Valorization of Biomass-Derived Platform Molecules via Photoredox Sustainable Catalysis

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Abstract

The conversion of biomass into valuable chemicals has promise for application in biorefineries. Light-driven photoredox catalysis, with the typical features of green route and operation under mild conditions, is considered a promising strategy for renewable biomass or biomass-derived intermediates conversion into high-value-added chemical feedstocks. In this review, we strongly emphasize the recent advances in photocatalytic valorization of lignin model compounds and biomass-derived alcohols. We briefly summarize the advances in photocatalytic cleavage of the $\beta$-O-4 bond or C–C bond into usable chemicals in the lignin model. On the other hand, we clarify not only the hybrid system for cooperative biomass-relevant alcohols oxidation and hydrogen ($H_2$) evolution but also the tunable accessibility to variation of the target products from the same alcohol reactant by catalyst design and optimization of reaction conditions. It is hoped that this review will inspire the rational design of photoredox catalysis-based systems toward efficient biomass-derived platform molecules valorization to obtain target-oriented valuable products.

Keywords  Biomass · Photoredox catalysis · Value-added chemicals · Selective oxidation

Introduction

Exploring the alternative and sustainable feedstocks has recently attracted substantial attention because of the shortage of fossil resources [1]. Biomass, which is available in large amounts and includes all animals, plants, and microorganisms, is an important part of renewable energy [2]. Thus, the efficient development and use of biomass will play a very vital role in resolving energy and ecological issues. Lignocellulose is a typical representative of biomass consisting of lignin, hemicellulose, and cellulose. Among these substances, lignin, which accounts for 30% of the total mass [3], is the only relevant renewable feedstock for aromatic compounds [2]. Although the traditional strategies for the transformation and fragmentation of lignin have made progress, the high energy consumption and harsh reaction conditions are always hazardous to the environment. In contrast, the bio-enzymatic degradation of lignin can occur under mild conditions, but the higher cost of enzyme increases the overall cost of the process [4]. Within this context, photocatalytic valorization of biomass has received substantial attention because of its low energy consumption and secondary-pollution-free operation [5–9].

Noticeably, the photoredox-catalyzed conversion of biomass-derived alcohols (such as 5-hydroxymethylfurfural, furfural alcohol and benzyl alcohol) has also attracted considerable attention in recent years (Fig. 1), because the produced carbonyl or C–C coupling compounds are important intermediates for the synthesis of high value-added fine chemicals [10, 11]. In particular, the visible-light-driven cooperative biomass-relevant alcohols oxidation and hydrogen ($H_2$) evolution can occur in one photoredox catalysis system, and in this scenario the photogenerated electrons and holes are simultaneously used to acquire upgraded chemicals as well as clean fuels [12–15]. Additionally, photocatalytic selective conversion of alcohols can be tuned to achieve different product distributions (e.g., aldehydes, ketones or C–C coupling compounds) [16, 17], which further promotes the flexibility of target-oriented product synthesis from biomass-derived alcohols.
In this review, we intend to concisely overview recent developments in the solar energy-driven photoredox-catalyzed transformation of biomass-derived compounds into useful chemical feedstocks, which includes selective fragmentation of lignin, biomass-derived alcohols oxidation, and dual-function selective alcohols conversion coupling with H₂ production in a cooperative manner. We also clarify the adjustable flexibility of switching the different target-oriented product distributions from the same alcohol substrate. Furthermore, the brief descriptions for the possible mechanism toward selective lignin model alcohols transformation or biomass-derived alcohols conversion are elaborated. Toward the end, the potential opportunities for future research toward biomass-derived platform molecules valorization are discussed, which we hope will contribute to the development of rational design of efficient photocatalysts for target solar-to-chemical transformations of biomass-derived platform molecules to target value-added feedstocks.

**Photocatalytic Conversion of Lignin Models**

Lignin, as a complex alcohol-containing molecule originating from lignocellulosic biomass, is considered the only relevant renewable feedstock for aromatic compounds [2]. The interlinkages of lignin primarily contain C–O bonds and C–C bonds, wherein the β-O-4 bond is greatest in content (Fig. 2a) [18, 19], and the β-O-4 bond can be broken down into chemically useful fragments [20]. Likewise, rupturing the C–C bond can yield valuable compounds, but cleaving the C–C bond of lignin is more difficult because of its relatively high bond dissociation energy (BDE) [21, 22]. Owing to the recalcitrance and complexity of raw lignin, many studies have used representative lignin model compounds to replace lignin macromolecules to explore the potential of lignin valorization into high-value chemicals. Figure 2b depicts a typical lignin model that can be converted into several low-molecular-mass aromatic compounds, which can be further used to synthesize other value-added chemicals. Although lignin model compounds minimize the challenges of lignin fragmentation, certain mild conditions still cannot meet the requirements of their efficient transformation. In this regard, semiconductor-based photoredox catalysis has emerged as an accessible approach to organic synthesis [23, 24], e.g., selective alcohol oxidation [25, 26] and nitroaromatic reduction reactions [27, 28]. To this end, solar energy-driven fragmentation of lignin models into useful aromatics has received considerable attention.
**β-O-4 Bond Cleavage**

The pivotal problem of lignin fragmentation is the scission of β-O-4 bonds. Catalytic cleavage of the β-O-4 linkage in lignin evidently undergoes a two-step continuous process (Fig. 3) [29]. In the first step, an oxidation catalyst is used to oxidize CαH–OH groups into Cα=O. Afterward, the oxidation catalyst is separated out, and then the Cβ–O bond is ruptured into aromatic monomers by addition of a hydrogenation catalyst. However, this method uses two catalysts that work independently under oxidizing and reducing conditions. To predigest the reaction steps and boost yields, a photocatalytic oxidation-hydrogenolysis tandem system for rupturing the β-O-4 bonds of the lignin model via a dual light wavelength switching strategy has been designed by Luo et al. [30], in which the α-C–OH moiety of the lignin model is oxidized to α-C=O by Pd/ZnIn2S4 under 455 nm wavelength light, and then the titanium dioxide-sodium acetate (TiO2-NaOAc) system is applied in rupturing the C–O bond neighboring the α-C=O bond by switching to 365 nm wavelength light (Fig. 4a). As a result, the C–O bonds are cleaved to yield phenols and acetophenone with high selectivity under aerobic conditions. Additionally, in the TiO2-NaOAc system, the (101) crystal facet of TiO2 is the most active crystal facet and the addition of alkali NaOAc enhances the catalytic activity with ethanol as the external hydrogen source. Control experiments suggest that photogenerated electrons play a unique role in this system.

![Fig. 3](image1.png) Calculated BDE of the two-step conversion strategy of the β-O-4 linkage in the lignin model. Reprinted with permission from Ref. [29]. Copyright 2018 American Chemical Society.

![Fig. 4](image2.png) a Illustration of the photocatalytic cleavage of β-O-4 alcohols into phenol and acetophenone by oxidation-hydrogenolysis via the dual-wavelength switching strategy. b Possible reaction mechanism for photocatalytic C–O bond cleavage of β-O-4 ketones over TiO2. Reprinted with permission from Ref. [30]. Copyright 2016 American Chemical Society. e The gas phase fragmentation of PP-ol and PP-ol-αCD detected using mass spectrometry. f Schematic illustration of photocatalytic cleavage of the β-O-4 bond via self-hydrogen transfer hydrogenolysis over ZnIn2S4. Reprinted with permission from Ref. [31]. Copyright 2017 American Chemical Society.
The electrons reduce Ti$^{4+}$ on the surface of TiO$_2$ to Ti$^{3+}$ instead of directly involving in cleaving the C–O bond, and then the Ti$^{3+}$ formed in situ adsorbs the substrate for weakening the C–O bond to directly participate in cleaving the C–O bond (Fig. 4b). This strategy not only integrates the oxidation-hydrogenation process in one pot by dual-wavelength variation but also obtains the aromatic monomers with superior yield.

Although this two-wavelength strategy achieves a one-pot process, two catalysts are still required, and an external hydrogen donor is necessary. When alcohol is used as the hydrogen donor, the alcohol dehydrogenation product (aldehyde) may condense with phenol, making the product separation more complicated. To overcome this obstacle, Luo and coworkers [31] have extracted phenolic products from the β-O-4 lignin model phenoxy-1-phenylethanol (PP-ol) over ZnIn$_2$S$_4$ to further integrate the dehydrogenation and hydrogenolysis reaction in one system, in which the alcoholic groups (CαH–OH) of PP-ol serve as hydrogen donors to form a “hydrogen pool” on ZnIn$_2$S$_4$ to produce phenolic compounds by self-hydrogen transfer. To ensure a pure self-hydrogen transfer mechanism, solvents without ability of hydrogen-donating (acetonitrile, acetone and tetrahydrofuran) are added to the reaction, obtaining a similar conversion (97–99%) and moderate yield of cleaving β-O-4 bond products (phenol and acetophenone), which suggests that the conversion of the lignin model into aromatic chemicals indeed occurs through self-hydrogen transfer. Additionally, the detections of H$_2$ and H$_2$S in the gas phase (Fig. 4c), which might originate from the hydrogen species of PP-ol, further prove the occurrence of self-hydrogen transfer hydrogenolysis. H$_2$ and HD are detected by adding deuterated PP-ol (PP-ol-$\alpha$CD) to the photocatalysis reaction (Fig. 4d), revealing that the H-abstraction of the Cα-H–OH groups in PP-ol occurs on the surface of ZnIn$_2$S$_4$. Based on the above results, a proposed mechanism for the fragmentation of the β-O-4 lignin model is outlined in Fig. 4e. PP-ol undertakes H-abstraction of α–OH groups via photogenerated holes, resulting in the formation of 1f radicals. Thereafter, H-abstraction of α–CH groups leads to a yield of the β-O-4 ketone intermediate. At the same time, the proton hydrogen adsorbed on the surface of ZnIn$_2$S$_4$ reacts with photogenerated electrons to form a “hydrogen pool.” When the β-O-4 ketone intermediate enters the “hydrogen pool”, it generates the products of C–O bond cleavage (phenol and acetophenone), while a small portion of the reduced hydrogen species combines to produce H$_2$ accompanied by the formation of byproduct. This study opens a doorway to exploit renewable biofeedstock as well as solar energy to acquire phenolic compounds.

To efficiently use all three components of lignocellulose, Wu et al. [32] achieved lignin-first fractionation from birch woodmeal while retaining the hemicellulose and cellulose. The results imply that the native lignin can be effectively converted into functionalized aromatic compounds using soluble colloidal CdS quantum dots (QDs) under mild conditions. Conversions of the solid residue, including cellulose and hemicellulose, are conducted after the separation of colloidalized CdS QDs. As disclosed in Fig. 5a, b, acidolysis is initially performed for the solid residue to obtain xylose under mild conditions (fraction I). Afterward, the remaining solid undergoes enzymatic hydrolysis to yield glucose (fraction II), demonstrating that the lignin-first concept for use of biomass can be integrated with the current biorefinery system without wrecking carbohydrate fractions. In addition, the mechanism for the fragmentation of the β-O-4 lignin model is different from those of the previous two-step photocatalytic systems. The control experiments using either holes or electrons scavenger exhibit significantly decreased yields of aromatic fragments, confirming that photoexcited holes and electrons participate in the generation of acetophenone and phenol. The reaction is almost inhibited by adding 5,5-dimethyl-1-pyrroline-N-oxide (DMPO), indicating that the reaction proceeds via a free radical intermediate. Furthermore, when the Cα hydrogen of benzyl group is replaced by a methyl group, the reaction barely occurs, suggesting that the presence of the Cα hydrogen is critical for the reaction. Moreover, the generation of Cα radicals has been confirmed via the spin trapping agent 5-diisopropoxypyrophosphoryl-5-methyl-1-pyrroline-N-oxide (DIPPMPO). Typically, free radicals can react with DIPPMPO to produce a nitroxide intermediate that undergoes disproportionation to form a correspondingly stable nitroxide and hydroxylamine as the ultimate products (Fig. 5c) [33]. Positive ion electrospray ionization mass spectrometry of the product mixture after adding DIPPMPO clearly proves the formation of Cα radicals (Fig. 5d) [34]. As depicted in Fig. 5e, a main signal at 28.7 ppm, which is index to Cα radical adducts [33], appears in the $^{31}$P nuclear magnetic resonance (NMR) spectrum, further demonstrating that the Cα radical is essential for this system. Based on the above analysis, an electron–hole coupled (EHC) mechanism is illustrated in Fig. 5f, in which the oxidative dehydrogenation (ODH) of the Cα–H bond triggered by holes first occurs to produce the Cα radical intermediate, and then this intermediate undertakes the cleavage of the β-O-4 bond driven by the electron.

**C–C Bond Cleavage**

Much effort has been devoted to cracking the C–O bonds in lignin into valuable aromatic products, but selective C–C bond cleavage in macromolecular biomass under mild conditions remains a challenge. When a C–C bond is a targeted linkage to be fragmented, Cα–H bond activation is necessary [35]. However, once the C–OH bond is transferred to a ketone by oxy-dehydrogenation, the BDE of the Cα–Cβ bond
bond increases from 294.7 to 354.2 kJ/mol, which makes the production of ketone more difficult (Fig. 6a). To minimize this dilemma, a hybrid CuO/Ceria/anatase nanotube (CuO/CeO2/A-NTs) has been synthesized using a deposition–precipitation method and applied to photocatalytic selective oxidation of C–C bonds of the β–1 lignin model into benzaldehydes [36], in which bifunctional CuO not only improves the catalytic activity but also suppresses the side reaction. Previous studies have shown that the photoactivity of CeO2 is related to the oxygen vacancy and Ce3+ concentration on the surface [37]. Thus, Raman spectroscopy is conducted to semi-quantify the concentration of surface oxygen vacancies [38–40]. As a result, the introduction of CuO clusters can enhance the concentration of oxygen vacancies, and the generated oxygen vacancies simultaneously reduce Ce4+ to Ce3+, thus showing a higher activity for the formation of benzaldehyde by the C–C bond cleavage of the β–1 lignin model. As depicted in Fig. 6b, the optical band energies of CeO2/A-NTs (2.72 eV) and CuO/CeO2/A-NTs (2.62 eV) are lower than that of blank CeO2 (2.86 eV), indicating that the presence of Ce3+ can narrow the band gap of CeO2. Furthermore, theoretical calculations demonstrate that the introduction of CuO clusters moves the valence band edge of titanium dioxide to a higher energy position, thereby inhibiting the side reaction of the oxygen dehydrogenation to enhance the selectivity of benzaldehydes.

Selective C–C bond cleavage in lignin affords useful and functional fine aromatic chemicals [41], and the interaction between the catalyst surface and lignin substrate also has a great effect on the conversion of lignin. Liu et al. [42] have used two-dimensional flexible graphite phase carbon nitride (g-C3N4) to provide an ideal surface for the adsorption of lignin models. The g-C3N4 prepared from melamine (C3N4-M) and urea (C3N4-U) is synthesized to investigate the effect on photoactivity with different precursors. Herein, C3N4-U exhibits higher C–C cleavage activity than C3N4-M, which is due to a larger surface area and the faster carrier transport of C3N4-U [43]. When the mesoporous graphitic
carbon nitride (mpg-C₃N₄) is used, the lignin model substrate is nearly completely transformed. The solid-state NMR techniques and density functional theory (DFT) calculations jointly verify π–π stacking interactions between the mpg-C₃N₄ and lignin models. As shown in Fig. 6c, the off-diagonal peak pair appears at (6.8, 15.9) ppm in the two-dimensional hydrogen double quantum magic angle spinning nuclear magnetic resonance (2D ¹H–¹H DQ MAS NMR) spectrum, indicating the spatial proximity of the hydrogen on the benzene ring of the substrate to the NHₓ group in mpg-C₃N₄ [44], which facilitates the transfer of photogenerated holes and promotes the photocatalytic reactions. DFT establishes a corrugated CₓNₓ surface with an adsorption energy of −0.92 eV and is parallel to two benzene rings of the substrate molecule (Fig. 6d). The distance between triazine rings and benzene rings is 3.2–3.9 Å, corresponding to the range of π–π stacking interactions, which contributes to the molecular activation [45, 46] and further promotes the conversion of the lignin model molecule.

In addition to semiconductor-based catalysts, quinones, as organic oxidants with excellent oxidation ability, are also used to promote the depolymerization of lignin. Mitchell and coworkers [47] have evaluated the photocatalytic C–C bond scission of lignin models using the simpler and cheaper 1,4-benzoquinone (BQ) with copper nanoparticles as the electron transfer mediator (ETM) under oxygen atmosphere, in which BQ is formed by in situ oxidation of 1,4-hydroquinone (BQH₂), and oxygen is used as a terminal oxidant. The cleavage of the C–C bond might occur by H-abstraction from the O–H bond by the photoexcited triplet quinone (³BQ) to form an alkoxy radical intermediate, and then this alkoxy radical is subjected to C–C homolytic fragmentation to form a new alkyl radical and aldehyde (Fig. 7). Finally, two carbonyl compounds are obtained by C–C cleavage reaction via a radical process. In addition, this methodology has been expanded to a range of β-O-4 lignin model compounds by effectively using visible light, leading to the production of a series of aromatic products.

**Photocatalytic Conversion of Biomass-Relevant Alcohols**

In addition to lignocellulosic biomass, photocatalytic transformation of biomass-relevant alcohols is highly appealing for obtaining usable chemicals [48–50], which are pivotal...
intermediates for the manufacture of pharmaceuticals, confectionery fragrances and biopolymers [10, 51]. Hence, we concisely summarize the recent overview of selective photocatalytic conversion of biomass-relevant alcohols in this section, including photooxidation of biomass-relevant alcohols, visible-light-induced hydrogenation of biomass-derived 5-hydroxymethylfurfural (HMF) and photoredox-catalyzed conversion of biomass-relevant alcohols integrated with hydrogen (H\textsubscript{2}) evolution.

**Mechanisms for the Photocatalytic Conversion of Biomass-Relevant Alcohols**

Photocatalytic conversion of biomass-relevant alcohols can be performed under aerobic or anaerobic conditions. In aerobic atmosphere (Fig. 8a), oxygen molecules are reduced to reactive oxygen species, such as the superoxide radical (\textsuperscript{•}\textsubscript{O}_2\textsuperscript{−}) and singlet molecular oxygen (\textsuperscript{1}\textsubscript{O}_2), by the photoexcited electrons in the conduction band (CB). These species subsequently participate in the transformation of biomass derivatives [48]. Meanwhile, holes accumulated in the valence band (VB) can oxidize biomass-derived alcohols to the corresponding carbonyl compound. By contrast, under anaerobic conditions (Fig. 8b), the photogenerated electrons are available for reducing protons to produce H\textsubscript{2}, while biomass-relevant alcohols can be oxidized by photoinduced holes, realizing the simultaneous use of electrons and holes.

### Photocatalytic Conversion of HMF

#### Selective Oxidation of HMF

A summary of the recent advances in selective photoredox-catalyzed biomass-relevant alcohols conversion is outlined in Table 1. HMF, as one of the representatives of biomass intermediates, can be oxidized to many species containing polymeric precursors, such as 2,5-furandicarboxylic acid (FDCA) and 2,5-diformylfuran (DFF) [52]. FDCA plays a key role in commerce, because it is the alternative to terephthalic acid in the production of polyesters, polyurethanes and polyamides [11]. For instance, Xu et al. [53] have constructed a photocatalyst of cobalt thioporphyrazine (CoPz) bonded on g-C\textsubscript{3}N\textsubscript{4} (CoPz/g-C\textsubscript{3}N\textsubscript{4}), which possesses a highly efficient photocatalytic performance for selective transformation of HMF into FDCA under aerobic conditions. Because of the powerful interaction between g-C\textsubscript{3}N\textsubscript{4} and CoPz, an optimal FDCA yield of 96.1\% is achieved after 22 h of irradiation in basic conditions. Although a high HMF (99.6\%) conversion is obtained on bare g-C\textsubscript{3}N\textsubscript{4} catalyst under the same conditions, no liquid products are detected, indicating that HMF is likely to mineralize into H\textsubscript{2}O and CO\textsubscript{2} on bare g-C\textsubscript{3}N\textsubscript{4}. Furthermore, the electron paramagnetic resonance (EPR) technique using 2,2,6,6-tetramethylpiperidine (TMP) and 5,5-dimethyl-1-pyrroline-N-oxide (DMPO) as spin-trapping regents is used to monitor oxidative species in the reaction. As disclosed in Fig. 9a, remarkable EPR signals for \textsuperscript{1}\textsubscript{O}_2-TMP adducts are observed over CoPz/g-C\textsubscript{3}N\textsubscript{4} irradiated by a pulsed laser and are assigned to the nitroxide produced by the reaction of TMP and \textsuperscript{1}\textsubscript{O}_2 [54]. However, a characteristic quartet signal of DMPO-hydroxyl radical (\textsuperscript{•}\textsubscript{HO}) adducts is detected over bare g-C\textsubscript{3}N\textsubscript{4} by adding DMPO to the reaction [55]. The above results further show that, compared with bare g-C\textsubscript{3}N\textsubscript{4}, CoPz/g-C\textsubscript{3}N\textsubscript{4} has a suitable oxidation capacity to selectively oxidize HMF into FDCA instead of undertaking photocatalytic mineralization.

![Fig. 7 Proposed mechanistic pathways of C–C bond cleavage in the lignin model via BQH\textsubscript{2}. Reprinted with permission from Ref. [47]. Copyright 2014 American Chemical Society](image)

![Fig. 8 Illustration of mechanisms for the photocatalytic conversion of biomass-relevant alcohols under a aerobic and b anaerobic conditions](image)
Table 1 An overview of the recent literature for biomass-relevant alcohols conversion

| Entry | Photocatalysts       | Substrate | Solvent | Atmosphere | Product       | Light source                      | Ref.     |
|-------|----------------------|-----------|---------|------------|---------------|-----------------------------------|----------|
| 1     | Nb$_2$O$_5$-800      | HMF       | BTF     | O$_2$      | DFF           | 300 W Xe lamp, > 400 nm           | [58]     |
| 2     | Pt/SGCN              | HMF       | H$_2$O  | N$_2$      | DFF and H$_2$ | LED lamp, > 400 nm               | [62]     |
| 3     | g-C$_3$N$_4$         | HMF       | H$_2$O  | N$_2$      | DFF and FDCA  | 15 W FL lamp, 340–420 nm         | [10]     |
| 4     | Ni/CdS               | HMF or FA | H$_2$O  | N$_2$      | DFF or Furfural, H$_2$ | Blue LED lamp, 450 nm | [13]     |
| 5     | Au-Ru/rGO           | HMF       | Toluene | O$_2$      | DFF           | 300 W Xe lamp                     | [57]     |
| 6     | Zn$_x$Cd$_{1-x}$-S-P | HMF       | H$_2$O  | Ar         | DFF and H$_2$ | White LED lamp                    | [84]     |
| 7     | Pt/g-C$_3$N$_4$      | HMF       | H$_2$O  | Air        | DFF           | 210 W Xe lamp, ≥ 420 nm          | [64]     |
| 8     | Pd/MIL-101(Fe)-NH$_2$| HMF       | CH$_3$CN| Air        | DFF           | 300 W Xe lamp, ≥ 420 nm          | [67]     |
| 9     | Ti$_3$C$_2$T$_x$/CdS | FA        | H$_2$O  | N$_2$      | Furfural and H$_2$ | 300 W Xe lamp, > 420 nm | [68]     |
| 10    | Co/CdS               | BA        | CH$_3$CN| N$_2$      | BAD and H$_2$ | 300 W Xe lamp, ≥ 420 nm          | [14]     |
| 11    | Ni$_x$S$_y$/ZnS      | BA        | H$_2$O  | N$_2$      | BAD and H$_2$ | 500 W Xe lamp, > 200 nm          | [85]     |
| 12    | Ni$_x$S$_y$/ZnS      | BA        | H$_2$O  | N$_2$      | BAD and H$_2$ | 300 W Xe lamp, ≥ 420 nm          | [86]     |
| 13    | CoP/CdS              | BA        | CH$_3$CN| N$_2$      | BAD and H$_2$ | 300 W Xe lamp, ≥ 420 nm          | [15]     |
| 14    | Carbon nitride aerogel| BA        | CH$_3$CN| O$_2$      | BAD           | 300 W Xe lamp, ≥ 420 nm          | [87]     |

HMF 5-hydroxymethylfurfural; BTF benzotrifluoride; DFF 2,5-diformylfuran; SGCN porous carbon nitride; FDCA 2,5-furandicarboxylic acid; FL fluorescent; FA furfural alcohol; rGO reduced graphene oxides; DHMF 2,5-dihydroxymethylfuran; BA benzyl alcohol; BAD benzyaldehyde

![Fig. 9](image_url)  
**Fig. 9** a EPR signals of (a) the DMPO-HO · adduct of bare g-C$_3$N$_4$ in water and (b) the TMP-1O$_2$ adduct of CoPz/g-C$_3$N$_4$ in water under light irradiation. b Possible mechanism for photocatalytic oxidation of HMF into FDCA with the CoPz/g-C$_3$N$_4$ catalyst. Reprinted with permission from Ref. [53]. Copyright 2017 American Chemical Society. Illustration of energy levels for Zn$_3$In$_2$S$_6$ and NiS c before and d after interfacing ($E_{vac}$, $E_F$, CB and VB denote the vacuum level, Fermi level, conduction band and valence band, respectively). e Schematic diagram of the dual-function photocatalytic reaction mechanism for selective oxidation of HMF to H$_2$ and DFF (e$^-$ and h$^+$ are photogenerated electrons and holes, respectively). Reprinted with permission from Ref. [59]. Copyright 2020 Elsevier
Figure 9b displays the photocatalytic mechanism of HMF conversion over CoPz/g-C$_3$N$_4$. Upon visible light irradiation, the CoPz dispersed on g-C$_3$N$_4$ transforms triplet O$_2$ ($^3$O$_2$) into 1O$_2$ by the energy transition, and the appropriate oxidation potential of 1O$_2$ can selectively convert HMF to the target product FDCA. HMF as a hygroscopic compound easily undergoes intermolecular etherification to form a HMF-dimer [56]. However, it can be used as a beneficial starting compound to yield usable and stable DFF. The target product DFF is widely used to manufacture the intermediates of pharmaceuticals, antifungal agents, ligands, and nematocides [57]. Nb$_2$O$_5$ catalyst calcined at 800 °C (Nb$_2$O$_5$-800) has been used for photocatalytic selective oxidation of HMF and shows an excellent DFF selectivity (90.6%) in HMF-containing benzo trifluoride (BTF) solvent under aerobic condition with visible light irradiation (Table 1, entry 1) [58]. Although the band-gap energy of Nb$_2$O$_5$ exceeds 3.2 eV, the selective HMF oxidation can occur under visible light, which might be attributed to the generation of visible-light-responsive alcoholate species between HMF and Nb$_2$O$_5$. To identify the main active species in this reaction, a benzoquinone scavenger for O$_2^-$ or isopropanol scavenger for OH is added to the reaction. The results show that no substantial effect on the performance is detected, revealing that neither O$_2^-$ nor OH is the dominant active species for the reaction. In contrast, the photocatalytic performance remarkably decreases with the addition of a hole scavenger (ethylenediaminetetraacetic acid), suggesting that the photogenerated hole has an important effect on the photocatalytic process.

HMF Oxidation Coupled with Photocatalytic H$_2$ Generation

A dual-function photocatalytic reaction for selective oxidation of biomass-relevant alcohols into clean energy and useful fine chemicals can be conducted in one pot to simultaneously use photogenerated electrons and holes. Noble metal-free 2D/2D-3D NiS/Zn$_3$In$_2$S$_6$ composites have been fabricated via a one-step approach for efficient transformation of HMF into DFF and H$_2$ with high selectivity under visible light [59]. As shown in Fig. 9c, the VB potential of Zn$_3$In$_2$S$_6$ is estimated to be 1.83 eV. The optical band-gap energy of Zn$_3$In$_2$S$_6$ is calculated to be 2.80 eV based on the Kubelka–Munk function, and thus the CB potential of Zn$_3$In$_2$S$_6$ is determined as −0.97 eV. The work function of NiS (5.5 eV) is higher than that of Zn$_3$In$_2$S$_6$ (3.6 eV) [60, 61]. Consequently, when Zn$_3$In$_2$S$_6$ and NiS contact intimately, a barrier region is established in the contact interface (yellow rectangle in Fig. 9d), so that the photoexcited electrons in Zn$_3$In$_2$S$_6$ can spontaneously transfer to NiS across the interface (Fig. 9e), thereby contributing to the separation and migration of photogenerated charge carriers. HMF can be converted to DFF and H$^+$ by holes of Zn$_3$In$_2$S$_6$, while the electrons can react with H$^+$ on the NiS surface to form H$_2$. Hence, a cyclic and efficient reaction toward solar-to-chemical energy transformation of biomass-relevant feedstocks is running. Additionally, Pt modified porous carbon nitride (SGCN) has been synthesized by a combined sol–gel/thermal condensation approach and applied to photoredox-catalyzed HMF oxidation coupled with H$_2$ production (Table 1, entry 2) [62]. Because of the appropriate VBM of SGCN, the unselective OH radicals cannot be generated, thereby inhibiting the over-oxidation or photocatalytic mineralization.

Photocatalytic Hydrogenation of HMF

Notably, hydrogenation of HMF can obtain various valuable fine chemicals, such as 2,5-dihydroxymethylfuran (DHMF), 2,5-dimethyltetrahydrofuran (DMTHF) and 2,5-dihydroxytetrahydrofuran (DHTMF) [52, 63]. Among them, DHMF has been used not only in the synthesis of drugs but also in the production of resins, polymers and artificial fibers [52]. In this scenario, Pt/g-C$_3$N$_4$ has been synthesized for photocatalytic hydrogenation of HMF to DHMF under visible light with trimethylamine (TEA) as a sacrificial electron donor (Table 1, entry 7) [64]. In this system, the doped Pt promotes H$_2$ evolution from water, and then the formed H$_2$ partakes in the reduction of HMF to yield DHMF. Compared with other metal-doped g-C$_3$N$_4$ (M/g-C$_3$N$_4$, M=Zn, Ni, Cu, Co, La, Ce), Pt/g-C$_3$N$_4$ exhibits the best performance for HMF hydrogenation. Because of its high work function, Pt can efficiently trap electrons from the CB of excited g-C$_3$N$_4$, which facilitates the separation of the photogenerated electron–hole pairs [65, 66], resulting in the enhanced photoactivity of HMF hydrogenation.

In another example, Pd nanoparticles supported on amine-functionalized iron-based metal–organic frameworks (Pd/MIL-101(Fe)-NH$_2$) have been prepared by Dong and coworkers [67] for visible-light-induced transfer hydrogenation of HMF to DHMF with triethylamine-formic acid (TEA-HCOOH) as the hydrogen source (Table 1, entry 8). In comparison, Pd deposited metal–organic frameworks samples without amine group modification exhibit insufficient activity toward visible-light-induced transfer hydrogenation reactions, indicating that the amine group plays a unique role in the photocatalytic hydrogenation. DFT studies demonstrate that the amine group not only stabilizes Pd nanoparticles in Pd/MIL-101(Fe)-NH$_2$ but also enhances the electron density of the Pd center upon light adsorption, thereby promoting biomass-based HMF hydrogenation.

Photocatalytic Conversion of Furfural Alcohol (FA)

Selective Oxidation of FA

FA is a sustainable candidate for biofuels and high value-added fine chemicals, because it can be used as a starting
feedstock for the synthesis of ethyl levulinate, furfural, furoic acid, pentanediol, etc. [68]. In this regard, \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) nanosheet provides an environmentally friendly and sustainable way toward conversion of FA to furfuraldehyde [69]. Monolayer \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) nanosheet is a lepidocrocite-type photocatalyst composed of a polyanion layer, which is constructed from a twisted TiO\(_6\) octahedral plate with rich defects. Because of the ultrathin structure of monolayer \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) nanosheet, many unsaturated surface metal sites (USMSs) are fully exposed on the monolayer nanosheet and serve as the Lewis acid sites [70, 71]. Consequently, the FA molecules are activated on \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) nanosheet because of the intimate interaction between USMSs and FA. Hydrogen nuclear magnetic resonance (\(^1\)HNMR) spectroscopy and in situ Fourier transform infrared spectroscopy (FTIR) results illustrate that \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) possesses more surface Lewis and Brønsted acid sites than the layered counterpart. Lewis acid sites serve as the active sites for effectively adsorbing and activating FA molecules, and Brønsted acid sites significantly improve photocatalytic performance. Based on the above results, a proposed photocatalytic mechanism for the photocatalytic oxidation of FA to furfuraldehyde over \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) is elucidated in Fig. 10. Under visible light, the surface Lewis acid sites of \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) effectively adsorb and activate FA molecules, and \( O_2 \) is captured by photogenerated electrons to form \( O_2^- \). Thereafter, the absorbed FA on \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) is deprotonated to generate the active \( C_5H_5O_2^+ \) intermediate, and this intermediate further releases a proton with the assistance of \( O_2^- \) to produce furfuraldehyde. Because of the weak interactions between \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) and furfuraldehyde, furfuraldehyde can successfully release from the surface of \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \), leading to a high furfuraldehyde selectivity.

**FA Oxidation Coupled with Photocatalytic H\(_2\) Generation**

To effectively use photogenerated electrons and holes, \( Ti_3C_2T_x \) MXene and cadmium sulfide (\( Ti_3C_2T_x/CdS \)) are integrated via a low-temperature wet chemistry approach for FA transformation integrated with H\(_2\) evolution. In the \( Ti_3C_2T_x/CdS \) composites, \( Ti_3C_2T_x \) serves as an “electron

![Fig. 10](image_url) Proposed reaction mechanism for the photocatalytic selective oxidation of FA over \( H_{1.4}Ti_{1.65}O_4 \cdot H_2O \) with \( O_2 \). Reprinted with permission from Ref. [69]. Copyright 2018 Elsevier

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sink" to trap photoexcited electrons from CdS, thereby leading to the moderate photoactivity of FA oxidation coupled with H₂ production (Table 1, entry 9) [68]. The possible mechanism shows that the photogenerated charge carriers are initially formed in CdS under visible light irradiation. The electrons easily migrate from the CB of CdS to the Ti₃C₂Tx because of the low Fermi energy level of Ti₃C₂Tx and the close interfacial connection between CdS and Ti₃C₂Tx [72, 73], and then the photogenerated electrons react with H₂O to yield H₂. Simultaneously, FA is oxidized into furfural by the photogenerated holes.

**Photocatalytic Conversion of Benzyl Alcohol (BA)**

**Selective Oxidation of BA**

BA as a biofeedstock model substrate can be converted into not only carbonyl compounds but also C–C coupled chemicals by using renewable solar energy, and these compounds are the essential precursors for the synthesis of fragrances, confectioneries and pharmaceuticals [17, 74]. DiMeglio and coworkers [75] have uncovered that nitrate mediators (nitrate salts of manganese, calcium, lithium and magnesium) improve the photocatalytic performance of BA oxidation to benzaldehyde (BAD) over CdS nanowires. To further understand the role of nitrate in this system, tetramethyl-ethylene (TME) as a nitrate radical trap is introduced to the photocatalytic BA oxidation reaction, in which the nitrate radical (NO₃⁻) and TME undergo an olefin epoxidation to liberate NO₂ (Fig. 11a), implying the generation of NO₃⁻. Compared with nitrate-free BA solution, a negligible difference in the photocatalytic reaction rate is observed in nitrate-free BA solution with TME. However, the photoactivity of BA oxidation is significantly decreased by adding TME to the LiNO₃-containing solution, suggesting that NO₃⁻ is oxidized to NO₃ over CdS, which then mediates BA conversion into BAD. Kinetic and trapping experiments suggest that selective oxidation of BA to BAD undergoes a mechanism mediated by NO₃ radicals instead of being directly oxidized by photoexcited holes. The O₂⁻ formed by photogenerated electrons acts as a base, and another NO₃ radical mediates the second electron transfer necessary for BAD production (Fig. 11b). In addition, nitrate-mediated unfunctionalized CdS nanowires also oxidize HMF into DFF, while no HMF oxidation is observed in the absence of the nitrate mediator. This study provides new opportunities for facile nitrate salts mediated biomass upgrading.

**BA Oxidation Coupled with Photocatalytic H₂ Generation**

Coproduction of value-added chemicals and H₂ can be obtained from organic feedstocks by making full use of photogenerated holes and electrons. CdS deposited anisotropic Pt-tipped Au nanorods (Au–Pt@CdS), a dual-function
photocatalyst, is applied to the transformation of BA to $\text{H}_2$ and BAD under Vis–NIR light irradiation [76]. The results show that the Au–Pt@CdS composite exhibits higher production of $\text{H}_2$ and BAD than CdS deposited Au nanorods (Au@CdS) and CdS deposited Pt-covered Au nanorods (Au@Pt@CdS), indicating that the spatial distribution of metal components and the composition of bimetal-semiconductor composites affect photocatalytic activity. In the Au@Pt@CdS composites, the unique surface plasmon resonance (SPR) of Au nanorods not only boosts light absorption but also enhances the local electric field, thus promoting the charge carrier generation and transfer of CdS. Pt nanoparticles anisotropically located at the tips of Au nanorods act as a cocatalyst and an electron reservoir to drive the directional charge transfer and promote the proton reduction. As a result, an excellent activity of coproduction of BAD and $\text{H}_2$ from BA is achieved. Mechanistic studies demonstrate that when the Au–Pt@CdS composite is simulated by Vis–NIR light (λ > 420 nm), the CdS is excited by short wavelength visible light (420 nm < λ < 520 nm) to yield charge carriers. Moreover, the SPR excitation of plasmonic Au–Pt bimetal by longer wavelength light offers an enhanced electric field to improve the photoabsorption, and facilitates the charge carriers generation and separation in CdS [77–79]. Afterward, the photogenerated electrons migrate from the CB of CdS to the anisotropic Au–Pt bimetal component, and then the electrons combine with protons to generate $\text{H}_2$ by diffusing through the porous and loose CdS thin layer [80, 81]. BA is selectively oxidized into BAD through the holes accumulated in the VB of CdS (Fig. 11c). The photogenerated charge carriers in ternary Au–Pt@CdS are spatially separated, and their recombination is remarkably inhibited, which leads to enhanced photocatalytic performance.

Recently, our group synthesized a flower-like nickel modified ZnIn$_2$S$_4$ (Ni:ZnIn$_2$S$_4$) composite via a photodeposition method to tune the selectivity of BA oxidation [82]. Compared to blank ZnIn$_2$S$_4$, Ni:ZnIn$_2$S$_4$ exhibits a higher BAD selectivity, because the Ni site facilitates the α-H abstraction to inhibit the formation of hydrobenzoin (HB). In addition, control experiments and the EPR technique jointly confirm that the CH(OH)Ph free radicals are the main active species in the reaction. Similarly, Ni-based compound (Ni$_x$P and NiS) modified ZnIn$_2$S$_4$ composites are also favorable to α-H abstraction, showing a photoactivity and selectivity trend analogous to that of Ni:ZnIn$_2$S$_4$. A possible mechanism for photocatalytic transformation of BA integrated with H$_2$ production over Ni:ZnIn$_2$S$_4$ is sketched in Fig. 11d. The ZnIn$_2$S$_4$ component in the Ni:ZnIn$_2$S$_4$ composite is initially excited to generate charge carriers under visible light. The electrons flow from the CB of ZnIn$_2$S$_4$ to the Ni sites, and then protons originating from water combine with the photogenerated electrons to yield H$_2$. Simultaneously, the proton of OH group in BA is adsorbed on the surface of ZnIn$_2$S$_4$ and the other proton abstracted from C–H is adsorbed by the Ni site, forming two abstracted protons and a CH(OH)Ph radical. Finally, BAD is formed from the CH(OH)Ph radical oxidation by holes, and abstracted protons are either reduced by photogenerated electrons to release H$_2$ or adsorbed on Ni:ZnIn$_2$S$_4$. This work offers a paradigm for tuning the selectivity of target-oriented biomass intermediates valorization to obtain fuels and high-value-added chemicals.

**Tunable Flexibility of Target Products from BA**

Interestingly, the different product distributions of BA oxidation can be tuned by photodepositing the low-valent Cd atoms onto the surface of the CdS quantum dots (QDs) modified with mercaptopropionyl ligands. Weiss’ group [16] has achieved a high selectivity of C–C coupling products (mainly hydrobenzoin) with the addition of Cd(ClO$_4$)$_2$ to the reaction to promote Cd$^0$ photodeposition. However, a high BAD selectivity (99%) is obtained by adding an electron scavenger (anthroquinone-2-sulfonate) to the same reaction to inhibit the photodeposition of Cd$^0$. Furthermore, the selectivity of C–C coupled products reaches 91% (68% hydrobenzoin, 15% benzil, and 8% 2-phenyl-acetophenone) by increasing the concentration of BA from 2 mmol/L to 5 mmol/L, because a higher α-hydroxybenzyl radical favors C–C coupling. On the contrary, the introduction of an electron scavenger not only effectively inhibits C–C coupling by preventing deposition of Cd$^0$ (Fig. 12a) but also removes the coulombic attraction of the hole in the CdS QDs, thereby the α-hydroxybenzyl radical intermediate can transfer to a photoexcited hole before coupling occurs.

In another example, Wang’s group [17] achieved controllable benzo (BZ) or deoxybenzoin (DOB) production from BA accompanied by $\text{H}_2$ evolution over Zn$_{0.6}$In$_2$S$_{3.6}$. The variation between DOB and BZ is related to the bottom potential of the CB of Zn$_{0.6}$In$_2$S$_{3.6}$, which can be manipulated by changing the relative contents of the indium and zinc precursors. In the ternary Zn$_{0.6}$In$_2$S$_{3.6}$, a higher Zn/In ratio corresponds to a lower CB bottom potential of the Zn$_{0.6}$In$_2$S$_{3.6}$ [83], leading to a high reduction capacity and contributing to the production of DOB. To understand the reaction pathway, the time distribution curve of the photocatalytic transformation of BA over Zn$_{0.6}$In$_2$S$_{3.6}$ was investigated. As depicted in Fig. 12b, BA is rapidly converted into HB and $\text{H}_2$ using blue LED illumination, and a maximum yield (62%) of HB appears after 4 h of irradiation. Afterward, the amounts of dominant product DOB, by-products BZ and trimers increase with decreasing HB, accompanied by the production of $\text{H}_2$. Additionally, control experiments were conducted to prove that HB is the intermediate in the production of DOB and BZ. As sketched in Fig. 12c, Zn$_{0.6}$In$_2$S$_{3.6}$ displays a moderate DOB yield (64%) and a relatively low yield of BZ (22%) under the illumination of blue LED. In comparison, the reaction over Zn$_{0.2}$In$_2$S$_{3.6}$.
affords BZ and DOB in 61% and 25% yield, respectively. The above results further confirm that the variation of target product over the Zn$_x$In$_2$S$_{3+x}$ can be tuned by altering Zn/In ratios. Based on the above analysis, a tentative reaction mechanism of photocatalytic fragmentation of HB into DOB is proposed in Fig. 12d. Under blue LED irradiation, the cleavage of the C–H bond in HB occurs by photogenerated holes to deliver the R$_{1b}$ radical. Afterward, the R$_{1b}$ radical transfers to the CB and reduces into 1,2-diphenylethenol (DPE) through photogenerated electrons. Finally, DPE tautomerizes to DOB.

**Conclusions and Outlook**

In summary, we have concisely recapped the recent progress in the photocatalytic upgrading of biomass-derived chemicals, including photocatalytic fragmentation of lignin models and photoredox-catalyzed biomass-relevant alcohols conversion. In particular, we offer the latest overview of visible-light-promoted rupture of β-O-4 bonds or C–C bonds of lignin models into valuable aromatic fragments. Then, we highlight the dual-function photoredox system for H$_2$ production integrated with alcohols oxidation and the photocatalyzed reaction toward controllable target-oriented product variation from the same substrates. Although considerable progress has been made in biomass-derived platform molecules conversion, it cannot meet the current requirements of industrial production.
and the selectivity of single C–C coupling chemicals remains unsatisfactory. Therefore, there remains a wide scope for improving the value of biomass-derived feedstocks for practical applications. In the end, we provide personal insights to further expand the manufacture of biomass-derived compounds molecule valorization under visible light.

First, the efficient strategies for developing and using the complicated and recalcitrant native lignin must be further explored. The viable catalytic tandem reaction systems, by effectively combining photocatalysis and traditional catalysis (thermocatalysis and enzyme catalysis), are anticipated to provide a more efficient and entire use of native lignin. Second, selectivity is a key issue for biomass-relevant derivatives conversion. In this regard, multifunctional semiconductor-based catalysts for tuning the selectivity of the target product are worthy of exploration. For example, surface modification can be used to alter a catalysts’ ability to adsorb or desorb intermediates, thereby tuning the selectivity of the target product. In addition, adjusting the band gap of photocatalysts can tailor the suitable oxidation capacity of photogenerated holes for alcohol conversion and different product distributions. Third, photocatalytic dual-function systems based on simultaneously using photoexcited electrons and holes need to be further established. For instance, assembling biomass-derived platform molecules (e.g., alcohols) oxidation and other reduction reactions, such as CO₂ reduction and N₂ fixation, offer a promising way to extend cooperative photoredox catalysis to synthesize the value-added chemicals and clean fuels together. Finally, the underlying mechanism of various photocatalytic biomass-derived chemical conversions remains to be investigated. In this context, in situ characterization techniques, such as in situ Fourier transform infrared spectroscopy (FTIR) and in situ electron spin resonance (ESR) spectroscopy, in combination with mass spectrometry and theoretical calculations, can be used to track the intermediate species during photocatalysis, which aids the understanding of the specific mechanistic pathway and explains the selectivity of products. It is hoped that this timely overview of the latest advances in photocatalytic biomass-derived compounds upgrading will inspire the rational design of photocatalysts toward selective organic transformations with tunable target-oriented product distributions.

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