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Nanoconfined tandem three-phase photocatalysis for highly selective  $\text{CO}_2$  reduction to ethanol

A noteworthy 94.15% ethanol selectivity is achieved *via* a nanoconfined tandem three-phase  $\text{CO}_2$  photoreduction system, which integrates hydrophobicity and the nanoconfinement effect with tandem reactions, reducing the energy barrier for  $^*\text{CO}-^*\text{CO}$  coupling, thereby significantly enhancing ethanol selectivity.

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## Introduction

The urgent need for sustainable development in response to the challenges posed by the greenhouse effect and dwindling fossil resources has led to the exploration of eco-friendly strategies.<sup>1,2</sup> Among these strategies, the photocatalytic CO<sub>2</sub> reduction reaction (PCRR) has emerged as a promising approach for converting CO<sub>2</sub> and H<sub>2</sub>O into chemicals and fuels.<sup>3,4</sup> During the PCRR, solar energy is transformed into chemical energy, and stored in the chemical bonds of resulting compounds containing one (C<sub>1</sub>) or multiple (C<sub>2+</sub>) carbon atoms. Importantly, the production of C<sub>2+</sub> products is particularly desirable due to their higher energy density and economic potential.<sup>5</sup> Furthermore, ethanol, a liquid C<sub>2</sub> fuel, has garnered significant research attention owing to its ease of storage and transport.<sup>6</sup> Nevertheless, ethanol production faces challenges in terms of low selectivity, largely attributed to the challenging C-C coupling process necessary for generating C<sub>2+</sub> products.<sup>7</sup>

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## Nanoconfined tandem three-phase photocatalysis for highly selective CO<sub>2</sub> reduction to ethanol<sup>†</sup>

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The conversion of CO<sub>2</sub> and H<sub>2</sub>O into ethanol with high selectivity via photocatalysis is greatly desired for effective CO<sub>2</sub> resource utilization. However, the sluggish and challenging C-C coupling hinders this goal, with the behavior of \*CO holding the key. Here, a nanoconfined and tandem three-phase reaction system is established to simultaneously enhance the \*CO concentration and interaction time, achieving an outstanding ethanol selectively of 94.15%. This system utilizes a tandem catalyst comprising an Ag core and a hydrophobic Cu<sub>2</sub>O shell. The hydrophobic Cu<sub>2</sub>O shell acts as a CO<sub>2</sub> reservoir, effectively overcoming the CO<sub>2</sub> mass-transfer limitation, while the Ag core facilitates the conversion of CO<sub>2</sub> to CO. Subsequently, CO undergoes continuous reduction within the nanoconfined mesoporous channels of Cu<sub>2</sub>O. The synergy of enhanced mass transfer, nanoconfinement, and tandem reaction leads to elevated \*CO concentrations and prolonged interaction time within the Cu<sub>2</sub>O shell, significantly reducing the energy barrier for \*CO-\*CO coupling compared to the formation of \*CHO from \*CO, as determined by density functional theory calculations. Consequently, C-C coupling preferentially occurs over \*CHO formation, producing excellent ethanol selectivity. These findings provide valuable insights into the efficient production of C<sub>2+</sub> compounds.

The C-C coupling process, which is sluggish and possesses a high energy barrier, is significantly influenced by the adsorbed carbon monoxide (\*CO) on the catalyst surface.<sup>7,8</sup> Typically, following the conversion of a CO<sub>2</sub> molecule to \*COOH, the resulting \*CO governs the direction of the reaction towards the production of either C<sub>1</sub> compounds (e.g., methanol) or C<sub>2+</sub> compounds (e.g., ethanol).<sup>9</sup> Specifically, the hydrogenation of \*CO into its derived intermediates, such as \*CHO, facilitates the formation of C<sub>1</sub> products, while the dimerization of \*CO or the polymerization of \*CO and its derived intermediates to proceed with C-C coupling initiates the C<sub>2</sub> pathway.<sup>9</sup> Furthermore, according to Le Chatelier's principle, an elevated concentration of \*CO is generally beneficial for the PCRR<sup>10</sup> and the C-C coupling process.<sup>11</sup> Since \*CO is a downstream intermediate of CO<sub>2</sub> protonation, researchers have strived to increase the surface \*CO concentration to promote the formation of C<sub>2+</sub> products by enhancing the local concentration of CO<sub>2</sub>.<sup>8,12</sup> However, in aqueous solutions, the PCRR encounters the mass-transfer limitation of CO<sub>2</sub> due to its poor solubility and diffusion coefficient, resulting in a 1:1300 ratio between CO<sub>2</sub> and H<sub>2</sub>O molecules at 1 atm pressure. This limitation further leads to a significant scarcity of diffusing CO<sub>2</sub> molecules reaching the catalyst surface.<sup>13</sup> Researchers have proposed that this mass-transfer limitation can be overcome by adjusting the catalyst surface to be hydrophobic, which can promote the enrichment of gaseous CO<sub>2</sub> molecules on the catalyst surface. This adjustment creates abundant three-phase (gaseous CO<sub>2</sub>, liquid water,

and solid catalysts) reaction sites for enhanced PCRR.<sup>13,14</sup> Our previous study has developed a hydrophobic cuprous oxide ( $\text{Cu}_2\text{O}$ ) hollow structure to mitigate the  $\text{CO}_2$  mass-transfer limitation, achieving enhanced ethanol production.<sup>5</sup> Nonetheless, its selectivity remains unsatisfactory. One possible reason is the limited increase in  $^*\text{CO}$  concentration, while another could be the insufficient interaction time between  $\text{C}_1$  intermediates necessary for effective C-C coupling, which could not align with the sluggishness of this process.<sup>15</sup>

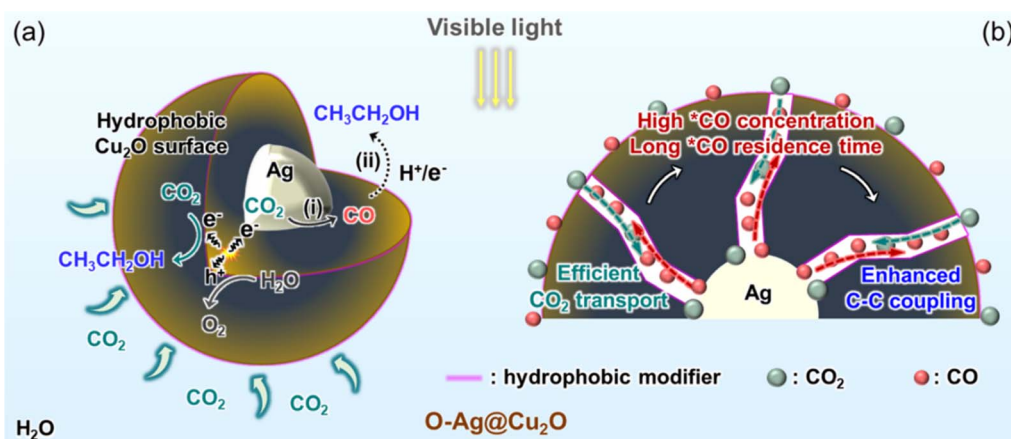
Herein, a nanoconfined and tandem three-phase reaction system is constructed, simultaneously improving the  $^*\text{CO}$  concentration and interaction time between  $\text{C}_1$  intermediates, thereby elevating the selectivity of ethanol production to 94.15%. Specifically, the tandem three-phase PCRR employs a tandem catalyst comprising silver (Ag) particles covered by a mesoporous shell of hydrophobic  $\text{Cu}_2\text{O}$ . It is well-known that Ag exhibits high activity and selectivity for CO formation in the  $\text{CO}_2$  reduction reaction,<sup>16,17</sup> with a relatively weak CO binding strength on its surface.<sup>18</sup> Moreover,  $\text{Cu}_2\text{O}$  contains  $\text{Cu}^1$  centers, which are considered effective catalytic sites for  $\text{CO}_2$  conversion,<sup>19</sup> facilitating CO adsorption and enhancing  $^*\text{CO}$  binding for subsequent coupling reactions with protons and electrons.<sup>20,21</sup> Thus, when this tandem catalyst is constructed in a core–shell structure, denoted as  $\text{O-Ag@Cu}_2\text{O}$ , the surface of Ag particles serves as the reactive sites for converting  $\text{CO}_2$  to CO, constituting the first step of the tandem reaction, as depicted in Scheme 1a.<sup>17</sup> Then, the generated CO desorbs from the Ag particles, diffuses along the mesoporous channels, and re-adsorbs onto the hydrophobic  $\text{Cu}_2\text{O}$  reactive sites for subsequent reduction, constituting the second step of the tandem reaction. During the reaction process, the hydrophobic  $\text{Cu}_2\text{O}$  surface facilitates the capture and enrichment of  $\text{CO}_2$  molecules, ensuring a concentrated source of CO for efficient transport to the reactive sites. Upon exposure to visible light, the high concentration of CO can undergo direct reduction on the  $\text{Cu}_2\text{O}$  surface. Moreover, CO<sub>2</sub> can diffuse through the

mesoporous  $\text{Cu}_2\text{O}$  shell to interact with the inner Ag particles, where it is converted to CO (Scheme 1a). As CO diffuses outward along the mesoporous channels (Scheme 1b), the nanoconfinement effect of these channels increases the likelihood of collisions between CO molecules and the channel walls, as well as among the CO molecules themselves.<sup>22</sup> This increased collision probability raises diffusion resistance, thereby extending the residence time of CO. This phenomenon not only leads to the aggregation of CO within the channels, increasing the  $^*\text{CO}$  concentration, but also provides sufficient time for  $^*\text{CO}$  molecules to interact with each other. Consequently, the designed system achieves a synergistic effect of mass transfer enhancement, nanoconfinement, and tandem reaction, resulting in high  $^*\text{CO}$  concentrations at active sites and prolonged interaction time. Density functional theory (DFT) calculations demonstrate that the elevated  $^*\text{CO}$  concentration significantly lowers the energy barrier for  $^*\text{CO}$ – $^*\text{CO}$  coupling compared to the formation of  $^*\text{CHO}$  from  $^*\text{CO}$ . This results in the preferential formation of  $\text{C}_{2+}$  products over  $\text{C}_1$  products, thereby enhancing ethanol selectivity. This strategy achieves effective  $\text{CO}_2$ -to-ethanol conversion with high selectivity and offers guidance for producing  $\text{C}_{2+}$  products efficiently.

## Results and discussion

### Morphological characterization

The synthesis of  $\text{O-Ag@Cu}_2\text{O}$  commences with the preparation of Ag particles derived from a silver chloride (AgCl) colloid. These prepared Ag particles are nearly spherical, with an average size of approximately 40 nm (Fig. S1†). Subsequently, the Ag particles are coated with  $\text{Cu}_2\text{O}$  to establish the core–shell structure of  $\text{Ag@Cu}_2\text{O}$  (Fig. S2a†). Following this,  $\text{Ag@Cu}_2\text{O}$  is hydrophobically modified using minute quantities of 1-dodecanethiol (DDT) to form the target catalyst  $\text{O-Ag@Cu}_2\text{O}$  (Fig. S2b†). To assess the impact of hydrophobicity on  $\text{O-Ag@Cu}_2\text{O}$ ,  $\text{Ag@Cu}_2\text{O}$  is utilized as a control sample.



**Scheme 1** Photocatalytic  $\text{CO}_2$  reduction reaction (PCRR) over  $\text{O-Ag@Cu}_2\text{O}$ . (a) PCRR mechanisms at the active sites of Ag and hydrophobic  $\text{Cu}_2\text{O}$  surfaces. The hydrophobic surface serves as a reservoir for  $\text{CO}_2$  molecules, where they can also undergo reduction. (i) and (ii) Refer to the first and second steps of the tandem reaction, respectively. (b) Gas transport processes within the mesoporous channels. The hydrophobic  $\text{Cu}_2\text{O}$  channels facilitate the transportation of  $\text{CO}_2$  to the inner Ag surface, where it is converted into CO. The nanoconfinement effect of the channels results in an extended residence time for CO.

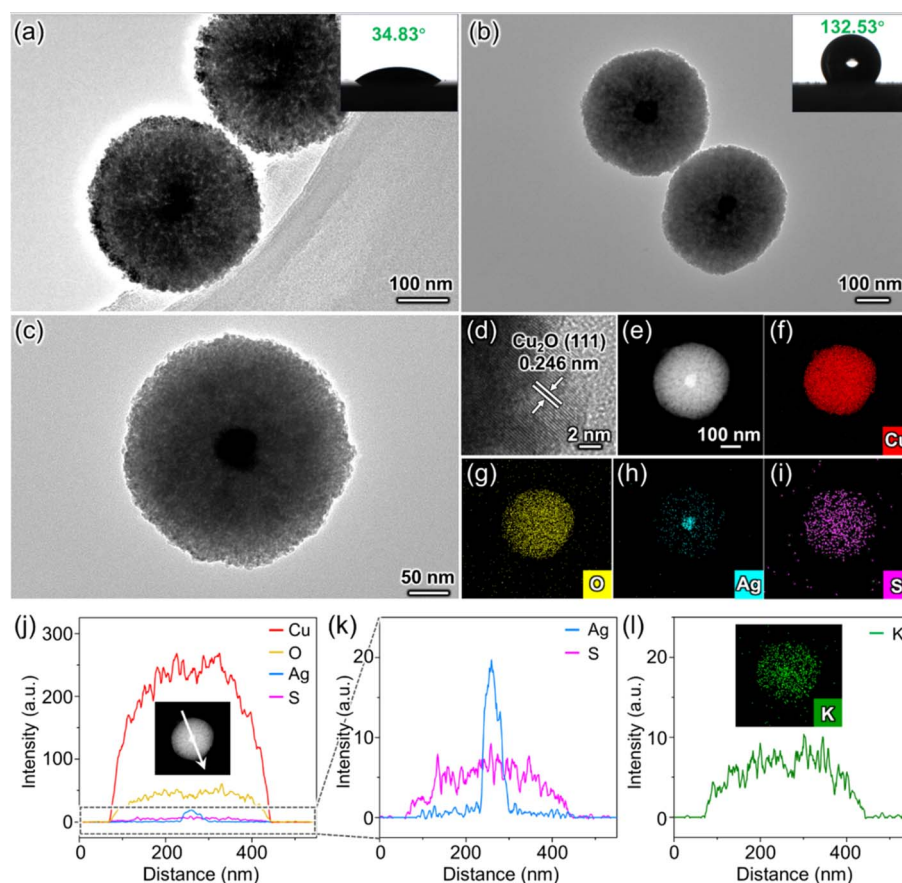


Additionally, to assess the role of Ag particles within the catalyst, bare  $\text{Cu}_2\text{O}$  (denoted as b- $\text{Cu}_2\text{O}$ ) and its hydrophobically modified counterpart (denoted as O-b- $\text{Cu}_2\text{O}$ ) are synthesized for comparison (Fig. S2c and d†). Furthermore, to investigate the impact of nanoconfinement effects in O-Ag@ $\text{Cu}_2\text{O}$ , the catalyst with Ag particles loaded onto the external surface of  $\text{Cu}_2\text{O}$  (denoted as Ag/ $\text{Cu}_2\text{O}$ ) and its hydrophobic variant (denoted as O-Ag/ $\text{Cu}_2\text{O}$ ) are prepared as control samples (Fig. S2e and f†). In this case, the spatial distribution of Ag particles on the external surface of  $\text{Cu}_2\text{O}$  means that the CO generated on the surface of Ag particles does not need to diffuse out through the mesoporous  $\text{Cu}_2\text{O}$ . Therefore, the nanoconfinement effect of the mesoporous channel does not impact the generated CO.

The successful synthesis of the Ag@ $\text{Cu}_2\text{O}$  structure is evident from the transmission electron microscope (TEM) images (Fig. 1a and S3†). The core-shell configuration of Ag@ $\text{Cu}_2\text{O}$ , with Ag serving as the core and mesoporous  $\text{Cu}_2\text{O}$  as the shell, exhibits a water contact angle (WCA) of  $34.83^\circ$ , indicative of its inherent hydrophilicity. Additionally, the mesoporous characteristic of the  $\text{Cu}_2\text{O}$  shell is substantiated by the  $\text{N}_2$  adsorption-desorption measurement, which displays a type IV curve with an H3 hysteresis loop (Fig. S4a†). Following DDT modification, the core-shell structure of O-Ag@ $\text{Cu}_2\text{O}$  remains intact. Notably,

a transition to hydrophobic behavior is observed, marked by a significant rise in the WCA to  $132.53^\circ$  (Fig. 1b). The magnified TEM image of O-Ag@ $\text{Cu}_2\text{O}$  further illustrates its mesoporous structure (Fig. 1c), featuring a pore diameter of approximately 19 nm (Fig. S4b†). Compared to microporous structures, which tend to impede molecular transport due to their smaller pore sizes ( $<2$  nm),<sup>23</sup> this remarkable mesoporous structure facilitates the mass transport of molecules.<sup>22</sup> Moreover, the lattice fringe of 0.246 nm shown in Fig. 1d aligns with the (111) plane of  $\text{Cu}_2\text{O}$ , suggesting the  $\text{Cu}_2\text{O}$  composition of the shell. Furthermore, the energy-dispersive X-ray spectroscopy (EDS) mapping and line scan results presented in Fig. 1e–k verify the spatial distribution relationship between Ag and  $\text{Cu}_2\text{O}$  within the core-shell structure of O-Ag@ $\text{Cu}_2\text{O}$ .

More importantly, the EDS line scan result of sulfur (S) element originating from DDT shows that DDT is evenly distributed throughout the entire mesoporous structure of O-Ag@ $\text{Cu}_2\text{O}$  (Fig. 1k). This observation suggests that DDT permeates the mesoporous channels to some extent, increasing the hydrophobicity of the channels, thereby facilitating the  $\text{CO}_2$  transport within them. To assess whether the presence of DDT in mesoporous channels impedes water infiltration, a potassium ion ( $\text{K}^+$ ) tracing experiment was conducted (for details see



**Fig. 1** (a and b) TEM images of Ag@ $\text{Cu}_2\text{O}$  and O-Ag@ $\text{Cu}_2\text{O}$ , respectively. Insets in (a and b) show the WCAs of the corresponding samples. (c) TEM image of a single O-Ag@ $\text{Cu}_2\text{O}$  particle. (d) HRTEM image of O-Ag@ $\text{Cu}_2\text{O}$ . (e) HAADF-STEM image of O-Ag@ $\text{Cu}_2\text{O}$  and corresponding EDS mapping profiles of Cu (f), O (g), Ag (h), and S (i). (j) EDS line scan of O-Ag@ $\text{Cu}_2\text{O}$ . The inset displays the position and direction of the line scan. (k) Enlarged view of the region outlined by the dashed box in (j). (l) EDS line scan and mapping results from a  $\text{K}^+$  tracing experiment. The line scan position and direction are the same as that in the inset of (j).



Experimental Procedures in ESI†).<sup>24</sup> The experimental procedure involved immersing the sample in a potassium bicarbonate solution and subsequently analyzing the distribution pattern of  $K^+$ . EDS mapping combined with EDS line scan results clearly illustrate that  $K^+$  is evenly distributed throughout the entire sphere, indicating that water is capable of penetrating the mesoporous channels. This observation ensures the accessibility of subsequent  $CO_2$  and water molecules into the sphere through the mesoporous channels, where they can react on the surface of Ag particles. To further elucidate that the presence of DDT on the surface of O-Ag@Cu<sub>2</sub>O does not hinder the migration of charge carriers to the catalyst surface, Ag<sup>+</sup> probe experiments for O-Ag@Cu<sub>2</sub>O and Ag@Cu<sub>2</sub>O are compared (Fig. S5 and S6,† for details see Experimental Procedures in ESI†).<sup>5</sup> The atomic percentages of Cu and Ag elements in O-Ag@Cu<sub>2</sub>O (Fig. S5†) are found to be comparable to those in Ag@Cu<sub>2</sub>O (Fig. S6†), indicating that the quantity of Ag resulting from the reduction of adsorbed Ag<sup>+</sup> on O-Ag@Cu<sub>2</sub>O surface by photo-generated electrons is analogous to that on Ag@Cu<sub>2</sub>O surface. Therefore, the surface hydrophobic modification does not significantly affect the interaction between photogenerated electrons migrating to the surface and reactants adsorbed on the surface. To further explore the impact of different synthesis scales on O-Ag@Cu<sub>2</sub>O and to preliminarily assess the potential for large-scale production, TEM and WCA characterizations were performed on samples obtained at various scales (Fig. S7†). The results reveal no significant differences in morphology or hydrophobicity across the different scales, indicating that the designed catalyst may be suitable for scaling up. This finding provides a basis for considering future large-scale production.

Additionally, control samples including Ag/Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, b-Cu<sub>2</sub>O, and O-b-Cu<sub>2</sub>O are successfully synthesized and confirmed through TEM and WCA characterizations (Fig. S8 and S9†). Moreover, the comparable specific surface areas and pore diameter distributions for the synthesized samples render the variation in specific surface areas and pore size negligible (Fig. S4 and S10†). In addition, the Ag contents (wt%) in O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O are quantified as 2.032% and 2.027%, respectively, using inductively coupled plasma-optic emission spectroscopy (Table S1†). This similarity in Ag content between the two samples enables a comparison of the effects of distinct spatial distributions of Ag particles without the complicating factor of varying amounts of Ag.

### Structural characterization

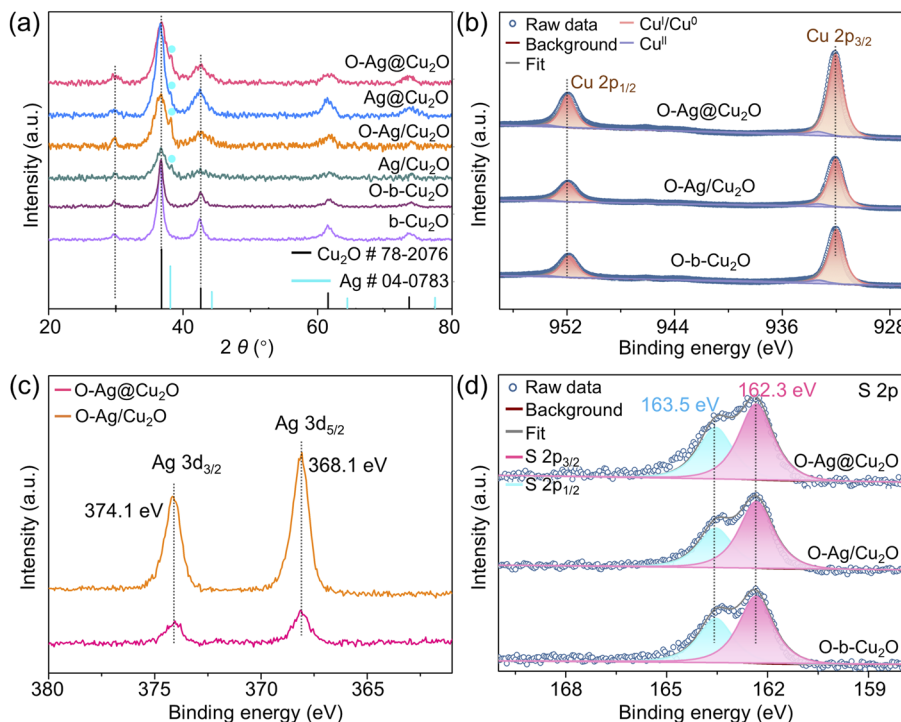
X-ray diffraction (XRD) analysis unveils the composition of the synthesized samples, showcasing a well-matched cubic phase Cu<sub>2</sub>O (JCPDS card no. 78-2076), alongside a weak peak corresponding to metallic Ag (Ag<sup>0</sup>) particles observed in O-Ag@Cu<sub>2</sub>O, Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, and Ag/Cu<sub>2</sub>O (Fig. 2a). X-ray photoelectron spectroscopy (XPS) further confirms the chemical composition of the synthesized samples. The high-resolution Cu 2p XPS spectra of O-Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, and O-b-Cu<sub>2</sub>O exhibit identical peak positions at 932.3 and 952.3 eV for Cu 2p<sub>3/2</sub> and Cu 2p<sub>1/2</sub>, respectively (Fig. 2b), indicating consistent valence

states of Cu (Cu<sup>I</sup>), as confirmed by Cu LMM Auger spectra (Fig. S11a†).<sup>25</sup> The weak peaks at 933.6 and 953.5 eV attributed to Cu<sup>II</sup> indicate the slight oxidation of the surface Cu<sub>2</sub>O. Similarly, their corresponding samples without hydrophobic modification show comparable Cu 2p XPS spectra, indicating their similar composition (Fig. S11b†). Additionally, the XPS surveys of O-Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, Ag@Cu<sub>2</sub>O, and Ag/Cu<sub>2</sub>O all exhibit discernible Ag signals (Fig. S11c and d†). In the cases of O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O, the high-resolution Ag 3d XPS spectra display consistent peak positions, indicating a uniform chemical state of Ag across these catalysts. Furthermore, the binding energies for Ag 3d<sub>5/2</sub> and Ag 3d<sub>3/2</sub> are located at 368.1 and 374.1 eV, respectively (Fig. 2c), consistent with the characteristic signals of Ag<sup>0</sup> and in agreement with the XRD results. In the cases of Ag@Cu<sub>2</sub>O and Ag/Cu<sub>2</sub>O, although the Ag 3d peak positions are consistent, indicating the uniform chemical state of Ag in these two catalysts, the binding energies are slightly shifted to 368.3 eV for Ag 3d<sub>5/2</sub> and 374.3 eV for Ag 3d<sub>3/2</sub> (Fig. S11e†). While the signals typical for Ag<sup>I</sup> are at 368.4 eV for Ag 3d<sub>5/2</sub> and 374.4 eV for Ag 3d<sub>3/2</sub>,<sup>26</sup> this suggests a coexistence of Ag<sup>0</sup> and Ag<sup>I</sup> on the surface of Ag particles in Ag@Cu<sub>2</sub>O and Ag/Cu<sub>2</sub>O,<sup>27</sup> implying a slight oxidation of the Ag particles for Ag@Cu<sub>2</sub>O and Ag/Cu<sub>2</sub>O. Additionally, as demonstrated in Fig. S12,† DDT exhibits the ability to interact with Ag particles. Consequently, when DDT is utilized to modify Ag@Cu<sub>2</sub>O and Ag/Cu<sub>2</sub>O, Ag<sup>I</sup> on the Ag particle surface may be reduced due to the reducibility of DDT,<sup>28</sup> leading to the conversion of the chemical state of Ag in O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O to Ag<sup>0</sup>. This finding further supports the notion that DDT can penetrate the mesoporous Cu<sub>2</sub>O shell in O-Ag@Cu<sub>2</sub>O, thereby accessing the surface of internal Ag particles.

Additionally, S signals are detected in the XPS surveys of O-Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, and O-b-Cu<sub>2</sub>O (Fig. S11d†), indicating the presence of DDT on the catalyst surface, consistent with observations from Fourier-transform infrared spectroscopy (FTIR) (Fig. S13†). Importantly, the high-resolution S 2p XPS spectra display consistent peak positions, suggesting a comparable state of DDT across the surfaces of all three samples (Fig. 2d and S11f†). Furthermore, the binding energy of S 2p<sub>3/2</sub> at 162.3 eV corresponds to surface thiolate species signals, implying the formation of -S-Cu by cleaving the sulphydryl group (-S-H) in DDT.<sup>29</sup> This observation indicates the occurrence of chemical adsorption between DDT and the Cu<sub>2</sub>O surface, which is further supported by Raman spectra (Fig. S14†). This chemical adsorption enables DDT to stably bind to the catalyst surface, preserving its stable hydrophobic properties essential for the catalytic reaction.

Subsequently, UV-vis diffuse reflectance spectroscopy was utilized to assess the light absorption ability and bandgap of the synthesized catalysts (Fig. S15†). All samples display a similar visible-light absorption range (Fig. S15a†), and the estimated bandgaps based on the Tauc plots are uniformly approximately 2.18 eV across all samples (Fig. S15b†). Additionally, after DDT modification, the absorbance intensity of the samples slightly decreases, which theoretically is unfavorable for the catalytic reaction.<sup>1</sup> However, subsequent photocatalytic performance tests demonstrate that the DDT-modified samples generally





**Fig. 2** Structural characterizations. (a) XRD patterns of b-Cu<sub>2</sub>O, O-b-Cu<sub>2</sub>O, Ag/Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, Ag@Cu<sub>2</sub>O, and O-Ag@Cu<sub>2</sub>O. (b) High-resolution Cu 2p XPS spectra of O-Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, and O-b-Cu<sub>2</sub>O. (c) High-resolution Ag 3d XPS spectra of O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O. (d) High-resolution S 2p XPS spectra of O-Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, and O-b-Cu<sub>2</sub>O.

perform better than their unmodified counterparts. This indicates that variations in the catalyst's absorbance intensity are not crucial factors contributing to the difference in performance. The charge separation efficiency is also compared by photoluminescence (PL) spectra and transient photocurrent response (Fig. S16†). As charge carrier recombination can release fluorescence, higher PL spectra intensity generally correlates with reduced charge separation efficiency.<sup>30</sup> Moreover, in transient photocurrent response, lower transient photocurrent intensity is indicative of inferior charge separation efficiency.<sup>30</sup> Remarkably, the PL and photocurrent intensity of O-Ag@Cu<sub>2</sub>O, Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, and Ag/Cu<sub>2</sub>O do not differ significantly. Following DDT modification, a slight increase in PL intensity indicates a slight decrease in carrier separation efficiency, aligning with the results of photocurrent testing, which might have had some detrimental effects on catalyst activity.<sup>31</sup> However, subsequent evaluation of photocatalytic performance reveals that the DDT-modified sample outperforms its unmodified counterpart. This discrepancy implies that the minor variation in carrier separation efficiency is not the primary determinant of the disparity in catalyst performance. Additionally, the decreased PL intensity and increased photocurrent intensity observed in O-Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, Ag@Cu<sub>2</sub>O, and Ag/Cu<sub>2</sub>O compared to O-b-Cu<sub>2</sub>O and b-Cu<sub>2</sub>O suggest enhanced efficiency in charge separation. This improvement can be ascribed to the presence of Ag particles, which can trap the photogenerated electrons.<sup>32</sup> As illustrated in Fig. S17,† when Ag particles come into contact with Cu<sub>2</sub>O, the difference in their Fermi levels induces a redistribution of

interfacial charges.<sup>33</sup> Under illumination, electrons in the conduction band of Cu<sub>2</sub>O at the interface ultimately transfer to the Ag particles, while the holes in the valence band remain within Cu<sub>2</sub>O.<sup>34,35</sup> Consequently, the presence of Ag particles suppresses the recombination of electrons and holes in Cu<sub>2</sub>O to some extent, thereby enhancing their separation efficiency.

### PCRR performance

The photocatalytic performance of O-Ag@Cu<sub>2</sub>O and relevant reference catalysts for CO<sub>2</sub> reduction with H<sub>2</sub>O was evaluated in an aqueous solution under visible light ( $\lambda > 420$  nm). As depicted in Fig. 3a, all Cu<sub>2</sub>O-based catalysts demonstrate activity in producing liquid alcohols. However, the C<sub>2</sub> product of ethanol is the main carbonaceous product for O-Ag@Cu<sub>2</sub>O, with an ethanol generation rate of 450.19  $\mu\text{mol g}^{-1} \text{h}^{-1}$  and an ethanol selectivity of up to 94.15%, as evident from the chromatogram of the original product detection (Fig. S18†). In terms of both ethanol selectivity and generation rate, this performance ranks among the top levels observed in photocatalytic CO<sub>2</sub> reduction systems over the past five years (a detailed comparison is presented in Table S2†). In contrast, methanol is the predominant carbonaceous product for the reference catalysts of Ag@Cu<sub>2</sub>O, O-Ag/Cu<sub>2</sub>O, Ag/Cu<sub>2</sub>O, O-b-Cu<sub>2</sub>O, and b-Cu<sub>2</sub>O. Specifically, O-Ag@Cu<sub>2</sub>O exhibits a 10.89-fold increase in ethanol selectivity compared to Ag@Cu<sub>2</sub>O. Additionally, the calculation of effective photogenerated electrons utilized in the PCRR reveals that O-Ag@Cu<sub>2</sub>O employs a 6.65-fold greater amount of electrons compared to Ag@Cu<sub>2</sub>O, with a significant portion contributing to ethanol formation (Fig. 3b). These

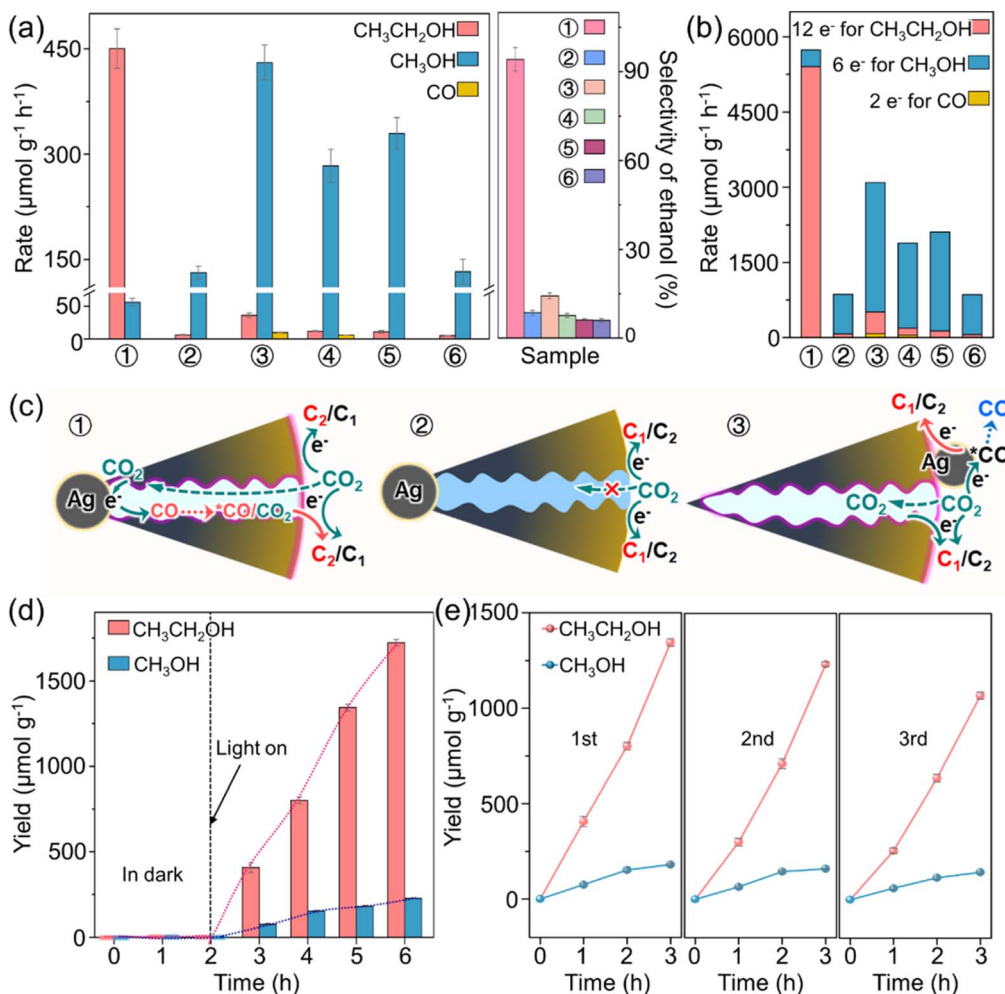


Fig. 3 PCRR performance. Generation rates of carbonaceous products and selectivity towards ethanol (a) and amount of effective photo-generated electrons utilized in the PCRR (b) for ① O-Ag@Cu<sub>2</sub>O, ② Ag@Cu<sub>2</sub>O, ③ O-Ag/Cu<sub>2</sub>O, ④ Ag/Cu<sub>2</sub>O, ⑤ O-b-Cu<sub>2</sub>O, and ⑥ b-Cu<sub>2</sub>O. The error bar in (a) represents the standard deviation of three measurements. (c) Schemes of the PCRR in the structures of ① O-Ag@Cu<sub>2</sub>O, ② Ag@Cu<sub>2</sub>O, and ③ O-Ag/Cu<sub>2</sub>O. The pink color on the inner wall of the pores and the outer surfaces of O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O represents DDT. (d) Ethanol and methanol yields over time for O-Ag@Cu<sub>2</sub>O. (e) Recycle experiments for O-Ag@Cu<sub>2</sub>O.

findings suggest the promotive effect of hydrophobicity on the PCRR. This effect can be further corroborated by comparing the photocatalytic performance of O-Ag/Cu<sub>2</sub>O and Ag/Cu<sub>2</sub>O, O-b-Cu<sub>2</sub>O and b-Cu<sub>2</sub>O, with more carbonaceous product generation and greater utilization of electrons for the hydrophobic ones. Moreover, compared to O-Ag/Cu<sub>2</sub>O with Ag particles loaded on the outer surface of Cu<sub>2</sub>O, O-Ag@Cu<sub>2</sub>O with Ag particles encapsulated within Cu<sub>2</sub>O exhibits a 6.59-fold increase in ethanol selectivity and a 1.86-fold increase in electron utilization. Additionally, no CO is detected for O-Ag@Cu<sub>2</sub>O, whereas a small amount of CO is observed for O-Ag/Cu<sub>2</sub>O. To further rule out the possibility that the significant performance difference between O-Ag/Cu<sub>2</sub>O and O-Ag@Cu<sub>2</sub>O arises from the different sizes of Ag particles in the catalysts, the Ag particle size in the comparative catalyst O-Ag/Cu<sub>2</sub>O was adjusted to match that in O-Ag@Cu<sub>2</sub>O. Performance tests reveal that even with identical Ag particle sizes, O-Ag/Cu<sub>2</sub>O still exhibits significantly lower performance than O-Ag@Cu<sub>2</sub>O (Fig. S19<sup>†</sup>). In addition, compared with O-b-Cu<sub>2</sub>O, both O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O

exhibit promoted CO<sub>2</sub>-to-alcohol conversion performance, closely related to the role of Ag particles.

In light of these experimental findings, potential reasons for the performance differences observed among various catalysts are proposed. As illustrated in Fig. 3c, the hydrophobic modification imparts hydrophobic characteristics to both the external surface and mesoporous channels of the catalyst, facilitating the enrichment and capture of CO<sub>2</sub>. Consequently, concerning the disparity between O-Ag@Cu<sub>2</sub>O and Ag@Cu<sub>2</sub>O, the hydrophobic properties of O-Ag@Cu<sub>2</sub>O enable CO<sub>2</sub> accumulation on both the inner and outer surfaces of O-Ag@Cu<sub>2</sub>O. Moreover, CO<sub>2</sub> can migrate towards the surface of the Ag particles enclosed within O-Ag@Cu<sub>2</sub>O, allowing Ag to actively participate in the PCRR.<sup>36</sup> Conversely, it is challenging for CO<sub>2</sub> to accumulate on the hydrophilic surface and enter the mesoporous channels of Ag@Cu<sub>2</sub>O, rendering the Ag particles ineffective. This observation is consistent with the PCRR results showing that the photocatalytic performance of Ag@Cu<sub>2</sub>O is similar to that of b-Cu<sub>2</sub>O (Fig. 3a), indicating the inefficacy of

the Ag particles (Fig. 3c and S20†). Furthermore, the difference between O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O stems from the distinct positions of the Ag particles within the catalyst. In the case of O-Ag@Cu<sub>2</sub>O, once the Ag particles inside the catalyst generate CO during PCRR, CO molecules need to diffuse out through the mesoporous channels. During this diffusion process, the nanoconfinement effect of the channels causes CO to aggregate within them. This phenomenon increases the concentration of \*CO on the active Cu<sub>2</sub>O sites and ample duration for \*CO molecules to engage in mutual interactions. This promotes the C-C coupling process and thereby accelerates ethanol production. In contrast, in the case of O-Ag/Cu<sub>2</sub>O, Ag particles are positioned on the external surface of the catalyst. Following CO generation, some of it may re-adsorb onto nearby active Cu<sub>2</sub>O sites, enhancing the PCRR, while the remainder may diffuse directly into the surrounding environment, leading to the production of gaseous CO. This direct overflow of CO limits the increase in \*CO concentration on the Cu<sub>2</sub>O active sites, resulting in a reduced ethanol production capacity compared to O-Ag@Cu<sub>2</sub>O.

To further evaluate the solar energy utilization efficiency of O-Ag@Cu<sub>2</sub>O, the solar-to-fuel conversion efficiency (STF) and apparent quantum efficiency (AQE) are calculated (Fig. S21 and Table S3 and S4†). The STF and AQE ( $\lambda = 420$  nm) are estimated to be 0.215% and 3.44%, respectively. These values surpass the typical records in the literature for liquid alcohol production, which range from 0.005% to 0.186% for STF and 0.37% to 1.5% for AQE.<sup>5,13,37–43</sup>

The exceptional photocatalytic performance of O-Ag@Cu<sub>2</sub>O can be further demonstrated by the time-dependent generation of ethanol and methanol (Fig. 3d). Under dark conditions, no discernible products are observed. However, upon exposure to visible light irradiation, the yields of ethanol and methanol progressively increase with prolonged reaction time, with ethanol exhibiting a notably higher yield compared to methanol. This nearly linear growth trend suggests the good stability of O-Ag@Cu<sub>2</sub>O, which can be further supported by recycling experiments. Notably, after three cycles, O-Ag@Cu<sub>2</sub>O retains approximately 80% of its initial activity (Fig. 3e), which is commensurate with the enhanced stability reported for Cu<sub>2</sub>O-based catalysts in previous studies.<sup>5,13,44–46</sup> Moreover, a comprehensive post-reaction characterization analysis of O-Ag@Cu<sub>2</sub>O reveals minimal alterations in its morphology and structure (Fig. S22†), providing preliminary evidence of its commendable stability.

To further investigate the long-term stability of O-Ag@Cu<sub>2</sub>O, the PCRR was extended to 16 hours under continuous operation (Fig. S23a†). As the reaction progresses, the production of ethanol and methanol continues to increase, indicative of the catalyst's sustained effectiveness, albeit with a gradually diminishing growth rate. This slowdown in production growth may be attributed to two primary factors. Firstly, while the primary Cu-containing component in O-Ag@Cu<sub>2</sub>O remains Cu<sub>2</sub>O after the 16 hours reaction, a portion of its surface is oxidized to Cu<sup>II</sup> by its photogenerated holes (Fig. S23b–e†),<sup>47</sup> leading to a slight decrease in photocatalytic performance. Secondly, the generated methanol and ethanol may undergo re-

oxidation by the photogenerated holes of Cu<sub>2</sub>O,<sup>48</sup> thereby reducing their net yield. The re-oxidation of methanol and ethanol allows them to act as sacrificial agents for hole consumption during the PCRR, thereby partially suppressing the self-oxidation of Cu<sub>2</sub>O in O-Ag@Cu<sub>2</sub>O. Overall, despite experiencing some oxidation, O-Ag@Cu<sub>2</sub>O exhibits commendable stability.

To further confirm that CO<sub>2</sub> is the carbon source for ethanol production, an isotope labeling experiment using <sup>13</sup>CO<sub>2</sub> was conducted. The gas chromatography-mass spectrometry (GC-MS) analysis of ethanol reveals a series of peaks at mass-to-charge ratios (*m/z*) ranging from 43 to 48 (Fig. S24a†), providing qualitative evidence for the formation of <sup>13</sup>CH<sub>3</sub><sup>13</sup>CH<sub>2</sub>OH. Specifically, the *m/z* values of 47 and 48 correspond to the fragment ion of <sup>13</sup>C<sub>2</sub>H<sub>4</sub>O<sup>+</sup> and the molecular ion of <sup>13</sup>C<sub>2</sub>H<sub>5</sub>OH<sup>+</sup>, respectively.<sup>49</sup> Additionally, the most intense peak at *m/z* = 32 is attributed to the cleavage of the C-C bond, forming <sup>13</sup>CH<sub>2</sub>OH.<sup>49</sup> Combined with the control experiments (Fig. S24b†), it can be concluded that the PCRR process is driven by light, with CO<sub>2</sub> serving as the carbon source. To further ensure the accuracy of ethanol selectivity, detailed analysis of the CO<sub>2</sub> reduction products was conducted using <sup>1</sup>H nuclear magnetic resonance (NMR) spectroscopy. This analysis aimed to identify any other carbon-containing compounds, such as formic acid. The result indicates that ethanol and methanol are the only carbon-containing products detected (Fig. S24c†).

Additionally, the PCRR conducted using only CO<sub>2</sub> and H<sub>2</sub>O as the reactants confirms the concurrent production of oxygen (O<sub>2</sub>). The band structure deduced from valence-band spectra (Fig. S25a†) and UV-vis diffuse reflectance spectroscopy (Fig. S15b†) provides evidence that the designed Cu<sub>2</sub>O-based catalyst is thermodynamically suitable for H<sub>2</sub>O oxidation reaction (Fig. S25b†). Furthermore, the oxidation of H<sub>2</sub>O to O<sub>2</sub> involves the transfer of four holes (2H<sub>2</sub>O + 4h<sup>+</sup> → O<sub>2</sub> + 4H<sup>+</sup>), which are generated within the valence band (VB) of the material. In Cu<sub>2</sub>O, the VB is derived from the hybridization of Cu 3d and O 2p orbitals.<sup>50</sup> Thus, both Cu and O atoms could theoretically serve as active sites for O<sub>2</sub> production. However, since Cu atoms are primarily recognized as the active sites for CO<sub>2</sub> reduction, it is more likely that O atoms serve as the active sites for O<sub>2</sub> evolution.<sup>48,51</sup> Additionally, the O<sub>2</sub> generation rate of O-Ag@Cu<sub>2</sub>O reaches 375.77 μmol g<sup>-1</sup> h<sup>-1</sup> (Fig. S25c†), which is much higher than that of other catalysts, indicating the excellent photocatalytic performance of O-Ag@Cu<sub>2</sub>O.

### Mechanism study

*In situ* FTIR technique was employed to identify reaction intermediates. To elucidate the differences caused by hydrophobicity, *in situ* FTIR spectra for O-Ag@Cu<sub>2</sub>O and Ag@Cu<sub>2</sub>O were compared (Fig. 4a and b). Furthermore, to investigate the influence of the spatial distribution of Ag particles, *in situ* FTIR spectra for O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O were analyzed (Fig. 4a and c). Notably, the signal at 1545 cm<sup>-1</sup>, corresponding to \*COOH,<sup>52</sup> indicates that CO<sub>2</sub> is chemically adsorbed on the catalyst surface and interacts with protons. Moreover, the signals for monodentate carbonate (m-CO<sub>3</sub><sup>2-</sup>) at 1458, 1508,

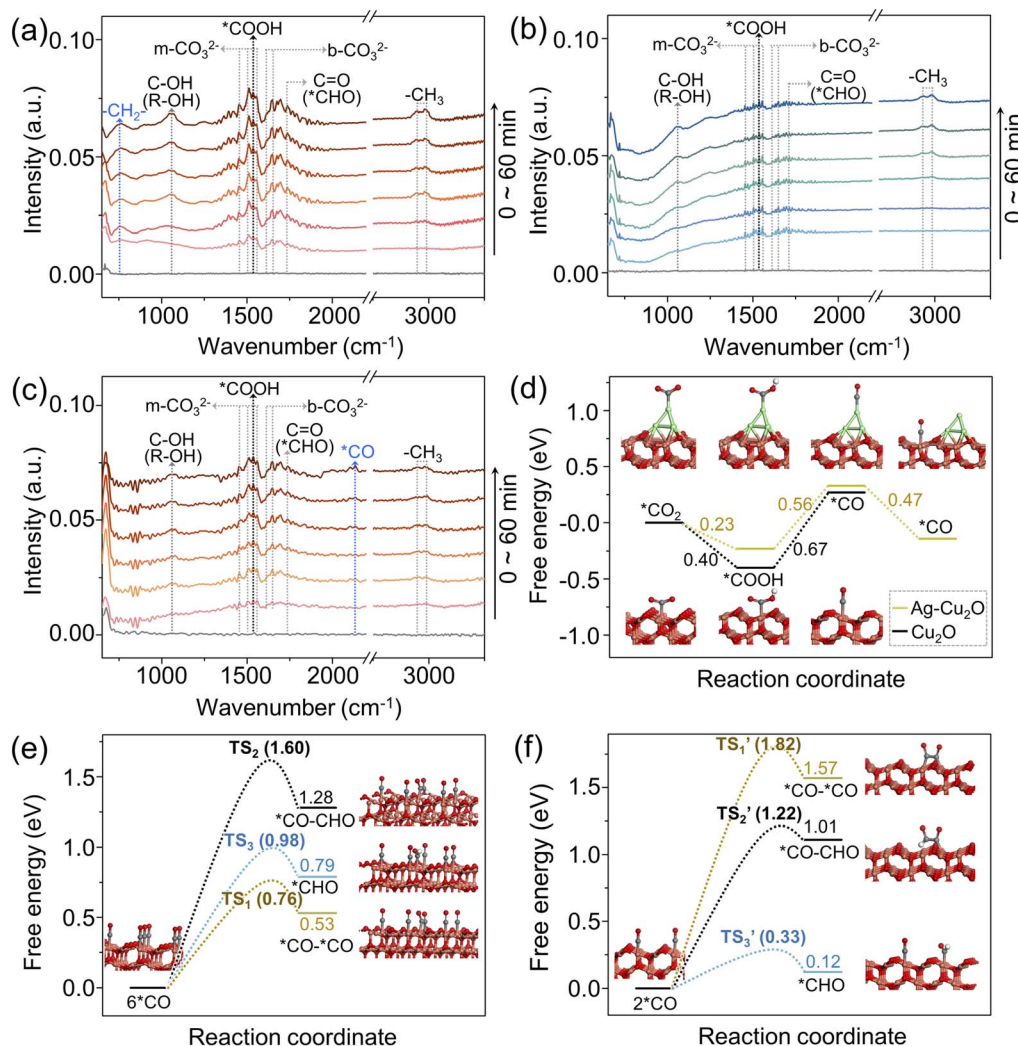


Fig. 4 (a–c) *In situ* FTIR spectra obtained under different irradiation times for O-Ag@Cu<sub>2</sub>O, Ag@Cu<sub>2</sub>O, and O-Ag/Cu<sub>2</sub>O, respectively. (d) Gibbs free energy diagrams for \*CO formation under the models of Ag-Cu<sub>2</sub>O and Cu<sub>2</sub>O. (e and f) Gibbs free energy diagrams for three potential reaction pathways after \*CO formation at elevated and low \*CO concentrations, respectively. The brown, gray, red, and green balls represent Cu, C, O, and Ag atoms, respectively. TS<sub>1</sub>, TS<sub>2</sub>, and TS<sub>3</sub> represent the transition states of \*CO-\*CO, \*CO-CHO, and \*CHO, respectively, at elevated \*CO concentration. TS<sub>1</sub>', TS<sub>2</sub>', and TS<sub>3</sub>' represent the transition states of \*CO-\*CO, \*CO-CHO, and \*CHO, respectively, at low \*CO concentration.

1538, and 1556 cm<sup>-1</sup>, and bidentate bicarbonates (b-CO<sub>3</sub><sup>2-</sup>) at 1620 and 1650 cm<sup>-1</sup>,<sup>53</sup> suggest interactions between the chemically adsorbed CO<sub>2</sub> and H<sub>2</sub>O molecules on the catalyst surface. Therefore, the presence of \*COOH, m-CO<sub>3</sub><sup>2-</sup>, and b-CO<sub>3</sub><sup>2-</sup> signals are indicative of effective CO<sub>2</sub> adsorption.<sup>9</sup> It is evident that the intensity of effective CO<sub>2</sub> adsorption for O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O is significantly higher than that for Ag@Cu<sub>2</sub>O, implying a greater accumulation of CO<sub>2</sub> molecules on the surfaces of O-Ag@Cu<sub>2</sub>O and O-Ag/Cu<sub>2</sub>O compared to Ag@Cu<sub>2</sub>O. This observation further demonstrates the enhanced CO<sub>2</sub> accumulation facilitated by hydrophobic properties. As the reaction progresses, the appearance of the stretching vibration peak of the aldehyde carbonyl group at 1733 cm<sup>-1</sup> implies the formation of \*CHO,<sup>54</sup> which is a crucial intermediate for C<sub>1</sub> compound formation. Moreover, the symmetric and antisymmetric stretching vibration peaks of -CH<sub>3</sub> at 2885 and 2975 cm<sup>-1</sup>,<sup>55</sup> along with the C-OH stretching vibration peak of

alcohols at 1060 cm<sup>-1</sup>,<sup>55</sup> reveal the formation of alcohols across all three catalysts. However, an evident peak at 760 cm<sup>-1</sup>, corresponding to the rocking vibration of -CH<sub>2</sub>-<sup>55</sup> is exclusively observed for O-Ag@Cu<sub>2</sub>O, aligning with the significant ethanol production associated with O-Ag@Cu<sub>2</sub>O. Additionally, for O-Ag/Cu<sub>2</sub>O, a discernible signal at 2134 cm<sup>-1</sup> can be attributed to \*CO,<sup>56</sup> demonstrating the formation of \*CO on the outer surface of O-Ag/Cu<sub>2</sub>O. This finding corroborates the proposed hypothesis that CO generated on Ag particles located on the Cu<sub>2</sub>O outer surface of O-Ag/Cu<sub>2</sub>O can re-adsorb onto the Cu<sub>2</sub>O outer surface. In contrast, CO generated on Ag particles located in the core of O-Ag@Cu<sub>2</sub>O experiences diffusion through the mesoporous channels within the Cu<sub>2</sub>O shell. Due to the prolonged residence time within the pores, \*CO intermediates interact with each other, resulting in a relatively low amount of CO reaching the outer surface, thereby no obvious signal being detected.

To gain a deeper insight into the enhanced ethanol selectivity observed in O-Ag@Cu<sub>2</sub>O, DFT calculations were conducted. For clarity, the CO<sub>2</sub>-to-ethanol conversion process is delineated and analyzed in three stages:<sup>57</sup> initially, the reduction of CO<sub>2</sub> molecules to \*CO; followed by the protonation of \*CO and the C-C coupling process; and finally, the subsequent proton-electron transfer leading to the formation of ethanol. To reveal the working principle of Ag, the model of the Cu<sub>2</sub>O (111) surface with Ag atoms atop (designated as Ag-Cu<sub>2</sub>O) and the model of the bare Cu<sub>2</sub>O (111) surface (designated as Cu<sub>2</sub>O) were compared (Fig. S26†). As shown in the configurations inserted in Fig. 4d, for Ag-Cu<sub>2</sub>O, when a CO<sub>2</sub> molecule is adsorbed onto the Ag atom, it undergoes protonation to form \*COOH, followed by further protonation to produce \*CO. Subsequently, the \*CO desorbs from the Ag atom and re-adsorbs onto the adjacent Cu atom. While for Cu<sub>2</sub>O, the adsorbed CO<sub>2</sub> molecule on the Cu atom is protonated into \*COOH and \*CO in order. The energy barrier of the uphill conversion of \*COOH to \*CO decreases by 0.11 eV over Ag-Cu<sub>2</sub>O, implying that the presence of Ag is conducive to \*CO formation. More importantly, the desorption of \*CO from Ag atoms and its subsequent adsorption onto Cu atoms is an energetically favorable, spontaneous process, suggesting that \*CO generated on Ag can readily re-adsorb onto nearby Cu atoms. Therefore, for O-Ag@Cu<sub>2</sub>O, aided by the hydrophobic property and nanoconfinement effect, the Ag core produces more CO, resulting in a greater accumulation of \*CO within the mesoporous channels of the Cu<sub>2</sub>O shell.

To investigate the impact of \*CO concentration on the Cu<sub>2</sub>O surface on ethanol formation, simulations were performed with six and two \*CO molecules adsorbed on Cu atoms, representing elevated \*CO concentration and low \*CO concentration scenarios, respectively (Fig. 4e and f). After \*CO formation, three potential reaction pathways were considered: the hydrogenation to form \*CHO, the dimerization of \*CO (\*CO-\*CO coupling), and the polymerization of \*CO and \*CHO (\*CO-CHO coupling). To gain deeper insight into the interaction of \*CO in these three different reaction pathways, the transition state energies of \*CO in these processes under different \*CO concentrations were quantified,<sup>58</sup> and the configurations of each transition state are presented (Fig. S27†). As depicted in Fig. 4e, at elevated \*CO concentration, the energy barrier for the transition state (TS<sub>1</sub>) associated with the \*CO-\*CO coupling process is the lowest among the three investigated pathways, with a value of 0.76 eV, reduced by 0.22 and 0.84 eV compared to those of \*CHO formation (TS<sub>3</sub>) and \*CO-CHO coupling (TS<sub>2</sub>), respectively. In contrast, at low \*CO concentration, \*CHO formation is the most favorable, with its transition state (TS<sub>3</sub>) energy barrier being the lowest, at 0.33 eV (Fig. 4f). These findings suggest that \*CO-\*CO coupling is preferred over \*CHO formation at elevated \*CO concentration, whereas \*CHO formation is favored over C-C coupling at lower \*CO concentration. Given that \*CHO is a key intermediate for C<sub>1</sub> compound formation and C-C coupling is essential for C<sub>2</sub> compound formation, increased \*CO concentration enhances ethanol selectivity. Combining these insights with possible CO<sub>2</sub>-to-ethanol and CO<sub>2</sub>-to-methanol conversion pathways proposed based on DFT calculations (Fig. S28†), elevated \*CO

concentration not only promotes ethanol formation but also methanol formation, albeit with a significantly greater effect on ethanol formation. Thus, increasing \*CO concentration enhances both PCRR efficiency and ethanol selectivity, corroborating experimental results.

## Conclusions

In summary, a nanoconfined and tandem three-phase reaction system has been developed to simultaneously address the limited \*CO concentration and inadequate interaction time between C<sub>1</sub> intermediates. Consequently, efficient ethanol production is achieved with a selectivity of 94.15% and a generation rate of 450.19  $\mu\text{mol g}^{-1} \text{h}^{-1}$ . This system is based on a tandem catalyst, O-Ag@Cu<sub>2</sub>O, comprising an Ag core covered by a mesoporous hydrophobic Cu<sub>2</sub>O shell. The hydrophobic Cu<sub>2</sub>O shell can alleviate the poor CO<sub>2</sub> mass transfer, ensuring sufficient CO<sub>2</sub> supply to the Ag core for conversion to CO. Subsequently, CO undergoes continuous reduction within the nanoconfined mesoporous channels of Cu<sub>2</sub>O. This process unfolds in three parts: (i) hydrophobicity guarantees ample CO<sub>2</sub> supply, enabling Ag to generate substantial CO; (ii) nanoconfinement of CO within the mesoporous channels of the Cu<sub>2</sub>O shell leads to its aggregation and prolonged residence time; and (iii) the tandem reaction of CO<sub>2</sub>-to-CO and CO-to-C<sub>2</sub> products is enhanced due to the substantial CO<sub>2</sub> and CO present during this process. The synergy of improved mass transfer, nanoconfinement, and tandem reaction results in high \*CO concentrations at active sites and prolonged interaction time, further promoting C-C coupling and aligning the reaction time with that of C-C coupling. Furthermore, the elevated \*CO concentration makes the energy barrier for \*CO-\*CO coupling much lower than that for \*CO forming \*CHO, thereby prioritizing the occurrence of C-C coupling. This results in outstanding photocatalytic performance for ethanol production, particularly in terms of ethanol selectivity. Additionally, the elevated \*CO concentration enhances the PCRR efficiency, achieving an STF of 0.215%. This strategy presents a promising avenue for the efficient production of C<sub>2+</sub> compounds.

## Data availability

The authors declare that the data supporting the findings of this study are available within the article and its ESI.†

## Author contributions

Conceptualization: Hailing Huo, Ang Li, formal analysis: Hailing Huo, Ting Hu, Chengxi Huang, Ang Li, funding acquisition: Erjun Kan, Ang Li, investigation: Hailing Huo, Zhiqing Zhong, Qiang Ju, Liang Zhang, methodology: Hailing Huo, Ting Hu, Cheng Zhan, Chengxi Huang, Fang Wu, Ang Li, Erjun Kan, supervision: Ang Li, Erjun Kan, validation: Hailing Huo, Zhiqing Zhong, Qiang Ju, Liang Zhang, visualization: Hailing Huo, writing – original draft: Hailing Huo, Ang Li, writing – review & editing: Hailing Huo, Ang Li, Erjun Kan.



## Conflicts of interest

There are no conflicts to declare.

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