



Molecular oxidation-reduction junctions for artificial photosynthetic overall reaction

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Constructing redox semiconductor heterojunction photocatalysts is the most effective and important means to complete the artificial photosynthetic overall reaction (i.e., coupling CO₂ photoreduction and water photo-oxidation reactions). However, multiphase hybridization essence and inhomogeneous junction distribution in these catalysts extremely limit the diverse design and regulation of the modes of photogenerated charge separation and transfer pathways, which are crucial factors to improve photocatalytic performance. Here, we develop molecular oxidation–reduction (OR) junctions assembled with oxidative cluster (PMo₁₂, for water oxidation) and reductive cluster (Ni₅, for CO₂ reduction) in a direct (*d*-OR), alternant (*a*-OR), or symmetric (*s*-OR) manner, respectively, for artificial photosynthesis. Significantly, the transfer direction and path of photogenerated charges between traditional junctions are obviously reformed and enriched in these well-defined crystalline catalysts with monophasic periodic distribution and thus improve the separation efficiency of the electrons and holes. In particular, the charge migration in *s*-OR shows a periodically and continuously opposite mode. It can inhibit the photogenerated charge recombination more effectively and enhance the photocatalytic performance largely when compared with the traditional heterojunction models. Structural analysis and density functional theory calculations disclose that, through adjusting the spatial arrangement of oxidation and reduction clusters, the energy level and population of the orbitals of these OR junctions can be regulated synchronously to further optimize photocatalytic performance. The establishment of molecular OR junctions is a pioneering important discovery for extremely improving the utilization efficiency of photogenerated charges in the artificial photosynthesis overall reaction.

artificial photosynthesis | heterogeneous photocatalyst | photocatalytic CO₂ reduction | polyoxometalate

To cope with the increasingly serious greenhouse effect, exploring green, economic, and sustainable CO₂ conversion and utilization methods to achieve carbon neutrality has become an important research goal of global development (1–7). As we all know, green plants can use solar energy to complete photosynthesis, that is, effectively reducing CO₂ to energy-rich organic carbon products along with oxidizing H₂O to O₂ (Scheme 1*A*) (8, 9). By simulating the synergistic mode of photosystem I (PSI) and PSII in natural photosynthesis, type-II/Z-scheme/S-scheme heterojunction photocatalyst combining two different kinds of semiconductor materials (photocatalyst I [PCI] and PCII), with the characteristics of improved separation efficiency of photogenerated electron hole and strong redox capability, has been proposed to realize the artificial photosynthesis (Scheme 1*B*) (10–13). However, heterojunction materials are difficult to achieve precisely structural adjustment on the oxidative and reductive components from the molecular level. Therefore, it is almost impossible to study the specific effect of the different connection modes and spatial orientations of oxidative and reductive moieties in catalysts systematically on photocatalytic performance. In contrast, inspired by the design concept of heterojunctions, we think that using oxidative and reductive structural motifs to construct periodically distributed molecular junctions, which can essentially create more charge separation and transfer pathways, represents a good choice to achieve the above objective.

In the proposed biomimetic oxidation–reduction (OR) junction, a specific electron transfer mode can occur between the oxidation part and reduction part under light irradiation. Hence, a single molecular OR junction can be regarded as a subminiature heterojunction. It is well known that classic heterojunction catalysts are composed of two or more kinds of semiconductor materials by bulk phase recombination; the oxidative and reductive components are concentrated in their respective regions (14–17). However, the oxidative and reductive catalytic moieties in molecular junctions are spatially periodically arranged. Consequently, the nature of molecular OR catalysts is quite

Significance

Developing photocatalysts that can complete CO₂ reduction and H₂O oxidation reactions simultaneously to realize artificial photosynthesis has great significance for energy and climate issues. Regulating the transfer mode of photogenerated charges in catalysts can greatly improve the efficiency of charge separation and photocatalytic performance. Here, three molecular oxidation–reduction (OR) junctions were constructed by connecting oxidation and reduction clusters in different ways and realized efficient artificial photosynthesis. These OR junctions served as a clear molecular model system to firstly explore the specific effects of different connection modes of oxidation and reduction parts on the migration of photogenerated charges. The specific design schemes of molecular OR junctions can further guide the subsequent design and synthesis of more-efficient catalysts for artificial photosynthesis.

The authors declare no competing interest.

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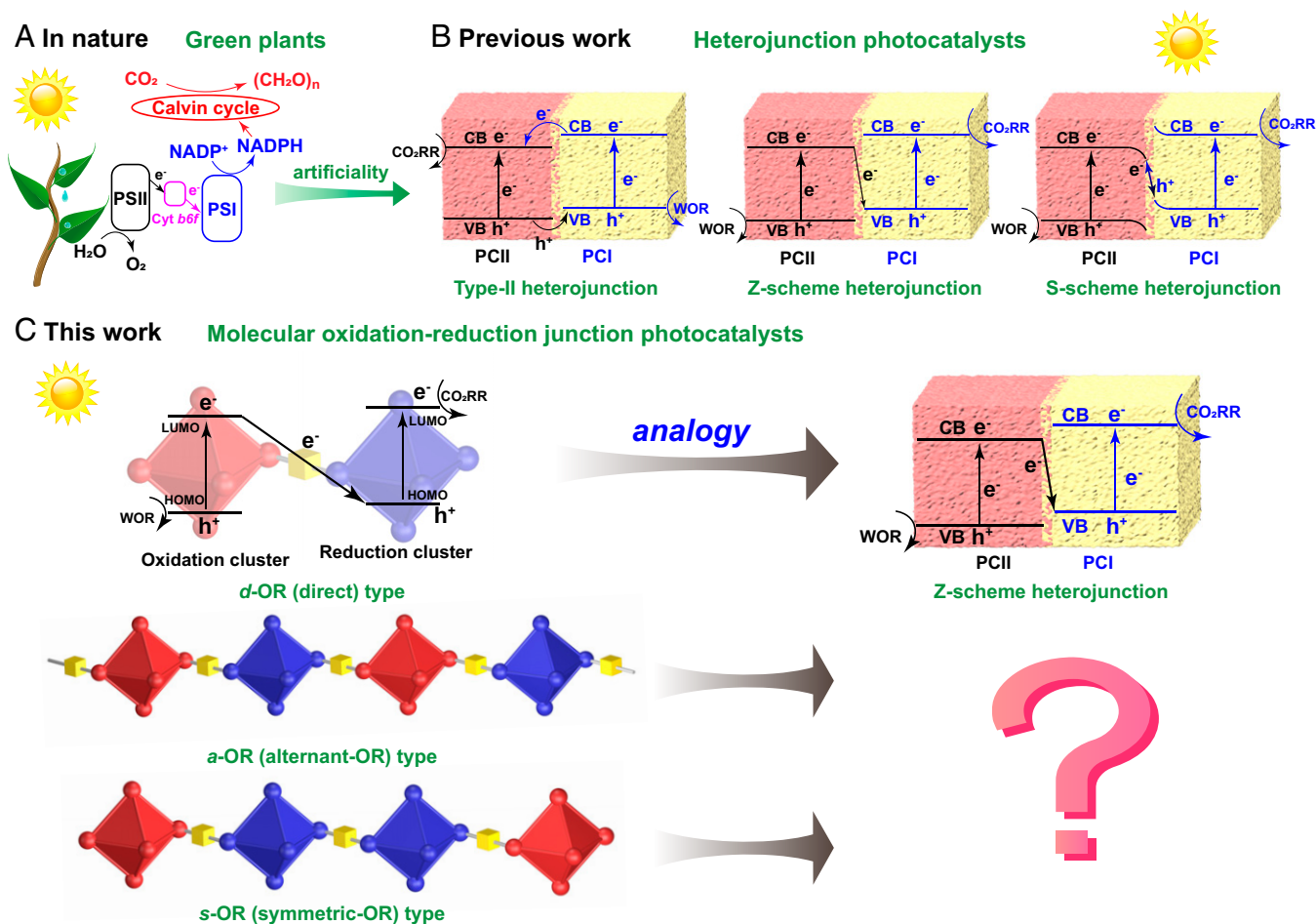
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Scheme 1. Photocatalytic mechanisms in different systems. (A) Operational mechanism of photosynthesis in green plants. (B) Operational mechanisms of artificial photosynthesis in heterojunction catalysts. The mentioned three kinds of heterojunction photocatalysts (type II, Z scheme, and S scheme) are all composed of oxidative part and reductive parts. The reason for their different categories is the distinction in charge transfer between the oxidative and reductive parts. (C) Operational mechanisms of artificial photosynthesis in molecular OR junctions.

different from the traditional heterojunction catalysts. More importantly, oxidation and reduction parts in the lattice of the molecular junctions can be arranged and connected in diversely spatial patterns (e.g., alternating or symmetrical manner), which means that more possibilities of charge separation and transfer pathway can be achieved in the finite or infinite structures of molecular junctions. These cases may also take place in the aggregative heterojunction, but it is difficult to carry out precise structural design and regulation in such a complicated multiphase hybrid system. In this regard, the well-defined crystalline structures and monophase periodic distribution of molecular OR junctions can provide a clear and intuitive research platform. In addition, the small size of the clusters (usually <3 nm) ensures the rapid charge migration between junctions (18).

Based on the above considerations, we designed and constructed three molecular OR junctions, $(\text{PMo}_{12}\text{O}_{40})\{\text{Ni}_5(\text{bzt})_6(\text{H}_2\text{O})_8(\text{C}_2\text{H}_5\text{O})(\text{py})_2(\text{NO}_3)\}$ (**d-OR**, 1H-bzt = 1H-benzotriazole, py = pyrazine), $(\text{PMo}_{12}\text{O}_{40})\{\text{Ni}_5(\text{bzt})_6(\text{H}_2\text{O})_3(\text{C}_2\text{H}_5\text{O})_5(\text{NO}_3)\}$ (**a-OR**), and $\{(\text{PMo}_{12}\text{O}_{40})\text{Ni}_5(\text{bzt})_6(\text{H}_2\text{O})_5(\text{C}_2\text{H}_5\text{O})(\text{py})_2(\text{NO}_3)\}_2(\text{py})$ (**s-OR**), which are assembled with oxidative (O) polyoxometallate $\text{H}_3\text{PMo}_{12}\text{O}_{40}$ (**PMo₁₂**) cluster and reductive (R) $\text{Ni}_5(\text{bzt})_6(\text{NO}_3)_4(\text{H}_2\text{O})_4$ (**Ni₅**) cluster in direct, alternant, and symmetric manners, respectively (Scheme 1 C). According to the energy-level distribution, X-ray photoelectron spectroscopy (XPS), and in situ XPS analysis, we found that the separation and transfer pathway of photogenerated charges between **PMo₁₂** and **Ni₅** clusters in

d-OR is similar to the PCI and PCII systems in Z-scheme heterojunction photocatalyst. Therefore, **d-OR**, **a-OR**, and **s-OR** represent an appropriate molecular model system to explore the specific influence of the different connection modes and spatial orientations of oxidation and reduction parts in catalyst on the photocatalytic performance. As expected, when **d-OR**, **a-OR**, and **s-OR** are used as photocatalysts, they all exhibit superior artificial photosynthesis performance in a mixed atmosphere of CO_2 and water vapor and are more efficient than independent **Ni₅** and **PMo₁₂**. In these three molecular junctions, **PMo₁₂** and **Ni₅** act as oxidative and reductive catalytic sites to convert H_2O and CO_2 into O_2 and CO. The bridging O atoms enable rapid charge transfer between **PMo₁₂** and **Ni₅**. After 10 h of light irradiation, the yield of produced CO is **s-OR** (238.68 $\mu\text{mol/g}$) > **d-OR** (152.70 $\mu\text{mol/g}$) > **a-OR** (112.19 $\mu\text{mol/g}$), indicating that the connection modes between the oxidation part and the reduction part indeed exert an important influence on photocatalytic performance. Through the comparison of the migration of photogenerated electrons and holes, we found that, compared with **d-OR**, the photogenerated charges of **s-OR** can transfer to the lower energy level, thereby decreasing the catalytic efficiency. Furthermore, density functional theory (DFT) calculations proved that the rearrangement of the orbitals leads to a decreased lowest unoccupied molecular orbital (LUMO) energy level of **Ni₅** in **d-OR**, which is more conducive to the occurrence of the catalytic reaction. In **a-OR**, the orbital distribution changes slightly relative to unconnected monomers **Ni₅** and **PMo₁₂**.

When *d*-OR is connected in a symmetric model to form *s*-OR, a new lower vacant orbital will be generated in the center of the *R*-*R* part to promote the photoexcited charge transfer process and inhibit the electron hole recombination of the *Ni*₅ part. This work discloses that the diversified connection modes and spatial arrangements of oxidative and reductive components in catalysts for artificial photosynthetic overall reaction can greatly alter the separation and recombination efficiency of photogenerated charges and then make an impact on the final photocatalytic performance. More importantly, the establishment of molecular OR junctions provides a very important platform for discovering and deeply understanding more migration types of photogenerated charges in photocatalysts.

Results

Structural Design and Fabrication of Photocatalysts. It is very important to select suitable oxidative and reductive clusters to successfully construct OR junction photocatalysts. The following points need to be considered. (1) These two kinds of clusters should have oxidizability and reducibility, respectively. (2) These two kinds of clusters should possess appropriate charge properties (such as anion and cation, respectively), which can make it easy to be connected through chemical bonds and crystallized. Following these ideas, we select a classical polyoxometalate (*PMo*₁₂) as the oxidative cluster unit and a Ni-based cluster (*Ni*₅) as the reductive cluster unit to assemble OR photocatalysts. *Ni*₅ is a compound with a tetrahedral configuration (19). Each metal ion at the vertex of the tetrahedron coordinates with a water molecule and a *NO*₃[−] ion, of which the *NO*₃[−] can be replaced by other anionic groups (20, 21) (*SI Appendix*, Fig. S1).

Three types of OR photocatalysts, *d*-OR, *a*-OR, and *s*-OR, were synthesized by a simple solvothermal method using *PMo*₁₂, *Ni*₅, and pyrazine as the raw materials (Fig. 1*A* and *SI Appendix*, Fig. S2). Single-crystal X-ray diffraction (SCXRD) analysis reveals that *d*-OR assembled directly with *PMo*₁₂ and *Ni*₅ clusters crystallizes in the triclinic space group *P*-1 (*SI Appendix*, Table S1). *Ni*₅ cluster keeps its original host tetrahedral configuration, but one of its original *NO*₃[−] ions is replaced by a terminal O atom from *PMo*₁₂ (Fig. 1*B*). The bridging O atom connecting *PMo*₁₂ and *Ni*₅ in a very short distance of 3.86 Å may allow for rapid photogenerated charge transfer. Intermolecular hydrogen bonds and π - π interactions between ligands make *d*-OR molecules form a three-dimensional supramolecular compound (*SI Appendix*, Fig. S3). *a*-OR crystallizes in the monoclinic space group *P*₂₁/*c* and two formula units (*Z* = 2) per unit cell (*SI Appendix*, Table S1) and is a one-dimensional chain structure by connecting *PMo*₁₂ and *Ni*₅ clusters alternately (Fig. 1*C*). The *a*-OR chains are further extended into a three-dimensional supramolecular array via hydrogen bonds (*SI Appendix*, Fig. S4). It's worth noting that all the vertical Ni ions in *d*-OR and *a*-OR coordinate with at least one solvent molecule that is easy to leave and makes exposed Ni ions as potential open sites for the adsorption and activation of substrates (*SI Appendix*, Figs. S5 and S6). That is to say, the number of catalytic sites is equivalent to the number of vertical Ni ions in *d*-OR and *a*-OR. The *s*-OR type molecule is built from a pyrazine group connecting two mirror-symmetric OR units (*O*-*R*-py-*R*-*O*). SCXRD analysis shows that *s*-OR crystallizes in the triclinic space group *P*-1 and one formula unit (*Z* = 1) per unit cell (*SI Appendix*, Table S1). The part of "OR" in *s*-OR has almost the same configuration as a

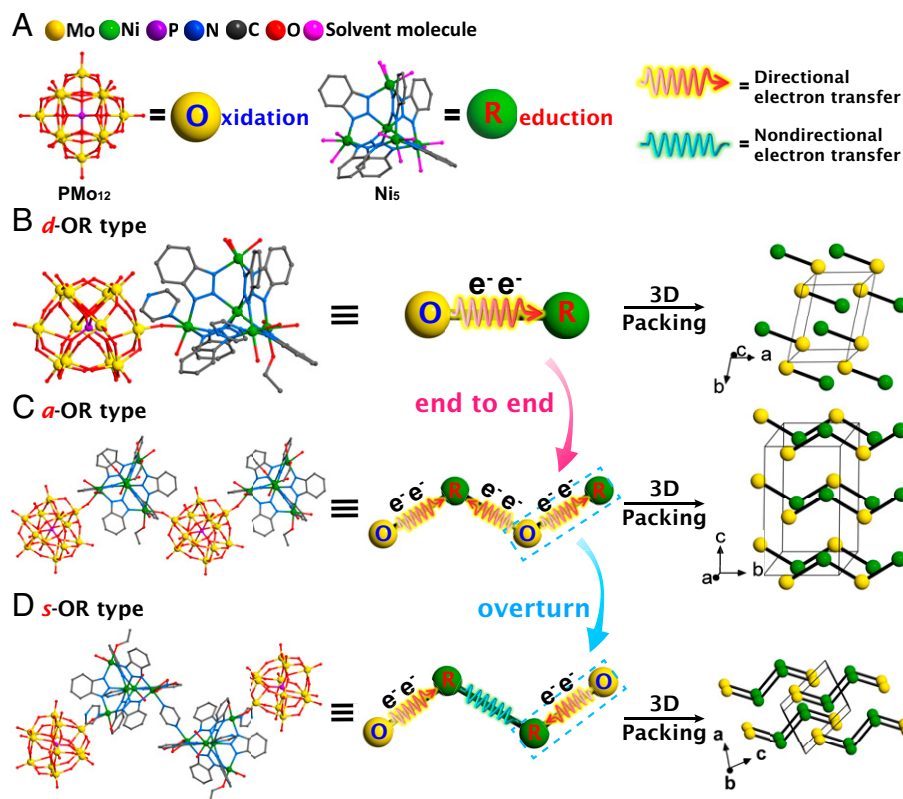


Fig. 1. The structure composition of photocatalysts. (A) Crystal structures and simplified balls of oxidative *PMo*₁₂ cluster and reductive *Ni*₅ cluster. The red arrows represent the directional photoelectron transfer between clusters. The cyan curves represent the nondirectional charge transfer between two clusters. (B–D) Crystal structures determined by SCXRD analysis, simplified structures, and packing diagrams of *d*-OR-, *a*-OR-, and *s*-OR-type photocatalysts. The simplified diagram in the *Middle* shows possible charge transfer between *PMo*₁₂ (O) and *Ni*₅ (R) clusters. All hydrogen atoms are omitted for clarity.

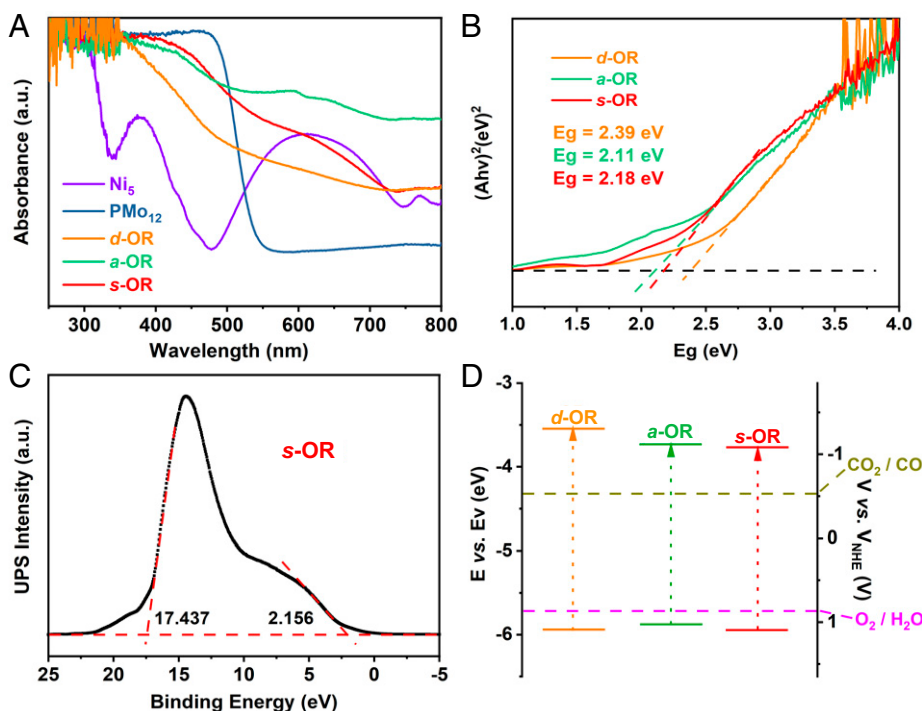


Fig. 2. Characterization of the electronic structures of *d*-OR, *a*-OR, and *s*-OR. (A) UV-Vis absorption spectra of Ni_5 (purple), PMo_{12} (blue), *d*-OR (yellow), *a*-OR (green), and *s*-OR (red) in the range of 250–800 nm. (B) Diffuse reflectance UV-Vis spectra of K–M function versus E_v (eV) of *d*-OR (yellow), *a*-OR (green) and *s*-OR (red). The intersection values of the tangents with the baseline are their corresponding values of band gaps. (C) UPS spectra of *s*-OR. The horizontal dashed red line is the baseline. The other two dashed red lines are the tangent of the curve. The intersection values of the tangents with the baseline are the edges of the UPS spectra. a.u., arbitrary unit. (D) Band structure diagram for *d*-OR (yellow), *a*-OR (green), and *s*-OR (red). All of the three molecular junctions can achieve artificial photosynthetic overall reaction thermodynamically.

single *d*-OR molecule (Fig. 1D). The coordination-saturated Ni linkers (Ni_3 and Ni_5) are encompassed by two O from chelate NO_3^- ions and four N atoms from three bzr and pyrazine ligands, which indicates these Ni ions in *s*-OR cannot serve as catalytic sites (SI Appendix, Fig. S7). The *s*-OR molecules are further stacked into three-dimensional supramolecular structures through multiple hydrogen bonds and π – π interactions (SI Appendix, Fig. S8).

Characterization of Photocatalysts. The purity of *d*-OR, *a*-OR, and *s*-OR was confirmed by the well-matched experimental and simulated powder X-ray diffraction (PXRD) patterns (SI Appendix, Fig. S9). The ultraviolet and visible (UV-Vis) absorption spectra of Ni_5 , PMo_{12} , *d*-OR, *a*-OR, and *s*-OR were characterized to compare their light-harvesting ability. As shown in Fig. 2A, PMo_{12} shows strong absorption in the range of 250–480 nm, while Ni_5 exhibits a broad absorption range of 250–700 nm. It is noted that, when PMo_{12} and Ni_5 were assembled into *d*-OR, *a*-OR, and *s*-OR, their light absorption was throughout the whole range of 250–800 nm. Obviously, these OR photocatalysts show better sunlight utilization than independent PMo_{12} and Ni_5 . The band gap energy (E_g) of *d*-OR, *a*-OR, and *s*-OR were further calculated to be 2.39 eV, 2.11 eV, and 2.18 eV, respectively, through Tauc plots (Fig. 2B). Moreover, UV photoelectron spectrometer (UPS) experiment was carried out to confirm their highest occupied molecular orbital (HOMO) positions (Fig. 2C and SI Appendix, Fig. S12) (22). The obtained HOMO values were -6.30 eV, -5.92 eV, and -5.84 eV (vs. vacuum level) for *d*-OR, *a*-OR, and *s*-OR respectively, which are all below the oxidation level (-5.67 eV vs. vacuum level) for H_2O to O_2 (23). Then the LUMO levels were determined at -3.91 eV (*d*-OR), -3.76 eV (*a*-OR), and -3.53 eV (*s*-OR) that are all above the reduction level for CO_2 -to-CO reduction (-4.34 eV vs. vacuum level) (24). Based

on the above results, the band structures of *d*-OR, *a*-OR, and *s*-OR are properly positioned to thermodynamically achieve CO_2 reduction reaction (CO_2RR) and water oxidation reaction (WOR) simultaneously, corroborating their potential to be catalysts for artificial photosynthetic overall reaction (Fig. 2D).

Artificial Photosynthetic Overall Reaction. Benefiting from the broad range of light absorption and appropriate band structures of *d*-OR, *a*-OR, and *s*-OR, they were utilized as catalysts to perform the photosynthetic overall reaction in a gas–solid system under illumination. The entire catalytic system runs only in the presence of photocatalysts, water vapor, and CO_2 . The results showed that all these photocatalysts can convert CO_2 and H_2O into CO and O_2 . After 10 h of light irradiation, independent Ni_5 and PMo_{12} clusters showed low CO yields of 71.27 $\mu\text{mol/g}$ and 46.95 $\mu\text{mol/g}$, respectively. In sharp contrast, *d*-OR, one of the assemblies composed of Ni_5 and PMo_{12} clusters, revealed a much higher performance, with the CO yield of 202.88 $\mu\text{mol/g}$. It's obvious that the connection of Ni_5 and PMo_{12} through a bridging O atom greatly promotes the efficiency of photocatalysis. Under the identical reaction condition, the CO yield (170.12 $\mu\text{mol/g}$) catalyzed by *a*-OR was slightly lower than *d*-OR, while *s*-OR revealed the highest CO_2 -to-CO reduction performance (271.25 $\mu\text{mol/g}$) among them (Fig. 3A). Moreover, H_2 as the most-common primary competitive product was not detected by gas chromatography (SI Appendix, Figs. S13 and S14).

A series of deletional control experiments were carried out to confirm the whole reaction, of which no photosynthetic products could be detected with the absence of photocatalysts or in the dark (SI Appendix, Table S5). When the reaction system was lacking water or performed under Ar atmosphere, only small amounts of CO were produced (ca. 48 $\mu\text{mol/g}$ for *d*-OR, 28 $\mu\text{mol/g}$ for *a*-OR, and 47 $\mu\text{mol/g}$ for *s*-OR, as entries 3 and

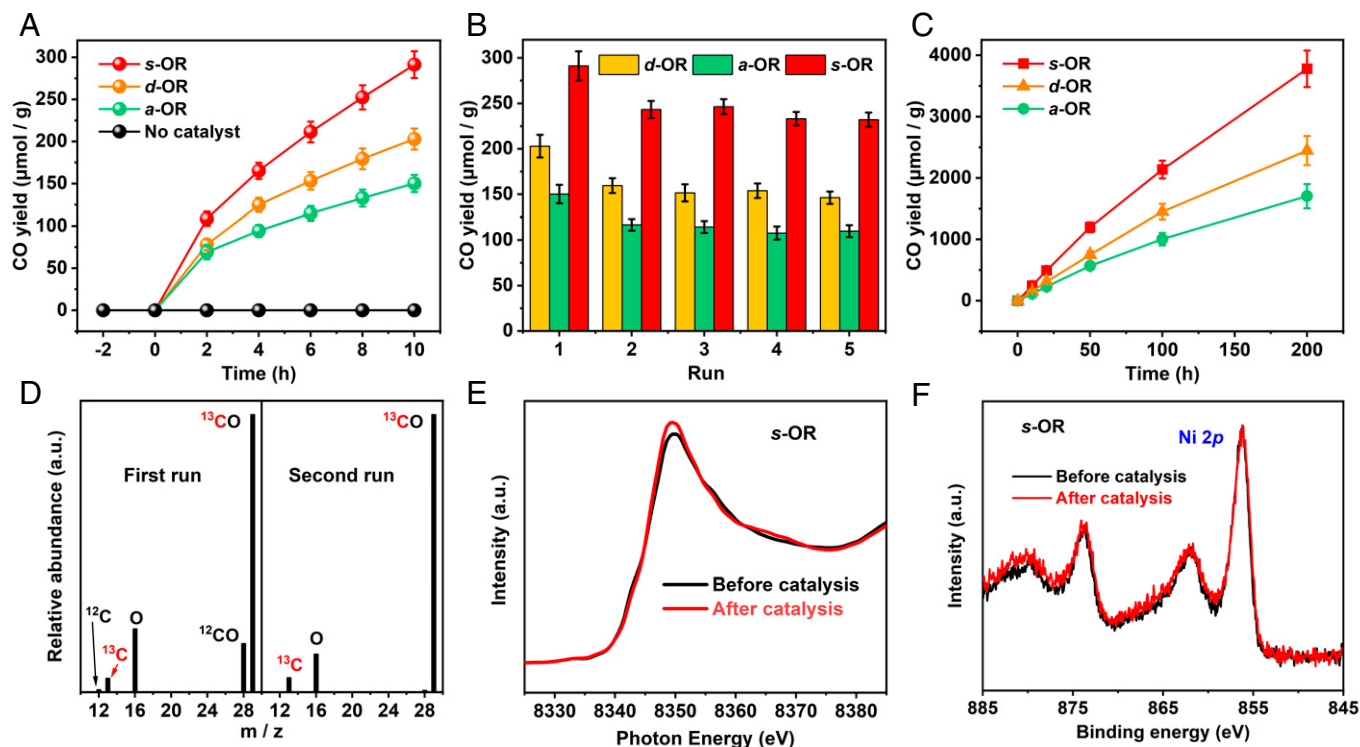


Fig. 3. Photocatalytic CO₂ reduction performance of *d*-OR, *a*-OR, and *s*-OR. (A) The yield of CO catalyzed by *d*-OR, *a*-OR, and *s*-OR. (B) Cycle performance of *d*-OR, *a*-OR, and *s*-OR. (C) Photocatalytic durability of *d*-OR, *a*-OR, and *s*-OR under continual irradiation. (D) Mass spectra extracted from gas chromatography–mass spectrometry analysis of CO product from ¹³CO₂ reduction. The molecular ion peaks appearing at *m/z* peak of 12, 13, 16, 28, and 29 are ascribed to ¹²C, ¹³C, ¹⁶O, ¹²C¹⁶O, and ¹³C¹⁶O, respectively. (E and F) Ni L-edge X-ray absorption near-edge spectroscopy spectra (E) and high-resolution Ni 2p XPS spectra (F) of *s*-OR before and after catalysis.

5 shown in *SI Appendix, Table S5*). It suggested that the photosynthetic system was operated by light, photocatalysts, CO₂, and H₂O together. The durability of photocatalysts was evaluated by refilling CO₂ and water after each 10-h run (see details in *SI Appendix*); it was found that the CO production catalyzed by *d*-OR, *a*-OR, and *s*-OR were dropped to 159.42, 136.78, and 223.23 μmol/g, respectively, on the second run. Then the yields of CO displayed negligible decay on the next three runs, suggesting the outstanding recyclability of these three photocatalysts (Fig. 3B). The catalytic stability was further assessed by keeping the reaction under continuous irradiation. The production of CO increased almost linearly in at least 200 h (Fig. 3C). Combined with the cycling experiments, *d*-OR, *a*-OR, and *s*-OR all revealed excellent catalytic durability. As for WOR, the production of O₂ was evaluated on an online test system. The molar ratio of produced CO and O₂ was close to stoichiometric 2:1, being in line with the theoretical value (*SI Appendix, Fig. S15*). Isotope-tracing experiments were carried out to prove these photocatalysts are indeed active for simultaneous CO₂RR and WOR rather than decomposition of photocatalysts. When using fresh *s*-OR as a representative photocatalyst under water vapor and ¹³CO₂ atmosphere to operate the overall reaction, only a trace amount of CH₄, C₂H₆, and acetaldehyde (CH₃CHO), as well as the main CO and O₂ product, could be detected by gas chromatography–mass spectrometry. By analyzing the *m/z* values of products, CH₄, C₂H₆, and CH₃CHO were not labeled by ¹³C (*SI Appendix, Fig. S16*), and two obvious signals at *m/z* 28 and 29 corresponding to ¹²CO and ¹³CO were shown in the CO peak with the ratio of absolute abundance of 1:5 (Fig. 3C, Left). The content of ¹²CO was close to the yield loss between the first and second runs of the cycling test. The unlabeled products might be derived from the conversion

of ethanol molecules in the catalysts, which caused the decline in CO yield during the first two runs of the cycling experiment. Then we reintroduce ¹³CO₂ into the reactor for the second run of illumination. The result showed that only CO could be detected, and the signal at *m/z* 28 was minuscule (<1% contrasted with *m/z* 29, Fig. 3C, Right). It can be concluded that CO production in the second run was all converted from CO₂. The O source in O₂ was verified by using H₂¹⁸O as the reactant; the obvious *m/z* peak at 36 revealed that the O₂ was indeed converted from H₂O (*SI Appendix, Fig. S17*). After photocatalysis, the unchanged X-ray absorption near-edge spectroscopy and XPS spectra indicated the structural integrity of the used catalysts (Fig. 3D and E and *SI Appendix, Fig. S18*). Obviously, *d*-OR, *a*-OR, and *s*-OR cannot only complete the artificial photocatalytic overall reaction like traditional heterojunction catalysts but can also establish a pioneering system to study the influence of the connection mode and spatial arrangement of oxidation and reduction components on the photocatalytic performance from their clear and periodically distributed structures, which has never been reported before. Therefore, in this work, we call these constructed molecular compound catalysts “molecular OR junctions”. They may be widely used to study the important effects of more-unknown photogenerated charge separation and transfer paths between redox components in catalysts on the photocatalytic performance of the overall reaction.

Discussion

It is well recognized that the efficiency of photocatalysis is mainly affected by light harvesting, charge separation, and charge utilization (25). Because of the similar light absorption and catalytic sites of *d*-OR, *a*-OR, and *s*-OR, the efficiency of

charge separation, in other words, electron hole recombination, may affect their performance of artificial photosynthesis mostly. We notice that the ratio of Ni_5 and PMo_{12} units are all 1:1 in *d*-OR, *a*-OR, and *s*-OR, so the connection modes and spatial orientations of Ni_5 and PMo_{12} clusters in these molecular OR junctions may be the key factors to impact their charge separation efficiency. From the analysis of the band structures of PMo_{12} and Ni_5 (SI Appendix, Figs. S19 and S20), the LUMO and HOMO positions of Ni_5 are all more negative than those of PMo_{12} , proving the oxidative capacity of PMo_{12} and reductive capacity of Ni_5 . To determine which transfer pathway the photogenerated charges between PMo_{12} and Ni_5 parts in these molecular junctions obeys, the direction of the internal electric field needs to be verified. Due to the larger work function of Ni_5 than that of PMo_{12} (26), electrons can transfer from Ni_5 to PMo_{12} when they link together through a coordination bond (SI Appendix, Fig. S21). XPS results provide further evidence for this fact (SI Appendix, Fig. S22). The negative shift in Mo 3*d* and the positive shift in Ni 2*p* of *d*-OR, *a*-OR, and *s*-OR indicate the electron transfer from Ni_5 to PMo_{12} , resulting in the formation of an internal electric field from Ni_5 to PMo_{12} , which is consistent with the inference of UPS. Importantly, high-resolution in situ XPS spectra for Ni 2*p* of *d*-OR were performed. Compared with the dark condition, the

negative shift in the Ni 2*p* binding energy under light irradiation indicates the increased electron density at the Ni sites. The results prove that the migration direction of photogenerated electrons is from PMo_{12} to Ni_5 (SI Appendix, Fig. S23). It follows that, under light irradiation, the photogenerated electrons in the LUMO of PMo_{12} tend to recombine with the photogenerated holes in the HOMO of Ni_5 , leaving photogenerated holes in the HOMO of PMo_{12} and photogenerated electrons in the LUMO of Ni_5 with strong redox capacity to complete WOR and CO_2RR (SI Appendix, Fig. S24) (27). That means PMo_{12} and Ni_5 link together and combine into *d*-OR to form a Z-scheme-like molecule.

When *d*-OR is under illumination, electrons are excited from HOMO levels of Ni_5 and PMo_{12} to their corresponding LUMO positions. The photogenerated electrons of PMo_{12} then transfer to Ni_5 under the internal electric field. Then the reserved electrons in the LUMO of Ni_5 and holes in the HOMO of PMo_{12} participate in the CO_2RR and WOR, respectively (Fig. 4A). The matched band structures and proper charge migration between Ni_5 and PMo_{12} inhibit the recombination of photogenerated electrons and holes and render *d*-OR to reveal better performance than Ni_5 and PMo_{12} themselves (SI Appendix, Fig. S24). The consequence is in accordance with the regulation of Z-scheme nanomaterial photocatalysts

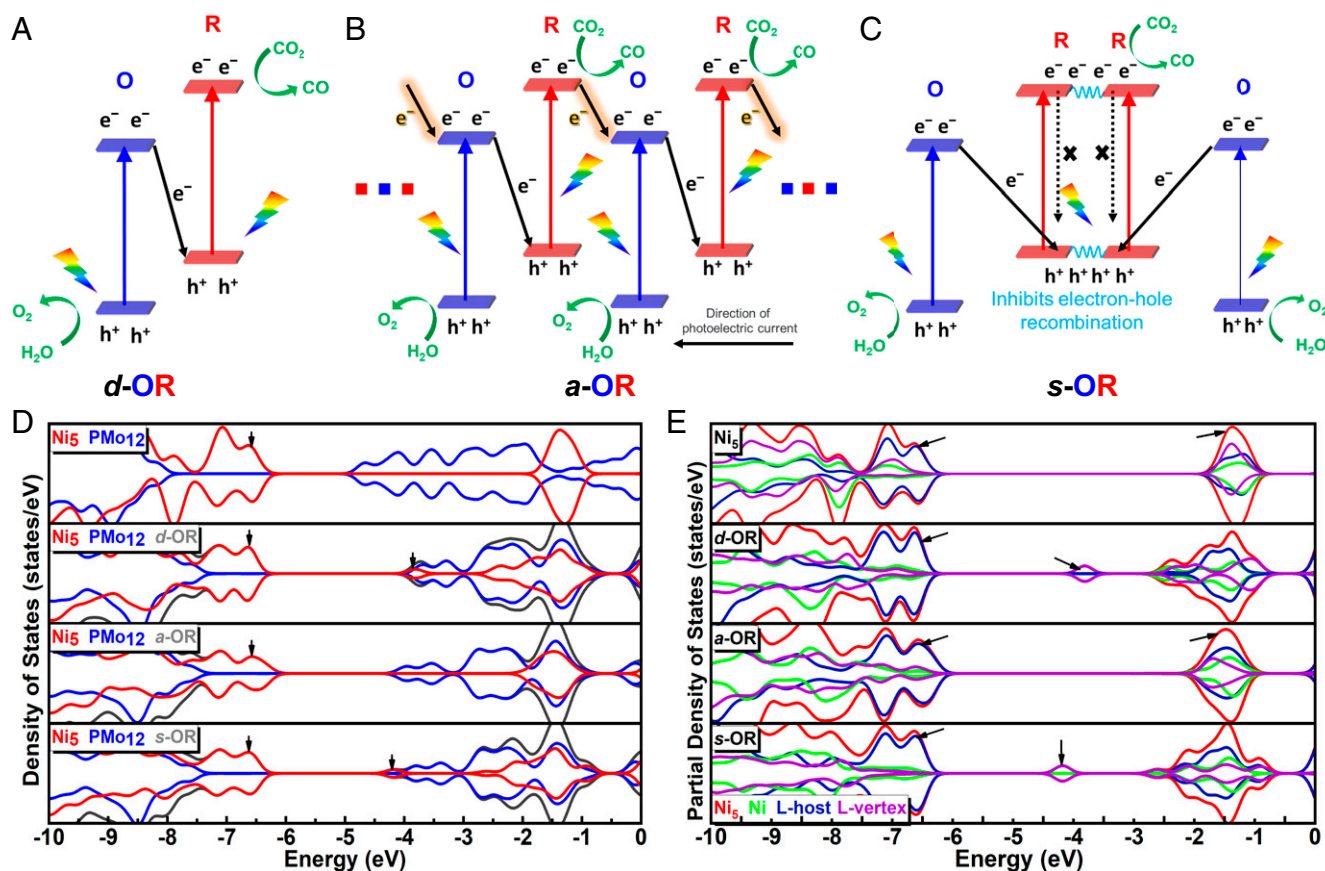


Fig. 4. Proposed charge separation and transfer pathways for *d*-OR, *a*-OR, and *s*-OR photocatalysts under light irradiation and corresponding evidence from total densities of state (TDOSs) and partial densities of state (PDOSs). (A–C) According to the calculated band structure and the results of XPS and in situ XPS analysis, the photogenerated charges of *d*-OR can migrate from PMo_{12} to Ni_5 under light irradiation. When *d*-OR extends to the infinite one-dimensional structure of *a*-OR, the photogenerated electrons can further transfer to the next cluster, which fundamentally weakens the reducing capacity. While *d*-OR changes into *s*-OR, presenting a mirror-symmetric O-R-R-O pattern, the symmetric charge-transfer pathway can extremely inhibit the recombination of photogenerated electron hole pairs, thus achieving the best photocatalytic performance. (D) TDOS and PDOS of isolated PMo_{12} , isolated Ni_5 , and PMo_{12} and Ni_5 moieties of *d*-OR, *a*-OR, and *s*-OR clusters. For the isolated Ni_5 and PMo_{12} , there is almost no orbital overlap and hybridization, and the probability of charge transfer is slight. For the *d*-OR, *a*-OR, and *s*-OR clusters, the unoccupied states' energy levels of Ni_5 moieties of *d*-OR and *s*-OR exhibit a significant decline. (E) PDOS of isolated Ni_5 and Ni_5 moiety of *d*-OR, *a*-OR, and *s*-OR clusters. **Ni** stands for the Ni moiety, **L-host** stands for the benzotriazole moiety, and **L-vertex** stands for the piperazine, H_2O , and NO_3 moieties.

(12, 28). Additional density of state analysis has also supported this point (Fig. 4D). Compared with the isolated Ni_5 , the LUMO levels of the Ni_5 moieties in $d\text{-OR}$ are depressed obviously, resulting in the smaller HOMO–LUMO gaps becoming more easily excited by visible light. And subsequent partial densities of state (PDOS) analysis (Fig. 4E) showed that in the unoccupied states of Ni_5 moiety in $d\text{-OR}$ existed polarization distribution, which further confirmed that the junction mode of $d\text{-OR}$ could inhibit electron hole recombination. When Ni_5 and PMo_{12} are connected alternately to be $a\text{-OR}$ through the coordination bonds, the greater orbital overlap between OR fragments would promote more interaction and charge transfer. However, the delocalized distribution of the unoccupied state similar to the isolated Ni_5 exists in $a\text{-OR}$, in which the local excited states would be produced and be quenched by fast electron hole recombination. On the other hand, when taking $a\text{-OR}$ under light irradiation, following the same regulation as $d\text{-OR}$, the excited electrons of PMo_{12} transferred to the adjacent Ni_5 . Nevertheless, because of its infinite one-dimensional structure, the electrons in the LUMO of Ni_5 can further migrate to the next PMo_{12} (Fig. 4B and SI Appendix, Fig. S25). Because some of the electrons transfer to the energy levels with weaker reducing capacity, it can decrease the catalytic performance of $a\text{-OR}$. As for $s\text{-OR}$, because the OR units are

joined together in a mirror-symmetrical manner, the direction of electron migration is far from the holes. It leads to the efficiency of electron-hole separation in $s\text{-OR}$ being greatly improved (Fig. 4C). Obviously, a stronger polarization distribution of the unoccupied states exists in $s\text{-OR}$, and the lowest unoccupied state is mainly contributed by **L-vertex** ligands (Fig. 4E). And due to the connection mode of Ni clusters in $s\text{-OR}$, the LUMO orbital is mainly distributed on the piperazine and NO_3 ligands of R-R center (SI Appendix, Fig. S26), which further reduces the overlap of HOMO and LUMO orbitals and inhibits the electron hole recombination of inner Ni_5 to improve the catalytic activity of Ni_5 . Equally important, the steady-state photoluminescence (PL) measurements of $d\text{-OR}$, $a\text{-OR}$, and $s\text{-OR}$ were carried out to support the above-mentioned points. The PL emission quenching of $a\text{-OR}$ and $s\text{-OR}$ is more enhanced and declined than $d\text{-OR}$, respectively, indicating the efficiency of electron hole recombination decreases from $a\text{-OR}$, $d\text{-OR}$, to $s\text{-OR}$ (SI Appendix, Fig. S27A) (29, 30). In addition, time-resolved fluorescence decay spectra were performed to evaluate the charge carrier dynamics of $d\text{-OR}$, $a\text{-OR}$, and $s\text{-OR}$ (SI Appendix, Fig. 27B and Table S6). The longest average lifetime of $s\text{-OR}$ implies that the photo-generated charges in $s\text{-OR}$ can survive the longest to participate in the catalytic reaction, in line with the catalytic efficiency.

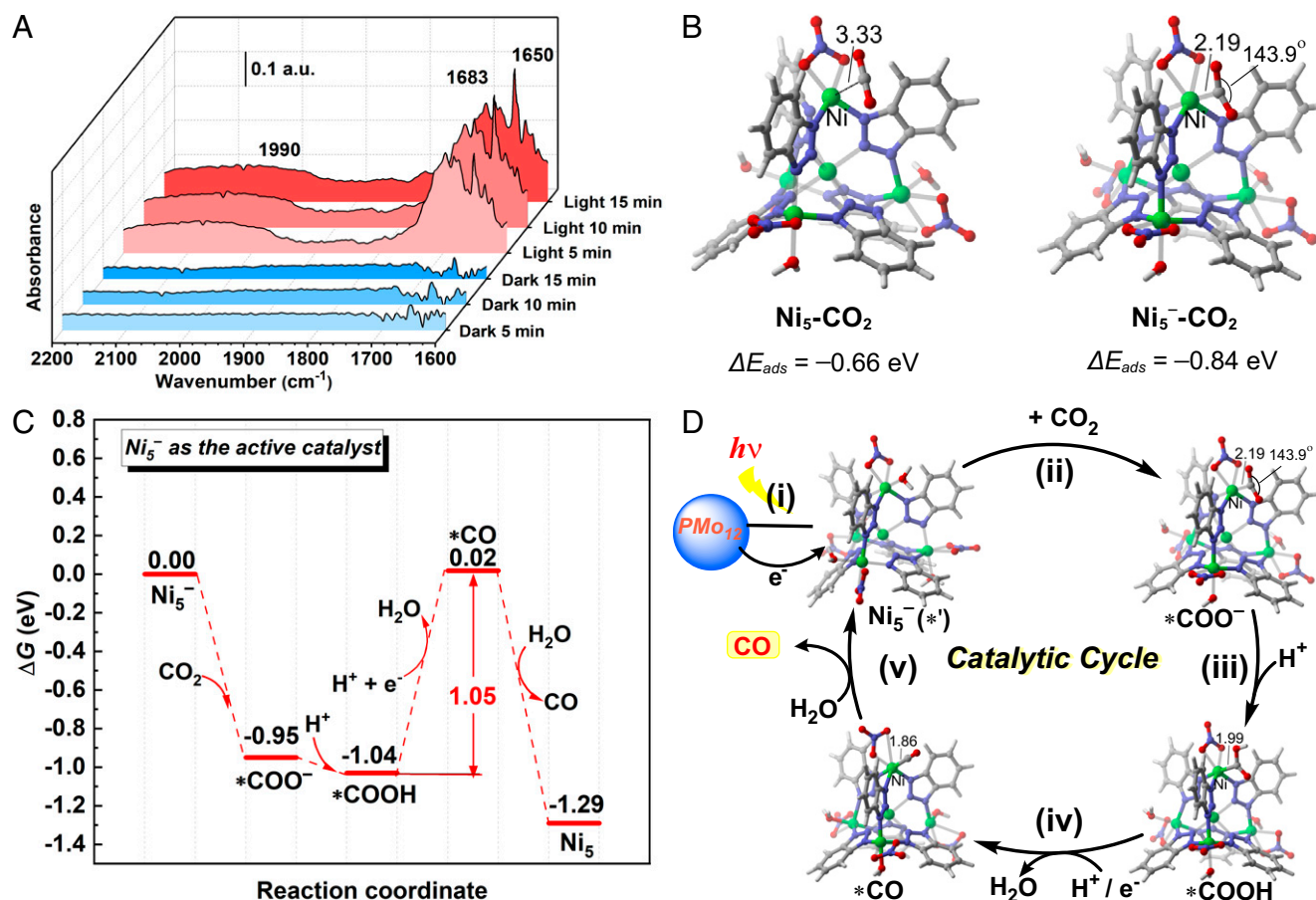


Fig. 5. Study on mechanism of artificial photosynthesis. (A) In situ diffuse reflectance infrared Fourier-transform spectroscopy under dark and light conditions. (B) Optimized structures of the neutral complex (Ni_5) and the charged state (Ni_5^-) after attracting CO_2 ; the bond lengths are given in Å. The carbon dioxide electron adsorption energy (ΔE_{ads}) of Ni_5 is -0.66 eV, and the adsorption energy of reduced state Ni_5^- is -0.84 eV. (C) Gibbs energy profile for the CO_2 reduction process to CO production with reduced state Ni_5^- as the catalytic center. For the intermediate species involved in the catalysis, we also investigated varieties of adsorption possibilities and selected the most-stable structures to evaluate the changes of potential energy surfaces (SI Appendix, Fig. S29). The potential determining step is the CO_2RR process with a barrier of 1.05 eV. (D) The proposed possible mechanism: (i) the electron transfer from PMo_{12} moiety to the Ni_5 moiety by photoexcitation; (ii) the activation of CO_2 to form $^*\text{COO}^-$ species; (iii) the protonation to obtain $^*\text{COOH}$ species; (iv) the proton-coupled electron transfer process to afford the reduced $^*\text{CO}$ species, and (v) the desorption of CO.

In order to capture the intermediate species during the reaction to study the underlying catalytic mechanism, in situ diffuse reflectance infrared Fourier-transform spectroscopy (DRIFTS) was carried out to understand the catalytic process in depth. As shown in Fig. 5A, in a mixed atmosphere of CO₂ and water vapor, there was no additional peak appearing after 15 min in the darkness. In sharp contrast, after 5 min of light irradiation, two peaks at 1,650 and 1,683 cm⁻¹ attributed to Ni-CO₂⁻ and Ni-CO₂H complex rapidly arose (31). The contemporaneous band around 1,990 cm⁻¹ can be assigned to Ni-CO species (32). In order to further understand the reaction mechanism of the CO₂RR process, DFT calculations have been carried out by employing Ni₅ as the model based on the above results (*SI Appendix*, Fig. S28). Firstly, we investigated the effect of electron injection on the CO₂ adsorption of Ni₅. The CO₂ adsorption on the neutral Ni₅ requires the electronic energy (ΔE_{ads}) of -0.66 eV, of which the CO₂ molecule remains almost linear and the C-Ni distance is shortened to 3.33 Å. In contrast, the ΔE_{ads} value of the reduced Ni₅⁻ complex becomes -0.84 eV with a further shortened C-Ni bond distance (2.19 Å) and a slightly curved CO₂ moiety (143.9 degree) (Fig. 5B). The calculation results indicate that the reduced Ni₅⁻ as the active site is more favorable to adsorbed CO₂ forming a carbonate *COO⁻ species than the neutral Ni₅. Therefore, Ni₅⁻ has potential advantages as an active catalyst of CO₂RR. Then, the *COO⁻ captures a proton to afford a stable *COOH (Fig. 5C and D). Such protonation process is exoergic of 0.09 eV. Subsequently, a proton-coupled electron transfer (PCET) process requires a moderate Gibbs free energy change (ΔG°) of 1.05 eV to obtain the *CO species with H₂O releasing, which serves as the potential determining step of the CO₂RR process. Finally, the target CO product is released via a spontaneous ligand exchange process with the ΔG° value of -1.31 eV. In comparison, the neutral Ni₅ complex would undergo two endoergic PCET processes to afford the *CO species, resulting in a large energy barrier of 1.19 eV (*SI Appendix*, Fig. S30). According to the above-mentioned experiment and calculation results, a favorable CO₂RR mechanism catalyzed by the reduced Ni₅⁻ has been illustrated in Fig. 5D.

Conclusion

In summary, we firstly proposed the concept of molecular OR junctions, and that established such three kinds of model molecular OR junctions to complete the artificial photosynthetic overall reaction. Through linking oxidative PMo₁₂ cluster and reductive Ni₅ cluster with a bridge O atom in direct, alternant, and symmetric manners, three molecular junctions, *d*-OR, *a*-OR, and *s*-OR, were successfully obtained. As expected, *d*-OR, *a*-OR, and *s*-OR can all realize outstanding CO₂-to-CO photoconversion (15.9, 13.7, and 22.3 μmol g⁻¹ h⁻¹ for *d*-OR, *a*-OR, and *s*-OR, respectively, >200 h) coupled with photo-oxidating H₂O to O₂, without any assistance of cocatalyst, photosensitizer, or sacrificial agent. Importantly, in virtue of the well-defined and periodic structures of these molecular OR junctions, we investigated the effect of diversified connection modes and spatial arrangements of oxidative and reductive components in overall

reaction catalysts on photocatalytic performance. Combined with the related experimental characterization and detailed DFT theoretical calculations, we find that the R-R part in *s*-OR reduces the overlap of HOMOs and LUMOs and inhibits the recombination of photogenerated electron hole pairs, resulting in the better catalytic activity of *s*-OR than *d*-OR and *a*-OR. Beyond the artificial photosynthetic whole-reaction system, these results may also have implications for other photocatalytic systems, like water splitting or redox organic conversion. Obviously, both of the specific design schemes of molecular OR junctions and the corresponding structure-performance relationship can further guide the subsequent design and synthesis of more-efficient catalysts for artificial photosynthesis.

Materials and Methods

All starting materials, reagents, and solvents used in experiments were commercially available, high-grade purity materials and used without further purification. Thermogravimetric analyses of the samples were performed on a PerkinElmer TG-7 analyzer heated from room temperature to 700 °C in flowing N₂/O₂ with a heating rate of 20 °C/min. Fourier transform infrared spectroscopy (FTIR) measurements were recorded in the range of 4,000–400 cm⁻¹ on a Mattson Alpha-Centauri spectrometer using the technique of pressed KBr pellets. PXRD measurements were recorded ranging from 5 to 50° at room temperature on a D/max 2500 VL/PC diffractometer (Japan) equipped with graphite monochromatized Cu Kα radiation (λ = 1.54060 Å). The UV-Vis-near infrared spectra were performed on a Varian Cary 5000. XPS and UPS were measured on an Escalab 250Xi. PL spectra were recorded by a FluoroMax-4 spectrofluorometer (HORIBA Scientific). In situ DRIFTS was carried out on a Bruker Tensor II FTIR NEXUS. In situ XPS tests were performed on a ThermoFisher Nexsa.

DFT Calculations. All of the geometry optimizations, vibrational frequency evaluations, and intrinsic reaction coordinate (IRC) (33) calculations were performed at the (U)M06 (34) functional with the def2-SVP (35) basis set. More accurate electronic energies on the potential energy surface (PES) were corrected by single-point energy calculations at the (U)M06-D3 (36) functional with the def2-TZVP (37) basis set. The above-mentioned quantum calculations were carried out with the Gaussian 09 (38). The partial density of states (PDOS) were performed for the crystal structure using Multiwfn (39) software with Gaussian broadening function and full width at half maximum (FWHM) of 0.3 eV. The three-dimensional molecular structures were performed by CYLview (40) program, and the relevant molecular orbitals were printed by Visual Molecular Dynamics (41) program.

Data, Materials, and Software Availability. All data are included in the manuscript and/or *SI Appendix*.

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