Structure—Activity Relationships of Hierarchical Three-Dimensional Electrodes with Photosystem II for Semiartificial Photosynthesis

Xin Fang,† Katarzyna P. Sokol,† Nina Heidary,† Tarek A. Kandiel,†,‡,† Jenny Z. Zhang,† and Erwin Reisner†,‡,†

†Department of Chemistry, University of Cambridge, Cambridge CB2 1EW, United Kingdom
‡Department of Chemistry, Faculty of Science, Sohag University, Sohag 82524, Egypt

Supporting Information

ABSTRACT: Semiartificial photosynthesis integrates photosynthetic enzymes with artificial electronics, which is an emerging approach to reroute the natural photoelectrogenetic pathways for sustainable fuel and chemical synthesis. However, the reduced catalytic activity of enzymes in bioelectrodes limits the overall performance and further applications in fuel production. Here, we show new insights into factors that affect the photoelectrogenesis in a model system consisting of photosystem II and three-dimensional indium tin oxide and graphene electrodes. Confocal fluorescence microscopy and in situ surface-sensitive infrared spectroscopy are employed to probe the enzyme distribution and penetration within electrode scaffolds of different structures, which is further correlated with protein film-photoelectrochemistry to establish relationships between the electrode architecture and enzyme activity. We find that the hierarchical structure of electrodes mainly influences the protein loading but not the enzyme activity. Photoactivity is more limited by light intensity and electronic communication at the biointerface. This study provides guidelines for maximizing the performance of semiartificial photosynthesis and also presents a set of methodologies to probe the photoactive biofilms in three-dimensional electrodes.

KEYWORDS: Photosystem II, inverse opal, indium tin oxide electrode, graphene electrode, semiartificial photosynthesis

Natural photosynthesis harvests sunlight to energize electrons and pump protons from water oxidation for carbon dioxide fixation, which is carried out by a series of enzymes that are orchestrated in thermodynamics and kinetics. Photosystem II (PSII) is the only enzyme known to photocatalyze the energy-demanding water oxidation reaction at a high turnover frequency (TOF) of \( \sim 100 \text{ s}^{-1} \), extracting electrons from water and initiating the vectorial electron flow to the terminal acceptor (Figure S1a). Photoelectrogenesis in PSII starts from the excitation of the reaction center chlorophylls (P680), followed by electron transfer from the excited P680* to the terminal acceptor plastoquinone B (Qb) via pheophytin and plastoquinone A (Figure S1b). In nature, the fully reduced Qb (QbH2) will dissociate from the reaction center complex and donate the electrons to photosystem I (PSI) via a cytochrome b6f complex. The electron transfer pathway from PSII to the cytochrome can be intercepted by exogenous acceptors such as molecular mediators and synthetic electrodes, which enables unique opportunities to redirect the photosynthetic electron flux for chemical synthesis and fuel production (Figure S1c).

Semiartificial photosynthesis combines strengths of both synthetic materials and biocatalysts to produce value-added chemicals with high selectivity and efficiency, which cannot be achieved by synthetic biology or artificial photosynthesis alone.9−11 Photosynthetic enzymes have been wired into many synthetic electrodes via self-assembled monolayers, metal oxide scaffolds, and redox polymer matrices, but their responsive photocurrent was limited by the low protein loading.12−18 To overcome this limitation, hierarchical three-dimensional (3D) electrodes with high surface area were produced to enhance protein loading and facilitate electron exchange at the protein-electrode interface (biointerface).19,20 For example, inverse opal−indium tin oxide (IO−ITO) electrodes allow proteins to penetrate through their interconnected macropores and have mesoporous skeletons with high surface area for protein binding and electronic communication (Figure 1a), thereby increasing the mediated photocurrent up to 1 mA cm\(^{-2}\). An attractive alternative to ITO are graphene materials as they can provide conductive and biocompatible surfaces for biocatalysts at low cost.25,28 In addition, the surface chemistry of graphene materials can be readily modified via physical/chemical approaches, which creates variable platforms to investigate
the interaction at biointerfaces. Planar graphene electrodes have been used to interface with isolated photoactive enzymes and intact photosynthetic apparatus, which usually produced submicroampere photocurrent that has no practical significance. A variety of 3D graphene structures have been applied for photovoltaics and electrocatalysis, but they have not yet been used to host photoactive enzymes for semiartificial photosynthesis. A major challenge facing semiartificial photosynthesis is that enzymes integrated in electrodes often exhibit reduced activity when stripped from their in vivo environment and that only a small fraction of enzymes at the biointerface are in an electroactive orientation (Figure S1d). Addressing these challenges requires an in-depth interrogation of bioelectrodes to pinpoint factors governing the photoelectrogenesis at the biointerface. Nevertheless, such investigations are lacking due to difficulties in characterizing the nanoscale proteins in complex 3D structures, which makes the protein-electrode interaction and its correlation with photoelectrochemistry largely unclear, and further challenges efforts to enhance the photocatalytic performance of bioelectrodes.

We prepared IO–ITO electrodes with different macro- and mesostructures using a co-assembly method with polystyrene (PS) beads and ITO nanoparticles, followed by annealing (Figure 1b). We employed 750 nm and 3 μm PS beads to create different macropores: larger PS beads will produce larger macropores and interconnecting channels after heat treatment for protein penetration but reduce the effective surface area for protein binding. To vary the mesostructure, we used polydispersed ITO nanoparticles (Sigma-Aldrich, SA) and synthesized monodispersed ITO nanoparticles with average sizes of approximately 10 nm (ITO-10), 20 nm (ITO-20) and 40 nm (ITO-40) (Figures S2 and S3 and Supporting Information). The IO–ITO electrode features interconnected macropores that enable protein penetration and integration. PSII can immobilize in the mesoporous scaffold that was composed of ITO nanoparticles. We employed confocal fluorescence microscopy and in situ infrared spectroscopy to visualize the protein distribution and monitor their penetration into the electrode scaffolds, respectively, and finally correlated with the enzyme activity that was quantified by protein-film photoelectrochemistry (PF-PEC). We find that the 3D hierarchical electrodes with smaller macropores and mesopores larger than the protein size enable higher loading capacity and better enzyme retention, whereas the individual enzyme activity is more sensitive to the light intensity and electronic communication at the biointerface, rather than the electrode morphology. Our results reveal the relationship between the electrode structure and enzyme activity and provide useful guidelines for the design and optimization of photoactive bioelectrodes.

Figure 1. Integration of PSII into IO–ITO electrodes for semiartificial photosynthesis. (a) Schematic representation of hierarchical structures of the IO–ITO electrodes. The PSII loaded IO–ITO electrode was used as the photoanode for PF-PEC. Platinum and Ag/AgCl were used as the counter and reference electrode, respectively. The IO–ITO electrode features interconnected macropores that enable protein penetration and integration. PSII can immobilize in the mesoporous scaffold that was composed of ITO nanoparticles. (b) SEM images of 750 nm IO–ITO electrodes and 3 μm IO–ITO electrodes made with different ITO nanoparticles. Scale bars: 500 nm (top); 2 μm (bottom).
The electrochemical properties of IO–ITO and IO–graphene electrodes were studied by cyclic voltammetry (CV). CV scans of both IO electrodes showed a typical electrical double layer capacitive behavior and electrochemical stability within the potential window of 0–0.5 V (vs SHE) at pH 6.5 (Figure S9a–c). As the capacitance of the electrical double layer capacitor scales proportionally with the electrochemically active surface area, we used the specific capacitance as a measure of the surface area accessible to electrolyte solution. The specific capacitance of IO–ITO electrodes decreased with increasing particle size as the smallest nanoparticles create a higher effective surface area (Figure 2a). The 750 nm IO–graphene electrodes displayed a higher capacitance than 3 μm electrodes. Increasing graphene will introduce more structural blockage for electrolyte penetration in 750 nm electrodes, thus reducing the capacitance, whereas it will provide more surface area instead of blocking the macropores in 3 μm electrodes (Figure S9d).

The light transmission of electrodes was measured by ultraviolet–visible (UV–vis) spectroscopy (Figure S10a–c). All the IO electrodes showed similar transmittance (<4%) regardless of materials, macro- and mesostructures (Figure S10d). Because of its wide bandgap (>4 eV), ITO has minimum light absorbance in the visible-light spectrum and therefore has been widely used for transparent electrodes. 

The IO–graphene electrodes were prepared by coassembling graphene oxide (GO) microsheets and PS beads (Figure S7a–c). PS beads were first mixed with a GO solution and dropcast on an ITO substrate. The GO–PS film was then annealed at 500 °C to remove the PS beads and reduce GO into graphene (reduced GO) (Figure S7d–g). The hydrophilicity of graphene can be further improved by ozone treatment (Figure S7g). The graphene electrode scaffold is composed of planar graphene sheets, which provides a large surface area but minor porosity and roughness compared to IO–ITO scaffolds (Figure S7h). We prepared IO–graphene electrodes with different morphologies by varying the size of PS beads (750 nm and 3 μm) and the concentration of GO solutions (Gr I, 4 mg mL⁻¹ and Gr II, 10 mg mL⁻¹) (Figure S8). In IO–graphene electrodes, the PS beads dictate the macroporosity, whereas the concentration of the GO solution mainly affects the surface area and pathways for enzyme penetration. More graphene sheets provide higher surface area for protein binding but may also block channels for protein integration and attenuate the light intensity. The major morphological difference with ITO counterparts is that macropores in IO–graphene electrodes were not interconnected. Proteins therefore can only penetrate into the electrodes via channels created by thermal decomposition of PS.

The 750 nm IO–ITO electrodes exhibited much lower PSII loading than IO–graphene electrodes. Despite comparable capacitance, IO–graphene electrodes exhibited much lower PSII loading than IO–ITO electrodes (Figure S12), which is likely due to weaker interaction and lack of effective pathways for protein penetration. At the center of electrode design for bioelectronics is creating large conductive surfaces to interface with redox-active proteins. However, this well-known principle should be complemented by considering the protein-electrode interaction to ensure the increased surface is accessible to the proteins.

Figure 2. PSII integration and distribution within IO–ITO electrodes. (a) Specific capacitance of IO–ITO electrodes obtained from CV scans (mean ± s.d., n = 3). (b) PSII loadings on IO–ITO electrodes obtained from UV–vis analysis (mean ± s.d., n = 3). (c,d) CLSM images of the PSII in ITO-SA 750 nm (c) and ITO-SA 3 μm (d) electrodes. (e,f) The 3D visualization of the PSII distribution within ITO-SA 750 nm (e) and ITO-SA 3 μm (f) electrodes. The 3D view was reconstructed from Z-stacking images that were acquired by scanning 20 μm downward from the electrode surface. Excitation: λex = 633 nm. Emission: λem = 650–750 nm. Scale bars in (c–f): 20 μm.
Despite numerous bioelectrodes developed for semiautomatic photosynthesis, PSII integration and distribution within electrodes has not been studied, which makes it difficult to evaluate the contribution of electrode structures. Here, we employed confocal laser scanning microscopy (CLSM) to visualize the spatial distribution of PSII within the 3D electrode scaffolds by its strongly fluorescent chlorophyll a (Chl a) molecules (see Figures S13–S15 for control experiments). The morphology of PSII films reflects the macroscopic structural features of electrodes such as macroporosity and cracks (Figure S16a). PSII uniformly distributed in 750 nm ITO–ITO electrodes whereas it formed aggregates in 3 μm ITO–ITO electrodes (Figure 2c,d). In both cases, PSII can penetrate throughout the entire ITO–ITO scaffold via channels interconnecting macropores (Figures 2e,f and S16b). The depth profile of fluorescence indicates that most PSII populated in the middle range of the electrode scaffold (Figure S17). Although the light intensity will decay along the electrode depth, the light scattering within the electrode will likely enable most PSII to access irradiation. In IO–graphene electrodes, CLSM images show that the enzymes could penetrate through the IO–graphene electrodes except Gr II-750 nm, where most PSII accumulated near to the surface due to excess graphene blocking pathways for protein penetration (Figure S18).

We further investigated the protein–electrode interaction by probing the dynamic process of protein adsorption and desorption within the 3D scaffold. This was possible by employing in situ attenuated total reflection-infrared (ATR-IR) spectroscopy to track the PSII penetration within IO–ITO scaffolds. ATR-IR spectroscopy features its surface-sensitivity to molecular vibration and can indicate protein unfolding or denaturation through changes in its spectral bands. The ATR-IR setup consisted of a 20 μm-thick IO–ITO scaffold deposited on a silicon prism (Figure S19a). As the evanescent IR wave will penetrate ~0.5 μm from the internal surface of the silicon prism, the monitored spectral features of protein amide bands can only stem from PSII at the bottom layer of the IO–ITO scaffold. Hence, the dynamic adsorption process of PSII can be monitored by its characteristic amide I and amide II bands centered at 1656 and 1546 cm⁻¹, respectively (Figure S19b).

When PSII was dropcast on the IO–ITO scaffold, the growing intensities of both amide I and amide II bands indicated an increased amount of PSII reaching the bottom of the IO–ITO structure, confirming the penetration of PSII through the entire 20 μm-thick film (Figure 3a,b). While positions of amide I and amide II bands remained unchanged during the PSII adsorption in both 750 nm and 3 μm ITO–ITO scaffolds (Figure 3c,d), their intensity followed a biexponential increase consisting of a fast and a slow kinetic process (Figure 3e,f). The first exponential growth was completed after 15 min for 750 nm and 1.8 min for 3 μm electrodes, which shows larger macropores favor fast protein penetration within the 3D scaffold and further suggests smaller macropores enable more mesoporous skeletons available for protein adsorption.

The nature of protein–electrode interaction was examined by a desorption experiment: the PSII-loaded IO–ITO scaffolds were washed with the PSII stock buffer solution (10% glycerol, 30 mM MgCl₂, 15 mM CaCl₂, 40 mM 2-(N-morpholino)-ethanesulfonic acid, pH 6.5) and buffer solutions with increasing KCl concentrations (0.5 M, 1.0 M, 3.0 M). In the 750 nm IO–ITO scaffold, the desorption process caused only minor changes in the amide bands intensities (Figure 3e), suggesting PSII was well-retained within the scaffold. In contrast, the amide bands increased steadily with KCl concentrations in the 3 μm IO–ITO scaffold (Figure 3f), indicating that an increased amount of PSII was desorbed from

Figure 3. ATR-IR spectroscopy of PSII-loaded IO–ITO scaffolds. (a,b) ATR-IR spectra of PSII adsorption at the bottom of ITO-SA 750 nm (a) and ITO-SA 3 μm (b) scaffolds. (c,d) The 2D visualization of the position and intensity of amide bands during the PSII adsorption in ITO-SA 750 nm (c) and ITO-SA 3 μm (d) scaffolds. The color bars indicate the IR absorbance (mOD). (e,f) Adsorption kinetics and desorption experiments of PSII loaded in ITO-SA 750 nm (e) and ITO-SA 3 μm (f) scaffolds. The biexponential fitting is shown as black lines in (e,f).
the scaffold by high ionic strength solutions and penetrated deeper into the IO scaffold thereafter. The difference in protein desorption can be attributed to the macroporosity, where desorbed proteins are more likely to be recaptured by the 750 nm IO–ITO scaffold but diffuse to the bottom in the 3 μm IO–ITO scaffold. The protein desorption induced by the high ionic strength solutions points to a dominating electrostatic interaction between PSII and the IO–ITO scaffold, which can be further verified by the desorption experiment on a planar ITO surface (Figure S20). The positive charge of ITO nanoparticles (ζ-potential: +32 mV at pH 6.5) enables PSII to interact with the IO–ITO scaffold via its negatively charged surfaces (Figure S21).59–61

The photoactivity of PSII-loaded electrodes was quantified by PF-PEC. Direct electron transfer (DET) photocurrent of IO–ITO electrodes correlated broadly with the PSII loading, except for ITO-40 750 nm that underperformed in DET compared with the ITO-SA 750 nm electrode (Figure 4a, S22 and Table S1). This may be due to thick PSII films that were inadequately wired by the conductive scaffold in ITO-40 750 nm and monodispersed ITO-40 nanoparticles that formed less contact sites with PSII compared with polydispersed ITO-SA nanoparticles. This problem was mitigated by introducing a diffusional redox mediator (2,5-dichloro-1,4-benzoquinone, DCBQ) that can relay electrons from proteins to the electrode, regardless of aggregates and orientation. Both ITO-40 and ITO-SA 750 nm electrodes showed a comparable mediated electron transfer (MET) photocurrent (Figure 4a). The MET photocurrent of 750 nm IO–ITO electrodes increased proportionally with the PSII loading, whereas that of 3 μm IO–ITO electrodes showed a major deviation from the PSII loadings, which may result from different mediator transport within protein films in different electrodes (Figure 4a). After PF-PEC measurements, less than 1% of PSII was released from the 750 nm IO–ITO electrode, whereas 22% were desorbed into the electrolyte solution from the 3 μm IO–ITO electrode (Figure S23), which is consistent with the results from ATR-IR spectroscopy that the 750 nm IO–ITO scaffold enabled better retention of PSII. Long-term chronoamperometry shows that the half-life of protein films inside the 750 nm IO–ITO electrode was ~4 min, significantly longer than that on a flat ITO electrode (<1 min), which manifests that the hierarchical structure is essential to secure the PSII within the electrode (Figure S24).

The TOF is calculated based on the photocurrent density and protein loading of the electrode (see Supporting Information for details), which represents the average photoactivity of PSII integrated in the electrodes (Figure 4b and Table S1). Despite large differences in photocurrent, 750 nm IO–ITO electrodes showed a similar TOFDET of 0.07 ± 0.02 s⁻¹. The highest TOFDET was achieved on the ITO-SA 750 nm electrode (0.10 ± 0.01 s⁻¹) (Figure 4b). As the MET current largely varied with the PSII loading for 750 nm IO-electrodes, TOFMET of 750 nm electrodes remained close to 2.8 ± 0.5 s⁻¹, which suggests the individual activity of PSII was less affected by the mesostructure of electrodes. Likewise, 3 μm IO–ITO electrodes showed a TOFDET of approximately 0.07 ± 0.01 s⁻¹ (Figure 4b). The comparable TOFDET for 750 nm and 3 μm IO–ITO electrodes despite their significant differences in photocurrent and PSII loadings suggests that the enzymatic activity is not dominated by the electrode morphology. When enzymes are integrated into electrodes, their TOFs are typically much lower than the average activity of purified PSII used in this study (TOF: ~20–30 s⁻¹) (Table S2), which indicates that most enzymes were not performing at their optimal rate. One limitation is the light intensity reaching PSII in electrode scaffolds. Both TOFDET and TOFMET increased with irradiation intensity up to 20 mW cm⁻² and decreased with electrode thickness (Figures 4c and S25). For example, despite having higher PSII loading, the 80 μm thick IO–ITO electrode had similar DET photocurrent with 40 μm thick electrode, indicating inefficient electronic communication within the thick electrode scaffold, which was improved with diffusional mediators (Figure S25). Further possible limitations may reside in the insufficient wiring of proteins by the electrode scaffold, due to protein aggregation and random
orientations. The adverse influence of protein orientation can be eliminated by modifying the surface chemistry of ITO to assist the electroactive orientation and by optimizing the surface mesoporosity of the electrode scaffold to multiply the contact sites to the enzyme. We also find that T0F_SMT would be affected by the mediator concentration, which may be due to the exogenous mediators quenching the excited antenna chlorophyll molecules (Figure S26).

Compared with their ITO counterparts, PSII in graphene electrodes exhibited less photoactivity (Figure S27 and Table S3), which is likely due to the strong absorption of graphene sheets attenuating the light intensity reaching PSII. Moreover, graphene electrodes had less protein binding capacity, which can be attributed to their macrostructures that do not have sufficient interconnected channels for protein penetration and their mesostructures that lack topographical roughness to physically bind protein films. In addition, hydrophilic interaction has been previously suggested between PSI and GO. We thereby speculate PSII might preferably interact with the thermally reduced graphene through a similar manner (Figure S28), which is weaker than the electrostatic interaction taking place in IO–ITO electrodes.

In this work, we conducted a systematic study on PSII-integrated 3D electrodes to reveal structure–activity relationships underlying photoelectrogenesis in photoactive bioelectrodes. (1) The protein integration in an electrode scaffold will be determined both by material and morphology. IO–ITO electrodes with small macropores and large mesoporosity tend to bind most proteins by physical/electrostatic interactions. (2) DET photocurrent arising from electronic communication between electroactive proteins and electrode surfaces correlates with the protein loading in the electrode. The photoactivity of PSII is not sensitive to the electrode morphology but will be governed by the light intensity and electronic communication at the biointerface. (3) MET photocurrent is convoluted by the interplay of protein loading and mediator diffusion within protein films and thus is dependent on the scaffold morphology. (4) IO–ITO electrodes are superior to IO–graphene as platforms for photoactive bioelectrodes, because their material and structures are more favorable for protein integration and light penetration (Figure 4d). Overall, our study cements the ITO-SA 750 nm electrode as the benchmark platform for PF–PEC due to the following advantages: (1) the hierarchical electrode architecture provides abundant conductive and hydrophilic mesoporous surfaces to secure proteins within the scaffold; (2) the polydispersed ITO nanoparticles create cavities with dimensions similar to PSII and enable numerous contact sites to interface with the protein; (3) ITO has low light absorbance, thereby allowing high light transmission and internal scattering in the 3D scaffold (Figure 4e).

Further studies are needed to elucidate the mass transport in protein films integrated in 3D electrode scaffolds. The PSII–3D electrode represents a model system in semiautomatic photosynthesis, and exemplifies the biotic–abiotic hybrid system that underlies a broad range of applications such as biosensing, biocatalysis, and biofuel production. This study advances our understanding of semiautomatic photosynthesis and presents a set of methodologies to probe protein–electrode interactions, which will underpin the ongoing efforts to reproduce the high bioactivity in artificial systems.

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