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Electrochemical sensing of estradiol benzoate using hydroxyapatite with three-dimensional channel frameworks†

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Ubiquitous estradiol benzoate (17β -estradiol-3-benzoate) in the environment and daily life has become a hot issue in society since it has been proven to be closely associated with metastatic tumour formation. In the current work, hydroxyapatite with three-dimensional channel frameworks has been successfully demonstrated to be highly sensitive for the electroanalytical sensing of this pollutant, with a detection limit of 24.4 ng L⁻¹ (0.065 nM), attributed to the enhanced electron transport capabilities and three-dimensional structure of the hydroxyapatite structure/surface.

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Introduction

Endocrine disrupting chemicals (EDCs) including estrogens, triazine, phthalates, etc. have gained global attention because of their well-known serious adverse effects. 1,2 As an important EDC, estrogens share the characteristics of EDCs, such as they may interact with the endocrine system of organisms and cause non-negligible adverse effects on the reproduction of wildlife and humans.³⁻⁵ One such example of an EDC is 17β-estradiol-3benzoate (E2B) which is a ubiquitous environmental agent. It is an estradiol analog commonly used in livestock for inducing weight gain and synchronising estrus cycles in heifers and cattle.6,7 However, as an analogue of EDC, E2B binds with estrogen receptors (ERs) which are expressed in human cells such as breast, bone and prostate, leading to relevant metastatic tumour formation.8,9 As early as 1988, the use of this kind of xenobiotic agent for improving the growth and feed conversion rates of food-producing animals has been banned in the European Union (EU) and no residues of these anabolic substances should be present in animal products.7 To enforce the prohibition on anabolic steroid abuse and prevent the illegal use of E2B, effective monitoring and detection methods have been developed.

For E2B determination, the most common techniques in use are high-performance liquid chromatography (HPLC), gas chromatography (GC), ¹⁰⁻¹² gas chromatography-mass spectrometry (GC-MS), ¹³ radioimmunoassay, ¹⁴ and liquid

chromatography-tandem mass spectrometry (LC-MS/MS).^{7,15} Although electrochemical methods have been demonstrated to exhibit many advantages over the traditional analytical methods, including rapid analysis, economical instrumentation, simple operation and high sensitivity and excellent selectivity, there are, to the best of our knowledge, no reports on the electrochemical determination of this specific endocrine disruptor.

In our previous research, hydroxyapatite (Ca₅(PO₄)₃OH, HAp) fabricated by template-assisted biomineralisation has been extensively applied in the field of chemically modified electrodes for biosensors, mainly due to its high biocompatibility and unique three-dimensional network structure.16-21 HAp materials with a hierarchical structure originating from a natural egg-shell have been successfully applied in the electrochemical sensing of bisphenol A (BPA) with a detection limit of 2.46 nM.18 On the other hand, it has been demonstrated that HAp materials manipulated by cation-exchange membraneassisted biomineralisation have significant electrochemical sensing capabilities attributed to their three-dimensional networks and beneficial electron transport process.16 In this work, we demonstrate proof-of-concept for the electroanalytical sensing of E2B using hydroxyapatite, which gives rise to highly sensitive analytical performances within aqueous solutions due to the material's inherent three-dimensional channel frameworks and useful electron transport capabilities.

Results and discussion

The fabrication of the hydroxyapatite (HAp) material/surface was performed as described in our previous report, ¹⁶ and for readers' convenience, these details are reported in the ESI.† In brief, the growth of HAp is performed in a self-designed crystallizer (Fig. S1†) at a pH of 11 in order to ensure the orientational growth of HAp crystals. After holding for 1 to 7 days, the crystals

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are scraped off the membrane and constructed into chemically modified carbon paste electrodes, denoted throughout as HAp-CPE. The crystalline phase and structure evolution during the 1 to 7 days' fabrication process (see the ESI†) are illustrated in Fig. 1. The corresponding PDF numbers are indicated along with the X-ray diffraction (XRD) patterns. Initially, the crystals are mainly composed of monetite (CaPO₃OH, PDF 09-0080) after 1 day. After extending the time of reaction from 3 to 7 days, the major constituents of the material are transformed to HAp with the variety of crystalline parameters such as the interplanar spacing (d) values of the strongest three diffraction peaks of the (211), (112) and (002) planes and lattice constants in each sample calculated by peak profile and cell refinement in order to reflect the transformation tendency, as shown in Table S1.† The values of d_{002} , d_{112} and c increase with the prolonged reaction time, while the others display a decrease. The variation of the unit cell for each crystalline structure with time is schematically shown in Fig. 2. The HAp structured family, which commonly display a P63/m hexagonal symmetry and have the ideal crystallochemical formula Ca₁₀(PO₄)₆(OH)₂, can be described as one-

dimensional ion conductors, where the [Ca₄^I][(PO₄)₆] framework

is constructed of face-sharing Ca^IO₆ metaprism columns linked

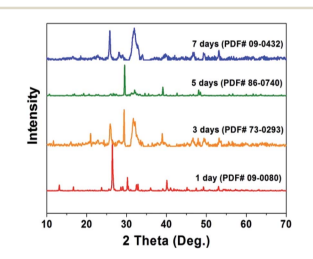


Fig. 1 X-ray diffraction patterns of the crystals after 1 to 7 days' fabrication time

through corner-connection to isolated PO4 tetrahedra. The resulting channels contain Ca^{II} and OH ions that are potentially ion conducting and exchangeable and the ion conduction is essentially electron transportation. It is likely that the variation of these lattice parameters will impact the sensing capability of the HAp material towards E2B; a suitable crystalline lattice will potentially favour electron transfer and provide a useful electrochemical surface.

The electrochemical behaviour of E2B at the HAp-CPE in different methanol: PBS ratios is illustrated in Fig. 3. It is readily evident that there is a clear electrochemical reduction peak facilitated by the HAp. Note that the response of the HAp-CPE in the presence and absence of E2B is presented in Fig. S2† indicating that the voltammetric signal is due to the electrochemical reduction of E2B and not any other electrochemical processes such as the supporting electrolyte degradation. The effect of changing the voltammetric scan rate is shown in Fig. 3 where the peak potential shifts with the scan rate and the shape of the voltammetric signatures indicates an electrochemically irreversible reaction. The underlying electrochemical process is likely the electrochemical reduction of the -C=O group to -C-OH via a one-electron and one-proton process. Analysis of the current as a function of the scan rate and square-root of scan and the peak potential (E_r) as a function of natural logarithm of scan rate $(\ln v)$ are presented in Fig. 3. In the case of using an electrolyte/solution of methanol: PBS ratio of 10:90, the peak current is found to be linearly dependent on the scan rate indicating that the electrochemical process is adsorptioncontrolled, while in the case of the methanol: PBS ratio of 90: 10, the peak current is linear in the case of the square-root of scan rate indicating a diffusion-controlled process; clearly, the solution/electrolyte has a key influence on the electrochemical process. Moreover, it was obtained that in both the mixtures, the natural logarithm of scan rate ($\ln \nu$) was linearly related to the peak potential (E_r) . Further analysis of the voltammetric data presented in Fig. 3 in the form of a log Ip vs. log scan rate produces a slope of 0.85 indicating a mixed adsorption and diffusional response in the case of the methanol: PBS ratio of 90:10. The electron transfer number (n) was deduced for the electrochemical reduction of E2B using the HAp-CPE and is presented in the ESI.†

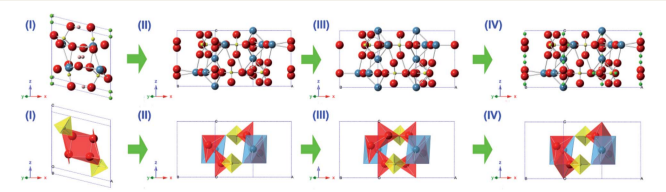


Fig. 2 Ball-stick models and polyhedral representations of crystalline structure evolution after (I) 1 day, (II) 3 days, (III) 5 days and (IV) 7 days' fabrications. All the angles of view are in the [010] plane (blue ball: Ca, red ball: O, yellow ball: P, green ball: H).

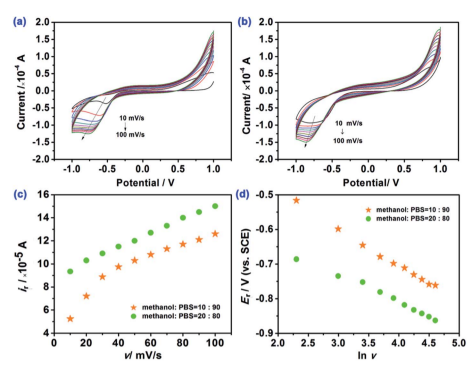


Fig. 3 Cyclic voltammetric responses of 1 μ g mL⁻¹ E2B using the HAp-CPE modified with 40% of 7 day HAp over the range of scan rates from 10 mV s⁻¹ to 100 mV s⁻¹ in methanol : PBS = (a) 10 : 90 (v/v) and (b) 90 : 10 (v/v). (c) Dependence of the reduction peak current (i_r) on the voltammetric scan rate (v). (d) The relationship between the reduction potential (E_r) and natural logarithm of voltammetric scan rate (v).

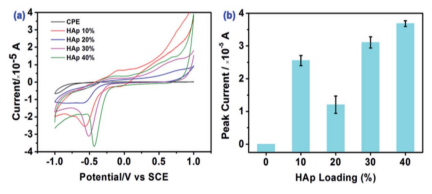


Fig. 4 Cyclic voltammetric curves of the 7 day HAp modified electrode in methanol : PBS = 10 : 90 (v/v) using different HAp loadings. E2B concentration: $1 \mu \text{g mL}^{-1}$.

The effect of reaction time (days) on HAp utilised within the HAp-CPE was also explored towards the magnitude of the electrochemical signal corresponding to the electrochemical reduction of E2B. It was found that the peak current corresponded to 0.6×10^{-5} A for 1 day fabrication time where the magnitude of the peak current was found to increase to 1.1×10^{-5} A for 7 days' fabrication time; consequently, the latter was chosen for further experiments. Next, loading (%) with the HAp-CPE was explored with the voltammetric signal monitored as a function of HAp loading from 10 to 40%. Fig. 4 shows the voltammetric signatures along with the effect on the peak current/signal. It is interesting to note that the voltammetric peak current shifts to less positive potentials as the % of HAp is increased along with a corresponding increase in the peak

height with the exception of the 20% HAp loading. The shape of the cyclic voltammogram also quantitatively changes from that of being largely diffusional to that of a thin-layer type response, which might be incorrectly assumed to be a transition to an electrocatalytic surface rather than true changes in the surface morphology and associated mass transport. Consequently, the largest voltammetric response/peak height is observed with a 40% HAp loading which is a combination of a useful electrochemical material being introduced into the electrode along with changes in the voltammetric response from having a contribution from a mixed diffusion and an adsorption electrochemical process (see earlier)/thin-layer voltammetry.

The analytical performance of the HAp-CPE towards the sensing of E2B was evaluated using cyclic voltammetry. Additions

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Fig. 5 Typical cyclic voltammetric responses and associated calibration plot obtained using the HAp-CPE towards the sensing of E2B. Solution composition; methanol : PBS = 10 : 90 (v/v). HAp: 7 day fabrication time; 40% loading of HAp in the HAp-CPE. Concentration of E2B: 1 ng mL $^{-1}$ ~400 ng mL $^{-1}$. Error bars are an average of three measurements and their associated standard deviation.

of E2B over the range of 1 ng mL⁻¹ to 400 ng mL⁻¹ were made into a methanol: PBS = 20:80 (v/v) solution as shown in Fig. 5. It is evident that at low E2B concentrations a single reduction wave is observed where at higher concentrations a pre-wave appears reflecting the adsorption of E2B on the HAp surface within the HAp-CPE. Analysis of the voltammetric response as a function of concentration, as shown in Fig. 5, shows a linear response at low E2B concentrations, which is observed to plateau at higher concentrations due to the saturation of E2B on the HAp surface. Analysis of the linear portion of the calibration plot reveals a linear response over from 1 to 200 ng mL⁻¹ ($i_p (\times 10^{-4} \text{ A}) =$ $0.00551 C_{E2B} (\text{ng mL}^{-1}) + 0.60681; R^2 = 0.98$). Based on this linear range, the detection limit (based on 3σ) was determined to correspond to 24.4 ng L⁻¹ (0.065 nM), which is much lower than the previous reports for the determination/sensing of E2B using GC $(0.15 \text{ mg L}^{-1})^{12}$ and is comparable with GC-MS/MS or LC-MS/ MS $(4.1 \mu g kg^{-1})$, indicating that this proposed method can potentially be used for the sensitive monitoring of low E2B concentrations. Note that this is the first reported example of the electrochemical sensing of E2B and hence no comparison to other electroanalytical methodologies can be made.

Conclusions

In summary, we have demonstrated a proof-of-concept for the electrochemical sensing method of the pollutant E2B within methanol/PBS mixtures. Sensitive analytical performances are observed using HAp as the basis of the electrochemical sensing platform due to its favourable electrochemical response and 3D framework with a possible detection limit of 24.4 ng $\rm L^{-1}$ (0.065 nM). Future work is recommended for exploring this material in real applications.

Conflicts of interest

There are no conflicts to declare.

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