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Bias-Free In Situ H₂O₂ Generation in a Photovoltaic-Photoelectrochemical Tandem Cell for Biocatalytic Oxyfunctionalization

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Supporting Information

ABSTRACT: Peroxygenases catalyze selective oxyfunctionalization of hydrocarbons with high conversion efficiencies using H₂O₂ as a key cosubstrate. Here, we report an unbiased photoelectrochemical (PEC) tandem structure consisting of a FeOOH/BiVO₄ photoanode, a Cu(In,Ga)Se₂ solar absorber, and a graphitic carbon nitride/reduced graphene oxide hybrid cathode for light-driven peroxygenase catalysis. Powered by sufficient photovoltage generated by the solar absorber, the PEC platform generates H₂O₂ in situ through reductive activation of molecular oxygen using water as an electron donor in the absence of external bias. The peroxygenase from Agrocybe aegerita catalyzed the stereoselective hydroxylation of



ethylbenzene to (R)-1-phenylethanol with total turnover numbers over 43 300 and high enantioselectivity (ee > 99%) in the unbiased PEC tandem system.

KEYWORDS: oxyfunctionalization, photovoltaics, photoelectrochemical cells, biocatalysis, peroxygenases

elective oxyfunctionalization of C-H bonds remains a major challenge in organic chemistry because it requires activation of kinetically inert C-H bonds.¹ Heme-dependent oxygenases are powerful oxygenation tools due to their inherent feature of balancing reactivity with high selectivity.² In particular, so-called "unspecific" peroxygenases (UPOs, EC 1.11.21) have gained much interest recently because they do not rely on nicotinamide cofactors and complex electrontransfer chains to generate the catalytically active ferryl-oxo heme radical cation (compound I). They catalyze selective oxygen transfer from H_2O_2 to diverse organic substrates having nonactivated C-H bonds under mild conditions.³ Nonetheless, UPOs suffer from oxidative degradation of the heme moiety at excess concentrations of H2O2.4 To circumvent the inactivation, careful control of H2O2 supply is required. Stepwise addition of H₂O₂ leads to significant dilution of the reaction mixtures; thus, in situ generation of H_2O_2 through O_2 reduction is desirable to mitigate this issue.⁵ To date, the most common method for the generation of H₂O₂ uses glucose oxidase,^{6,7} which is not promising due to poor atom-economy (i.e., one equivalent of H_2O_2 from one equivalent of glucose) and practical issues such as high viscosity of glucose solutions. To resolve this problem, several different strategies have been proposed for in situ generation of H₂O₂ using more efficient electron donors (e.g., methanol, formate, electricity) through

electrochemical,^{8,9} photochemical,¹⁰⁻¹⁴ or enzymatic methods.15-17

Here, we report a bias-free, photoelectrode-photovoltaic (PV) dual absorber tandem photoelectrochemical (PEC) platform for solar-powered in situ H₂O₂ generation coupled with peroxygenase catalysis, as depicted in Figure 1. The PEC reduction of oxygen is an economically and ecologically promising approach to produce H₂O₂ using abundant solar energy and water, avoiding the accumulation of byproducts. Moreover, physical separation of anodic and cathodic reactions in a PEC system can prevent oxidative degradation of redox enzymes and H_2O_2 oxidation at the anode.^{18,19} The unassisted PEC system reduces O₂ and simultaneously generates H₂O₂ by using abundant solar energy and water in the absence of external bias. The Cu(In,Ga)Se₂ (CIGS) thin-film solar cell²⁰ in the PEC tandem device supply sufficient photovoltage, fulfilling the thermodynamic requirement for spontaneous PEC reaction while maximizing the fraction of solar energy collected. For solar water oxidation, we employed n-type bismuth vanadate $(BiVO_4)$ that exhibits a suitable band gap (ca. 2.4 eV) and superior photochemical stability in aqueous

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Figure 1. External bias-free, photoelectrochemical generation of H_2O_2 using water and solar energy for peroxygenase catalysis. *AaeUPO* catalyzes chemo- and stereospecific conversion of ethylbenzene to (*R*)-1-phenylethanol. In situ provision of H_2O_2 is achieved through the reduction of oxygen using FeOOH/BiVO₄/CIGS solar cell in tandem and CN/rGO film electrode under illumination.

environments.²¹ We have paired BiVO₄ with CIGS solar cells to construct a PEC-PV tandem device. As a counter electrode in the PEC tandem cell, we explored the feasibility of graphitic carbon nitride (CN) and reduced graphene oxide (rGO) hybrid film²² as an effective cathode material for O_2 reduction using electrons extracted from water oxidation at the BiVO₄ photoanode. CN is an attractive electrocatalyst for O2 reduction because of its high nitrogen content, easily tunable structure, and cost-efficient production.²³ To facilitate charge transfer and increase the electrochemical active surface area, rGO is used as a supporting layer for CN. As depicted in Figure 1, the proposed tandem system generates H_2O_2 through O₂ reduction under zero applied bias, which then takes part in selective hydroxylation of ethylbenzene to (R)-1-phenylethanol by the recombinant peroxygenase from Agrocybe aegerita (AaeUPO).

To synthesize the CN/rGO hybrid, we casted cyanuric acidmelamine supramolecular assembly and GO aqueous dispersion blended in ethylene glycol onto a fluorine-doped SnO₂ (FTO) glass using a doctor-blade technique, followed by heating at 550 °C under a nitrogen atmosphere for 4 h according to the literature.²² The resulting hybrids are denoted as CN/rGO-n, where *n* represents the weight % of GO in the precursor paste. During the calcination process, GO was thermally reduced, and a dark gray colored electrode was formed (Figure S1a). Scanning electron microscopy (SEM) (Figures S1b,c) revealed that the CN/rGO-0.2 hybrid film exhibited a porous morphology consisting of well-interconnected sheets with a thickness of approximately 40 μ m. In addition, transmission electron microscopy (TEM) analysis revealed a layered nanostructure consisted of rGO and CN sheets (Figure S2). We investigated the composition and chemical structure of CN/rGO using X-ray photoelectron spectroscopy (XPS). The high-resolution N 1s spectrum of CN/rGO-0.2 (Figure S3a) was deconvoluted into three typical peaks for pyridinic N (C–N = C) at 398.7 eV, tertiary N (N– $(C)_3$) at 400.1 eV, and amino groups (C-N-H) at 401.4 eV, confirming the formation of CNs.²⁴ The minor peak centered at 404.2 eV corresponded to the charging effect (or π excitation) in the heterocycles.²⁵ With regard to the C 1s spectrum of CN/rGO-0.2, there were mainly two carbon states including C-C bonds at 284.6 eV and sp²-bonded carbon coordination (N-C=N) at 288.5 eV (Figure S3b). Besides, two minor peaks at 285.4 and 286.1 eV corresponded to C-OH (from rGO) and C-O-C/C-O-N, respectively. C-O-

N bonds play a role in the covalent bonding between the amine groups from CN and the hydroxyl (or carboxyl) groups from GO.²² The coupling between CN and rGO was further confirmed by Fourier transform infrared spectroscopy (FTIR) (Figure S4). The breathing mode of the triazine unit shifted from 810 to 812 cm⁻¹ because of the strong interaction between CN and rGO.

We evaluated CN/rGO-*n* hybrid's electrocatalytic activity for O₂ reduction by cyclic voltammetry (CV) in N₂ or O₂saturated 0.1 M KPi buffers with a scan rate of 50 mV s⁻¹ (Figure 2a, Figure S5). All CN/rGO-*n* films exhibited



Figure 2. (a) CV curves of the CN/rGO-0.2 electrode measured in N₂- or O₂-saturated 0.1 M KPi buffer (pH 7.0) at a scan rate of 50 mV s⁻¹. (b) Nyquist curves of CN/rGO-*n* electrodes measured in a 0.1 M KPi buffer (pH 7.0) at an applied bias of -0.3 V (vs Ag/AgCl). All electrodes had a geometrical surface area of 2 cm².

significantly increased cathodic currents in O2-saturated solutions compared to the currents generated in a N2-saturated solution suggesting cathodic O₂ reduction. Increasing the amount of GO increased the peak current density of CN/rGO*n* from -0.2 mA cm^{-2} (for CN/rGO-0.2) to -0.53 mA cm^{-2} (for CN/rGO-0.5) (Figure S5). However, the increase in the reduction current from CN/rGO-0.4 to CN/rGO-0.5 was not clearly observed. This tendency is attributed to the presence of excess rGO that could block the sites available for O_2 adsorption. Similarly, double-layer capacitance increase was observed with the increasing GO content. As shown in Figure S6, the estimated double-layer capacitance value of CN/rGO-0.5 was 29 times higher than that of CN/rGO-0.1. The sheer increase in double-layer capacitance indicates that the incorporation of rGO enhanced the electrical conductivity and the specific surface area of CN/rGO hybrid.²⁶ We investigated the effect of rGO on charge separation efficiency by performing electrochemical impedance spectroscopy (EIS) measurements for CN/rGO-n in 0.1 M KPi buffer (Figure 2b, Figure S7). The charge transfer resistance of CN/rGO-n film was evaluated by the semicircle diameter in the high frequency region of the Nyquist plots. The arc radius of the semicircles decreased drastically with higher loading of GO from 0.1 to 0.5 wt %, indicating the decrease in charge transfer resistance. The result shows that rGO acts as an electron conductive channel to facilitate efficient electron transfer to the solution.

We assembled a FeOOH/BiVO₄ photoanode and a CIGS solar cell in a tandem manner to drive unassisted H_2O_2 generation under illumination. We prepared a nanoporous BiVO₄ film (active area: 4 cm²) on a FTO substrate according to the literature,²⁷ then photoelectrochemically deposited FeOOH as a cocatalyst on the BiVO₄ photoelectrode to promote the PEC activities of BiVO₄ toward water oxidation. BiVO₄ formed a nanostructure where particles of approximately 200–300 nm diameter were interconnected each other

(Figure S8a). The band gap energy for the nanostructured FeOOH/BiVO₄ was estimated to be approximately 2.5 eV (Tauc plot, Figure S8b). In contrast, the CIGS solar cell possessed a band gap energy of around 1.1 eV;²⁸ thus, it can utilize the longer-wavelength (<1.13 mm) photons that are transmitted through the front FeOOH/BiVO₄ photoanode. To confirm the existence of FeOOH layer, we carried out XPS analysis on BiVO₄ and FeOOH/BiVO₄ (Figure S9). After the photoelectrodeposition of FeOOH, Fe 2p peaks and a strong peak for hydroxyl group at 530 eV in the O 1s spectrum appeared, consistent with the literature.²⁹

The J-V performances of the FeOOH/BiVO₄ photoanode and the CIGS solar cell under filtered illumination via the FeOOH/BiVO₄ are shown in Figure 3a. The theoretical



Figure 3. (a) J-V curves for CIGS PV cell with FeOOH/BiVO₄ placed in front (black) and FeOOH/BiVO₄ photoanode (red) under illumination (AM 1.5G, 100 mW cm⁻²), where the currents are based on actual areas of each components without normalization. The intersection point of two curves indicates a theoretical operating current of the tandem device. (b) Linear sweep voltammetry curves of the FeOOH/BiVO₄ photoanode (black) and the FeOOH/BiVO₄/CIGS tandem device (red) measured in 0.1 M KPi buffer (pH 7.0) at a scan rate of 50 mV s⁻¹ under illumination.

operating current of the tandem system was estimated to be 1.35 mA from the photocurrent at the intersection of individual J-V curves of the photoanode and the solar cell. This value is close to the actual photocurrent (1.34 mA) measured from the linear sweep voltammetric analysis of the assembled tandem device in a three-electrode configuration (Figure 3b).

We applied the CN/rGO cathode to the tandem system consisting of the FeOOH/BiVO₄ photoanode and the CIGS solar cell for in situ H₂O₂ generation driven by solar energy using water as an electron donor. To prevent possible oxidation of H2O2 at the anode, we utilized a two-chamber setup in which FeOOH/BiVO₄/CIGS in tandem and CN/ rGO cathode were placed in different reaction vessels connected by a salt bridge. Under this configuration, the photocurrent was reduced to 40% of the value obtained from the one-chamber setup due to salt bridge's ionic resistance (Figure S10).³⁰ The photocurrent generated by the tandem system in a two-compartment setup was stabilized at 0.46 mA for ~14 h under illumination. To facilitate mass transfer of dissolved O2, we gently bubbled O2 gas into the electrolyte solution of the cathodic compartment during the experiments.³¹ As shown in Figure 4a, we observed H₂O₂ generation by the PEC tandem platform, which reached a plateau of 6.5 mM after 2 h, which is 8.9-fold higher than a control experiment without CIGS solar absorber. The result indicates that an efficient production of H₂O₂ at CN/rGO electrode can be achieved in the absence of external bias by employing CIGS



Figure 4. (a) Bias-free photoelectrochemical in situ H_2O_2 formation over CN/rGO-0.3 cathode with FeOOH/BiVO₄/CIGS tandem device or FeOOH/BiVO₄ photoanode under O₂- or N₂-purging. (b) H_2O_2 formation rates and current efficiencies by the tandem system under O₂-purging as a function of GO loading amount (wt %). H_2O_2 production rates are divided by the geometrical surface area of the electrodes.

PV cell that provides sufficient photovoltage to reduce O₂. In the absence of oxygen, no H₂O₂ generation was observed consistent with the proposed reduction of O_2 . To evaluate the effect of the rGO content in the cathode on H₂O₂ generation, we measured initial production rate and current efficiency by varying the loaded GO amount from 0.1 to 0.5 wt % (Figure 4b). Consistent with the EIS and CV analysis, an increase in H_2O_2 production rate was observed with the increasing amount of GO from 0.1 to 0.4 wt %. The CN/rGO-0.4 electrode exhibited the highest H₂O₂ production rate of 2.8 $mM h^{-1} cm^{-2}$ with a current efficiency of 92%. Higher loadings of GO (i.e., CN/rGO-0.5) did not improve the O₂-to-H₂O₂ reduction rate (1.9 mM h⁻¹ cm⁻²) showing a slight decrease of current efficiency (73%), which is attributed to the kinetic limitation by oxygen mass transfer to the cathode surface.³² In addition, the surface nitrogen heteroatom of CN has been considered to be active sites for O2 reduction,³³ which could be blocked by the formation of rGO at high content of GO,³⁴ decreasing overall H₂O₂ production rate.

We conducted *Aae*UPO-catalyzed stereospecific hydroxylation of ethylbenzene to (*R*)-1-phenylethanol as a model reaction using the PEC tandem platform (Figure 5a). With the unassisted H_2O_2 generation in the PEC tandem device, we observed a (*R*)-1-phenylethanol formation at a rate of 0.89 mM h⁻¹, which was 4.2 times faster than in the absence of CIGS solar cell. The optical purity of the product exceeded 99% enantiomeric excess (ee) in all cases, indicating the conversion was catalyzed by the enzyme. Note that (*R*)-1-



Figure 5. (a) Time profile of ethylbenzene conversion to (*R*)-1phenylethanol by *Aae*UPO in the tandem configuration and in the absence of CIGS solar cell. (b) Influence of GO concentration on initial TOF and TTN of *Aae*UPO-catalyzed ethylbenzene hydroxylation driven by the proposed PEC tandem system. Reaction conditions: 0.1 M KPi buffer (pH 7.0, 1 mL) containing 200 nM *Aae*UPO and 100 mM ethylbenzene; 100 mW cm⁻²; 25 °C.

phenylethanol was not produced at all in the absence of any key component (e.g., light, oxygen, or AaeUPO). We further examined the effect of rGO content in the CN/rGO cathode on the performance of AaeUPO-catalyzed oxyfunctionalization in terms of turnover frequency (TOF) and total turnover number (TTN). In agreement with the H₂O₂ production rate, the initial TOF_{AaeUPO} and TTN_{AaeUPO} also increased with the incorporation of rGO. As shown in Figure 5b, CN/rGO-0.4 exhibited the best performance in terms of reaction rate with a $\text{TOF}_{Aae\text{UPO}}$ of 12.9 s⁻¹ and had the highest $\text{TTN}_{Aae\text{UPO}}$ of 43 300. The TOF value is much higher compared with other PEC platforms for in situ H2O2 generation such as flavinhybridized carbon nanotube photocathode (TOF: 1 s⁻¹).³⁵ Additionally, a further increase of the GO loading lowered the biocatalytic conversion; initial TOF_{AaeUPO} of 12.2 s⁻¹ and the TTN_{AaeUPO} of 29 700 were observed for CN/rGO-0.5. The decrease in the TTN with CN/rGO-0.5 may be caused by excess electrons provided by the highly conductive CN/rGO film, and possibly, hydroxyl radicals originating from H₂O₂ reduction $[E (H_2O_2/OH) = 0.38 \text{ V vs NHE, pH 7}]$ cause oxidative inactivation of AaeUPO (Figure S11). According to the literature,³⁶ hydroxyl radicals among reactive oxygen species can directly attack the active site of AaeUPO. Using the CN/rGO-0.5 cathode, product formation ceased within 4 h (Figure S12), which we ascribed to biocatalyst deactivation. The results suggest that careful control of rGO amount is essential to sustain activity of AaeUPO, while suppressing undesired side reactions. We further explored the substrate scope of photoelectroenzymatic hydroxylation in the tandem PEC system. As listed in Table S1, a broad range of aromatic compounds were converted into their corresponding alcohols. Taken together, the deliberate integration of the PEC-PV tandem device with AaeUPO can produce fine and specialty chemicals without any external energy supply in pursuit of a higher efficiency, while the previously reported PEC method³⁵ requires an additional bias of -0.4 V (vs Ag/AgCl) in addition to the light energy to promote optimized UPO reactions. We envision that a further increase of turnover numbers can be achieved by enhancing enzyme stability through the use of hydroxyl radical scavengers and physical separation of AaeUPO from the electrode.

In summary, we successfully demonstrated peroxygenasecatalyzed oxyfunctionalization of C-H bonds through unbiased photoelectrochemical H₂O₂ generation using a PEC tandem cell structure consisted of a FeOOH/BiVO₄ photoanode, a CIGS solar cell, and a CN/rGO cathode. The singlepass tandem configuration allows for the well-managed light harvesting and provides sufficient energy to drive water oxidation at the FeOOH/BiVO4 and simultaneous O2 reduction at the cathode under no external bias. The CN/ rGO cathode having significantly enhanced electrical conductivity and effective surface area owing to rGO allowed for high oxygen reduction activity. By integrating the CN/rGO-0.4 cathode to the PEC tandem cell, H₂O₂ was generated at a rate of 2.8 mM h⁻¹ cm⁻² under visible light in the absence of external bias. Finally, we successfully achieved photoelectroenzymatic hydroxylation of ethylbenzene to enantiopure (R)-1-phenylethanol with a TTN of 43 300 and a TOF of 12.9 s^{-1} in the tandem platform. Our approach suggests that careful integration of unbiased PEC system with peroxygenases can promote C-H oxyfunctionalization using solar-powered electrons extracted from water.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscatal.9b04454.

Experimental procedures, SEM, TEM, XPS, FTIR, and electrochemical characterization (PDF)

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Notes

The authors declare no competing financial interest.

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