Cerium oxide-chitosan based nanobiocomposite for food borne mycotoxin detection

Ajeet Kaushik,1,2 Pratima R. Solanki,1 M. K. Pandey,1 Sharif Ahmad,2 and Bansi D. Malhotra1,3,4

1Department of Science and Technology Centre on Biomolecular Electronics, National Physical Laboratory, New Delhi 110012, India
2Department of Chemistry, Materials Research Laboratory, Jamia Millia Islamia, New Delhi 110025, India
3Centre for Nano-Bioengineering and Spintronics, Chungnam National University, Daejeon 305-764, Republic of Korea

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Cerium oxide nanoparticles (NanoCeO2) and chitosan (CH) based nanobiocomposite film deposited onto indium-tin-oxide coated glass substrate has been used to coimmobilize rabbit immunoglobulin (r-IgGs) and bovine serum albumin (BSA) for food borne mycotoxin [ochratoxin-A (OTA)] detection. Electrochemical studies reveal that presence of NanoCeO2 increases effective electro-active surface area of CH-NanoCeO2/indium tin oxide (ITO) nanobiocomposite resulting in high loading of r-IgGs. BSA/r-IgGs/CH-NanoCeO2/ITO immunoelectrode exhibits improved linearity (0.25–6.0 ng/dl), detection limit (0.25 ng/dl), response time (25 s), sensitivity (18 μA/ng dl−1 cm−2), and regression coefficient (r2 ~ 0.997). © 2009 American Institute of Physics. [doi:10.1063/1.3249586]

Electrochemical immunosensors due to accuracy, sensitivity, selectivity and cost-effectiveness have been extensively used to detect proteins, biomarkers, biological toxins, and biological-warfare agents in critical situations, food, environment, pharmaceutical chemistry, and clinical diagnostics.1–3 The choices of nanomaterials as appropriate matrices due to unique optical, electrical, and molecular properties and in particular high reactivity and beneficial chemically tailored physicochemical properties have been utilized for antibodies immobilization with proper orientation that help to obtain sensitive, compact, and stable immunosensor.2,3

Among nanomaterials, metal oxides have been found to exhibit high surface-to-volume ratio, high surface reaction activity, high catalytic efficiency, and strong adsorption ability that make them potential candidate materials for the fabrication of biosensor.4–5 Among these, nanostructured CeO2 due to high mechanical strength, oxygen ion conductivity, biocompatibility, oxygen storage capacity, nontoxicity, high chemical stability and high electron transfer have aroused much interest for development of implantable biosensors. Moreover, high isoelectric point (IEP) of CeO2 (~9.2) can be helpful to immobilize desired biomolecules of low IEP via electrostatic interactions.2,5–13

Nanostructured CeO2 has been used for various biosensor applications such as immunosensor for mycotoxin,2 to immobilize cholesterol oxidase/glucose oxidase to detect cholesterol/glucose,5–7 as an insoluble oxidant to minimize interferences,8 for hydrogen peroxide detection via immobilizing horseradish peroxidase,9,10 and DNA hybridization detection.11 However, there is a considerable scope to improve the biosensing characteristics of NanoCeO2 by dispersing these in electroactive biopolymer e.g., chitosan (CH) to fabricate nanobiocomposite. CH due to its excellent film-forming ability, mechanical strength, biocompatibility, nontoxicity, susceptibility to chemical modification, cost-effectiveness, etc., has been explored for biosensor development.3 Moreover, amino groups of CH provide a hydrophilic environment compatible with the biomolecules.3,14

In this context, nanoporous CH-NanoCeO2 composite has been used to fabricate DNA and cholesterol biosensor.12,13

We report results of studies relating to coimmobilization of r-IgG and BSA onto CH-NanoCeO2 nanobiocomposite film onto indium-tin-oxide coated glass substrate for OTA detection. BSA/r-IgGs/CH-NanoCeO2/indium tin oxide (ITO) immunosensor shows improved sensing characteristics. OTA is produced by several species of Aspergillus and Penicillium and occurs in a variety of food commodities that causes nephrotoxin, hepatotoxin, teratogen, and carcinogen (group 2B, possibly by induction of oxidative DNA damage).2,3

All chemicals of analytical grade were purchased from Sigma Aldrich. 1 gm of cerium ammonium nitrate was dissolved in distilled 20 ml de-ionized water. NanoCeO2 (10 mg), synthesized using co-precipitation13 method is dispersed in 10 ml of CH solution [prepared by dissolving 50 mg CH in 10 ml of acetate buffer (0.05M, pH 4.2) solution] by constant stirring at room temperature (25 °C). The nanobiocomposite films are fabricated by uniformly spreading 10 μl solution of CH-NanoCeO2 nanobiocomposite onto an ITO (0.25 cm2) surface and dried at room temperature for about 12 h and then washed with deionized water to remove any unbound particles. OTA (Aspergillus ochraceus) solution is prepared in phosphate buffer (PB, 50 mM, pH 7.0) with 10% methanol. Solution of r-IgGs is prepared in PB. The immobilization of r-IgG has been carried out by spreading 10 μl solution onto the CH-NanoCeO2 nanobiocomposite electrode. BSA (98%) dissolved in PB is used as the blocking agent for nonspecific binding sites. Both the solutions containing 0.015M NaN3 as a preservative are aliquoated and stored at −20 °C.
X-ray diffraction studies (XRD) (Rigaku, miniflux 2) have been used to confirm formation of CeO2 nanoparticles. Scanning electron microscope (SEM) (LEO-440), Fourier transform infrared (FTIR) (Perkin Elmer) and Autolab (Potentiostat/Galvanostat) are used to characterize electrodes and immunoelectrodes. The electrochemical studies [differential pulse voltammetry (DPV) and cyclic voltammetry (CV)] are carried out in phosphate buffer saline (50 mM and 0.9% NaCl) containing 5 mM [Fe(CN)_6]^{3-}/4- at pH 7, at this pH biomolecule retains its natural structure and activity.

XRD pattern [Fig. 1(a)] of the CeO2, synthesized using coprecipitation method exhibits characteristic diffraction planes corresponding to CeO2 fluorite structure. However, the observed reflection planes are broad due to small particle size (~10 nm) estimated using Scherrer’s formula. The incorporation of NanoCeO2 in CH matrix has been confirmed using SEM wherein, regular planer morphology of CH and immunoelectrodes. The bands at 1643 and 1560 cm⁻¹ observed in FTIR spectra of CH-NanoCeO2 nanobiocomposite (circle, curve iii) correspond to amide II band of r-IgGs (β-sheet, main secondary structure element of IgGs), indicating immobilization of r-IgGs. However, presence of sharp band at 1647 cm⁻¹ in BSA/r-IgGs/CH-NanoCeO2/ITO immunoelectrode (circle, curve iv) corresponds to amide II band of BSA indicating immobilization of BSA. It is known that,
in the CH-NanoCeO₂ nanobiocomposite, CH with positively
NH₃⁺ group and surface charged NanoCeO₂ (IEP ~ 9.2) bind with carboxyl terminated group [fragment crystalizable (F⁺)]
of r-IgGs via electrostatic interactions and the free amino
terminated site [fragment antigen binding (Fab)] part is free
to bind with carboxylic group of OTA.

In the DPV studies [Fig. 2(a)], the magnitude of current
response for CH-NanoCeO₂/ITO nanobiocomposite electrode (curve ii) is higher than that of CH/ITO electrode
(curve i) revealing that presence of NanoCeO₂ results in increased electro-active surface area leading to increased adsorption of redox species [Fe (III)/(IV)]. The presence of NanoCeO₂ in nanobiocomposite results in improved electronic and ionic transport due to uniformly distributed NanoCeO₂ within CH matrix rather than clustered matrix (SEM analysis) resulting in three-dimensional electron conductive network extended throughout the ion-conductive matrix of CH. The surface concentration of redox species onto CH-NanoCeO₂/ITO electrode (2.2 × 10⁻⁶ mol cm⁻³) is higher than that of CH/ITO (1.4 × 10⁻⁶ mol cm⁻³) suggesting that incorporation of NanoCeO₂ increases electroactive surface area for diffusion of electrons resulting in high electrocatalytic properties of nanobiocomposite that may be responsible for improved sensing characteristics due to high loading of r-IgGs.

The magnitude of current for r-IgGs/CH-NanoCeO₂/ITO nanobiocomposite electrode (curve iii) further increases due to available free NH₃⁺ group on r-IgGs resulting in increased adsorption of redox species. The magnitude of current decreases for BSA/r-IgGs/CH-NanoCeO₂/ITO immunoelectrode (curve iv) revealing that blocking of non-binding sites of r-IgGs and pores of CH-NanoCeO₂/ITO electrode due to immobilization of BSA hinder electron transfer between the medium. The values of diffusion coefficient (D) for CH/ITO electrode, CH-NanoCeO₂/ITO electrode, r-IgGs/CH-NanoCeO₂/ITO immunoelectrode and BSA/r-IgGs/CH-NanoCeO₂/ITO immunoelectrode have been calculated as 2.2 × 10⁻⁷, 4.6 × 10⁻⁷, 7.8 × 10⁻⁷, and 6.0 × 10⁻⁷ cm² s⁻¹ respectively, using Randles–Sevcik equation.

The results of CV studies conducted on both electrodes and immunoelectrode indicate similar behavior [inset, Fig. 2(a)]. Moreover, CV studies of BSA/r-IgGs/CH-NanoCeO₂/ITO immunoelectrode [Fig. 2(b)] as a function of scan rate (10–100 mV/s) show that magnitude of current response exhibits linear relationship with square root of scan rate [inset, Fig. 2(b)], revealing a diffusion-controlled process. The observed linear dependence of potential difference as a function of scan rate reveals the facile electron transfer and follows Eqs. (1)–(3), respectively.

\[ I_{pa}(\text{BSA/r-IgGs/CH-NanoCeO}_2/\text{ITO})[A] = 3.7 \times 10^{-5} [A \text{ s/mV}] \text{ scan rate(mV/s)}^{1/2} \quad R^2 = 0.984, \]  
\[ I_{pa}(\text{BSA/r-IgGs/CH-NanoCeO}_2/\text{ITO})[A] = 1.2 \times 10^{-4} [A \text{ s/mV}] \text{ scan rate(mV/s)}^{1/2} \quad R^2 = 0.978, \] 
\[ \Delta E_{pa}(\text{BSA/r-IgGs/CH-NanoCeO}_2/\text{ITO})[V] = 0.164 [V] \quad \text{with} \quad R^2 = 0.99. \]  

It is observed that magnitude of the electrochemical response current of BSA/r-IgGs/CH-NanoCeO₂/ITO immuno electrode increases on addition of OTA (Fig. 3) due to formation of antigen-antibody complex between OTA and r-IgGs on nanobiocomposite surface that acts as electron transfer-accelerating layer. A linear relationship (inset, Fig. 3) between magnitude of current and OTA concentration has been observed and follows Eq. (4).

\[ I(A) = 2.21 \times 10^{-4} (A) + 17.5 \times 1.2 \text{ Ad L ng}^{-1} \times [\text{OTA concentration (ng dl}^{-1}] \quad R^2 = 0.997. \]  

The high value of the association constant (Kₐ, 4 × 10⁻¹² mol⁻¹) reveals high affinity of r-IgGs toward OTA due to favorable conformation of r-IgGs onto nanoporous CH-NanoCeO₂ nanobiocomposite. The CH-NanoCeO₂/ITO based immunosensor exhibits improved sensing characteristics such as low detection limit of 0.25 ng dl⁻¹, high sensitivity of 17.5 µA/ng dl⁻¹ cm⁻², reproducibility of 20 times and long term stability 80 days in comparison to BSA/r-IgGs/CH/ITO and BSA/r-IgGs/ NanoCeO₂/ITO based immunosensor, that are summarized in the supplementary table.¹⁷

In summary, NanoCeO₂ increases electroactive surface area of CH-NanoCeO₂ nanobiocomposite for the immobilization of r-IgGs resulting in enhanced electron transport. The BSA/r-IgGs/CH-NanoCeO₂/ITO immunosensor represents a new immunosensor platform for the detection of OTA and exhibits improved sensing characteristics than that of the sol-gel derived NanoCeO₂ and CH based immunoelectrode and may find potential application for detection of other clinically important antigens such as aflatoxin, citrinin, etc.

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¹⁷See EPAPS supplementary material at http://dx.doi.org/10.1063/1.3249586 for a supplementary table.