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Design and Preparation of Nanostructured Prussian Blue Modified Electrode for Glucose Detection

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1. Introduction

High sensitivity and anti-interference ability are critical parameters in glucose biosensor design, because of the complexity of blood composition. Prussian blue (PB) is a hexacyanoferrate with two different iron valences (+2 and +3) and was initially developed as a blue pigment in the 1700s (Bartoll, 2008). Recently researchers found that PB is an excellent material for use in the fabrication of glucose biosensors, because of its non-toxicity, high electrocatalytic activity, and low overpotential detection (Ricci & Palleschi, 2005; Karyakin et al., 2007; Wang, 2008). During glucose detection, hydrogen peroxide ($H_2O_2$) is produced via the enzymatic oxidation of glucose. Subsequently, PB reduces the $H_2O_2$ and transfers the electrons to generate a current response on the electrode surface. Thus, PB is the mediator of electron transfer in the detection process, making the electron sensitivity of PB - a key component of biosensor performance.

Chemical deposition and electrodeposition are the main processes used to synthesize a PB film on the electrode surface (Itaya et al., 1982; Zakharchuk et al., 1995). However, the sensitivity of the electrode modified with a PB film cannot be controlled, because of difficulties in film construction. Millward et al. (2001) used a self-assembly approach to deposit a PB film on a gold electrode, which greatly accelerated the development of PB-based biosensors. The advantages of applying nanostructured materials for improving biosensor performance have been recognized in recent years, especially for improving sensitivity (Cella et al., 2010; Cao et al., 2010). The utilization of nanoparticles can enhance electrode performance, even if the same materials are used. We developed a PB nanostructure to produce a high sensitivity biosensor for glucose detection, which can be directly grown on the electrode surface.

We previously synthesized PB nanoparticles on a platinum (Pt) electrode surface using a self-assembly approach (Liu et al., 2009). We found that the size and quantity of PB particles greatly affected $H_2O_2$ detection performance. After immobilization of the glucose enzyme, this type of biosensor was sensitive to trace concentrations of glucose in solution (Liu et al. 2009). However, it was difficult to form a regularly structured PB particle film, when synthesis was conducted via self-assembly. Thus, we further developed the self-assembly approach to deliver a novel method for obtaining a regular film of PB crystals, are thereby improving biosensor sensitivity and effectiveness.
2. Mechanism of H\textsubscript{2}O\textsubscript{2} detection

PB is typically produced from a reaction between two compounds, K\textsubscript{4}[Fe(CN)\textsubscript{6}] and FeCl\textsubscript{3} (see equation 1).

\[
3K_4[Fe(CN)_6] + 4FeCl_3 \rightarrow Fe_4[Fe(CN)_6]_3 + 12KCl
\]
\[
K_4[Fe(CN)_6] + FeCl_3 \rightarrow KFe[Fe(CN)_6] + 3KCl
\]

KFe[Fe(CN)\textsubscript{6}] and Fe\textsubscript{4}[Fe(CN)\textsubscript{6}]\textsubscript{3} are soluble and insoluble PB, respectively, and both species are required. The unit cell of PB is a face-centered cubic structure with lattice parameters, a = b = c = 10.143 Å. It is interesting that neighboring iron ions have different valences. The Fe\textsuperscript{2+} ion, in the Fe(CN)\textsubscript{6}\textsuperscript{4+} complex, is readily reduced to the Fe\textsuperscript{3+} ion. The signal generated by electron transfer from Fe(CN)\textsubscript{6}\textsuperscript{4+} is generally used in electrochemical detection to characterize the properties of the PB film.

Fig. 1. (a) The structure of the PB unit cell. (b) Mechanism of H\textsubscript{2}O\textsubscript{2} detection.

PB has two functions when used as a material in biosensor fabrication, i.e., electrocatalysis and electron transfer: When a potential (typically - 0.05 V) is imposed on a PB modified electrode, the PB will acquire electrons and move into its reduced state, Prussian White (PW). If H\textsubscript{2}O\textsubscript{2} is present in the detection system, PW rapidly reduces H\textsubscript{2}O\textsubscript{2} to OH\textsuperscript{-}. After donating electrons, PW reverts back to PB (Fig. 1). The direction of electron transfer during this process is from the electrode surface to H\textsubscript{2}O\textsubscript{2}. Thus, electrocatalysis and electron transfer determine overall electrode performance.

Regular crystalline structure is a morphological feature that has been widely demonstrated to enhance catalytic ability (Liu et al., 2010; Prevot et al., 2011). However, it is difficult to control regular PB crystal growth using conventional preparative methods, because of its rapid reaction rate during crystal formation. Consequently, novel methods are required to facilitate the regular crystal structure growth in the PB film.

3. PB crystal formation using a self-assembly approach

Self-assembly is a common approach for production of PB-modified electrodes. Scheme 1 shows the process of PB synthesis, which starts with a pre-coating layer of polyelectrolyte, to provide electrostatic adsorption sites, followed by dipping the electrode into a reactive solution for PB formation. Poly(diallyldimethylammonium chloride) (PDDA) is typically
applied as the cation polyelectrolyte layer (Hornok & Dékány, 2007; Li et al., 2008). This layer provides a foundation for further layer deposition, which means that its properties are essential for regular PB growth. The regular structure of PB is not readily formed via self-assembly, but this approach has been successfully applied to regular crystal growth for other materials (Wang et al., 2009; Guang et al., 2009). The PB self-assembly problem might be attributable to first layer selection, and temperature can greatly affect polymer properties. We determined the optimum method for regular crystal formation in a PB film by controlling PDDA load temperature.

Scheme 1. Production of a PB-modified electrode using a self-assembly method.

We investigated the properties of PDDA layer with respect to changes in temperature, prior to the self-assembly of PB. Electrochemical impedance spectroscopy (EIS facilitates observation of the structure and charge transfer behavior of polymer films. We compared the impedances of bare Pt and the electrodes loaded by PDDA at different temperatures. Electron transfer is essential for sensitivity in the detection of $\text{H}_2\text{O}_2$. However, PDDA is not an excellent mediator for electron transfer, but we were more concerned with the resistance provided by the PDDA layer. Fig. 2 shows the results of impedance experiments, illustrating the electrode property at high frequencies due to the electron transfer kinetics of redox species at the electrode surface. The linear region is dominated by the mass transfer of probe species from the bulk solution to the interface region. We found that increasing the adsorption temperature led to a reduction in interface resistance. We compared the exact resistance values using a classical Randles circuit (inset of Fig. 2), the results of which were applied to fitting the EIS curves. The values of $R_\text{ct}$ which resulted in electron transfer ability were calculated as 4.741, 5.932, 9.722, 11.47, and 13.03 $\Omega$ respectively, for the bare Pt electrode, and the electrode with PDDA adsorbed at 35, 30, 25, and 20 °C. The resistance of the unmodified Pt electrode was the lowest, and the addition of PDDA gradually decreased the electron transfer ability. The impedance was enhanced by a reduction of the PDDA adsorption temperature. The fluidity of the polymer was very poor at low temperatures, which led to severe surface aggregation of the substrate with detrimental effects on PB assembly. This prompted the investigation of surface coverage. The fractional electrode coverage rate ($\theta$) of PDDA was estimated according to equation 2.
$\theta = 1 - \frac{R_{ct}^{bare}}{R_{ct}^{PDDA}}$ (2)

Where $R_{ct}^{bare}$ is the charge transfer resistance value for the bare Pt electrode, and $R_{ct}^{PDDA}$ is the same value for the PDDA modified electrode at different adsorption temperatures. The calculated results were 0.2062, 0.5123, 0.5867, and 0.6361 for 35, 30, 25, and 20 °C, respectively. By combining these results with the resistance results, we determined why resistance did not decrease as a function of temperature increase. PDDA did not cover the whole Pt surface and electrons preferred to transfer in areas without polymer because of the lower resistance. Local preferences for electron transfer resistance were consistent with variations in polymer coverage.

Fig. 2. The EIS diagram for the bare electrode and the electrodes loaded with a PDDA layer at 20, 25, 30, and 35 °C. The inset shows the equivalent circuit used for PDDA modified electrodes, where $R_{ct}$ is the charge transfer resistance, $R_s$ is the solution resistance, $Z_w$ is the Warburg impedance due to the mass transfer from the redox probe to the electrode, and $C_{dl}$ is the double layer capacitance.

Thus, the improvement of electron transport, demands that PDDA is loaded as sparingly as possible. This is because a high loading of PDDA further hinders assembly of PB, which means that a correct loading balance must be achieved.

The affect of PDDA adsorption temperature on PB morphology, and performance, were evaluated to determine the optimum production temperature, and provide a higher electrode performance via the synthesis of a regular PB structure. We investigated whether a regular PB morphology could be achieved by adjusting the PDDA adsorption temperature. Fig. 3 (a) to (d) shows the great differences in surface morphology of PB on modified electrodes, when the PDDA layer was adsorbed at different temperatures. When the PDDA was deposited at 20 °C, the film surface was covered with irregular particles, but the particles grew bigger as the temperature increased to 25 °C. As the temperature increased, the whole surface changed and, instead of irregular particles, an abundance of nanocubes was observed on the electrode surface, with each nanocube measuring ~100 nm. These
regular crystals formed the film and although some nanocubes remained on the surface, the amount decreased. We also measured the PB concentrations on the electrodes. Fig. 3 (e) shows that the PB concentration was 6.059, 6.771, 7.03, and 4.827 nmol cm$^{-2}$ at temperatures of 20, 25, 30, and 35 °C, respectively. At the low temperature, the amount of PDDA covering the Pt surface was so large that it caused the aggregation of PB particles. Thus, an excess of crystal nuclei might lead to imperfect growth of PB crystals.

Fig. 3. The FESEM images of PB modified electrodes, where PDDA were adsorbed at different temperatures: (a) 20 °C; (b) 25 °C; (c) 30 °C; (d) 35 °C. (e) Shows the PB concentrations on electrodes at different PDDA adsorption temperature.

The film was comprised of small PDDA particles, which provided adsorption sites for PB assembly. The PB concentration was low at high temperature, because of low PDDA coverage. We showed that a PDDA layer adsorbed at low temperature resulted in a strong
impedance to electron transfer, so we selected 30 °C as the optimal temperature for PDDA adsorption. According to the performance detection procedure, the sensitivity of an electrode prepared at 30 °C was 1179.6 mA M⁻¹ cm⁻², which was much higher than electrodes prepared with no control over PDDA adsorption temperature (Liu et al., 2009). This improved self-assembly approach to PB film formation successfully produced nanocubic PB crystals and enhanced the sensitivity of H₂O₂ detection. This study demonstrated the constraints on electron transfer impedance imposed by the PDDA layer so we next determined the factors affecting the regular growth of PB in the absence of a polyelectrolyte.

4. Aerosol deposition (AD) for synthesis of PB nanocubes

4.1 Synthesis mechanism

Synthesis of a PB film with a regular morphology, in the absence of polyelectrolyte, demands control of the formation rate during regular PB film growth. Current approaches require PB formation in reactive solutions. However, the concentration of molecules in solution is very high, and PB is produced in seconds when solutions of K₄[Fe(CN)₆] and FeCl₃ are mixed. Thus, it is desirable to achieve PB formation with lower concentration solutions. Aerosols are composed of tiny droplets, which are so small that they are readily suspended in the air. There has been a recent growth in interest in the application of aerosols (Boissiere et al., 2010). The advantages of aerosols are as follows: small volumes lead to micro-reactions and the production of small particles; fewer molecules per unit are required in controlled reactions; the suspension of aerosols avoid the influence of gravity on preparation and promotes uniform growth. We aimed to produce regular PB crystals, without polyelectrolyte, by adopting the AD approach instead of conducting the reaction in solution.

If aerosols are to be used to deposit PB directly onto a Pt electrode, it was essential that an interaction occurred between Pt and reactive aerosols. Before we conducted the experiments, we confirmed that aerosols can be deposited on the electrode surface by virtue of Van der Waals force between the carbon and Pt atoms, according to the results of a Density Functional Theory (DFT) simulation (Chu et al., 2009). This simulation helped us to define the deposition order, i.e., the K₄[Fe(CN)₆]₃ aerosol should be initially attracted onto the Pt surface, because of the carbon conformation. The deposition time of the two aerosols was the same.

4.2 Morphology control

As shown in Fig. 4, aerosols of Fe(CN)₆⁴⁻ and Fe³⁺ were deposited on the Pt surface to form the PB film. The deposition time was a controlled parameter that determined the concentration of aerosols. The small size of liquid used meant that aerosols were more readily affected by environmental temperature, which affected the growth of crystals. Thus, deposition time and temperature were the main controlled factors in the whole process (Chu et al., 2010).

Cyclic voltammetry (CV) experiment showed that the PB concentration on the modified electrode directly correlated with deposition time. Fig. 5 shows the maximal and minimal current values of each curve for the redox peaks of Fe²⁺/Fe³⁺, caused by electron transfer from PB to PW and PW to PB. PB concentrations can be calculated using the data by following equation 3 (Grundler, 2007).
\[ \Gamma_T = \frac{Q}{nF \Delta E} \]  

Where \( \Gamma_T \) represents the concentration of PB, \( Q \) is a single peak (either a reduction or oxidation peak), \( n \) is the average transfer of electrons calculated by \( \frac{57}{\Delta E} \) (\( \Delta E \) is the difference in potential between the redox potentials), \( F \) is the Faraday constant, and \( A \) is the area of the electrode.

Fig. 4. PB film preparation using the aerosol deposition method.

Fig. 5. The CV results for PB-modified electrodes produced for 2, 3, 3.5, 4, 5, and 6 hours. The preparation temperature was kept at 25 °C. The inset shows the diagram of calculated PB concentrations.
We investigated the PB crystal morphology for electrodes prepared at 25, 35, and 45 °C. Fig. 6 shows that regular nanocrystals formed on the electrode surface at 25 and 35 °C. When the film was prepared at 25 °C, the shape of crystals was cuboid-like with a size of ~500 nm, whereas at 35 °C, the crystals were smaller and possessed a more obviously cubic structure. Thus, it was determined that 35 °C was appropriate for the synthesis of larger volumes of crystals with a regular structure. A further increase of the temperature to 45 °C, led to the surface being populated with small particles with an undesirable morphology. Higher temperature also caused serious cracks in the film due to differences of thermal expansion between the PB film and the metal of the Pt electrode. Fig. 7(d) shows an HRTEM image of the growth of crystals produced in one direction, where the d-spacing of the lattice fringes was measured as 0.500 nm, which is consistent with the (200) lattice plane of PB in the PDF card (JCPDS card no.73-0687).

Temperature strongly affects PB morphology, because deposition of aerosols can reduce the amount of molecules available to participate in the reaction. Thus, the driving force of chemical equilibrium is reduced compared to reactions in solution, thereby slowing down the reaction product formation rate. High temperature might accelerate the formation of nuclei for further growth, but the availability of more growth cores could lead to an increase in the amount, but a decrease in the size, of PB crystals. Further problems occur if the temperature is increased further, because the interaction between PB and Pt is governed by the van der Waals force, which is a weak force. Thus, if the temperature is too high molecules will escape from the restraint of interaction due to high thermal motion activity. This can lead to inadequate and imperfect PB crystal growth.
Regular crystals with perfect edges lead to higher sensor performance, because they increase the catalytic activity of biosensors. Thus, a PB electrode with nano-cubic structure is expected to show enhanced sensitivity in the detection of H$_2$O$_2$.

### 4.3 Performance in H$_2$O$_2$ detection

Chronoamperometry is often used in the performance measurement of electrochemical biosensor detection performance. We tested the detection performance of three electrode systems; the Pt electrode we developed as described above, a counter Pt electrode, and an Ag/AgCl electrode. The process was conducted in a phosphate buffer solution (pH = 6.5) to ensure system stability. The potential imposed on the PB modified electrode was - 0.05 V. The stirring was provided to ensure the uniform dispersion of H$_2$O$_2$. After running the baseline, 10 μM H$_2$O$_2$ was repeatedly added into the buffer solution at a standard time interval.

![Fig. 7. Current response during H$_2$O$_2$ detection. The inset shows a simulation diagram of current versus H$_2$O$_2$ concentration.](image)

Fig. 7 shows the change in current after the introduction of H$_2$O$_2$. After 19 treatments of H$_2$O$_2$ addition, the step of the current signal remained clear and stable. This shows that the PB construction maintains good stability with an increase of H$_2$O$_2$ concentration. The current response data had a linear relationship with the H$_2$O$_2$ concentration, which we simulated. The slope of this linear function indicated the H$_2$O$_2$ detection sensitivity of the PB modified electrode. The best sensitivity was calculated as 1163 mA M$^{-1}$ cm$^{-2}$. This performance of an electrode prepared using the AD method was higher than traditional preparation methods. The enhancement of sensitivity was attributed to the formation of nanocubic PB crystals. The nanocubes possessed perfect edges, which provided strong electrocatalytic sites for the H$_2$O$_2$ reduction. High crystallinity might also provide more PB sites to participate in detection per unit area. Electrons transfer from Fe$^{2+}$ and Fe$^{3+}$ proceeds along a linear pathway $Fe - C \equiv N - Fe$ in the PB unit cell. Thus, the synthesized cube structure matches the PB cell construction of a face-centered cubic lattice, which means that we produced a regular path for electron transport that might decrease the resistance, in terms of geometry.
This production method for performance enhancement was based on the formation of regular structures by decreasing the reaction rate. A common approach for performance improvement depends on the introduction of other compounds with high catalytic ability or favorable conductivity. We confirmed that the AD approach can produce the regular nanocubes of PB, but we further investigated whether the incorporation of other compounds during the production process might further increase the sensitivity. The inclusion of other compounds might affect the growth of PB, and the structure of the film, and further sensitivity improvements might accrue from cooperation between the two materials.

5. Codeposition of a PB-TiO$_2$ composite film using the AD method

Material selection during biosensor design demands attention to several critical requirements, i.e., non-toxicity, biocompatibility, stability, and cost. Codeposited compounds must also be compatible with PB. Titanium dioxide (TiO$_2$) is a common material, which satisfies all the required properties and is particularly suited to nanoparticle production, because it is very light. This allows the mixing of powdered TiO$_2$ with PB aerosols prior to suspension in the air. Synergistic reaction between semiconductors can increase performance, and both of PB and TiO$_2$ are semiconductors, with energy gaps of 1.58 and 3.2 eV, respectively. The main catalytic function of TiO$_2$ results from electron excitation of its valence bond to a conduction bond. However, 3.2 eV is a large gap which requires UV light to provide the energy for electron transit. The gap of PB is much lower than that of TiO$_2$. Combination of these two semiconductors means that they can pad each gap and make excitation easier (Yamamoto, 2009). Thus, doping TiO$_2$ nanoparticles into the PB film could improve the catalytic ability. This composite design might enhance the H$_2$O$_2$ detection sensitivity of the prepared PB electrode.

We next tested the effect of TiO$_2$ nanoparticle doping into deposition aerosols.

![Fig. 8](image-url)

Fig. 8. (a) The comparison of CV diagrams for electrodes with only PB and PB doped with TiO$_2$. The electrode coating was conducted for 6 hours, at 35 °C. (b) Comparison of calculated PB surface concentrations for both electrodes.

We added 0.1 mmol TiO$_2$ to the 0.01 mol/L FeCl$_3$ solution to achieve co-deposition of TiO$_2$ during the preparation process. The mixed solution was converted to aerosols using an ultrasonic nebulizer. The concentration of TiO$_2$ was very low because of its poor solubility. Higher amounts would interfere with the ultrasonic method used to produce aerosols. Fig. 8
(a) shows that redox peaks for the electrode with a composite film was lower than that with a PB film. After doping with TiO₂, the difference between the potentials of oxidation and reduction was larger. The PB concentration was reduced approximately four-fold, according to the calculation results shown in Fig. 8 (b). The deposition of TiO₂ particles in the reactive aerosols, led to them becoming highly packed on the electrode surface, which meant they occupied many growth sites that might have been occupied by PB, after electrode surface adsorption. The interaction between Pt and TiO₂ was very weak. Thus, without the packing of PB particles, TiO₂ particles might easily fall off and detach PB crystals growing around them. This would reduce the concentration of PB on the electrode surface. The widening of the distance between peaks was caused by an increase in electron transfer resistance. Electron transfer through TiO₂ is much harder compared with PB, due to the larger bond gap. The change of concentration might also affect the surface morphology so we investigated the micro-scale surface morphology using FESEM. Fig. 9 (a) shows that the film did not grow fully over the surface. There were numerous flaws on the film surface, making the film discontinuous. The lack of adequate PB prevented correct growth. Compared to the surface morphology shown in Fig. 6(b), the ground film was smooth instead of rough, under the same preparation conditions. However, we found that some nanocubes grew on the ground film, which was very different from the PB-only modified electrode. Nanocubes only grew in the ground film, when the film was only formed of PB crystals, but because of the large size of TiO₂ particles, the PB crystals rapidly aggregated around TiO₂. PB might struggle to form a regular shape, because of interference from TiO₂ particles, and regular crystals only formed freely on the ground film. The low PB concentration and defects in the film limits any synergistic functionality of PB with TiO₂, thereby decreasing electrode performance. We prolonged the deposition time in order to form a uniform film and, after increasing the time to eight hours, surface concentration was found to be 5.4 nmol cm⁻². Fig. 9 (b) shows that the ground film was uniform and contained no defects. More nanocubes were apparent on the surface and the size increased to ~2 μm. This was an obvious improvement on electrodes prepared for less than six hours. We compared H₂O₂ detection sensitivity with an electrode prepared under 35 °C for six hours.

![Fig. 9. FESEM images of PB-modified electrodes prepared at 35 °C for (a) 6 h, and (b) 8 h.](image-url)

The introduction of TiO₂ decreased the PB surface concentration and resulted in significant changes in film morphology, particularly the PB crystal shape. This meant that the performance of the prepared PB-TiO₂ composite film was expected to be no improvement on the PB-only film. However, based on the results of H₂O₂ detection (Fig. 10), the
sensitivity was calculated to be 1300 mA M$^{-1}$ cm$^{-2}$, which was a little higher than the PB-only film. This sensitivity is based on a PB concentration that was half that of the PB-only film. The catalytic function of TiO$_2$ was confirmed. The reductive ability of PB was weakened due to lower PB coverage, but TiO$_2$ particles contribute to overall catalytic function.

Fig. 10. H$_2$O$_2$ detection sensitivity comparison of electrodes with PB-only film and PB-TiO$_2$ composite film.

This preliminary work confirms that this approach can be applied to the production of regular structures PB and improve the performance of electrodes. Further research includes adjusting the ratio of the doping content and identifying other compounds with better solubility to substitute for TiO$_2$.

6. Conclusions and future prospects

Regular nanostructured PB can be synthesized using three methods: self-assembly with temperature controlled polyelectrolyte adsorption, the aerosol deposition approach, and the aerosol co-deposition approach. The regular structured PB film increased detection sensitivity of electrodes. Electron transfer direction in solution is from H$_2$O$_2$ to the electrode, and provision of a regular structured pathway for electron transport enhanced the detection efficiency of the modified electrode. This might require the production of oriented structures in the PB film, such as nanotubes or nanowires. However, if this assumption is to be satisfied a template or external force must be applied to set the PB growth direction. An alternative method is to identify compounds to dope the PB film and exploit their synergistic functionality, as we found with TiO$_2$. Such methods are not confined to PB fabrication and they may be appropriate in other fields requiring regular crystal synthesis.

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A biosensor is a detecting device that combines a transducer with a biologically sensitive and selective component. Biosensors can measure compounds present in the environment, chemical processes, food and human body at low cost if compared with traditional analytical techniques. This book covers a wide range of aspects and issues related to biosensor technology, bringing together researchers from 12 different countries. The book consists of 20 chapters written by 69 authors and divided in three sections: Biosensors Technology and Materials, Biosensors for Health and Biosensors for Environment and Biosecurity.

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