Supporting Information

Three-Phase Photocatalysis for the Enhanced Selectivity and Activity of CO$_2$ Reduction on a Hydrophobic Surface

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Experimental Procedures

Materials

Acetone, Al₂O₃ basic, 2-bromopropionyl bromide (98%), cyanic acid (98%), ethanol, ethyl acetate (99.8 %), hexane (95%), melamine (99%), N,N,N′,N″-pentamethyldiethylentriamine (PMDETA, ≥98%), 1H,1H,2H,2H-perfluorodecanethiol (pFDe, ≥97%), pyridine (99.8%), silica gel (for column chromatography), tetrahydrofuran (THF, ≥99.9%, inhibitor-free) and lithium hydroxide (LiOH, 99.9%), Palladium(II) acetylacetonate (Pd(acac)₂, 99%), Copper(II) acetylacetonate (Cu(acac)₂, ≥99%) and ¹³C labeled sodium bicarbonate (NaH¹³CO₃, 98 atom% ¹³C) were purchased from Sigma Aldrich. Hydrochloric acid (1 mol L⁻¹) was purchased from Merck KGaA. Platinum(II) bis(acetylacetonate) (Pt(acac)₂) was purchased from Tokyo Chemical Industry UK Ltd. Na₂S (≥98.0) and Na₂SO₃ (≥97.0) were purchased from Tianjin Fuyu Fine Chemical Co. Ltd. KHCO₃ (99.5%) was purchased from Macklin. Deionized water (H₂O, 18.25 MΩ-cm) supplied by an UP Water Purification System was used in the whole experimental processes. Copper (I) bromide (Cu(I)Br, 98%, Sigma Aldrich) was purified by stirring in glacial acetic acid overnight, filtered, and washing with absolute ethanol. Glycidyl methacrylate (GMA, 97%, Sigma Aldrich) was passed through a basic alumina column prior to use. All other chemicals were obtained from commercial suppliers and used without further purification.

Characterization

Crystalline structures were evaluated by X-ray diffraction (XRD) analysis using a Bruker D8 Advance instrument with Cu Kα radiation. The Transmission Electron Microscope (TEM) is a double-corrected Jeol ARM200F, equipped with a cold field emission gun. The acceleration voltage was set to 200kV and the emission was put to 10 µA. Room temperature photoluminescence (PL) spectra were monitored at a Perkin Elmer Luminescence Spectrometer (LS 50 B) with excitation wavelength of 350 nm. The specific surface areas were calculated from the isotherms using the BET method. The pore distribution and the cumulative volumes of pores were obtained by the BJH method from the desorption branch of the nitrogen adsorption isotherms. Inductively Coupled Plasma Optical Emission Spectrometer (ICP-OES) was performed on a Vista MPX ICP. The infrared (IR) spectrum was recorded by HIDEN QIC-20 mass spectrometer. The optical properties of the powders were examined by a Shimadzu UV 2600 instrument. Here we use the absorbance pattern to detect the light absorption within the whole range of UV and visible light. For solid samples, the light absorbance is detected based on diffuse reflection light. BaSO₄ is used as the reference sample. The reflection light when BaSO₄ is used is considered to have an absorbance of 0, which is used as the reference light. The absorbance is the part of light less than reference light. The value of the absorbance is calculated by the built-in program of the instrument.

Methods

Section 1. Synthesis of polymeric carbon nitride (PCN). The carbon nitride used in present work was synthesized via cyanic acid and melamine as precursors. A 1:1 molar ratio of cyanic acid and melamine were mixed in water and shaken for 24 hours at ambient temperature. Then the obtained milky suspension was centrifuged and washed with water three times, and dried at 50 °C under vacuum. Afterwards the precursor powder was annealed in capped crucibles at 550 °C for 4 h at a heating rate of 2.3 °C/min in nitrogen atmosphere. After cooling down the yellow carbon nitride powder was collected without any further treatment.[1]

Section 2. Modification the surface to synthesis hydrophobic PCN (o-PCN). The o-PCN was obtained through 3 steps.[2] (1) Synthesis of ene-PGMA. To obtain PGMA with double bond end modification, an ene-initiator (undecenyl 2-bromopropionate) was synthesized firstly according to the literature.[3] Then GMA was polymerized using atom transfer reaction polymerization (ATRP) with Cu(I)Br as catalyst, PMDETA as ligand and ene-initiator. Specifically, Cu (I) Br (15 mg) was added to a 25 mL round bottom flask in an ice bath, the flask was sealed with a rubber septum and flushed with Argon for 30 min. Subsequently, deoxygenated GMA (5.4 g) and PMDETA (17 mg) were added via syringe. After the solution turned light green, the ene-initiator (64 mg) was added via syringe. The mixture was stirred at 25 °C for 30 min. Thus, PGMA with ene-functionalization was prepared (ene-PGMA, Mₙ=25.9 kg/mol). (2) Grafting of ene-PGMA onto PCN was realized by mixing 300 mg ene-PGMA with 30 mg PCN in 15 mL THF in a round bottom flask, the flask was sealed with a rubber septum and flushed with Argon for 30 min. The mixture was put between two 50 W LED daylight sources (20 cm distance between light sources) to initiate the grafting. After 48 h reaction, the mixture was washed 2 times with THF and dried under vacuum, to obtain ene-PGMA/PCN. (3) Hydrophobic modification of ene-PGMA/PCN. In a 15 mL glass vial, 2 mL THF, 7 mg of ene-PGMA/PCN and 300 µL pFDe were mixed and cooled with an ice bath. 3 mg LiOH was added slowly, the cooling was removed and the resulting reaction mixture was stirred at ambient temperature for 1 hour. Afterwards, the mixture was diluted with THF and washed 3 times with 3 mL THF and centrifuged. The precipitate was dried under vacuum to obtain o-PCN.

Section 3. Pt loading on PCN to synthesize Pt/PCN. 0.3 g PCN was dispersed in an aqueous solution containing 20 mL H₂O and 10 mL ethanol under sonication. Then 40 µL H₂PtCl₆ solution (the concentration of the H₂PtCl₆ solution is 8 wt%) was added with sonication. After stirring for 30 min, the mixture was moved to a sealed transparent container which was set under a pressure of 0.13 bar. Subsequently, the mixture under 0.13 bar was irradiated under visible light supplied by the visible light source for 18 h. Subsequently, the as-prepared sample was separated by centrifugation and washed three times with H₂O. Then the samples were vacuum dried at 80 °C overnight to finally form Pt/PCN as a powder.
Section 4. Pt loading on o-PCN to synthesize Pt/o-PCN. 0.3 g o-PCN was dispersed in the mixture of 20 mL acetone and 10 mL ethanol under sonication. Then 0.03 g Pt(acac)2 was added to the mixture under sonication. After stirring for 30 min, the mixture was moved to a sealed transparent container which was set under a pressure of 0.13 bar. Subsequently, the mixture under 0.13 bar was irradiated under visible light supplied by the visible light source for 18 h. Then the as-prepared sample was separated by centrifugation and washed three times with acetone. Then the samples were vacuum dried at 80 °C overnight to finally form Pt/o-PCN as a powder.

Section 5. Cu and Pd loading on o-PCN to synthesize Cu/o-PCN and Pd/o-PCN. 0.3 g o-PCN was firstly dispersed in the mixture of 20 mL acetone and 10 mL ethanol under sonication. Then 0.020 g Cu(acac)2 or 0.023 g Pd(acac)2 was added to the mixture under sonication. After stirring for 30 min, the mixture was moved to a sealed transparent container which was set under a pressure of 0.13 bar. Subsequently, the mixture under 0.13 bar was irradiated under visible light supplied by the visible light source for 18 h. Then the as-prepared sample was separated by centrifugation and washed three times with acetone. Then the samples were vacuum dried at 80 °C overnight to finally form Cu/o-PCN or Pd/o-PCN as a powder. The TEM and HRTEM images of Cu/o-PCN and Pd/o-PCN are shown in Figure S9, in which the Cu or Pd particles loaded on o-PCN can be clearly observed, and the lattice spacing can prove the composition.

Section 6. Photocatalytic CO2 reduction. The home-made photocatalytic reactor consisted of a sealed chamber, an embedded window made by quartz glass and a liquid sampling port sealed by silicone pad. The reactor was connected to a gas circulation system with a ten-port valve (VICI) for on-line sampling to a gas chromatograph (Ruimin GC 2060, Shanghai). The gas circulation system was primarily made of stainless steel tubing and a home-made gas pump for the circulation of the gas. A mechanical pump was connected into the system to exhaust the carrier gas of the gas chromatograph when switch back the ten-port value. A pressure gauge was also connected into the system to monitor the gas pressure. The total volume of the gas in the circulation system after filling the reactor with solvent was 80 mL.

At the start of this reaction, 0.01 g catalysts were mixed with 40 mL KHCO3 (0.1 M), Na2S (0.1 M) and Na2SO3 (0.1 M) aqueous solution in the reactor. The solution was then placed in a water bath at 80 °C for 15 min under stirring. While stirring, NaOH was added to the solution to adjust the pH value to 11. After that, the gas was switched into an 110 W xenon lamp which was placed into the reactor for 0.5 h. After that, pure CO2 gas was introduced into the reactor for 1 h. Then the gas circulation system was started to circulate the gas at a flow rate of 200 ml/min. The gas was detected by TCD and FID with higher sensitivities. The role of visible light was introduced to the reaction space by the irradiation of a 300 W xenon lamp. The light was introduced to the reaction space by a quartz window placed into the reactor. The selected visible light wavelength was longer than 400 nm (λ > 400 nm). After the reaction of 1 h, the gas and liquid products was detected by the analysis system.

The analysis of the gaseous reaction mixtures containing CO, CH4, H2, was carried out using a gas chromatograph, which was equipped with a TCD, FID and a methanizer which contained a Ni catalyst. Argon (≥ 99.9999%) was used as the carrier gas. The back channel of GC was equipped with two packed columns, TDX-01 and Molsieve 5 Å, and two gas switch valves. During the analysis, 1.0 mL of gas sample in the sample loop of ten-port value was introduced to the TDX-01 column which was separated from the other gases due to its longer retention time. The rest of the gases after the TDX-01 column was further separated by the Molsieve 5 Å column. The gas product of H2 was detected by TCD and CH4, CO were further detected by FID with higher sensitivities. The role of methanizer was to convert CO to CH4 for FID analysis. The liquid product of CH3OH was injected into HP-5 capillary column (30 m × 0.32 mm ID × 0.5 μm, Agilent) and detected by FID after the reaction. No CH3OH can be detected in this work. Other liquid products such as HCOOH and HCHO possibly formed during the reaction were also analyzed by high-performance liquid chromatography (HPLC 7080B, Agilent Co., USA), but no such products were detected under our reaction conditions. Cycling test was operated using the same method, and each cycle continued 3 h. After each cycle, the catalysts were separated by centrifugation, dried under vacuum at 80 °C for 6 h and then reused in the next cycle.

Section 7. Calculation of selectivity of carbon derivatives. The selectivity of CO2 reduction to carbon derivatives (CO, CH4) on an electron beam (8 eV for CH4, 2 eV for CO and H2) has been calculated using the following equation:[7]

\[
\text{Selectivity of carbon derivatives (\%) = \left( \frac{[\text{CH}_4]}{[\text{CO}] + [\text{CH}_4]} \right) \times 100\%}
\]

Where n(CO), n(CH4) and n(H2) are the numbers (moles) of CO, CH4, and H2 formed within a certain period of time.

Section 8. in situ IR test. A diffuse reflection in situ IR was used to investigate the effective CO2 adsorption on the surface. Samples are loaded in the sample holder with a flat surface. A cover was fixed on the sample holder to form a reaction space, which was then cleaned by argon. Then the IR data was collected as background. Subsequently, visible light was introduced to the reaction space by observation window, and CO2 was introduced through container filled with water which make the gas bring water into the reaction space. Then IR data was collected every 20 min, and the change based on IR can be recorded.
Section 9. Stability of the polymer-modified surface during visible irradiation and during the reaction. The polymer modified surface of o-PCN is very stable under visible light irradiation and during the photoreduction, which can be proved by several evidences. (1) As shown in inset of Figure S7, after the photoreaction under visible light, the contact angle of water on o-PCN is 91.5°, which is very close to that before photoreaction (89.7°), almost remaining unchanged. Because the hydrophobic properties of o-PCN are provided by the polymer, it can be deduced that the polymers remain stable under visible irradiation and during the photoreduction. (2) It has been explained in details in section 6 of methods and Figure S7 that the carbon source of carbon derivatives is CO₂, which means the carbon skeleton of polymer did not react during photoreaction. (3) As described in section 2 of methods, the polymer was connected to PCN by visible light with 48 h, and after that the hydrophobicity of surface was greatly improved. It can be deduced that if the polymer is unstable under visible light, it will be decomposed totally in 48 h and would not improve the hydrophobicity of the surface.

Section 10. The calculation of turnover number (TON) over Pt/o-PCN. The turnover number (TON) was calculated using the following equation:^[6]

\[
\text{TON} = \frac{\text{moles of products evolved}}{\text{moles of active components on photocatalyst}}
\]

Moles of products evolved: In this work, only CO and CH₄ can be detected as the products of CO₂ reduction. We detected the amount of both CO and CH₄ base on the method described in section 6 of methods in the SI. The Pt/o-PCN can remain relatively stable during 9 h. After the reaction of 9 h with 0.01 g Pt/o-PCN, the amounts of CO and CH₄ were detected to be 12.751 and 6.594 μmol, respectively.

Moles of active components on photocatalyst: the loading amount of Pt was determined to be 2.61 wt% by the test of ICP. Thus, the moles of active components is \( (0.01 \text{ g} \times 2.61\%) / 195.1 \text{ g mol}^{-1} = 1.338 \times 10^{-6} \text{ mol} = 1.338 \mu\text{mol} \).

Thus, TON = (12.751 + 6.594) μmol / 1.338 μmol = 14.458. The TON value is obviously more than 1, reflecting the catalytic nature of the reaction.^[9]

Section 11. Investigation of carbon source with isotope labeled CO₂. The reaction system of the isotope experiment is the sealed circulatory system mentioned in section 6 of methods in SI. 1 g NaHCO₃ (98 atom% ¹³C, Sigma Aldrich) was added into 32.2 mL Na₂S (0.1 M) and Na₂SO₃ (0.1 M) aqueous solution in the reactor to provide H¹³CO₃⁻. Then after adding 0.01 g Pt/o-PCN, the system was sealed. The system was evacuated to remove air firstly, and then the suspension in the reactor was purged with argon for 1 h to remove air. Then 7.8 mL hydrochloric acid (1 M) was injected through the liquid sampling port with silica gel pad for the in situ generation of ¹³CO₂ via the reaction H⁺ + H¹³CO₃⁻ = H₂O + ¹³CO₂. According to the stoichiometric ratio, the amount of hydrochloric acid added can ensure the remaining of 4 mmol H¹³CO₂ in these 40 mL mixture, making the concentration of H¹³CO₂ to be 0.1 M. Then after a circulation for 1 h, visible light was provided for the photocatalytic reaction. After the reaction of 5 h, the gas sample was detected by a gas chromatography-mass spectrometer (GC-MS) instrument.

The gas chromatograph (GC) is the same one that we used to detect reaction activity (described in section 6 of methods in SI) with the columns of TDX-01 and Molsieve 5 Å. The difference is that the gas products separated by GC was introduced into a mass spectrometer (HIDEN QIC-20) instead of Flame Ionization Detector (FID). We monitored the characteristic mass-charge ratios (m/z) of 28, 29, 16 and 17 to detect the ¹²CO, ¹³CO, ¹²CH₄ and ¹³CH₄ in the product, respectively, and results are shown in Figure S6. According to the results of GC recorded by FID in the former experiment when detecting activity (section 6 of methods in SI), the retention time of CO and CH₄ is 4.990 and 9.234 min, respectively. Thus, because the same GC was used for GC-MS, the signal appeared in similar retention recorded by MS were the time is reasonable to be recognized as the signal of CO and CH₄. The retention time of carbon derivatives with ¹²C is slightly larger than that with ¹³C, which also have been observed by other researchers.[10] Figure S6b shows that ¹²CO can be detected when ¹³CO₂ and H¹³CO₃⁻ are provided, and ¹³CO can be detected when ¹²CO₂ and H¹³CO₃⁻ are used as the carbon source.[11] Figure S6c shows that ¹²CH₄ can be detected when ¹²CO₂ and H¹⁵CO₃⁻ are provided, and ¹³CH₄ can be detected when ¹³CO₂ and H¹³CO₃⁻ are used as the carbon source.[12] These results proved that the photocatalytic products indeed originate from CO₂ reduction.[13]

Section 12. The action spectra of photocatalytic CO₂ reduction reaction (CRR). The action spectra is a graph of the rate of an activity plotted against the wavelength of light.[13] To detect the activity of CRR with Pt/o-PCN under the irradiations with different wavelength, we use a similar method which was described in section 6 of methods in the SI. The only difference is the incident light (λ > 400 nm) to monochromatic light with a specific wavelengths, which were provided by a 300 W xenon lamp (Beijing PerfectLight Co.) and various bandpass filters (Beijing China Education Au-light Co., Ltd.). The CRR activities under monochromatic irradiations with wavelengths of 365, 380, 420, 435, 525 and 700 nm are listed in Table S1, According to which we can get the action spectra (Figure S8).

Section 13. Apparent quantum efficiency (AQE) for photocatalytic CRR. The AQE is defined by the equation:[13]

\[
\Phi_a = \frac{\int (d[x]/dt) \, dt}{\int [hv]^{inc} \, dt}
\]

(2)
where $\frac{d[x]}{dt}$ is the rate of change of the concentration of the reactant (or product) and $\frac{d[hv]}{dt}$ is the total optical power impinging on the sample. Generally, for the convenience of measurement and calculation, researchers usually use the integral form of equation (2):[14]

$$\text{AQE}(\%) = \frac{\text{number of the electrons taking part in reaction}}{\text{number of incident photons}} \times 100\% \quad (3)$$

The number of electrons taking part in the reaction is calculated according to the action spectra, which was described in section 12 of methods in the SI. In this work, the products of CRR are only CO and CH$_4$. Thus, the AQE of CRR should be calculated according to the action spectra of CO and CH$_4$. Normally, two electrons are needed to get one molecule of CO, and eight electrons are needed to get one molecule of CH$_4$. Thus, equation (3) can be transformed into:[14b]

$$\text{AQE}(\%) = 2 \times \frac{\text{number of evolved CO molecules} + 8 \times \text{number of evolved CH}_4\text{ molecules}}{\text{number of incident photons}} \times 100\% \quad (4)$$

The numerator and denominator of equation (4) are divided by unit time simultaneously, we can get:[15]

$$\text{AQE}(\%) = \frac{(2 \times \text{number of evolved CO molecules} + 8 \times \text{number of evolved CH}_4\text{ molecules})}{\text{unit time}} \times 100\% \quad (5)$$

We denote the numerator of equation (5) as $N_e$, and the denominator of equation (5) as $N_p$. Thus, $N_e$ means the amounts of electrons taking part in CRR during the unit time, and $N_p$ means the amounts of incident photons during the unit time, and equation (5) can be written as:[15]

$$\text{AQE}(\%) = \frac{N_e}{N_p} \times 100\% \quad (6)$$

Thus, we can calculate the AQE as long as we can get the value of $N_e$ and $N_p$ under the incident light with a specific wavelength. Here we demonstrate the process with the example of AQE calculation under irradiation with the wavelength of 420 nm.

$N_e$: The amount of CO and CH$_4$ generated by unit catalyst at the unit time was detected under the irradiation with the wavelength of 420 nm (as described in section 12 of methods in the SI). As shown in Table S1, the generation rate of CO and CH$_4$ under 420 nm is 0.531 and 0.229 μmol h$^{-1}$, which can be transferred to be $1.475 \times 10^{-10}$ and $6.361 \times 10^{-11}$ mol s$^{-1}$, respectively. Thus, $N_e = 2 \times 1.475 \times 10^{-10} + 8 \times 6.361 \times 10^{-11} = 8.039 \times 10^{-10}$ mol s$^{-1}$.

$N_p$: The energy of single photon (denoted as $E_s$) at $\lambda = 420$ nm can be calculated to be $4.736 \times 10^{-19}$ J according to $E_s = (hc)/\lambda$, where $h$ and $c$ are the Planck constant ($6.626 \times 10^{-34}$ J·s) and the speed of light ($2.998 \times 10^8$ m s$^{-1}$), respectively. The light intensity (denoted as $I_o$) was measured to be 0.35 mW cm$^{-2}$ by an irradiatometer (Photoelectric Instrument Factory of Beijing Normal University). The diameter (denoted as $d$) of illumination window of the reactor was measured to be 5 cm, and thus the irradiation area (denoted as $S_o$) can be calculated to be $19.63$ cm$^2$ according to the equation $S_o = \pi \times (d/2)^2$. Thus, the $N_p$ can be calculated by divide the total energy provided in unit time by the energy of single photon: $N_p = (I_o \times S_o) / E_s = 2.408 \times 10^{-08}$ mol s$^{-1}$.[15]

Thus, according to equation (6), AQE (%) = $N_e / N_p \times 100\% = 8.039 \times 10^{-10}$ mol s$^{-1} / 2.408 \times 10^{-08}$ mol s$^{-1} \times 100\% = 3.337\%$.

In addition, the AQE of CRR at $\lambda = 365, 380, 435, 525$ and 700 nm can be calculated by the same process. Results are listed in Table S1, according to which we can get the AQE spectra as shown in Figure S8.
Figure S1. Nitrogen adsorption isotherms and pore size distribution (inset) of a) bulk carbon nitride and b) Pt/o-PCN.

Figure S2. The mechanism of in situ photo-loading method to synthesis Pt/o-PCN and Pt/PCN. a) The progress of photo-loading to load Pt particles on o-PCN. S represents the hole sacrificial agent, and S⁺ represents the sacrificial agent which are already oxidized by holes. Here ethanol is used as the sacrificial agent. b) The formation of Pt/o-PCN. c) The progress of photo-loading to load Pt particles on PCN. S represents the hole sacrificial agent, and S⁺ represents the sacrificial agent which are already oxidized by holes. Here ethanol is used as the sacrificial agent. d) The formation of Pt/PCN.
**Figure S3.** a) TEM images of Pt/PCN. b) Pt particle size statistics of Pt/PCN based on image a. Average particle size: 2.18 nm. c) TEM images of Pt/o-PCN. d) Pt particle size statistics of Pt/o-PCN based on image c. Average particle size: 2.21 nm. Pt particles taking part in statistics are selected randomly.

**Figure S4.** Nitrogen atoms in different chemical surroundings with different bonding mode. A: C-N=C. B: N-(C)₃. C: C-N-H.
Figure S5. Stability along with time and recycling test of Pt/o-PCN.

Figure S6. Signal of GC-FID and GC-MS. a) The raw data of the signal of GC-FID which was used to investigation the activity using the method described in section 6 of methods in the SI. b) The GC-MS signal monitored by m/z = 28 and m/z = 29 for the detection of $^{12}$CO and $^{13}$CO, respectively. c) The GC-MS signal monitored by m/z = 16 and m/z = 17 for the detection of $^{12}$CH$_4$ and $^{13}$CH$_4$, respectively.
Figure S7. To investigate the carbon source of CRR with o-PCN, control experiment was operated by replacing CO\textsubscript{2} to N\textsubscript{2}. Result shown that without CO\textsubscript{2} there are almost no carbon derivatives can be detected. A few carbon derivatives in products can be attribute to the KHCO\textsubscript{3} solution, which is described in detail in section 6 of methods. Insert: Photographs of contact angles of water on o-PCN before and after photoreaction.

Figure S8. The action spectra and AQE spectra with the catalysts of Pt/o-PCN. This figure is constructed according to the data shown in Table S1. Under the irradiation with different wavelengths, the ration of CO and CH\textsubscript{4} is variable. The highest AQE within the visible region is 3.337, which is achieved under the irradiation of 420 nm.

Figure S9. The TEM and high-resolution TEM (HRTEM) images HRTEM images of Cu/o-PCN and Pd/o-PCN and their CO\textsubscript{2} photoreduction activities. a) The TEM images of Cu/o-PCN. Insert: HRTEM image of Cu particles. The lattice spacing is 0.212 nm, which is in good agreement with the (111) planes of Cu. b) the TEM images of Pd/o-PCN. Insert: HRTEM image of Pd particles. The lattice spacing is 0.138 nm, which is in good agreement with the (220) planes of Pd. c) the CO\textsubscript{2} reduction activities and selectivity over Cu/o-PCN and Pd/o-PCN, and the comparison with Pt/o-PCN. Only CO and CH\textsubscript{4} can be detected as the products of CO\textsubscript{2} reduction.
Figure S10. The quantitative analysis of effective mass transfer. Because the adsorption of CO$_2$ as the form of COOH$^*$ is a result of the effective mass transfer of gas phase to the surface, the rate of increase of the main IR signal peaks of COOH$^*$ may be used as the quantitative parameter to reflect the effective mass transfer. The main IR peak of COOH$^*$ is the one at the wavenumber of 1728 cm$^{-1}$ (Figure 4d). The average increasing rates can be calculated based on the slope of fitting lines, which determined to be 7.040 × 10$^{-3}$ min$^{-1}$ and 1.859 × 10$^{-3}$ min$^{-1}$ for Pt/o-PCN and Pt/PCN, respectively.

Table S1. The data of action spectra and AQE under incident light with different wavelengths.[4]

| Wavelength / nm | Generation rate / µmol h$^{-1}$ | N$_{p}$[3] / mol s$^{-1}$ | E$_{p}$[3] / J | I$_{p}$[3] / mW cm$^{-2}$ | N$_{p}$[3] / mol s$^{-1}$ | AQE$^{[3]}$ / % |
|-----------------|---------------------------------|--------------------------|-------------|-------------------------|--------------------------|-----------------|
| 365             | 0.539                           | 0.291                    | 9.461 × 10$^{-12}$ | 5.449 × 10$^{-12}$ | 0.54                      | 3.229 × 10$^{-08}$ | 2.930          |
| 380             | 0.674                           | 0.225                    | 8.744 × 10$^{-12}$ | 5.234 × 10$^{-12}$ | 0.41                      | 2.553 × 10$^{-08}$ | 3.426          |
| 420             | 0.531                           | 0.229                    | 8.039 × 10$^{-12}$ | 4.736 × 10$^{-12}$ | 0.36                      | 2.408 × 10$^{-08}$ | 3.337          |
| 435             | 0.124                           | 0.051                    | 1.822 × 10$^{-10}$ | 4.572 × 10$^{-10}$ | 0.29                      | 2.067 × 10$^{-08}$ | 0.882          |
| 525             | 0.002                           | 0                       | 1.111 × 10$^{-12}$ | 3.789 × 10$^{-12}$ | 0.38                      | 3.269 × 10$^{-08}$ | 0.003          |
| 700             | 0.003                           | 0                       | 1.666 × 10$^{-12}$ | 2.841 × 10$^{-12}$ | 0.19                      | 2.179 × 10$^{-08}$ | 0.008          |

[a] the action spectra and AQE are investigated using the target catalyst Pt/o-PCN. Incident light of different wavelengths was provided by a 300 W xenon lamp (Beijing PerfectLight Co.) and various bandpass filters (Beijing China Education Au-light Co., Ltd). [b] N$_{p}$ means the amounts of electrons taking part in CRR during the unit time. [c] E$_{p}$ means the energy of a single photon, which is calculated according to E$_{p}$ = (hc) / λ, in which the h, c and λ are the Planck constant, the speed of light and the wavelength of the photon. [d] I$_{p}$ means the intensity of incident light, which is measured by an irradiatiometer (Photoelectric Instrument Factory of Beijing Normal University). [e] N$_{p}$ means the amounts of incident photons during the unit time. [f] AQE was calculated according to the equation with integral form AQE = N$_{p}$/ N$_{0}$. 

References

[1] M. Shalom, S. Inal, C. Fettkenhauer, D. Neher, M. Antonietti, J. Am. Chem. Soc. 2013, 135, 7118-7121.
[2] Q. Cao, B. Kumru, M. Antonietti, B. V. K. J. Schmidt, Macromolecules 2019, DOI: 10.1021/acs.macromol.10b00894.
[3] a) K. Matyjaszewski, P. J. Miller, N. Shukla, B. Immaraporn, A. Geiman, B. B. Luokala, T. M. Siclovan, G. Kicikelbick, T. Vallant, H. Hoffmann, T. Pakula, Macromolecules 1999, 32, 8716-8724; b) S. De, A. Khan, Chem. Commun. 2012, 48, 3130-3132.
[4] S. Zhu, B. Jiang, W. B. Cai, M. Shao, J. Am. Chem. Soc. 2017, 139, 15664-15667.
[5] M. Dunwell, G. Lu, J. M. Heyes, J. Rosen, J. G. Chen, Y. Yan, F. Xiao, B. Xu, J. Am. Chem. Soc. 2017, 139, 3774-3783.
[6] J. L. White, M. F. Barudth, J. E. Pander H., Y. Hu, I. C. Fortmeyer, J. E. Park, T. Zhang, K. Liao, J. Gu, Y. Yan, T. W. Shaw, E. Abelev, A. B. Bocarsly, Chem. Rev. 2015, 115, 12888-12935.
[7] a) S. Neau, J. A. Macia-Aguilo, P. Concepcion, H. Garcia, J. Am. Chem. Soc. 2014, 136, 15969-15978; b) Q. Zhai, S. Xie, W. Fan, Q. Zhang, Y. Wang, W. Deng, Y. Wang, Angew. Chem. Int. Ed. 2013, 52, 5776-5779; c) S. Xie, Y. Wang, Q. Zhang, W. Deng, Y. Wang, ACS Catal. 2014, 4, 3644-3653.
[8] S. Wan, M. Xu, X. Wang, Y. Wang, Y. Zeng, J. Ding, S. Zhang, Q. Zhong, Dalton Trans. 2019, DOI: 10.1039/C1039DT02507C.
[9] a) S. Wang, Z. Ding, X. Wang, Chem. Commun. 2015, 51, 1517-1519; b) T. Takayama, H. Nakanishi, M. Matsu, I. Iwase, A. Kudo, J. Photoch. Photobiol. A 2018, 352, 416-421.
[10] P. Xia, M. Antonietti, B. Zhu, T. Heil, J. Yu, S. Cao, Adv. Funct. Mater. 2019, 29, 1900093.
[11] M. Jiang, Y. Gao, Z. Wang, Z. Ding, Appl. Catal. B: Environ. 2016, 198, 160-188.
[12] S. Bajpai, G. Bhat, Plant Physiol. 1970, 46, 1-5.
[13] a) J. M. Buriak, P. V. Kamat, K. S. Schanze, ACS Appl. Mater. Inter. 2014, 6, 11815-11816; b) M. Liang, T. Borjigin, Y. Zhang, H. Liu, B. Liu, H. Guo, ACS Appl. Mater. Inter. 2018, 10, 34123-34131.
[14] a) W. Dai, H. Xu, J. Yu, X. Hu, X. Luo, X. Tu, L. Yang, Appl. Surf. Sci. 2015, 356, 173-180; b) L. Ye, D. Wu, K. H. Chu, B. Wang, H. Xie, H. Y. Yip, P. K. Wong, Chem. Eng. J. 2016, 304, 376-383.
[15] A. Li, X. Chang, Z. Huang, C. Li, Y. Wei, L. Zhang, T. Wang, J. Gong, Angew. Chem. Int. Ed. 2016, 55, 13734-13738.