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Sacrifice and valorization of biomass to realize energy exploitation and transformation in a photoelectrochemical way

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Photoelectrochemistry (PEC) transfers photoexcited electrons from a photocatalyst to a cathode through a wire, to avoid, to a considerable extent, the recombination of charge carriers in solitary photocatalysis. The moderate bias also consumes less energy than electrocatalysis, rendering PEC a green and promising transformation approach. PEC biomass conversion can be performed at a lower bias than PEC water splitting, and biomass holds great potential for valorization as the world's largest renewable resource. In recent years, there has been escalating interest in the study of PEC valorization from the sacrifice of biomass, but only a few reviews have systematically and specifically addressed it. This review discusses the conversion mechanism of organics, the catalyst and system configuration, and the effect of other variables in detail based on different structures and types of biomass substrates. A brief outlook on the development of the field is outlined in the end as well, and PEC is expected to be an alternative low-carbon, energy-saving, multi-objective route to biomass valorization.

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1. Introduction

Except for a few resource-based countries, there is a burgeoning concern about the reserves of non-renewable fossil resources around the world. The waste and harmful gases generated by the consumption of fossil resources threaten the global ecological environment significantly. The search for alternative green raw materials and more environmentally sound and efficient technological conditions is currently a global trend. In sustainable development, solar energy is regarded as a key technology due to its abundance and worldwide availability for collection and conversion.^{1,2} Biomass, derived from plant photosynthesis, is an energy carrier of solar energy and is considered one of the most promising alternatives to combat the energy crisis and environmental problems of fossil fuels.³ Carbohydrates and/or aromatic ring-rich fractions in biomass can be converted into biofuels or chemicals by fermentation.⁴

Biomass is mainly composed of cellulose, hemicellulose, and lignin, and the relative content of each component varies with the type of biomass. Taking lignocellulose, about 100 billion tons of production per annum globally, as an example,

the content of cellulose and hemicellulose is about 40–50 wt% and 20–30 wt%, respectively, and this is mainly a C5–C6 carbohydrate polymer. In contrast, the content of lignin is slightly lower, about 15–25 wt%.⁵ The structures also differed to some extent between the different components (Fig. 1). Cellulose is a homopolymer composed of β -D-glucopyranose units linked by a β -glycosidic bond that can be deconstructed into glucose monomers.⁶ Hemicelluloses are amorphous branched polymers with a low polymerization degree (~200). Hemicellulose is more readily hydrolyzed than cellulose, which allows it to be removed under more benign reaction conditions. Hemicellulose contains mainly pentoses, hexoses, and glyoxylates. Pentoses (xylose and arabinose) and hexoses (glucose, galactose, mannose, rhamnose, and fucose) are sources of sugars that can be valorized to biofuels and renewable chemicals.⁶ Lignin, on the other hand, contains the world's largest renewable aromatic monomer and is an integral part of the cell wall. However, the utilization of lignin is not satisfactory. The pulp and paper industry produces about 5 million metric tons of lignin each year.⁷ However, 95% of lignin remains abandoned or is burned during the biorefining process due to its complex and irregular structure.^{7,8} Lignin has an amorphous three-dimensional network structure and the different aromatic monomers are interconnected by aliphatic C–C and C–O bonds. The polymers contain many hydroxyl and methoxy groups and the three major monomeric units of lignin can be distinguished according to the number of methoxy groups on

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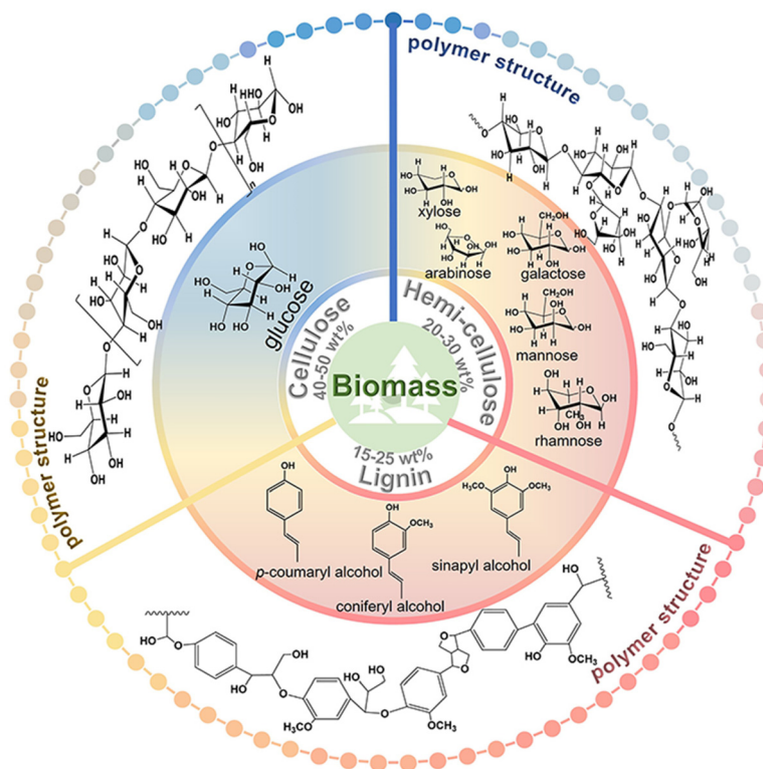


Fig. 1 The polymer structures and depolymerized monomers/monomeric units of the three major components of biomass.

the benzene ring: *p*-coumaryl alcohol, coniferyl alcohol, and sinapyl alcohol.⁹ The abundant aromatic content and the variety of oxygen-containing functional groups enable lignin to have gigantic potential for upgrading.

Biomass feedstock is commonly treated as a “fuel” source and converted into a fuel form through pyrolysis, saccharification, and fermentation, such as H₂, alcohols, and polysacchar-

ides.¹⁰ These transforming processes burden the overall process and lead to unwanted energy consumption, so it would be better to utilize (hemi-)cellulose or lignin directly as a “fuel”.¹¹ This purpose can be achieved through photocatalytic, electrocatalytic, and photoelectrochemical (PEC) systems under mild conditions (Fig. 2). For simple photocatalysis, illumination of light with a specific wavelength range or

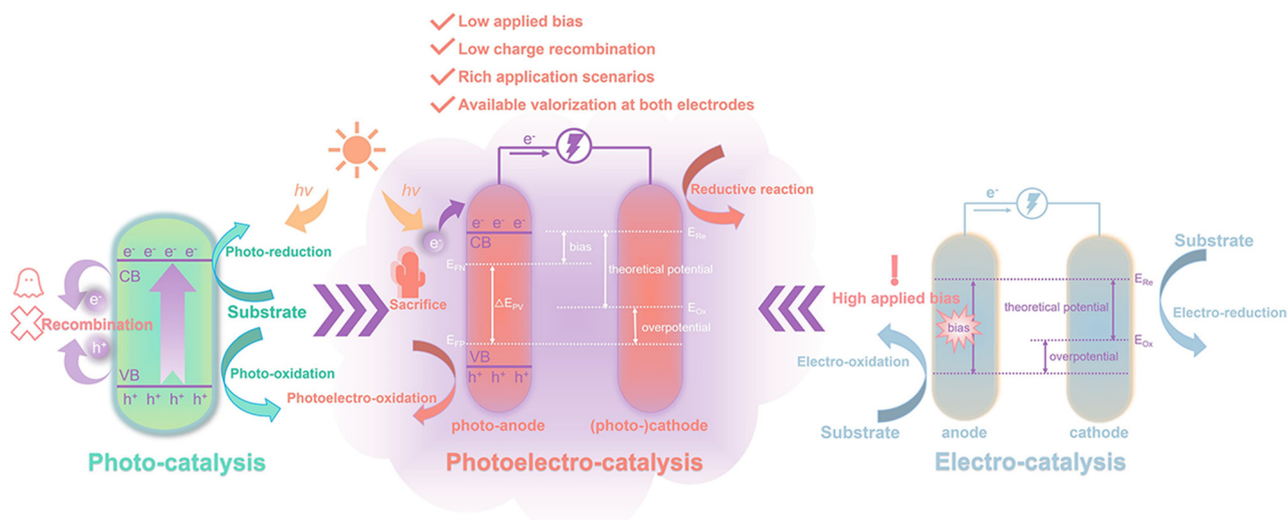


Fig. 2 Comparison of photo-, electro-, and photoelectro-catalysis. E_{FP} and E_{FN} are the quasi Fermi levels of holes and electrons, respectively; E_{Ox} and E_{Re} are the oxidative and reductive potentials of the substrate, respectively.

visible light (collectively referred to as incident light) onto a semiconductor catalyst excites electrons (e^-) in the valence band (VB) to the conduction band (CB), while the same number of holes (h^+) is generated in the VB. h^+ is strongly oxidizing and can carry out the oxidation of the substrate at the VB. Correspondingly, reduction reactions can be performed at the CB. Shang *et al.*¹² reviewed in detail the study of bifunctional photocatalytic reaction systems in which photogenerated electrons and holes can be exploited simultaneously, with the organics acting as sacrificial reagents to eventually produce high value-added products. However, the notable drawbacks of photocatalysis are low reaction rates and low quantum efficiency due to the recombination of photogenerated charges.¹³ At the same time, the insolubility of biomass hinders the supply of “fuel” in the liquid phase. The effective photodegradation of solid substrates is also challenging since a large contact area between the substrate and the photocatalyst is essential in photocatalysis.

Electrocatalysis is able to efficiently convert electrical energy into chemical energy and the electrical energy can be driven by sustainable energy sources (*e.g.* solar photovoltaic, wind, and hydrogen).^{14,15} In an electrocatalytic system, all electrons are induced by an external bias, which determines the current density of the circuit. The actual applied voltage is often higher than the theoretical energy requirement to compensate for overpotential or other energy losses.¹⁶ In contrast, semiconductor-based PEC systems that absorb and convert solar energy are capable of alleviating the dependence on pure electricity.

2. Working configurations of PEC systems

Photoelectrochemical cells use solar and electrical energy to promote target chemical reactions. In 1972, Honda *et al.*¹⁷ achieved PEC water splitting by coupling a crystalline n -TiO₂ photoanode and a platinum cathode under ultraviolet (UV) radiation. Since then, tremendous applied advancements in the field of PEC using a range of photoanodes have been emerging.¹⁸ Numerous studies have spotlighted the application of water splitting, which is a thermodynamically uphill reaction.^{19,20} During water splitting, the hydrogen evolution reaction (HER) and oxygen evolution reaction (OER) occur simultaneously at the cathode and anode, respectively. The reaction kinetics of the OER is slower than that of the HER because the OER is a four-electron transfer process accompanied by the formation of O–O bonds, whereas the HER is only a two-electron transfer process. In other words, the OER is the rate-limiting step for water cleavage.^{21–23} The oxidation of biomass and its derivatives is a promising alternative to the OER with a low oxidative potential.^{24–26} The substrate being oxidized, which can be photoconverted into high-value organic products, is usually waste or pollutants left over from industrial processes and production.²⁷ The whole process is simple and efficient; hence PEC decomposition of biomass-

derived waste/pollutants is also recognized as one of the most promising solutions.^{20,28–32}

2.1. Reaction mechanism of PEC

As shown in Fig. 2, photogenerated electrons on the VB are excited to the CB under illumination, and the same number of photogenerated holes is generated in the VB. Due to the presence of an external electric field, the highly oxidizing holes are transferred to the semiconductor surface to oxidize the substrate, and the photogenerated electrons are transferred along the external circuit to the cathode for the reduction reaction. The photogenerated carriers transported in the circuit create the photocurrent. The photovoltage generated at the photoanode can be determined by the potential difference (ΔE_{PV}) between the quasi Fermi energy levels of electrons and holes. For a particular photoelectrochemical reaction and semiconductor, the photocurrent density of the PEC system normally depends on the incident light intensity and the applied external bias. In terms of the operation mechanism, the photogenerated charge resulting from light illumination onto the catalyst is the essential reason for the catalytic activity of the catalytic system. Therefore, the incident light intensity plays a prominent role and the maximum current density of the system depends on it. If the incident light intensity is constant, the current density will increase with a higher bias.¹⁶

During the PEC oxidation of biomass derivatives, the active species driving the substrate in the reaction show more variations compared to water splitting. In addition to the direct contact of photogenerated holes with organic substrates, the hydroxyl radicals ($\cdot OH$) generated by the splitting of water and the superoxide radicals ($\cdot O_2^-$) generated by the reduction of soluble oxygen exhibit equally strong oxidative properties, although the oxidation selectivity of $\cdot OH$ is relatively low.⁵ Whether a single reactive species or multiple reactive species play a decisive role in PEC activity needs to be analyzed on the basis of the specific catalytic system. Therefore, the investigation of the specific active species in biomass PEC systems seems to be valuable and indispensable.

2.2. Comparison of PEC, photocatalysis, and electrocatalysis

It is hardly possible to make a photocatalyst undergo both oxidation and reduction half-reactions under the premise of reduced recombination of photogenerated electrons and holes, but the PEC system can be employed for separate studies of cathodes and anodes and allows better testing and rational development of catalysts for oxidation reactions.³³ For both electrocatalysis and PEC, electrocatalysis can produce a higher current density than PEC at a high bias. In the case of water splitting, light illumination of several sun powers is required for PEC if a current density of 100 mA cm⁻² is to be produced.³⁴ Oxidation reactions of many organics are thermodynamically and kinetically easier to be initiated than those of water, so PEC systems are capable of achieving higher current densities at a lower bias than electrocatalysis.³⁵ A common characteristic of photocatalysis and electrocatalysis is the generation of electrons and the following participation and pro-

motion of subsequent chemical reactions. In addition to photons and electrons induced by photocatalytic and electrocatalytic processes, there are a number of generated radical intermediates that are other common active components, such as carbon-centered radicals, $\cdot\text{OH}$, $\cdot\text{O}_2^-$, and chemically generated reactive oxygen species. These components are also applicable in the PEC process, and for more details on the radical induction mechanism, please refer to the review by Wu *et al.*⁵

As the applications of PEC systems continue to expand, more and more studies are shifting from water splitting to the exploration of the valorization of biomass and its derivatives as sacrificial agents. The oxidation of biomass at the anode builds the bridge of energy transfer for the whole system as a process of electron extraction. The biomass itself is either consumed at the anode or escalated simultaneously with the cathode to valorized products, which is of great research interest. However, a review of the literature that systematically summarizes and explores this research area is currently unavailable. This work will survey the PEC high-value applications of (hemi-)cellulose, lignin, and their derivatives in the last five years. Meanwhile, future research perspectives are proposed based on the discussion on targets of valorization, designs of systems, catalysts, and other impact factors.

3. Valorization application in PEC systems through biomass-based substrates

The majority of current bio-organic substrate conversions take place employing photoelectro-oxidation. Based on the lower oxidative potential of biomass-based substrates compared to water, a large number of different types of biomass and their derived monomers are used for a wide variety of high-value purposes. Cathode sides are being tried for more and more reductive applications beyond efficient H_2 production. From the perspective of substrate conversion, the valorization applications at the anode and cathode will be explored separately in detail.

3.1. Photoelectro-oxidation at the anode

The oxidation performance can be characterized using line scan voltammetry (LSV) curves. The photooxidation of organic substrates can be visualized with lower onset potentials and surging current densities compared to catalytic systems without organic substrates (sometimes can be directly interpreted as water splitting systems), proving that the PEC of organic compounds is thermodynamically and dynamically preferable. The photocurrent can be enhanced to varying degrees by different types of organic compounds.³⁷ However, in contrast to (hemi-)cellulose, lignin has a more complex structure and darker color, which makes direct photocatalysis of the liquid phase system more difficult. Therefore, there are also some differences in the PEC application for (hemi-)cellulose and lignin at the anode.

3.1.1. (Hemi-)cellulose and its derivatives. Glucose, the most common depolymerized product of cellulose, is worthy of exploration for its PEC conversion as a substrate. At 1.23 V vs. RHE, the photocurrent of $\text{nanoFe}_2\text{O}_3/m\text{-CuO}$ containing glucose was 15 times higher than that of $\text{nanoFe}_2\text{O}_3/m\text{-CuO}$ without glucose. The Faraday efficiency (FE), also known as the current efficiency, is defined as the proportion of the passing charge assigned to the formation of the target product. The system also exhibited high stability and high FE ($97.3 \pm 2.8\%$) at 1 V vs. RHE (Fig. 3), which demonstrates the outstanding role of glucose in enhancing the performance.³⁶ When single-atom-anchored defective TiO_2 nanorod arrays (NRAs) are employed as photoanodes, glucose can be selectively oxidized to high-value glucaric acid (Fig. 4). Glucaric acid and its derivatives are widely utilized in the production of a range of health-care commodities, including cholesterol reduction and cancer chemotherapy.³⁸ In this system, holes react with adsorbed water to generate $\cdot\text{OH}$ radicals,³⁹ which extract hydrogen atoms to form $\text{C}1=\text{O}$ bonds.^{40,41} The $\text{C}5=\text{O}$ bond is cleaved by hydrolysis and desorbed from the photoanode to produce gluconic acid.⁴⁰ The $\text{C}6$ -position is finally oxidized through the highly selective oxidation of the primary alcohol by Pt and the binding of the $\cdot\text{OH}$ radical to yield *L*-glucuronic acid. The $\text{C}6=\text{O}$ bond and re-formed $\text{C}6\text{-OH}$ activated by holes as well as $\cdot\text{OH}$ will be regarded as the $\text{C}1$ -position and follow the same pathway to eventually produce glucaric acid. The conversion of gluconic acid into *L*-glucuronic acid is the rate-control-

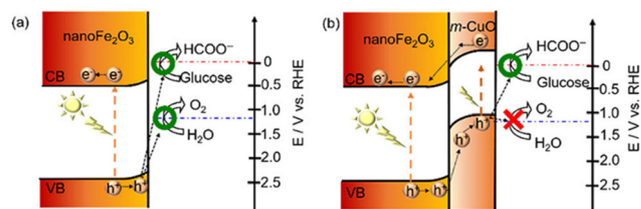


Fig. 3 Schematic representation of the electronic band structures of (a) $\text{nanoFe}_2\text{O}_3$ and (b) $\text{nanoFe}_2\text{O}_3/m\text{-CuO}$. Adapted from ref. 36 with permission from the Royal Society of Chemistry, copyright 2022.

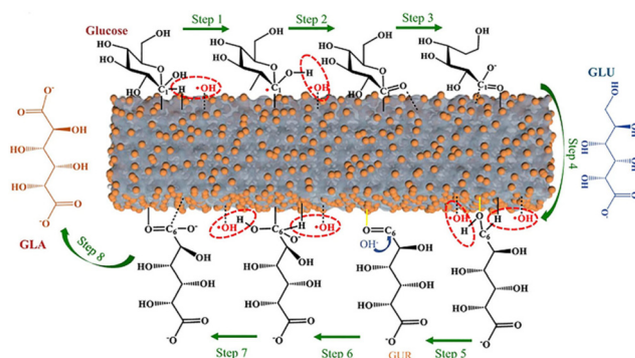
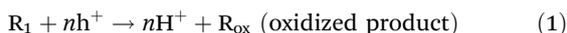


Fig. 4 Possible reaction pathway from glucose to glucaric acid over the Pt/def- TiO_2 photoanode. Adapted from ref. 38 with permission from Springer Nature, copyright 2023.

ling step and Pt plays a key role in the reaction with adsorbed $\cdot\text{OH}$ radicals.³⁸ The nanostructured semi-transparent WO_3 photoanode is capable of oxidizing glucose to gluconic acid and glucaric acid with acidic sulfate under simulated sunlight irradiation. The $2e^-$ oxidation pathway of glucose in this system is a direct transfer of photogenerated holes to the adsorbed organics, followed by the injection of a single-electron reaction intermediate to the CB to form gluconic acid.⁴² Pan *et al.*,⁴³ on the other hand, exploited the $\text{NiOOH}/\alpha\text{-Fe}_2\text{O}_3$ photoanode to oxidize glucose to formate (FA), where aldose is the main intermediate in the conversion process (Fig. 5a). Glucose was first oxidized to arabinose and FA by C1–C2 bond cleavage. Arabinose underwent similar successive C1–C2 bond ruptures to form the corresponding aldose and finally FA. This catalyst is also highly versatile for real biomass substrates. The PEC performance of the substrates obeys the following order: glucose > (hemi)cellulose > raw biomass.⁴³

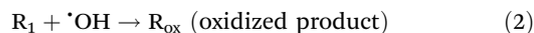
The other common biomass-derived substrate is glycerol. Glycerol is a by-product of the biodiesel production process and comprises roughly one-tenth of the biodiesel mass.⁴⁴ Glycerol is considered a promising sacrificial agent due to its non-toxic and trihydroxy structure.⁴⁵ In addition to being sacrificed to CO_2 for hydrogen evolution at the cathode, glycerol can be converted into many high-value chemicals such as dihydroxyacetone (DHA), glyceric acid (GLA), glycolic acid (GLCA), lactic acid, acetic acid, and FA.⁴⁶ The photo-oxidation of glycerol occurs through two main pathways:

(i) Reaction with photogenerated holes directly:



The photogenerated holes can activate glycerol molecules and efficiently cleave the C–C bonds of C2 and C3 intermediates. Consequently, highly selective FA^{50,51} and other C1 products are easily available through this catalytic pathway. For example, in the photoelectro-transformation of biomass for hydrogen production, most of the C is eventually converted into CO_2 .^{52,53} The N-doped WO_3 photoanode, however, is able to reform C from glycerol to CO, since CO itself is an important fuel and chemical that is widely exploited in the synthesis of olefins and aromatics.^{54,55} During the oxidation of this system, glyceraldehyde is the main liquid product, accompanied by the other liquid product FA. Aldehydes prefer to produce CO, while FA mainly liberates CO_2 . Glycerol is first directly oxidized by photogenerated holes to glyceraldehyde, which is further de-carbonylated to CO and glycolaldehyde. Glycolaldehyde is then de-carbonylated to CO and formaldehyde. In contrast, aldehyde intermediates such as formaldehyde are readily over-oxidized to acids thereby leading to CO_2 (Fig. 6c).⁴⁹

(ii) Reaction with some oxidizing species indirectly, taking $\cdot\text{OH}$ as an example:



DHA has been widely applied as a high-value raw material in the cosmetic, pharmaceutical, fine chemical, and food

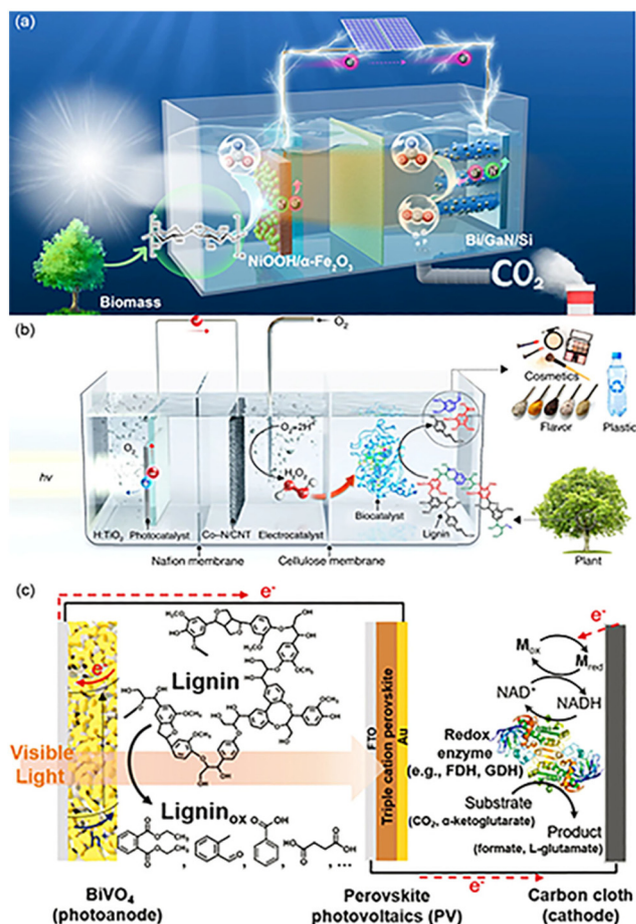


Fig. 5 Several designs of PEC configurations. (a) Schematic illustration of the tandem PEC cell for FA production from biomass and CO_2 . The PEC reactor consists of two compartments of photoanodic biomass oxidation (left-hand side) and photocathodic CO_2 reduction (right-hand side) for concurrent FA production. Adapted from ref. 43 with permission from Springer Nature, Copyright 2023. (b) Schematic illustration of the three-compartment photo-electro-biochemical reactor. The photo-electro-biochemical reactor consists of three separate compartments. The first compartment (left-hand side) is the anodic part of the photo-electrochemical cell where water oxidation takes place. The photoanode is connected to the Co-N/CNT cathode through a copper wire for *in situ* H_2O_2 generation in the second compartment (middle part). A Nafion proton exchange membrane separates the anodic and cathodic parts of the photo-electrochemical cell. The schematic of the second and third compartments shows the gradual increase in the concentration of photogenerated H_2O_2 and continuous flow to the biocatalytic system (third compartment), which are ideal conditions for selective lignin conversion. Adapted from ref. 47 with permission from Springer Nature, copyright 2019. (c) Schematic illustration of the lignin-fueled photoelectrochemical platform for solar-driven redox biotransformations. The BiVO_4 photoanode promotes the photoelectrochemical oxidation of lignin. The triple-cation perovskite PV forms a tandem structure with the BiVO_4 photoanode, providing supplementary photovoltage to drive reactions under zero external bias. The carbon-cloth cathode reduces NAD^+ to NADH for the reductive conversion of substrates using redox enzymes. Adapted from ref. 48 with permission from the Royal Society of Chemistry, copyright 2020.

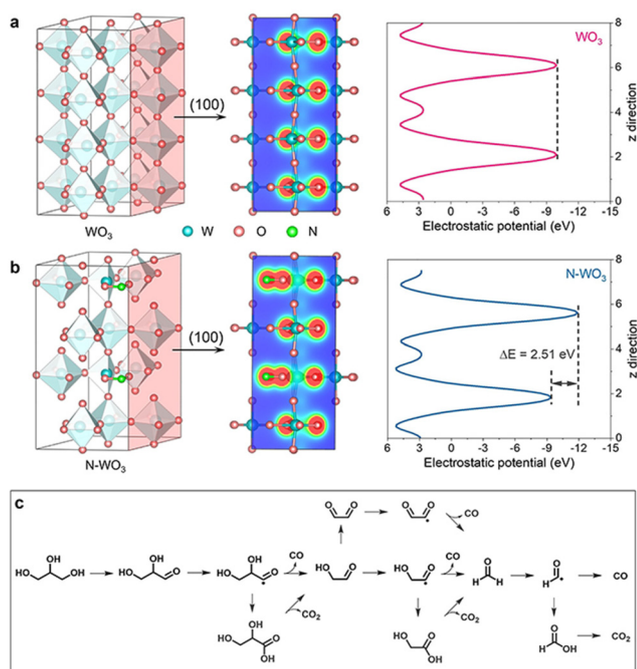


Fig. 6 Photoelectronic properties. Electron density isosurface (100) and electrostatic potentials of (a) WO_3 and (b) N-WO_3 . (c) Proposed reaction route for PEC glycerol reforming to CO. Adapted from ref. 49 with permission from John Wiley and Sons, copyright 2022.

industries. The price of DHA is \$150 per kg, while the price of raw and purified glycerol is only \$0.11 and \$0.66 per kg, respectively, which makes the catalytic production of DHA economically attractive.^{56,57} The $\text{Bi}_2\text{O}_3/\text{TiO}_2$ photoanode was able to PEC oxidize glycerol to DHA with 75.4% selectivity, and the conversion of the system could reach 50%, which is 160 times higher than that by photocatalysis under the same light conditions and 7 times higher than that by electrocatalysis at the same voltage (1 V vs. RHE). The PEC glycerol oxidation is an $\cdot\text{OH}$ radical-mediated process, and photogenerated holes play an essential role in the activation of H_2O by electron spin resonance (ESR), isotope labeling, fluorescent probe experiments, liquid chromatography-mass spectrometry (LC-MS), and free radical quenching experiments. PEC glycerol oxidation is accompanied by a lattice oxygen oxidation process.³⁹ Hence, the catalytic system follows two main parallel reaction mechanisms: (i) oxidation of glycerol by electrophilic $\cdot\text{OH}$ generated by water oxidation and (ii) dehydrogenation of glycerol by lattice oxygen followed by oxidation of lattice hydroxyl groups by photogenerated holes. The PEC configuration integrated with the Pt-perovskite photocathode and the $\text{Cu}_{30}\text{Pd}_{70}$ oxidation catalyst for the transition of three model compounds namely glycerol, ethylene glycol, and glucose revealed selectivities of $79 \pm 4\%$, $95 \pm 2\%$, and $65 \pm 7\%$ for glyceric acid, glycolic acid, and gluconic acid, respectively.⁵⁹ The slightly lower selectivity of gluconic acid may be due to the base degradation, isomerization and other side reactions of glucose over time.⁶⁰

Tartaric acid ($\text{C}_4\text{H}_6\text{O}_6$), a biomass derivative rich in hydroxyl and carboxyl groups, is not only adopted as an organic fuel

but is also able to complex with many metal ions, such as Bi^{3+} , Cu^{2+} , Fe^{2+} , etc.^{61,62} Thus, tartaric acid is able to establish a strong complexation with BiVO_4 , and this complexation also serves as a bridge for charge and energy transfer (Fig. 7b). It not only improves the photoelectric conversion efficiency of BiVO_4 but also effectively converts the chemical energy of biomass into H_2 .⁵⁸ A current density of 13.54 mA cm^{-2} can be achieved at 1.23 V vs. RHE, which is 1.8 times the theoretical maximum current density of BiVO_4 . Organic fuels containing only hydroxyl or carboxyl groups do not maximize PEC performance, while hydroxy-carboxyl-based biomass with other strong complexation capabilities can also contribute to PEC performance. Besides, Lv *et al.*³⁷ found FA, diethanolamine and triethanolamine as hole sacrificial agents, while methanol, glucose, and urea were $\cdot\text{OH}$ radical sacrificial agents after PEC oxidation of different kinds of organic compounds using Mn-doped CdS quantum dot-supported TiO_2 nanotube arrays (NTAs). Incident photon-to-current efficiency (IPCE) tests indicate a current doubling effect for methanol, where the $\cdot\text{OH}$ radical (which generates the first photoelectron) leads to a highly reduced α -hydroxymethyl radical.⁶³ The radical then injects a second electron into the photoanode to form formaldehyde.^{63–65} Each photon produces two photoelectrons, so the theoretical IPCE is up to 200%. In practice, only 130% is achieved at 320 nm due to the limitation of net charge separation and transfer. The three alcohol moieties of triethanolamine show the same current doubling effect as an effective electron donor, with about 160% at 320 nm. 5-Hydroxymethylfuran (HMF) was oxidized to 2,5-furandicarboxylic acid (FDCA) by a 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO)-mediated photo-electronic system, which consumed a charge of 55 C. HMF was ultimately oxidized to FDCA through the intermediates of furfuraldehyde and furfuraldehyde acid. The yield and FE of FDCA were 100% at 0.7 V vs. RHE.⁶⁶

In addition to model compounds, PEC oxidation of biomass feedstocks is being progressively explored. Cellulose films deposited on porous TiO_2 are used as a fuel and configured with the Pt cathode to decompose cellulose to CO_2 .¹¹ Carbonyl carbohydrate chains such as gluconic acid, oxalic acid, acetic acid, and FA are intermediate products dissolved

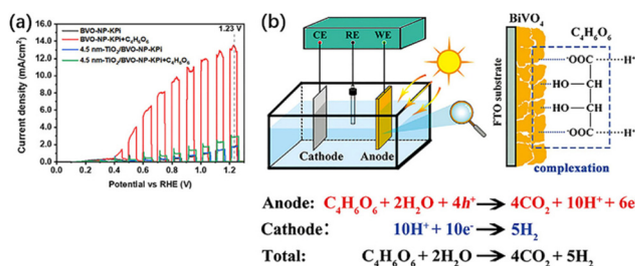
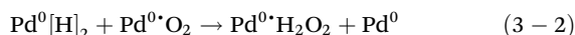
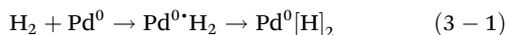


Fig. 7 (a) Current density versus potential curves of BVO-NP and 4.5 nm $\text{TiO}_2/\text{BVO-NP}$ photoanodes in potassium phosphate (KPi) and $0.5 \text{ M C}_4\text{H}_6\text{O}_6$ solutions under AM 1.5G illumination or in the dark. (b) Schematic illustration of the detailed reaction process in the BiVO_4 PEC fuel cell system. Adapted from ref. 58 with permission from John Wiley and Sons, copyright 2022.

in the liquid phase during PEC. In the decomposition process, cellulose is first cleaved to short carboxylic acids, which are further decomposed to CO₂. Pre-treated polyethylene terephthalate (PET) powder, real-life commercial PET bottles, and microcrystalline cellulose similarly underwent PEC conversion using the Cu₃₀Pd₇₀ catalyst with the Pt-perovskite photocathode. After 10 h in both compartments, the selectivities of glycolic acid from PET powder and PET bottles were 94% and 91%, respectively, and that of gluconic acid from microcrystalline cellulose was 67%.⁵⁹ Microalgal biomass can generate renewable energy because it does not need to occupy arable land for growth and is characterized by high lipid yields per unit, short growth cycles, and high environmental adaptability.⁶⁷ However, the thick cell wall and selectively permeable cell membrane structure make the extraction of lipids difficult.⁶⁸ Based on this, the PEC system consisting of the N-doped TiO₂ nanotube photoanode, a bipolar membrane, and the P-Pd cathode utilizes eventually ·OH radicals, which are produced from the H₂O₂ generated from H₂ and O₂ at both electrodes^{69,70} by the strong adsorption and activation ability of Pd-based catalysts, to realize this purpose:⁷¹



·OH radicals can attack the hydrogen bonds among the carbon chains between cellulose and pectin molecules in the algal cell wall.⁷² ·OH radicals can also attack diacylglycerides in the cell membrane, leading to peroxidation that breaks carbon chains of unsaturated fatty acids as well as lipids from saturated fatty acids.⁷³ After the cell wall and cell membrane rupture, organic solvents are then allowed to diffuse into the cell to extract the lipids. At the same time, ·OH also breaks single and double bonds in chlorophyll, which contributes to the decolorization of crude oil.⁷⁴

3.1.2. Lignin and its derivatives. As the predominant ether bond in the lignin structure, the β-O-4 bond is the most studied object in lignin model compounds, together with the real lignin being mainly oxidized by holes. C-H and O-H bonds in organic molecules can act as electron donors to consume photogenerated holes and produce H⁺. The opaque solution of dissolved lignin prevents light from reaching the photocatalyst, so the problem can be effectively solved *via* back irradiation, the addition of redox mediators, or light avoidance in the anode compartment.^{47,75,76}

Although there are many studies on the PEC conversion of β-O-4 bonds, different conversion outcomes can be achieved. There are four main conversion pathways as follows: (i) oxidation of β-O-4 alcohol to β-O-4 ketone; (ii) C_α-C_β cleavage of the β-O-4 bond; (iii) C_{Ar}-C_α cleavage of the β-O-4 bond; and (iv) C_β-O cleavage of the β-O-4 bond (Fig. 8). Pathway (i) can be achieved by dye-sensitized photoelectrochemical (DSPEC) systems. The C-O bond of β-O-4 can significantly be weakened after alcohol oxidation of the lignin skeleton, which contrib-

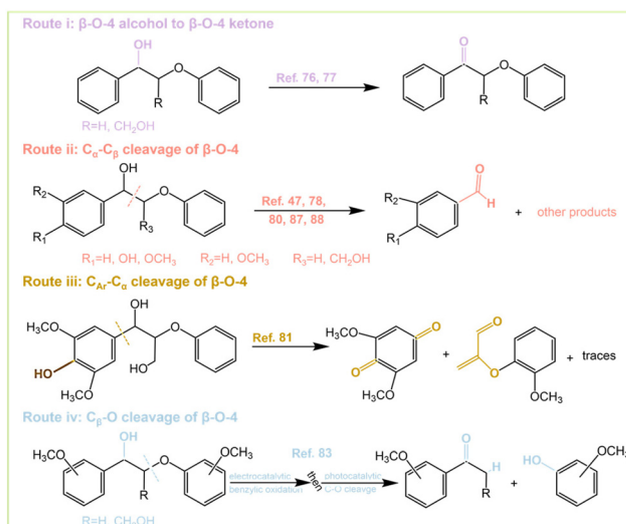
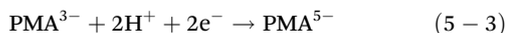


Fig. 8 Four routes of PEC β-O-4 transformation.

utes to the subsequent cleavage.^{76,77} Kim *et al.*⁷⁸ explored the oxidative reaction species of Kraft lignin using α-Fe₂O₃ (hematite) photoanodes and found that photogenerated holes play a key role in lignin depolymerization. Lignin depolymerization can be accomplished by oxidation of C_α-OH to C_α=O and subsequent oxidative cleavage of C_α-C_β. It can be expected that the depolymerization proceeds through pathway (i) and pathway (ii) successively. BiVO₄ facilitates the direct transfer of photogenerated holes to lignin, thus weakening the C-C bond and causing the decomposition as well as oxidation of lignin to organic acids.⁷⁹ Path (ii) is the most common way of breaking the β-O-4 bond and the typical product is the formation of benzaldehyde types of molecules. β-O-4 bonds are successfully converted by depositing V₂O₅ on the BiVO₄ photoanode under a bias of 2 V. The depolymerization products are mainly *p*-anisaldehyde, phenol, and FA.⁸⁰ The presence of hydroxymethyl at the β-position of β-O-4 significantly increases the decomposition activity, and the presence of methoxy at the *para*-position of both benzene rings further improves the conversion. Route (iii) was also carried out by a DSPEC system, where the active hydrogen atom transfer (HAT) mediator was 4-acetamido-2,2,6,6-tetramethylpiperidine-1-oxyl (ACT). Oxidation of HAT-DSPEC was able to afford 2,6-dimethoxybenzoquinone and the corresponding other cleavage product after C_{Ar}-C_α bond severing (Fig. 8 Route iii). Model compounds that do not contain *p*-hydroxyl groups are unable to undergo C-C bond fracture.⁸¹ The selectivity of the reaction depends on the reaction time and the presence of light. The gradual decrease of the anodic current with prolonged illumination time is associated with the irreversible depletion of the β-O-4 model compound and degradation of the photocatalyst.⁸² Route (iv) can be achieved by a two-step one-pot method of β-O-4 bonds (Fig. 8 Route iv and Fig. 11). This method is subject to electrocatalytic oxidation followed by photocatalytic cleavage to finally obtain cracked ketones and guaiacol.⁸³ The whole cata-

lytic system does not strictly belong to the scope of PEC, which is a combination of photocatalysis and electrocatalysis but has a strong significance. The reactions were performed on coniferyl- and sinapyl-derived dimers, and even natural lignin with good yields, and the investigation of lignin depolymerization revealed that the electrocatalytic oxidation–photocatalytic cleavage process not only cleaves the β -O-4 bond efficiently but also leaves the other components intact with high selectivity. The extraction of benzyl hydrogen during the reaction may be the rate- or turnover-determining step.

Oxidation of lignin occurred more readily than water oxidation when using a tandem structure of the BiVO_4 photoanode with a perovskite PV for the conversion of real biomass feedstocks, but other non-lignin biocomponents (e.g., cellulose and xylose) had negligible effects on photocurrent generation.⁴⁸ Oxidation of lignin occurs mainly at bonds with lower dissociation energies such as β -O-4 ($54\text{--}72\text{ kcal mol}^{-1}$) and β -5 ($54\text{--}63\text{ kcal mol}^{-1}$) rather than at bonds with higher dissociation energies such as β - β (81 kcal mol^{-1}).⁸⁴ Further extension of the overoxidation time enables decomposition to the simplest form, i.e. CO_2 . The phosphomolybdic acid (PMA) PEC biomass oxidation system also proved to have a more outstanding performance than PEC water oxidation. Compared to the high voltage ($1.5\text{--}1.6\text{ V vs. RHE}$) required to reach 10 mA cm^{-2} for the aqueous oxidation system,^{85,86} the biomass system required only 0.95 V vs. RHE at the same current density.⁸⁷ Comparison of the relative contents of β -O-4, β -5, and β - β structures of lignin before and after the reaction demonstrated that the main selective rupture of the oxidation reaction was that of the β -O-4 bond. The main products of lignin depolymerization are vanillin and CO (Fig. 9c), where CO is mainly formed due to the presence of dissolved oxygen molecules. According to Arrhenius kinetics, the activation energies for PMA reduction by oxidation of lignin, cellulose, hemicellulose, and lignocellulosic biomass are $24, 102, 79, \text{ and } 47\text{ kJ mol}^{-1}$ ($0.249, 1.057, 0.819, \text{ and } 0.487\text{ eV}$), respectively, indicating that lignin is the main source of electrons and is more readily depolymerized than other components.⁸⁸ Lignin was selectively removed from the lignocellulosic biomass after reacting with PMA, resulting in the formation of a hierarchical porous microstructure. The general reaction mechanism is as follows:



In addition, there is the PEC phenol oxidation process catalyzed by SnO_2 and Sb photoanodes coated on TiO_2 nanotubes in order to provide electrons for electrochemical CO_2 reduction. The current density under visible light irradiation is 12 times higher than that under dark conditions when phenol is added to Na_2SO_4 solution.⁸⁸ The reactions at the anode are as follows:



100% selective benzyl alcohol to benzaldehyde conversion was achieved by the $\text{Bi}_2\text{MoO}_6/\text{TiO}_2$ NTA photoanode and the $\text{C}/\text{Cu}_2\text{ONW}$ photocathode.⁹⁰ Benzaldehyde is a raw material that is widely employed in pharmaceutical, chemical, printing, and dyeing areas.^{91,92} The estimated flat-band potential (E_{FB}) of the photoanode is -0.34 V , which is more electronegative than that of O_2/O_2^- (-0.28 V vs. NHE).⁹³ The photoanode can adsorb O_2 via photogenerated electrons and further activate it to the O_2^- radical. Benzyl alcohol can be directly accelerated and converted into benzaldehyde through such radicals. Therefore, O_2 is very critical for the solution system.⁹⁴ The conversion of benzyl alcohol into benzaldehyde is a 2e^- process, and the conversion of benzaldehyde into benzoic acid is also a 2e^- process. The conversion of benzyl alcohol into benzaldehyde is a little easier, but the conversion of benzaldehyde into CO_2 is more difficult because the reaction is a 30e^- process. In the investigation for the effect of different substituents (e.g., $-\text{OCH}_3$, $-\text{CH}_3$, $-\text{F}$, and $-\text{NO}_2$) on the conversion of benzyl alcohol at the *para*-position, it was found that $^*\text{C}_6\text{H}_5\text{--CH}_2\text{OH}^+$ is the major intermediate. $-\text{OCH}_3$ has both an electron-withdrawing induction effect and an electron-donating conjugation effect. However, $-\text{OCH}_3$ is an electron-donating functional group because the lone pair of oxygen plays a dominant role in the electron-donating conjugation effect. The electron-donating functional groups are more helpful for the formation of $^*\text{C}_6\text{H}_5\text{--CH}_2\text{OH}^+$ intermediates, so the conversion of 4-methoxybenzyl alcohol is higher than that of benzyl alcohol. In contrast, the strong electron-withdrawing induction effect of the $-\text{NO}_2$ group reduces the electron cloud density of benzyl intermediates. Therefore, the higher the electron-donating ability, the higher the conversion. The electron-donating ability of different substituents is: $-\text{OCH}_3 > -\text{CH}_3 > -\text{H} > -\text{F} > -\text{NO}_2$. Benzoic acid is a precursor of phenol, nylon, and benzoate plasticizers in the chemical industry, and is also known to be applied as a food preservative, mold inhibitor, and pharmaceutical additive.^{95,96} After oxidation by the nanowire-structured $\text{Mo--BVO--TiCoNO--CoFeO}_x$ photoanode, benzyl alcohol was oxidized to benzoic acid (97.7% yield) in water, accompanied by an H_2 yield of $136.8\text{ }\mu\text{mol cm}^{-2}$.⁹⁷ Benzyl alcohol adsorbed on the photoelectrode is oxidized by photo-generated h^+ , where the alcohol hydroxyl group reacts with hydroxide in the electrolyte to form H_2O , which is stripped off from the substrate, leaving the remaining carbonyl radicals. Subsequently, the carbonyl radical interacts with the hydroxyl radical in the electrolyte to form benzoic acid.

Through oxidation in the anode cell, many valuable chemicals or fuels are delivered from the biomass derivatives. But what has to be mentioned is that many substrates are oxidized to low-value CO_2 in the anode cell in order to garner hydrogen or other valorized products efficiently in the cathode cell. The simultaneous valorization at both cathode and anode is also an ideal concept.

3.2. Reductive valorization at the cathode

When electrons accumulated on the CB at the anode are transferred to the cathode, a large amount of H^+ can be reduced to

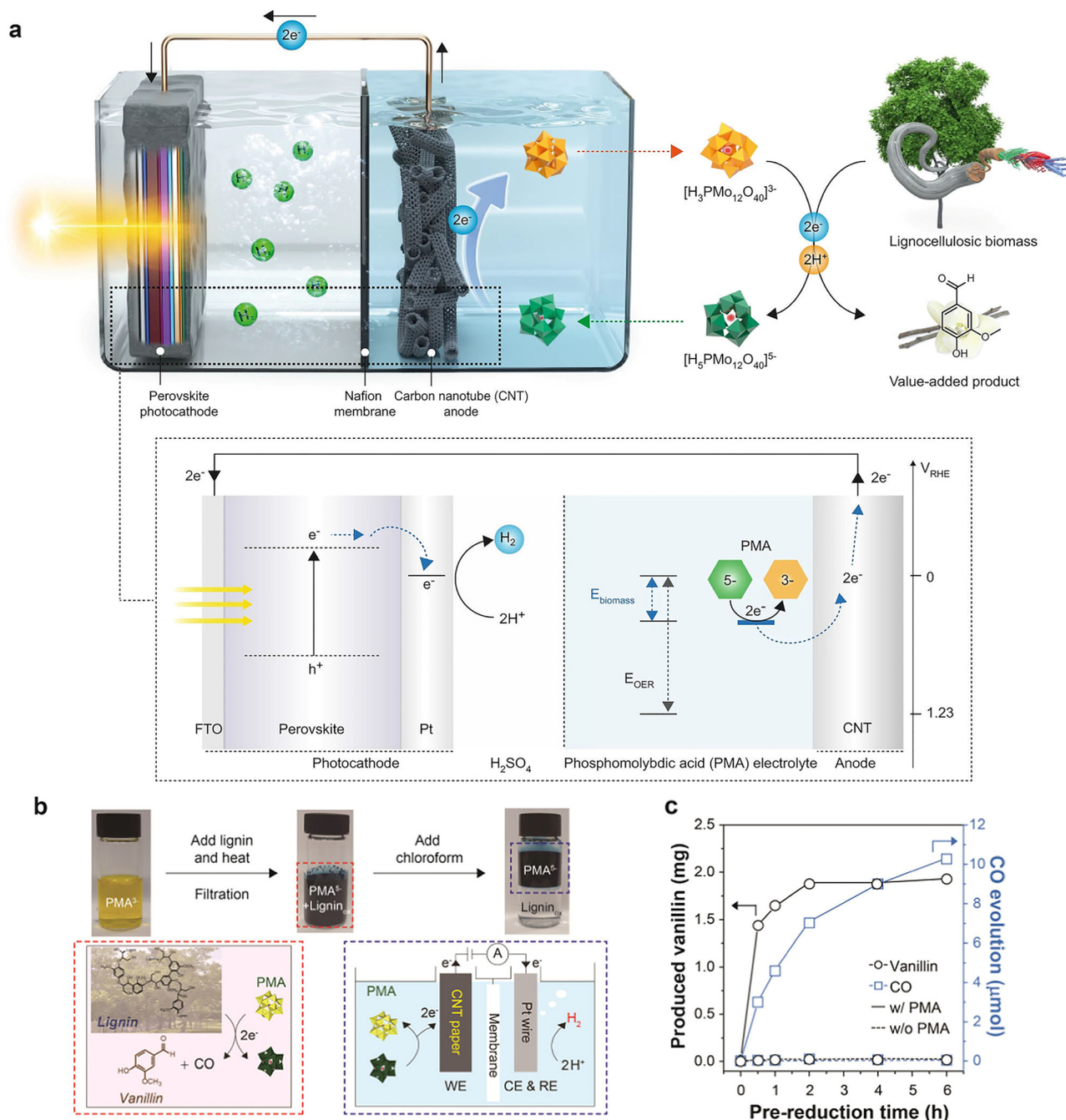


Fig. 9 (a) The perovskite photocathode with the Pt catalyst and the electron extraction system from biomass are combined for bias-free hydrogen evolution. Adapted from ref. 88 with permission from Springer Nature, copyright 2022. (b) Photographs and reaction scheme showing the pre-reduction of PMA upon lignin oxidation and separation of oxidized by-products from lignin using chloroform. (c) Concentration profile of CO and vanillin upon oxidation of lignin using PMA at 50 °C. Adapted from ref. 87 with permission from the American Chemical Society, copyright 2020.

H₂. H₂ is green source of energy and is recognized as one of the most promising alternatives among the existing forms of energy (coal, petroleum, fossil fuels) in society.^{98,99} Therefore, for most PEC conversions in order to achieve efficient hydrogen production at the cathode side, biomass-derived waste is often used as a hole sacrificial agent, and the performance of PEC for H₂ production depends entirely on the photoelectrotransformation of the photo-electrode.¹⁰⁰ The oxidation of PEC

furfuryl alcohol with the TiO₂ nanotube anode and the Ni foam cathode yielded only a small amount of furfural, attributed to the competitive reaction between furfural and water oxidation. The O₂ from H₂O oxidation diffuses to the cathode, which in turn affects the rate of hydrogen production.¹⁰¹ Kong *et al.*⁴⁹ achieved an H₂ evolution rate of 237 mmol m⁻² h⁻² with purity >99.99% using the N-doped WO₃ photoanode and the Pt cathode. When PMA is utilized as an oxidation catalyst,

the p-type photocathode Pt-black silicon can be used for efficient hydrogen production.⁸⁷ After the replacement of Pt-black silicon by a perovskite photocathode, a maximum current density of 19.8 mA cm⁻² was achieved without bias under 1 sun illumination. The hydrogen production efficiency reached 512 μmol cm⁻² h⁻¹, which is a record-breaking high value for the solar hydrogen production system reported so far, regardless of the type of device. At the same time, the system permits the continuous production of high-value chemicals through biomass depolymerization at night and solar hydrogen production conversion in the daytime.⁸⁸

There is a relative shortage of research on photocatalytic CO₂ reduction, and Shang in his review had great expectations for the reduction of CO₂.¹² The application of CO₂ reduction based on the sacrifice of organics in PEC systems has also been developed in recent years. The system for PEC phenol oxidation starts from electrochemical CO₂ reduction using a Cu nanowire cathode to reduce CO₂ to CO, accompanied by the production of H₂ and small amounts of carbohydrates such as CH₄, C₂H₄, and C₂H₆.⁸⁹ The amount of available CO can be increased when the system uses NaHCO₃ instead of Na₂SO₄, because the bicarbonate increases the CO₂ concentration in the solution thus facilitating the reduction of CO₂.¹⁰² PEC along with biocatalysis similarly achieved the CO₂ to FA conversion process. CO₂ was reduced to FA using nicotinamide cofactor analog (NADH)-dependent formate dehydrogenase from *Thiobacillus* sp. (TsFDH). The synthesis rate of FA was 4.5 times higher than that based on water oxidation.⁴⁸ The concentration of lignin at the anode was proportional to the formation rate of FA. The oxidation of lignin, during biocatalytic CO₂ reduction, is the rate-controlling step. This system also achieved selective amination of α-ketoglutarate to L-glutamate using L-glutamate dehydrogenase in addition to CO₂ reduction (Fig. 5c).

Moreover, the application of PEC in combination with biocatalysis seems to be even broader. The regenerated NADHs activate ene-reductases of the old yellow enzyme family, which can catalyze the enantioselective reduction of α,β-unsaturated carbohydrates. The conversion of 2-methyl-2-cyclohexen-1-one into enantiopure (*R*)-2-methylcyclohexanone was achieved together with a silicon photovoltaic mesoporous indium tin oxide (Si/*meso*ITO) photocathode.⁷⁸ Lignin peroxidase isozyme H8 (LiPH8) from white rot bacteria is capable of excellent selective splitting of β-O-4.¹⁰³ However, the high concentration of H₂O₂ is decisive for the system's stability. Photocatalytic production of H₂O₂ is a straightforward, clean, and ambient operable process, exploiting photogenerated electrons for O₂ reduction to produce H₂O₂. Based on this idea, divided three-compartment photo-electro-biochemical systems were designed for the valorization of lignin without additional bias and sacrificial agents (Fig. 5b). H₂O is oxidized by photogenerated holes on the VB to O₂, and electrons are transferred to the cathodic cell for the reduction of O₂ to harvest H₂O₂. H₂O₂ penetrates through the size-selective cellulose membrane and is then bio-catalytically valorized for lignin.⁴⁷ The main product of β-O-4 cleavage is 3,4-dimethoxybenzaldehyde, the

conversion of which is mainly limited by the concentration of H₂O₂. The bioenzyme was replaced with the horseradish peroxidase (HRP) biocatalyst in the presence of H₂O₂ for biopolymer synthesis as well. The conversion and biopolymer yield can be up to 98.3% and 73.3%, respectively, using coniferyl alcohol as the feedstock. The polymers consisted of common bonding types like β-O-4, β-β, and β-5. The number average molecular weight was 1103. Chitin is the second largest source of carbohydrates for the production of biofuels and chemicals due to its efficient and sustainable transformations. But destroying the crystalline structure similar to cellulose and producing soluble oligosaccharides and monosaccharides is one of the challenges in the development of chitin.¹⁰⁴ An inorganic biohybrid platform by integrating the Si photocathode and lytic polysaccharide monoxygenase (LPMO)-based enzyme systems can cascade catalyze α-chitin to soluble oxidized oligosaccharides. 2,6-Dimethyl-1,4-benzoquinone was chosen as a connecting agent among a host of quinones due to the best electrochemical reversibility and suitable redox potential. In the presence of light, the quinone mediator is reduced on a silicon photocathode. Capitalizing on the reducing power provided by the mediator, α-chitin is then degraded to soluble products by oxygen activation on LPMO. At the same time, the mediator is recovered to its initial state to complete the cycle. All components are indispensable.¹⁰⁴

Whether in the anodic or cathodic cell, there are many more well-known model compounds, bonding types, and even pristine biomass in the field of PEC biomass that can be explored, accompanied by conversion mechanisms that are challenging to fulfill by photocatalysis, electrocatalysis, and even thermo-catalysis. These exciting results cannot be achieved without the design and preparation of catalysts. Intelligent design of catalysts at both electrodes combined with appropriate catalytic configurations can lead to endless possibilities for PEC biomass conversion.

4. Design and exploitation of catalysts and PEC configurations

Similar to plain photocatalysis, a prerequisite for a PEC system to function well is the formation of an internal electric field (IEF) in the semiconductor photocatalyst/system. Aside from the direct application of acquainted photocatalysts, there are many ways to increase the catalytic activity, for example by creating defects such as oxygen vacancies, atomic doping, and atomic or compound modifications (Table 1). Nevertheless, a unique configuration is particularly vital when special conversion outcomes need to be achieved or the applied bias of the whole system needs to be brought down.

4.1. The modulation of catalysts

The main types of semiconductor catalysts utilized in biomass PEC systems are currently centered on TiO₂-based, WO₃-based, BiVO₄-based, perovskite, Fe₂O₃-based, and Si-based catalysts. Most types are already well known in photocatalysis, and there

Table 1 Summary of current valorization via PEC transformation using biomass and derivatives as substrates

Substrates	Anode	Cathode	Illuminants (mW cm ⁻²)	Bias (V vs. RHE)	Major products	Current density (mA cm ⁻²)	Entry ^{Ref.}
Glycerol	N-WO ₃	Pt foil	100	1.2	CO + H ₂	1.4	1 ⁴⁹
Tartaric acid	BiVO ₄	Pt plate	100	1.23	H ₂	13.54	2 ⁵⁸
Glucose	nanoWO ₃	Pt grid	100	1.23	Gluconic + glucaric acids or erythrose + arabinose	6.5	3 ⁴²
Glucose	nanoFe ₂ O ₃	<i>m</i> -CuO	100	1.0	FA	0.4	4 ³⁶
Glucose	Pt/def-TiO ₂ NRAs	Pt foil	100	0.6	Glucaric acid	1.91	5 ³⁸
Glycerol	CeO ₂ /Ce-doped ZnO ^a	Pt coil	150 W Xe lamp	1	H ₂	0.91	6 ¹⁴²
Glycerol	Bi ₂ WO ₆	Pt mesh	120	1.3	FA + H ₂	0.69–0.8	7 ¹³
Glycerol	Bi ₂ O ₃ /TiO ₂	Pt foil	100	1	DHA	0.27	8 ³⁹
Furfuryl alcohol	TiO ₂ NTs	Ni foam	Six UVA fluorescent lamps (<365 nm, 8 W)	0.5 V vs. Ag/AgCl	Furfuraldehyde + H ₂	0.002	9 ¹⁰¹
HMF	NiFe-LDH/BiVO ₄	Pt	100	0.7	FDCA	3.54	10 ⁶⁶
Glycerol	WO ₃	Pt wire	350 (LED UV lamp)	0.5–2	Glyceraldehyde + DHA	0.8	11 ¹¹⁹
Glycerol	CoNiFe-LDH/Ta ₃ N ₅	Pt plate	100	1.23	FA + H ₂	3.59	12 ⁵¹
Glycerol	BVO/TANF	Pt wire	100	1.23	FA	5.26	13 ⁵⁰
Glucose-CO ₂	NiOOH/α-Fe ₂ O ₃	Bi/GaN/Si	100	0	FA	1.25	14 ⁴³
β-O-4 alcohol	FTO TiO ₂ -RuC + NHPI/lutidine	Pt wire	200	0.75 V vs. SCE	β-O-4 ketone	0.13	15 ⁷⁶
β-O-4	FTO TiO ₂ NRAs RuC + ACT	Pt wire	100	0.1 V vs. Ag/AgCl	C _{aryl} -C _α bond cleavage products + 2,6-dimethoxybenzoquinone	0.69	16 ⁸¹
β-O-4 alcohol	FTO 1-TiO ₂ NRAs RuC + ACT	Pt wire	100	0.45 V vs. SCE	β-O-4 ketone	0.38	17 ⁷⁷
β-O-4	V ₂ O ₅ /BiVO ₄	Glassy carbon	100	2	<i>p</i> -Anisaldehyde + phenol + FA	2.8	18 ⁸⁰
β-O-4	H:TiO ₂	Co-N/CNT	100	0	3,4-Dimethoxybenzaldehyde	—	19 ⁴⁷
Phenol	Sb/SnO ₂ -TiO ₂ NTs	Cu nanowire	—	1.55 V vs. Ag/AgCl	CO + H ₂	0.17	20 ⁸⁹
Benzyl alcohol	Bi ₂ MoO ₆ @TiO ₂ NTA	C/Cu ₂ ONW	300 W Xe lamp	0	Benzaldehyde + H ₂	0.54	21 ⁹⁰
Benzyl alcohol	Mo-BVO-TiCoNO-CoFeO _{xy}	Pt	100	1.48	Benzoic acid + H ₂	1.31	22 ⁹⁷
Cellulose	TiO ₂	Pt	300 W Xe lamp (<500 nm)	0.65	CO ₂	3.8	23 ¹¹
Cellulose/PET bottle/PET powder	Cu ₃₀ Pd ₇₀	Pt-perovskite	100	0	Gluconic acid + glycolic acid + H ₂	3.7–5.1	24 ⁵⁹
<i>Chlorella</i>	N-Doped TiO ₂	P-Pd	100 W UV-Vis LED	—	Lipid	1.62	25 ⁷¹
α-Chitin	Pt	C/Si + LPMO	Xe lamp (>420 nm)	0	Oxidized oligosaccharides	0.085	26 ¹⁰⁴
Kraft lignin	α-Fe ₂ O ₃	Si/ <i>meso</i> ITO NADH	100	2.1 V vs. Ag/AgCl	Terephthalic acid + TME ^b + MHB ^c + IPM ^d + diethyl phthalate + EMC ^e	≈1.5	27 ⁷⁸
Kraft lignin	TiO ₂	Pt	25	2.3	Oxidized lignin	0.26	28 ⁷⁵
Alkali lignin/lignosulfonate-CO ₂	BiVO ₄ + perovskite	Carbon cloth	100	0	Oxidized lignin + FA	—	29 ⁴⁸
Kraft lignin	PMA	Pt black silicon	100	0.95	Vanillin + CO	10	30 ⁸⁷
Alkali lignin/lignocellulosic biomass	PMA	CsFAMA perovskite	100	0	Vanillin/vanillin + acetovanillone	19.8	31 ⁸⁸

Several catalyst preparation methods: oxygen vacancy defect (yellow), atom doping (blue), element/compound decoration (pink), or heterojunction construction (green). ^aThe catalyst was prepared by both CeO₂ decoration and Ce doping. ^bTME = 1-(2,3,4,5-tetramethoxy-6-methylphenyl)ethenone. ^cMHB = methyl 2-hydroxybenzoate. ^dIPM = (4-isopropylphenyl)methanol. ^eEMC = enantiopure (*R*)-2-methylcyclohexanone.

are discrepancies in the conversion from catalysts due to different band gaps.

4.1.1. TiO₂-based catalysts. TiO₂ is the most widely deployed n-type semiconductor photocatalyst but is confined by a narrow wavelength absorption range. To maximize the light absorption range, a series of approaches have been pursued to enhance the activity, such as the calcination of rutile under hydrogen to improve the charge transfer properties of TiO₂.⁴⁷ One class of systems that employs direct bandgap absorbers,

i.e., immobilized monolayer surface molecular dye coatings, to increase the light absorption range of TiO₂-based electrodes is called DSPECs.^{106,107} The design of DSPECs is strongly based on the design of dye-sensitized solar cells, which was first reported in 1991.¹⁰⁸ Combining the concept of mediated electrocatalysis (especially in lignin C–C or C–O bond rupture) and DSPECs, aminoxyl-mediated DSPECs for biomass conversion or alcohol oxidation in a heterogeneous medium are proposed.^{76,77,81} Three main components are involved: a primary catalyst com-

posed of TiO₂ and polypyridine Ru(II) complexes, aminoxyl-mediated co-catalysts, and sacrificial agents. Phosphonic acid or carboxylic acid groups on polypyridine Ru(II) complexes can be anchored onto TiO₂ and rely on aminoxyl compounds for charge transfer (Fig. 10a and b). Lignin and its model compounds are important sacrificial media in the DSPEC system.^{76,81} The likely reaction pathways are as follows: (i) photoexcitation of surface-bound Ru(II) photocatalysts; (ii) generation of electrons (TiO₂(e⁻)) and holes (Ru(III)) after electron injection into TiO₂ NRAs; (iii) electron transfer from the aminoxyl radical mediator (ARM) to Ru(III); (iv) formation of oxidized ARM⁺ that activates HAT-mediated oxidative cleavage of lignin substrates; and (v) relying on the HAT mediator, intramolecular proton transfer to direct C–C and/or C–O bond cleavage, and ARM-catalyzed dehydration to produce aromatic compounds¹⁰⁵ (Fig. 10c). C_α-OH of β-O-4 is able to be oxidized to C_α=O by mesoporous TiO₂ combined with (bis-2,2'-bipyridine) (2,2'-bipyridine-4,4'-dicarboxylic acid) Ru(II) (RuC) as a catalyst

and *N*-hydroxyphthalimide (NHPI)/lutidine as a co-catalyst.⁷⁶ RuC could catalyze the oxidation of NHPI to the *N*-oxy radical (PINO) form. The presence of lutidine is necessary to facilitate the charge transfer from photoexcited RuC* to NHPI. Beforehand, Bosque *et al.*⁸³ achieved the scission of the ether bond in a one-pot method using the electro-oxidation of β-O-4 by NHPI/lutidine followed by photocatalysis. The one-pot is featured in the fact that photocatalysis could be directly performed after the oxidation with the simple addition of the [Ir(ppy)₂(dtbbpy)](PF₆) photocatalyst, diisopropylethylamine (DIPEA), and FA to the reaction vessel. The photocatalytic performance is guaranteed even in the presence of additives from electrocatalytic oxidation in the system. NHPI is first oxidized to PINO at the anode, which can be facilitated by the proton-coupled electron transfer process of 2,6-dimethylpyridine (Fig. 11). HAT regenerates NHPI from the substrate and produces α-hydroxy radical 1' intermediates that should be prone to β-O-4 fracture. Polypyridine Ru(II) complexes have been

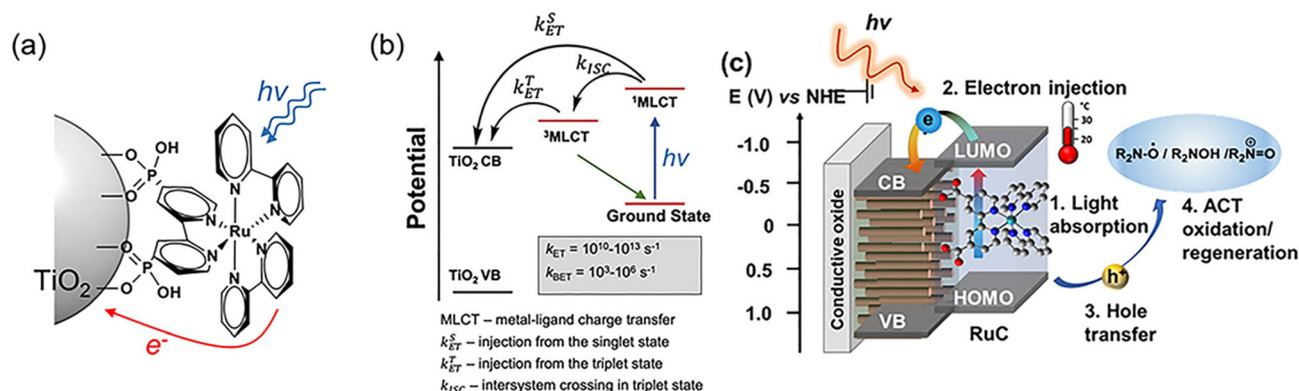


Fig. 10 (a) Illustration of a Ru-based chromophore and semiconductor assembly. (b) Photodynamics of the Ru(II) chromophore on a TiO₂ surface, electron injection from a singlet state and a triplet state into the conduction band, and back electron transfer to oxidized Ru³⁺. (c) Schematic for the visible light induced charge transfer between surface bound RuC and ACT in a PEC system. Adapted from ref. 105 with permission from the Royal Society of Chemistry, copyright 2023.

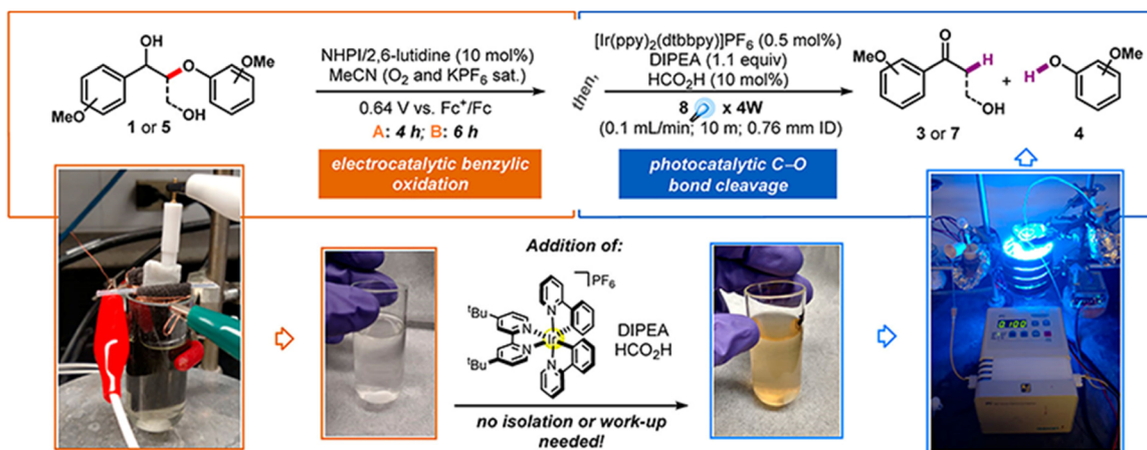


Fig. 11 Schematic for the one-pot electrocatalytic oxidation–reductive cleavage of β-O-4. Adapted from ref. 83 with permission from the American Chemical Society, copyright 2017.

widely applied in water splitting and lignin depolymerization due to their wide visible absorbance and durable excited state lifetimes.^{109,110} After replacing the mesoporous TiO₂ grown on the surface of fluorine-doped tin oxide (FTO) electrodes with TiO₂ NRAs (denoted as FTO|*t*-TiO₂|RuC, *t* is the thickness of TiO₂ nanorods (μm)), it was found that the length of TiO₂ NRAs has a significant effect on the photocatalytic performance; the shorter the NRAs the better the performance. The >99% yield of β-O-4 2° alcohol to ketone conversion and 88% FE could be achieved *via* the FTO|1-TiO₂|RuC anode with the shortest NRAs on the surface after 4 h of reaction.⁷⁷ ACT served as a photocatalyst to efficiently quench RuC* emission, indicating the charge transfer from photoexcited RuC* to ACT. This quenching efficiency is 3 times more amplified than that of the NHPI/base pair and does not require a base.⁸¹

Single-atom anchored defective TiO₂ NRAs as photoanodes can selectively oxidize glucose to high-value glucaric acid.³⁸ The oxygen vacancy-induced defect structure can modulate the charge carrier dynamics and energy band structure. The band gap of def-TiO₂ generated by oxygen vacancy defects drops from 2.99 eV to 2.70 eV. The narrower band gap is attributed to: (i) additional transitions due to incremental Ti³⁺ states; (ii) oxygen vacancies; (iii) hydroxyl functional groups; and (iv) other defects at different energy levels.¹¹¹ The prepared Pt/def-TiO₂ NRA catalyst is a core-shell structure made up of a rutile crystalline core with a disordered shell layer. The disordered shell layer furnishes a large number of active sites for the anchoring of single-atom Pt. The loading of Pt improves the selectivity for glucaric acid and the facilitation of Pt induces fewer holes to be involved in C-C bond breaking. In the system of PEC glycerol oxidation to DHA with 75.4% selectivity using the Bi₂O₃/TiO₂ photoanode, the p-n junction is formed by the Fermi energy level potential difference between the n-type semiconductor TiO₂ and the p-type semiconductor Bi₂O₃,¹¹² which generates a strong IEF. This p-n junction effectively inhibits the recombination of photogenerated carriers.³⁹ Bi₂O₃ prefers to be in contact with the middle -OH of glycerol, which facilitates the selective oxidation of the substrate. At the same time, DHA, whereas, is tough to be adsorbed on Bi₂O₃, avoiding further oxidation and improving selectivity. An analogous case is the Bi₂MoO₆@TiO₂ NTA photoanode and the C/Cu₂ONW photocathode, which are n-type and p-type semiconductors, respectively. The Fermi level of the Bi₂MoO₆@TiO₂ NTA photoanode is more negative than that of the C/Cu₂ONW photocathode, which causes an internal bias and drives the electrons of the photoanode to transfer and combine with the holes of the photocathode.⁹⁰ The tubular structure of the TiO₂ NTA can further enhance the refraction of light and improve light utilization. Native quantum dots (QDs) are very prone to recombination, and combining QDs with wide band gap acceptors into type-II heterojunctions can effectively extract charge and prevent recombination.¹¹³ In the TiO₂ NTA supported Mn-doped CdS QD photoanode, Mn doping increases the photoelectron lifetime of the composite photoanode and is more effective in enhancing the oxidation of the ·OH radical sacrificial agent, and there may be a correlation between both

of them.³⁷ The nominal doping of Mn at 15% has the longest photoelectron charge lifetime. Further doping results in the dominance of doping-induced defects and conduces the recombination of photoelectron charges.

Apart from the exploitation of UV light directly, a variety of methods to interact with TiO₂ to extend the light absorption of the catalytic system appear to be a more advisable way out for TiO₂-based catalysts. Its cost and performance advantages will keep it from being obsolete for a long time.

4.1.2. WO₃-based catalysts. WO₃ has the advantages of good light absorption properties, high stability and harmlessness under acidic and oxidative conditions, and the ability to modulate the morphology on demand.¹¹⁴⁻¹¹⁷ Nevertheless, the faster recombination gives rise to slower charge carrier transfer. The presence of sub-stoichiometric WO_{3-x} on the catalyst surface plays an instrumental role in the PEC performance in the oxidation of glycerol to glyceraldehyde and DHA by monoclinic WO₃. Oxygen vacancies are conducive to the formation of trap sites and facilitate the separation of photogenerated charges, but an appropriate quantity of oxygen defects is the clincher.¹¹⁸ There are four key points in this reaction system: (i) light irradiation as an initiator: serving to initiate the PEC glycerol oxidation; (ii) polarization by the external electric field as an accelerator to accelerate the charge separation; (iii) surface oxygen vacancies: serving as electron trapping sites and boosting the oxidation of water; and (iv) catalyst effect on the electro-oxidation of water: consuming ·OH radicals to prevent the indirect oxidation of glycerol.¹¹⁹ Upon this, glycerol mainly follows a direct oxidation path with photogenerated holes. WO₃ photoanodes also possess a remarkable tendency to oxidize acidic electrolyte anions,⁴² in some cases (HClO₄, H₂SO₄) even accompanied by the formation of peroxide species.^{120,121} The photo-oxidation products of acidic electrolyte anions, including the corresponding radical species, are considered intermediates of PEC reactions containing organics, especially when their concentration is low.¹²² It was found that ·HSO₄ radical species instead of S₂O₈²⁻ were formed for sulfates, and hypochlorite for chlorides when PEC glucose oxidation was performed with different electrolytes *via* nanostructured semi-transparent WO₃ photoanodes.¹²³ These substances improve the proton-to-current transition efficiency of the system. N-Doped catalysts exhibit enhanced current densities due to the reduced band gap and red-shift in the range of absorption energy bands.⁷¹ Structural changes are induced by N-doping in the WO₃ photoanode, hence the formation of IEFs,⁴⁹ specifically because an uneven charge distribution between two adjacent slices is imposed, creating an electrostatic potential difference. This asymmetric distribution of charge is capable of forming IEFs (Fig. 6a and b). The built-in electric field diminishes the trap depth and favors charge separation and transfer.¹²⁴ The WO₃/BiVO₄ complex can be adapted for hydrogen production from PEC of biomass. A Z-type heterojunction structure, which retains the CB (-0.1 eV) of BiVO₄ and the VB of WO₃ and enables the transformation of H⁺ into H₂ at 0 eV, is constituted.¹²⁵ BiVO₄-based catalysts also play an indispensable role in the biomass PEC process.

4.1.3. BiVO₄-based and perovskite catalysts. As mentioned above that BiVO₄ could form a strong complexation with tartaric acid, monoclinic BiVO₄ is regarded as a promising photoanode due to its low cost, narrow band gap, strong oxidation ability, and high theoretical maximum photocurrent (7.5 mA cm⁻²) and solar-to-hydrogen conversion efficiency (~9.1%) under 1 sun illumination.^{126–129} Strong adsorption of reaction intermediates and organic fuels on the electrode surface is a prerequisite for anodic reactions.¹³⁰ The deposited thin TiO₂ layer significantly reduced the PEC performance of the photoanode containing tartaric acid solution (Fig. 7a). Correspondingly, a BVO/TANF photoanode comprising BiVO₄ with phenolic ligands (tannic acid) synergistic with Ni and Fe ions (TANF) is also available. PEC glycerol oxidation in a 0.1 M Na₂SO₄/H₂SO₄ (pH = 2) electrolyte achieved more than 94% FE, where the selectivity of FA was more than 85%. The co-presence of Ni and Fe in the metal phenolic network for the achievement of a high current density is essential.⁵⁰ Unlike the mechanism of selective adsorption of glycerol middle-OH by Bi₂O₃ to oxidize to DHA, the selective primary-OH adsorption by BVO/TANF is dominant, but instead, the adsorption of secondary-OH was inferior. The presence of Ni in the metal phenolic network plays a principal role in the adsorption of glycerol primary-OH, and the addition of Fe²⁺ can promote the formation of a thickened network by more immobilized Ni²⁺ on BVO/TANF. The TEMPO-mediated NiFe-LDH/BiVO₄ photoanode also exhibits outstanding oxidation performance. The TEMPO mediator is highly selective for FDCA.⁶⁶ When the TEMPO concentration was increased to 5 mg mL⁻¹, the onset potential was significantly negatively shifted accompanied by a remarkable photocurrent enhancement, especially below 0.7 V vs. RHE. Under illumination, TEMPO was readily oxidized to TEMPO⁺ by the photogenerated holes on the VB of BiVO₄, and the TEMPO⁺ was recovered to TEMPO after reacting with HMF. As the content of NiFe salts was elevated, the morphology of NiFe-LDH was smaller and more uniform with a higher density.⁶⁶ The Mo-BVO-TiCoNO-CoFeO_{xy} photoanode produced by Wu *et al.*⁹⁷ through electrodeposition, drop-coating, and multiple thermal treatments has a nanowire structure where the length of the nanowire is multiplicatively correlated with the half-wavelength of the simulated sunlight, known as resonance, and this specialized effect is more receptive to solar radiation. This is the antenna effect of the non-quantum semiconductor structure, which is able to absorb more photons compared to other structures at the same volume.¹³¹ The TiCoNO layer is a protective layer for Mo-BVO because the photocurrent of Mo-BVO decays rapidly in strong bases due to the precipitation of V⁵⁺. The electrons in the non-d orbitals (s and p) supplied by Fe interact with Co to form Co-Fe. At the same time, the Fe species is the main adsorption site (*i.e.*, the catalytic site), and Co serves as a stabilizer to inhibit the precipitation of the Fe species in the strong bases. The CoFeO_{xy} layer has a better charge accumulation ability, and the Fe element acts as a charge provider in the charge redistribution process. A large amount of charge convergence in the CoFeO_{xy} layer will form a potential difference with the underlying elec-

trode, triggering a spontaneous charge flow and forming a built-in electric field.⁹⁷ Hence it has superior benzyl alcohol oxidation capability. β-O-4 can also be decomposed *via* the BiVO₄ photoanode under simulated sunlight and a 2.0 V bias. A 64% conversion was achieved after 20 h, and an elevated 10% conversion was obtained through the deposition of a V₂O₅ layer on BiVO₄ as well.⁸⁰ Although a bias of 2 V is applied, light is the key to achieving high activity. BiVO₄ acts as both a photoanode material and a heterogeneous catalyst for organic conversion in the system. V₂O₅ can only act as an electrocatalyst and has a synergistic effect with BiVO₄. The radius of the arc of the electrochemical impedance spectroscopy (EIS) Nyquist point with organics is normally much smaller than that without organics, proving its favorable electrical conductivity and interfacial charge transport.⁵⁸ In contrast, V₂O₅/BiVO₄ has a greater resistance, indicating that the presence of V does not increase the electrical conductivity. The reason for the higher conversion is that the surface roughness is increased with the participation of V. The corresponding perovskite PV cathode is deemed as a green option due to its excellent photovoltage output to drive the bias-free PEC reaction and the facile synthesis method that does not require an energy-intensive vacuum deposition process. The BiVO₄ potential can be further upgraded by PV through the remaining visible light transmitted through the photoanode in a tandem structure.⁴⁸ Meanwhile, the biocatalytic process at the cathode cannot happen in the absence of the perovskite PV, indicating the utmost significance of the solar absorber for bias-free PEC (Fig. 5c). There are also Pt-perovskite photocathodes which are composed of hybrid organic-inorganic halide perovskite interlayers between the electron and hole transport layers, together with the electrocatalyst CuPd alloy to reform polymeric waste. A selectivity of 60–90% yield and a product generation rate of about 70–130 μmol cm⁻² h⁻¹ were achieved, whose performance is 10²–10⁴ times more active than conventional photoreforming systems.⁵⁹ Cu possesses the best oxophilicity which assists in the adsorption of OH⁻, releasing active Pd sites for readily available substrate uptake.¹³² OH_{ad} further aids the oxidation process and removes the adsorbed intermediate species to reduce catalyst poisoning.^{132,133} The transport of the substrate to Pd sites is also limited by the high concentration of OH_{ad} species.¹³² The performance of many semiconductors in the cathode cell is incommensurate with that in the anode cell, but perovskites are always able to achieve excellent reduction performance due to their rich structural tunability. Therefore, perovskites are currently reported to be frequently deployed as high-value catalysts at the cathode.

4.1.4. α-Fe₂O₃- and Si-based catalysts. Catalytic systems coupled with Fe₂O₃ photoanodes and Si-based photocathodes tend to convert biomass without bias. Hematite (α-Fe₂O₃) has the advantages of a suitable band gap for absorption up to about 600 nm, low cost, non-toxicity, earth crust abundance, and high photochemical stability.¹³⁴ When glucose was added, the photocurrent was not much higher than that of PEC water, demonstrating the sluggish kinetics of glucose oxidation on the surface of unmodified α-Fe₂O₃. The NiOOH modification

had the best PEC performance in terms of the photocurrent density and starting potential, with a six-fold higher current density than that of the unmodified $\alpha\text{-Fe}_2\text{O}_3$.⁴³ Other catalysts such as $\alpha\text{-Fe}_2\text{O}_3$, $\text{FeOOH}/\alpha\text{-Fe}_2\text{O}_3$, and $\text{CoOOH}/\alpha\text{-Fe}_2\text{O}_3$ all exhibited poor selectivity (<50%) for the product FA, and only $\text{NiOOH}/\alpha\text{-Fe}_2\text{O}_3$ showed a unique performance. The coupled Si-based photocathode Bi/GaN/Si showed the lowest reported overpotential for achieving a high FE to FA (0 V), attributed to the coupling effect of strong absorption by p-n Si (1100 nm), efficient electron extraction and light trapping by GaN nanowires, and fast surface reaction kinetics of Bi photocatalysts (Fig. 5a). The system realized an average formate FE of up to 160%, where photoanodes and photocathodes accounted for 90% and 70%, respectively. The oxidation of lignin and the asymmetric hydrogenation of C–C bonds can be implemented in an additional unbiased PEC system constructed from hematite and Si-based electrodes. The electrode material is mesoporous indium tin oxide (*meso*ITO), which provides a more electrochemically active surface area than planar ITO, implying the ability to augment the rate of redox reactions.¹³⁵ The photoactivated Si PV is then able to supply an electron-driven more negative potential.⁷⁸ Both the photosynthetic components of the system (including the Rh-based electron mediator, *Thermus scotoductus*, and NAD^+) and the organics are imperative for C=C asymmetric hydrogenation.

4.1.5. Other catalysts. Some other catalysts with outstanding performance are everywhen applied in PEC systems. PMA has been relied upon for its unique characteristics of high acidity, reversible redox behavior at low potential, color change with redox reaction, and high solubility and stability under acidic conditions as a catalyst for the oxidative depolymerization of biomass¹³⁶ (Fig. 9a). Despite the low solubility or insolubility of many types of biomass in water, the color change from yellow to dark green was clearly observed in the PMA solution with biomass after overnight incubation at room temperature, indicating the reduction of PMA^{3-} to PMA^{5-} (ref. 87 and 88) (Fig. 9b). Ta_3N_5 is an n-type semiconductor with a narrow band gap of 2.1 eV that permits visible light absorption in the wavelength range up to 600 nm.^{137–139} But it has disadvantages such as weak carrier transport during water splitting, fast charge recombination, and high onset potential and overpotential.¹³⁹ The Ta_3N_5 catalyst synthesized by Wang has a typical sandwich structure: Ta_3N_5 nanotube film/ Ta_2N interlayer/ Ta substrate.⁵¹ The Ta_2N interlayer has high electrical conductivity, which allows for the transfer of Ta_3N_5 photoexcited charges to the Ta substrate.¹⁴⁰ In the meantime, Ta_2N behaves as a connecting bridge to ensure the bonding of Ta_3N_5 to the backing substrate, lowering the resistance to interfacial charge transfer. In response to the surface of Ta_3N_5 NTAs being susceptible to oxidative corrosion during PEC, the modification of two-dimensional trimetallic CoNiFe layer double hydroxide nanosheets (CoNiFe-LDH NSs) effectively inhibited oxidation by promoting the extraction and utilization of holes. The average lifetime of the photogenerated charges of the modified catalysts ($\tau = 1.16$ ns) was longer than that of the unmodified ones ($\tau = 0.64$ ns) as well. ZnO offers the advan-

tages of high adsorption properties, low cost, and excellent stability in alkaline media.¹⁴¹ Nonetheless, the drawbacks of a large band gap (3.2 eV) and low charge separation restricted its application. Since Ce^{4+} and Zn^{2+} have similar radii, it is possible to dope Ce without ruining the ZnO structure.¹⁴² What's more, through the chemical valence shift ($\text{Ce}^{4+} \rightarrow \text{Ce}^{3+}$), Ce doping generates a large number of oxygen vacancies for efficient charge separation. The modification of CeO_2 nanoclusters is contributive to the charge transport and adheres better to the substrate in the electrolyte than bare ZnO.¹⁴³

4.1.6. Prominent properties of diverse catalysts. Among semiconductor catalysts, it is particularly pertinent to select a suitable catalyst substrate depending on the cost, pH, band gap, and stability, apart from the fabrication of heterojunctions and structural modifications. Prominent semiconductor materials like TiO_2 , BiVO_4 , and perovskites are destined to be intensively investigated due to their respective non-negligible characteristics. TiO_2 has a simple preparation process and strong stability, while BiVO_4 has a low cost and relatively narrow band gap, and both are capable of forming strong complexation with conductive substances to realize catalyst-mediated photoelectric transformation. The multi-structural perovskites with tunable bandgaps provide an ultra-wide range of wave absorption and superb stability, which enables them to exert excellent activity as cathode materials. Besides, WO_3 and ZnO exhibit high stability in acidic and alkaline media, respectively, which coupled with suitable self-designed semiconductor photoelectrode materials makes specific conversion paths available.

4.2. The design of PEC configurations

Apart from the fabrication of highly active catalysts or complexes by designing vacancy defects, atomic doping, and elemental or compound modifications, the synergistic settlement of catalysts at both photoanodes can likewise fulfill specific conversion targets. In the $\text{NiOOH}/\alpha\text{-Fe}_2\text{O}_3\text{-Si}/\text{mesoITO}$ complex system, the long wavelength photons (>600 nm) passing through the front $\alpha\text{-Fe}_2\text{O}_3$ photoanode are available to the rear Si-based photocathode, which allows the hybrid apparatus to encompass most of the spectrum⁴³ (Fig. 5a). There is also a Pt-perovskite photocathode with an electrocatalyst CuPd alloy for the PEC system. This system is a highlight in photocatalysis as it applies dark mixed waste solution directly on the anode side without blocking the light in the cathode cell. A similar system is the three-compartment photo-electro-biochemical system (Fig. 5b). In order to avoid the underperformance from the dark lignin in conjunction with photocatalysis, the lignin is arranged in the biocatalytic cell for transformation. Since the lignin and photocatalyst are separated, the light absorption efficiency of the dark lignin is not decreased.⁴⁷ To resolve the contact problem between the reactants and the catalyst, cellulose films are deposited directly on the TiO_2 photoanode. The original porous and particulate TiO_2 surface is completely renewed to a uniform and smooth surface after cellulose support. The PEC reaction takes place near the interface of TiO_2 and cellulose.¹¹ Under solar illumina-

nation, nano-Fe₂O₃ can drive both water and glucose oxidation. Yet the FE of oxidation of glucose is only 60.8 ± 1.5% at 1.0 V vs. RHE and the selectivity of FA is not satisfactory. *m*-CuO could not only act as an electrocatalyst, but the p–n heterojunction composed of *m*-CuO and nanoFe₂O₃ could modulate the oxidation of photoinduced holes. The valence band edge of *m*-CuO is similar to its flat-band potential (1.15 eV), so the photogenerated holes of this heterojunction can only oxidize glucose. As shown in Fig. 6, photoelectric water oxidation is almost eliminated by the passivation of *m*-CuO. Moreover, the photocurrent of nanoFe₂O₃|*m*-CuO containing glucose at 1.23 V vs. RHE is 15 times higher than that of glucose-free nanoFe₂O₃|*m*-CuO.³⁶

With the aim of minimizing the bias, it is common to employ n-type and p-type semiconductors for the anode and cathode, respectively, apart from preparing catalysts with a high light absorption range and high activity, or by means of bio-catalysis.⁹⁰ Compensation for the potential difference can also be implemented by the series structure of the photoanode and PV. In the PEC lignin oxidation system with BiVO₄ and cathodic biocatalytic hydrogenation, there are three major components: (i) the BiVO₄ photoanode for PEC lignin oxidation; (ii) the tandem structure of the tri-cation perovskite PV with BiVO₄ and providing supplementary photovoltage to drive the redox reaction at 0 bias; and (iii) the carbon-cloth cathode to reduce NAD⁺ to NADH for enzyme redox transformation.⁴⁸ The tandem structure compensates for the potential difference through Z-schematic adsorption, resulting in a significant decrease in the molecular weight of lignin (Fig. 5c). Luo *et al.*³⁹ established a self-powered PEC system based on a two-electrode setup, consisting of a PV plate (~2 V) that provides a constant bias and a homemade PEC flow cell comprising a Bi₂O₃/TiO₂ photoanode with a 7 cm² irradiated area and a Pt wire cathode. A DHA yield of 1.04 mg cm⁻² h⁻¹ (selectivity >70%) and an H₂ yield of 0.32 mL cm⁻² h⁻¹ were obtained for this system.

The semiconductor materials with suitable band gaps could be prepared by modulating the catalyst structure with abundant elements and designing suitable PEC systems. The techniques combining the advantages of electrocatalysis or bio-catalysis could ultimately realize low-bias set-up and high-value conversion. This is the main feature of the current PEC area but it still holds infinite research potential. The flexibility of photoelectric conversion, which allows simultaneous valorized conversion at both anode and cathode sides or even in multiple compartments, is its greatest advantage. During the implementation of the system, some components or variables can also have an impact on the final catalytic performance.

5. The effect of related variables on PEC systems

There is no doubt that a suitable preparation of the photocatalyst and a smart configuration can directly determine the specific PEC conversion. At the same time, some factors such as solution pH, pretreatment, polarization, membrane, and

substrate concentration can affect the optimal operating conditions or the selectivity of the target product. A summary of some representative works is presented here.

(i) Solution pH: The nanocrystalline Bi₂WO₆ photoanode is capable of converting glycerol into FA at different pH values, but the performance of the transformation varies greatly. In acidic media, the selectivity increases to 88% but the photocurrent decreases to 50%. In alkaline media, glycerol oxidation is more convenient and the photocurrent increases to 0.8 mA cm⁻², but the selectivity of the product decreases.¹³ The reason for this is that at lower pH, there is stronger adsorption of glycerol to the photoanode, making the reaction from photogenerated holes to further oxidation easier, and thereby more selective for formic acid. Benzyl alcohol can be oxidized to benzoic acid in strong bases without organic solvents, while it is only oxidized to benzaldehyde in neutral organic-free solvents or just in organic solvent systems.⁹⁷ Except that, a huge pH difference at both sides of the membrane even leads to the leakage of polluting ions.⁷⁵

(ii) Pretreatment: For polymers, the structural properties of the substrates vary substantially depending on the processing methods in the industry. In the system of PEC microalgal biomass cleavage for lipid extraction, PEC pretreated *Chlorella* achieved a lipid extraction efficiency of 96%, roughly 10 times higher than that of non-pretreated *Chlorella*.⁷¹ The pretreatment step of biomass in the PEC biomass depolymerization system plays a key role in FA production. The FA yields of untreated samples were one to two orders of magnitude lower than those of pretreated ones.⁴³ The PEC conversion of PET powder, real-life commercial PET bottles, and microcrystalline cellulose by the CuPd alloy was likewise found to achieve even negligible selectivity of glycolic acid and gluconic acids from non-pretreated feedstocks.⁵⁹ The oxidation product of β-O-4 by PMA is mainly vanillin, while those of lignocellulose are mainly vanillin and acetovanillone. The differences in the outputs may be attributed to distinctions in the molecular structure of the various raw materials and diverse chemical treatment methods.⁸⁸ The tandem structure of BiVO₄ with a perovskite is able to degrade the molecular weight of alkali lignin from 51 000 *M_w* to 15 000 *M_w*, and the molecule keeps its aromatic ring intact. In contrast, the aromatic rings are converted into carbonyl or carboxyl groups from the decomposition of lignosulfonate. The properties of lignin oxidation can be differentiated from various sources and pretreatment methods.⁴⁸

(iii) Polarization: The production of C2 and C1 chemicals was retarded and the production of C3 was simulated under a sufficiently high potential (1.5 V vs. Ag/AgCl) in the PEC glycerol oxidation system by monoclinic WO₃, as the polarized oxygen evolution reaction consumes ·OH, further avoiding the excessive oxidation of ·OH.¹¹⁹ In the system with Kraft lignin as a reactant, the *J*-*V* curve from the voltage scanning from -1 to 3 V laid above that from 3 to -1 V. This hysteresis is attributed to the low ion transfer rate and the long distance between the electrodes, bringing about polarization near the electrode surface.⁷⁵

(iv) Membranes: For a given PEC system, membranes are capable of selectively permeating target ions or components. Different types of membranes have different functions and should be treated precisely. In a three-compartment photo-electro-biochemical system, Nafion and cellulose membranes are used to separate the anode cell, the cathode cell, and the biocatalytic cell, whereby the bioenzyme system is protected from reactive oxygen species ($\cdot\text{OH}$ and $\cdot\text{O}^{2-}$), shear stress from oxygen bubbles, high concentrations of hydrogen peroxide, and UV radiation⁴⁷ (Fig. 5b).

(v) Substrate concentration: Adequate contact between the reactants and the catalyst surface is one of the key factors for efficient PEC conversion. A moderate range of substrate concentrations ensure the uninterrupted contact between the catalytically active site and organic molecules, thereby maintaining the reaction rate and current density. Instead, a too high substrate concentration would affect the adsorption on the electrode surface due to diffusion limitation or agglomeration, and eventually, the current density would drop abruptly.

6. Conclusion and prospect

To be frank, the energy utilization efficiency of biomass electrocatalysis is high, and many of these systems could elevate the FE of organic substrates to about 100%. Inevitably, the selectivity of products in some fine chemistry areas is cut down because certain sensitive or highly active reaction processes are too vigorous. The applied potential itself that sustains electrocatalysis is a form of energy that needs to be injected into the system at the beginning of the reaction. Photocatalysis takes advantage of specific wavelengths of light or sunlight to perform selective conversion of substrates with the benefit of semiconductor catalysts. The PEC system is capable of transferring electrons to the cathode through wires to avoid charge recombination and to realize valuable applications with smooth parallel oxidation and reduction reactions. Biomass, the world's largest renewable resource, has the merits of degradation at a lower bias than water. By preparing suitable semiconductor catalysts and setting up a subtle photoelectrode device and a system model, biomass can be sacrificed for a range of valorizations.

Currently, however, most studies have concentrated on the conversion of glucose, glycerol, and β -O-4. These substrates are derived compounds that best represent the structure of the biomass. Upon attack by h^+ , $\cdot\text{OH}$, or other reactive species, the substrates are consumed or decomposed into small molecule products. The generation of glucaric acid and the $\text{C}_{\text{Ar}}\text{-C}_{\alpha}$ bond cleavage of β -O-4 exemplify the extremely strong oriented transformation ability of the PEC system. Mechanistic investigation of the conversion and exploration of energy formats for more types of model compounds are not clear yet, which is detrimental to the adequate utilization of biomass components. The combination of PEC valorization pathways in multiple realms is also not merely circumscribed to electrocatalysis and biocatalysis. The flexibility of PEC systems leads to more possi-

bilities for valorization of biomass. More and more studies are gradually looking at electrocatalysis and photocatalysis for inspiration and attempting breakthroughs as well as innovations in PEC. Beyond improving the selectivity of the aimed products, the modulation of semiconductors with different band gaps alters the redox intensity, while the arrangement of configurations concerning cathodes and anodes reduces the bias and achieves multi-path conversion. It is noteworthy that there is a lack of reductive valorization pathways for biomass substrates at the cathode and photoreduction of biomass will make PEC more universal like other catalytic approaches.

The development of PEC, similar to pure photocatalysis, is still at a theoretical level. The low optical transmittance and insolubility of polymeric raw materials are the main challenges. The upgrading of PEC needs to go through a procedure like structural clarification of biomass feedstock, efficiency improvement of energy harnessing, scale-up of reaction apparatus, and cost economization of catalysts and systems. These problems are being addressed step by step with the cooperation of multidisciplinary scientific research.

Conflicts of interest

There are no conflicts to declare.

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