of four independent fabricated PbO-NPs sensors are displayed in Fig. S2 in the ESI.† Thus, it suggests the high stability and reproducibility of the sensor.

### 3.3 Chronoamperometric measurement of PbO-NPs sensor

To investigate the selectivity of PbO-NPs sensor, the chronoamperometry (CA) method was used for studying the relationship between response of reduction current and time at fixed potential. The paraquat is a well-known herbicide. Thus,

the other popular herbicides such as glyphosate, glufosinate-ammonium and butachlor-propanil have been selected for interference effects in this study. The PbO-NPs sensor was applied with a potential of  $-0.8$  V for 240 s by addition of each interfering solution of 100 µL (10 mM) at every 60 s. The recovery percentage can be calculated as following eqn (6).

$$\% \text{ recovery} = \left( \frac{I_s - I_U}{I_s} \right) \times 100 \quad (6)$$

where  $I_s$  (µA) = response current for paraquat detection,  $I_U$  (µA) = response current for interfering compounds detection.



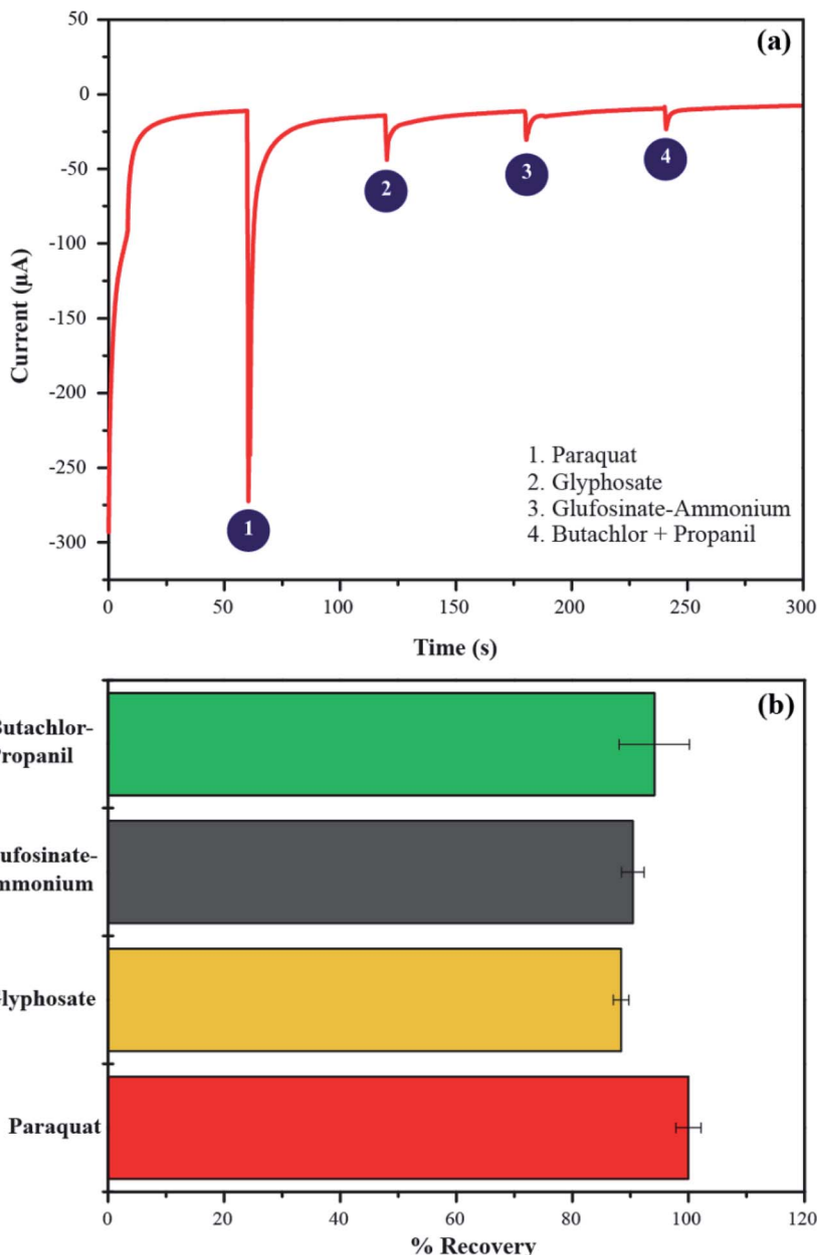


Fig. 10 (a) Dynamic chronoamperometric current responses of PbO-NPs sensor to various popular herbicides and (b) their % recovery of the PbO-NPs sensors (the error bar is SD from the currents of CA with three repeated measurements).

Table 1 Determination of paraquat in samples

Sample	Added concentration (mM)	Found concentration (mM)	Recovery (%)	RSD (%)
Juice	10	10.11	101.10	5.95
Milk	10	8.09	80.90	4.33

A real-time dynamic chronoamperometric plot is displayed in Fig. 10a. It reveals that the PbO-NPs sensor exhibits high selectivity to paraquat over other popular herbicides. The

measurements were performed using three PbO-NPs sensors. The average percentage of recovery is calculated to be 89–95% as shown in Fig. 10b. The sources of the error bars are SD from the currents of CA with three repeated measurements (four independent PbO/SPE sensors). It should be noted that the peak current of 10 mM paraquat as shown in Fig. 10a is higher than the peak current as shown in Fig. 9. The different currents come from different measurement methods between CV and CA technique. The CV experiment applies multiple potential cycles while the CA uses a fixed square-wave potential to apply the working electrode that may have some current fluctuation leading to difference currents between the two methods. In



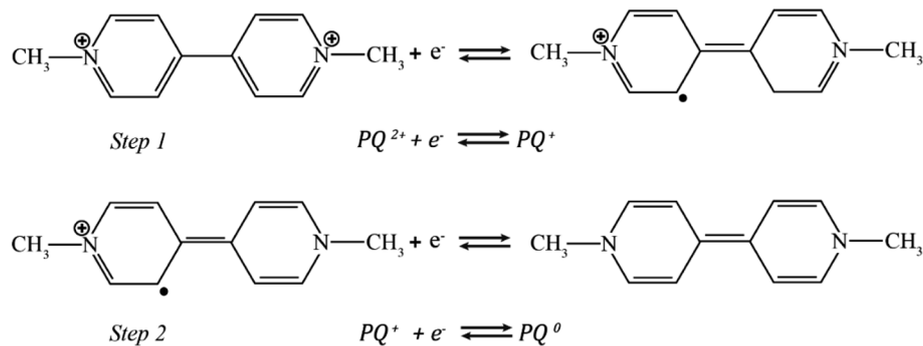


Fig. 11 Paraquat sensing mechanism for electrochemical behavior.

In addition, the measurement of current response in the CA method was applied with a steady potential ( $-0.8$  V) and times (every 60 s). The accumulation of charges between a conductive surface of electrode and electrolyte in double-layer can be formed and increased a current response over CV measurement. However, it's not make any effects on selectivity tests. The currents obtained from interactions of different herbicides can be compared with the similar method.

To further evaluate the sensitivity and selectivity of PbO-NPs SPE sensor, the CVs responses to glyphosate, glufosinate-ammonium and butachlor-propanil are shown in Fig. S3 in the ESI.† Due to its superior reaction sites of PbO and strong electrostatic interactions between negatively charged PbO-NPs/SPE and positively charged paraquat ions, the PbO-NPs SPE sensor exhibits high sensitivity and selectivity to paraquat at a potential of  $-0.8$  V. However, the PbO-NPs SPE sensor is also sensitive to glyphosate, glufosinate-ammonium and butachlor-propanil at different applied potentials with different electrochemical behaviors. Therefore, the selectivity of electrochemical sensors can be considered with a fixed applied potential matching with chemical reduction/oxidation between sensing materials and targeted chemical species.

### 3.4 Paraquat sensing mechanism

The electrochemical response occurs from interactions between paraquat and the PbO-NPs on a SPE. The concept of the sensing mechanism is described by charge transfer. The structure model of the metal oxide electrode in aqueous solution is generally divided into two distinct regions: a compact layer and a diffuse layer. The compact layer includes an inner Helmholtz layer (IHP) and an outer Helmholtz layer (OHP).<sup>49,50</sup> Therefore, when the aqueous solution contains paraquat molecules and water molecules under the electrostatic force, the hydrogen, or hydroxyl ions (the water dissociation), the positively charged paraquat are absorbed onto the negatively charged surface of the PbO-NPs resulting in charge transfer at the IHP of the compact layer. The interaction of the PbO-NPs and paraquat occurs through the  $\pi$ - $\pi$  stacking of aromatic ring forming charge transfer complexes. The negatively charged surface of the PbO-NPs will help to promote and attract the accumulation of paraquat leading to high sensitivity and selectivity of sensor.

In addition, the paraquat detection can be monitored by the two-reduction process which transfers one electron in each step. Therefore, the possible mechanism of paraquat detection may be described in the equation of Fig. 11 as follows<sup>51-53</sup>

Step 1: the first reduction state involves to the formation of the radical paraquat cation ( $PQ^{\bullet+}$ )

Step 2: the second reduction state involves to the transformation of  $PQ^{\bullet+}$  into neutral molecules  $PQ^0$ .

### 3.5 Detection of paraquat contamination in samples

Practical application of the PbO-NPs sensor for paraquat determination was estimated in samples such as milk and instant mixed vegetable-fruit juice. The 200  $\mu\text{L}$  of paraquat solution was added into the 200  $\mu\text{L}$  of each sample (10 mM paraquat concentration). The mixture was stirred for 1 min. The 100  $\mu\text{L}$  of mixture was dropped on PbO-NPs SPE sensor and examined by using CV analysis under optimized conditions (pH 7, 100  $\text{mV s}^{-1}$ ). The summarized data of recovery percentage and the relative standard deviation obtained is presented in Table 1. The PbO-NPs sensor can determine the paraquat concentration in mixed vegetable-fruit juice and milk with % recovery of 101.10% and 80.90%, respectively, referring well identify the paraquat contamination in juice. In case of milk, it consists of fat, proteins, lactose, etc. that can strongly affect on interaction between PbO-NPs surface and paraquat molecules resulting to underestimate paraquat concentration. This PbO-NPs sensor may prefer to be used in samples with low fat, proteins and lactose.

$$\% \text{ recovery} = \left( \frac{C_F}{C_A} \right) \times 100 \quad (7)$$

where  $C_F$  = found concentration of paraquat in spiked sample (mM) that obtained from cathodic peak current of CVs results and then calculated to paraquat concentration according to calibration linear regression plot as shown in Fig. 7b.  $C_A$  = added concentration (mM).

## 4. Conclusion

In summary, the electrochemical sensor for the paraquat detection has been successfully developed by modifying with



the PbO-NPs onto a screen-printed Ag working electrode surface. The PbO-NPs were synthesized *via* sparking method that offers a simple, rapid and low-cost method for producing nanoparticles under room temperature and normal pressure. The synthesized nanoparticles are spherical shape with the average size of  $22.95 \pm 5.46$  nm. The electrochemical behaviors have been investigated using the cyclic voltammetry. Under the optimal experimental conditions (scan rate of  $100 \text{ mV s}^{-1}$ ,  $0.1 \text{ M}$  Tris buffer (pH 7)), the PbO-NPs/SPE sensors demonstrate the redox reaction of  $\text{PQ}^{2+}$  by showing two cathodic peak currents around  $-0.8 \text{ V}$  and  $-1.1 \text{ V}$ . The PbO-NPs/SPE sensor exhibits very good linear response with concentrations in range from  $1 \text{ mM}$  to  $5 \text{ mM}$  ( $R^2 = 0.996$ ) with high sensitivity of  $204.8 \mu\text{A mM}^{-1} \text{ cm}^{-2}$ . The LOD of the PbO-NPs/SPE sensor is  $1.1 \text{ mM}$  ( $0.28 \mu\text{g L}^{-1}$ ) that is lower than maximum permissible limit for total pesticides in drinking water ( $0.5 \mu\text{g L}^{-1}$ ), for surface waters ( $1\text{--}3 \mu\text{g L}^{-1}$ )<sup>54</sup> and for paraquat residues in drinking water that varies from  $1\text{--}30 \mu\text{g L}^{-1}$  in different countries.<sup>55</sup> Moreover, the PbO-NPs sensor shows high selective to paraquat over other popular herbicides such as glyphosate, glufosinate-ammonium and butachlor-propanil. The developed sensors can be integrated into a smartphone for on-site field testing. Based on the presented methods and results, it is hoped that this work can be an alternative way to develop the electrochemical paraquat sensor for commercialization in future.

## Conflicts of interest

There are no conflicts to declare.

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