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Copper-doped perylene diimide supramolecules combined with TiO₂ for efficient photoactivity†

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Designing organic–inorganic hybrid semiconductors is an effective strategy for improving the performance of the photocatalyst under visible light irradiation. In this experiment, we firstly introduced Cu into perylenediimide supramolecules (PDIs) to prepare the novel Cu-doped PDIsm (CuPDIsm) with one-dimensional structure and then incorporated CuPDIsm with TiO₂ to improve the photocatalytic performance. The introduction of Cu in PDIsm increases both the visible light adsorption and specific surface areas. Cu²⁺ coordination link between adjacent perylenediimide (PDI) moleculars and H-type π – π stacking of the aromatic core greatly accelerate the electron transfer in CuPDIsm system. Moreover, the photo-induced electrons generated by CuPDIsm migrate to TiO₂ nanoparticles through hydrogen bond and electronic coupling at the TiO₂/CuPDIsm heterojunction, which further accelerates the electron transfer and the separation efficiency of the charge carriers. So, the TiO₂/CuPDIsm composites exhibit excellent photodegradation activity under visible light irradiation, reaching the maximum values of 89.87 and 97.26% toward tetracycline and methylene blue, respectively. This study provides new prospects for the development of metal-doping organic systems and the construction of inorganic–organic heterojunctions, which can effectively enhance the electron transfer and improve the photocatalytic performance.

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1. Introduction

Photocatalytic technology is environmentally friendly for organic pollutants degradation, hydrogen evolution and photo-oxidation synthesis under sunlight irradiation.^{1,2} It is well known that semiconductor photocatalysts play a key role to improve the photocatalytic efficiency, implying that the pursuit of high-quality photocatalysts remains a research priority in the field of photocatalysis.^{3–5}

Perylene diimide (PDI) organic materials and their derivatives, often synthesized by solution method under N₂ or Ar inert atmosphere conditions, have many merits, such as good electron-accepting character,^{6,7} high mobility,⁸ wide light response wavelength⁹ and outstanding photochemical stability,¹⁰ so they are the potential candidates for photocatalytic applications.^{11–13} PDI as a n-type organic semiconductor can be used as an independent catalyst in photocatalytic process, but its photocatalytic efficiency is low due to its electron-deficient π -conjugated cores, rapid charge recombination and inefficient charge transport molecular system.^{14,15} To solve these problems, some effective strategies

were employed to enhance the photocatalytic activity, for instance, designing PDI molecule,¹⁶ adjusting supramolecular configuration,^{6,17} coupling with other photocatalysts or cocatalysts,^{18–21} etc.

In fact, incorporating PDI assemblies with other photocatalysts or cocatalysts is the usual method, which constructs powerful photocatalytic systems with improved photoinduced electron transfer. For example, PDI/g-C₃N₄ showed the enhanced electron delocalization effect and the interfacial charge transfer with the formation of n–n heterojunction system.¹⁰ PDI coupling with WO₃ (ref. 22), BiOCl²³ and Ag₃PO₄ (ref. 24) had high photoactivity arisen from the efficient carriers separation and the formation of Z-scheme heterojunction. PDI/TiO₂ composites showed the improved photoactivity due to the electrons transporting from PDI to the conduction band of TiO₂.^{15,25,26} PDI/Pd composites had high electron transfer efficiency due to two opposite types of electron transfer pathway.⁷ PTCDI self-assembled with C₆₀ lowered the position of valence band, narrowed the band gap and promoted the transfer of the photogenerated carriers.²⁷ In PDI/Graphene system, graphene quantum dots (GQDs) contributed to the electrons delocalization *via* π – π action with PDI and promoted electron transfer from GQDs to PDI.²⁸

Recently, it is found that the doping of Zn and Co can facilitate fascinating modification of PDI supramolecules through π – π stacking of the organized PDI rings and the metal

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bridging between the side chains of adjacent PDI molecules.^{29,30} The synergistic effects between the PDI arrays and metal sites provide the promising way for developing ideal photocatalysts toward the better utilization of solar energy.³¹ However, to our knowledge, the incorporation of the metals into PDI supramolecules except Zn and Co have not been involved and how to introduce more types of metals into PDI assemblies remain a great challenge.

In this work, we firstly designed a simple solvothermal method to prepare PDI, different from the conventional synthesis method of PDI under N₂ or Ar inert atmosphere conditions. To keep the consecutive and rapid photo-induced electron transfer process, we prepared Cu-doping PDI supramolecular (CuPDIsM) by the self-assembly of PDI molecules in copper salt solution and then combined CuPDIsM with TiO₂ nanoparticles by solvothermal synthesis method, aiming for designing a novel CuPDIsM/TiO₂ photocatalysts with low-cost and excellent visible-light photocatalytic performance. Furthermore, we systematically elucidated the effect of Cu-bridging and TiO₂ coupling on the electron transfer process and the photocatalytic activity of the PDI supramolecules.

2. Experimental

2.1 Materials

Perylene-3,4,9,10-tetracarboxylic dianhydride, glycine, imidazole, tetracycline (TC), methylene blue (MB), and methyl orange (MO) were purchased from Aladdin. Triethylamine, hydrochloric acid, copper nitrate trihydrate, tetrabutyl titanate, glacial acetic acid, sodium hydroxide and *N,N*-dimethylformamide (DMF) were purchased from Sinopharm Chemical Reagent Co. All reagents were analytically pure and could be used without further purification.

2.2 Synthesis of crude PDI

Different from the conventional preparing method of PDI under N₂ or Ar inert atmosphere conditions, we have designed a solvothermal method to prepare PDI, which has many merits like convenient operation, good effect and low cost. Firstly, 1.4 g of perylene-3,4,9,10-tetracarboxylic dianhydride and 2.6 g of glycine were well mixed and then 25 g of imidazole was added as the solvent. The obtained mixture was put in a reactor and heated at 110 °C for 5 h. After cooling, 50 mL of anhydrous ethanol and 80 mL of 3 M hydrochloric acid were sequentially added with continually stirring. Finally, the precipitate was collected through centrifugation and washed with ethanol and deionized water for 3 times, respectively. After being freeze-dried, crude PDI powder with bright red color was obtained.

2.3 Synthesis of PDIsM and CuPDIsM

200 mL of deionized water, 0.834 mL of triethylamine and 1 mL of 2 M NaOH solution were uniformly mixed. Then 0.54 g of crude PDI product was added with continually stirring for 1 h until PDI powders were fully dissolved and subsequently 27.3 mL of 4 M hydrochloric acid was added with rapidly stirring. Soon, flocculent precipitates were collected with

centrifugation and washed with ethanol to get the dark red PDIsM products.

400 mg of crude PDI sample, 300 mg of Cu(NO₃)₂·3H₂O and 480 mg of NaOH were homogeneously mixed, then 40 mL of DMF and 110 mL of deionized water were added with stirring. The mixture was put into a reactor and heated at 110 °C for 3 days. After cooling, centrifugation, washing and freeze-drying, CuPDIsM powders were obtained.

2.4 Synthesis of TiO₂/PDIsM and TiO₂/CuPDIsM

Firstly, 10 mL of tetrabutyl titanate was thoroughly mixed with 40 mL of anhydrous ethanol, followed by the addition of 4 mL of glacial acetic acid. Then 10 mL of ethanol solution ($V_{\text{water}}/V_{\text{ethanol}} = 2:3$) was slowly added with continuous stirring at 40 °C for 1 h to obtain TiO₂ precursor solution. Subsequently, 1 wt% PDIsM or CuPDIsM suspensions were mixed with the TiO₂ precursor solution with continuously stirring at 70 °C for 4 h, followed by the aging at room temperature for 12 h. Finally, the mixture was transferred into a reactor and heated at 180 °C for 5 h. After cooling, centrifugation, washing and freeze-drying, TiO₂/PDIsM and TiO₂/CuPDIsM composites were obtained.

2.5 Characterization

The crystal structure of the as prepared samples was analyzed by X-ray diffraction (XRD, Rigaku, Smartlab diffractometer). The specific surface area was evaluated using N₂ adsorption-desorption experiments with a Micromeritics SSA-7000 instrument. The morphology of the samples was examined by emission-scanning electron microscopy (SEM, Hitachi SU8010) equipped with X-ray energy dispersive spectrometer and transmission electron microscopy (TEM, JEOL2100F). UV-visible absorption spectra and diffuse reflectance spectra (DRS mode) were obtained using the UV-2600 spectrophotometer. The structure analysis was studied by a Fourier transform infrared spectrometer (FTIR, Nicolet iS50, America) and X-ray photoelectron spectroscopy (XPS, Thermo Fischer ESCALAB Xi+). Photoluminescence spectroscopy (PL) measurements were conducted on a Hitachi F-4500 fluorescence spectrometer and zeta potential measurements were carried out in neutral aqueous solution using Zetasizer Nano ZS90 device. Electrochemical analysis was performed on a CHI 660D workstation with 0.1 M Na₂SO₄ aqueous solution as the electrolyte solution.

2.6 Photocatalytic performance evaluation

Tetracycline, methylene blue and methyl orange were selected as target pollutants and the photocatalytic activity of photocatalyst samples was evaluated under visible light irradiation. A 300W xenon light lamp was used as an irradiation source with a distance of 10 cm between the solution and the lamp. 20 mg of the prepared catalyst was dispersed in 40 mL of a solution containing TC (10 mg L⁻¹), MO (50 mg L⁻¹) or MB (50 mg L⁻¹ or 150 mg L⁻¹). Prior to irradiation, dark adsorption treatment was first carried out by magnetically stirring for 30 min to establish the adsorption-desorption equilibrium. Then 3 mL of the suspension was collected and filtered at given intervals of time during the light illumination. The filtrates were analyzed by UV-



vis spectrophotometer for TC at 358 nm, MO at 463 nm and MB at 664 nm.

3. Results and discussion

3.1 The organization and morphology investigation

The π - π stacking and crystal structure of self-assembled PDI supramoleculars can be revealed from UV-vis absorption spectra and XRD patterns, respectively.^{31,32} The UV-vis absorption spectra of PDI monomer solution, PDIsM and CuPDIsM suspensions are shown in Fig. 1a. The spectrum of PDI dissolved in ethanol solution ($V_{\text{water}}/V_{\text{ethanol}} = 1:1$) shows three pronounced peaks in the range of 400–600 nm, which is attributed to the 0–0, 0–1 and 0–2 electronic transitions of monomeric PDI molecules.^{33,34} Compared with PDI monomer, PDIsM dispersion shows broaden peaks with a blue-shift, indicating the H-type stacking between the PDI skeletons. CuPDIsM suspensions show an obvious blue-shift of the maximum peak as comparison with PDI monomer, so it also has the H-type stacking structure. Moreover, in the spectrum of CuPDIsM, the absorbance intensity of the 0–1 vibronic transition peak at 504 nm is higher than the 0–0 transition peak at 543 nm, which is a typical indication of H-type aggregation.^{35,36} Therefore, the as prepared PDIsM and CuPDIsM have H-type π - π stacking structures, different from other metal-PDI supramoleculars with J-type aggregation structure in the literature.^{29,30}

Fig. 1b shows the XRD patterns of PDIsM, CuPDIsM, TiO₂ and their composite samples. For the obtained PDIsM, the diffraction peaks between 24°–28° are generally considered to be the typical characteristic peaks of π - π stacking structures

between PDI skeletons molecules, with the stacking layer spacing of 3.27–3.71 Å. This ordered π - π stacking contributes to long-range electron delocalization and the migration of photogenerated electrons along the π - π superposition direction,^{33,36} which can enhance the electronic coupling^{34,37} and charge mobility.³⁸ For the as prepared TiO₂, the peaks at $2\theta = 25.3^\circ$, 37.8° , 48.1° , 54.9° and 62.7° are corresponding to the (101), (004), (200), (211), and (204) planes of anatase phase, respectively. For TiO₂/PDIsM composites, all the peaks can be detected in the spectra of PDI and TiO₂, implying that PDI supermolecules are integrated with TiO₂ without changing the crystal structure. For CuPDIsM, the intensity of the diffraction peaks is low as comparison with the spectrum of PDIsM, indicating that Cu doping hardly changes the crystal phase, but lowers the crystallinity of the PDI aggregates.

The morphology of the samples was analyzed by SEM and TEM. In Fig. S1a,† it can be seen that TiO₂ nanoparticles with the particle size of about 20–30 nm show heavy agglomerations due to the strong surface polarity. PDIsM and CuPDIsM show a clustered nanorod-like morphology with the diameter of less than 50 nm (Fig. S1b and c†). The nanorod shape of CuPDIsM is also observed in TEM image (Fig. 1c). This one-dimension structure originates from the self-assembly of carboxyl-containing perylene diimide molecules through intrinsic π -stacking of the aromatic core along the long axis and the hydrogen bond link between the carboxylic side chains along the short axis.^{14,33} The hydrogen bond between the side chains of the adjacent PDI molecules is replaced by metal-ligand interaction in CuPDIsM. For TiO₂/PDIsM and TiO₂/CuPDIsM (Fig. S1d and e†), TiO₂ particles are successfully distributed on the surface of PDIsM and CuPDIsM nanorods. Element

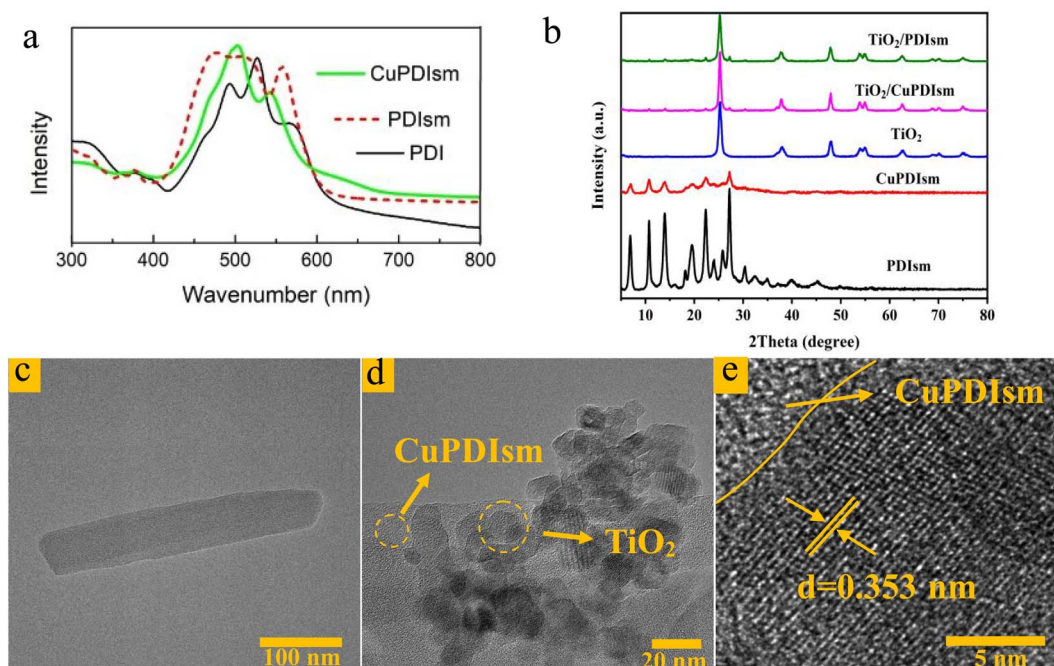


Fig. 1 UV-vis absorption spectra of PDI monomer solution and PDI supramolecular suspensions (a), XRD patterns of TiO₂, PDIsM, CuPDIsM and their composite samples (b), TEM images of CuPDIsM (c) and TiO₂/CuPDIsM (d), HRTEM of TiO₂/CuPDIsM (e).



mapping images (Fig. S2†) show that the presence of Cu element identifies the successful doping of Cu in PDIsm. The homogenous distribution of C, N, O and Ti indicates the good incorporation of TiO₂ with CuPDIsm. In addition, TEM images of TiO₂/CuPDIsm composites (Fig. 1c–e) show that TiO₂ nanoparticles are immobilized on the surface of CuPDIsm nanorods and the crystal plane spacing of 0.353 nm is corresponding to the highly reactive (101) plane of TiO₂ crystals. The close contact of TiO₂ with PDIsm and CuPDIsm indicates the formation of heterojunction interface, offering a direct pathway for photo-electron transport.

N₂ adsorption-desorption isotherms were employed to evaluate the specific surface area of the samples (Fig. S3a–c†). The isotherm plots of all samples belong to the IV type in the classification. Compared with PDIsm, CuPDIsm has much more mesopores and higher specific surface areas, which may be responsible for its lower crystallinity degree as discussed above in XRD results. The incorporation of PDIsm and TiO₂ further increases the mesopores and specific surface areas, so both Cu-doping and TiO₂-coupling can enhance the adsorption and reduce the migration distance of electrons from inside to the surface.

Fig. 2a shows FTIR spectra of the samples. The spectrum of CuPDIsm is similar to that of PDIsm, so the doping of Cu hardly affects the chemical structure of PDI supermolecules. The typical characteristic peak of PDIsm at 1695 cm⁻¹ is corresponding to the stretching vibrations of the C=O bond of the PDI molecule,²¹ which is attributed to the presence of carboxyl group at the side chains of PDI molecular. But this peak slightly shifts to 1697 cm⁻¹ in the spectrum of CuPDIsm, which may be ascribed to the electrostatic interaction between carboxyl and Cu²⁺.²⁵ Such blue shift verifies that PDI molecular interconnects each other through hydrogen bonds for PDIsm and Cu bridging for CuPDIsm along the short axis of the one-dimension supra-molecules. The Cu bridging is often in the way of metal-ligand interaction with one Cu²⁺ ion coordinated by three monodentate carboxylic groups from three deprotonated PDI ligands and one water molecule,²⁵ as shown in Fig. 2b. In the spectra of TiO₂/PDIsm and TiO₂/CuPDIsm, C=O peak also shifts to 1697 cm⁻¹, which may be due to the formation of hydrogen bonds between carboxyl group of the PDI molecule and -OH

group of TiO₂ surface.^{26,34,42–45} To further illustrate the interaction between Cu and PDI, Raman spectrum measurements were performed. As shown in the Fig. S4,† the characteristic bands in PDIsm are observed at 1292 cm⁻¹ and 1369 cm⁻¹ corresponding to C–H bending vibrations, and at 1572 cm⁻¹ corresponding to C–C/C=C stretching vibration in perylene core.^{39–41} Compared with Raman spectrum of PDIsm, these bands with stronger intensity in CuPDIsm slightly shift to the left, which may be attributed to Cu bonding between adjacent PDI units.

XPS characterization was used to study the surface chemistry of materials (Fig. 3a–f). In Fig. 3a, compared with PDIsm, CuPDIsm has a small new peak corresponding to Cu 2p signal, identifying the presence of Cu in this composite. The C 1s spectrum of PDIsm shows four peaks at 284.7, 286.1, 287.8 and 289.2 eV, representing C=C, C–N, C=O bonding and π -electron excitation, respectively, suggesting the formation of a self-assembled π -stacking structure. The O 1s spectrum of PDIsm shows three characteristic peaks at 531.6, 532.5 and 533.3 eV, related to C=O bond, C–O bond and adsorbed oxygen, respectively. Compared with PDIsm, CuPDIsm show a slight shift of C=O and C–O peaks to high binding energy, which is attributed to the formation of C–O–Cu interaction between adjacent PDI moleculars, further implying that CuPDIsm has a linear edge-to-edge organization through metal-ligand interaction along the short axis. An obvious blue shift of C=O and C–O peaks is detected in TiO₂/CuPDIsm, which may be due to the formation of the –C=O–H hydrogen bond and C–O–Ti bond at the heterojunction interface.²⁷ Such hydrogen bond and electronic coupling arisen from the interaction between surface carboxyl groups at PDI molecular side chain and the surface hydroxyl groups and unsaturated dangling bonds on TiO₂ particle, providing the fast passage for the electron transfer, as shown in Fig. 4.

In Ti 2p spectra (Fig. 3d), TiO₂ has two peaks at about 529.36 and 530.5 eV, corresponding to Ti–O bond and the surface hydroxyl groups (Ti–OH bond), respectively.⁴⁶ Notably, these two peaks in TiO₂/CuPDIsm shift to lower binding energy position, indicating that the surface electron density of TiO₂ increases.^{24,47} This increasing surface electrons of TiO₂ originates from the transfer of electrons from CuPDIsm to TiO₂ through hydrogen bonds and C=O–Ti at the heterojunction

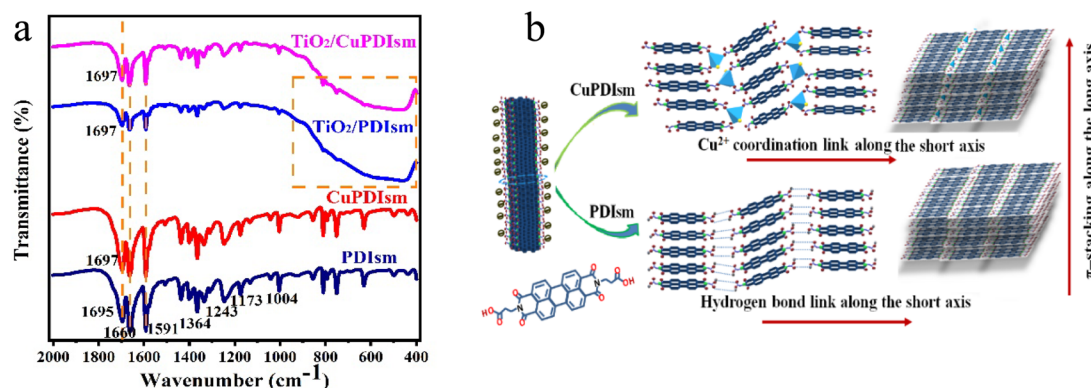


Fig. 2 FTIR spectra of the samples (a), self-assemble illustration of adjacent PDI molecular *via* hydrogen bond or tetra-coordinated interactions (b).

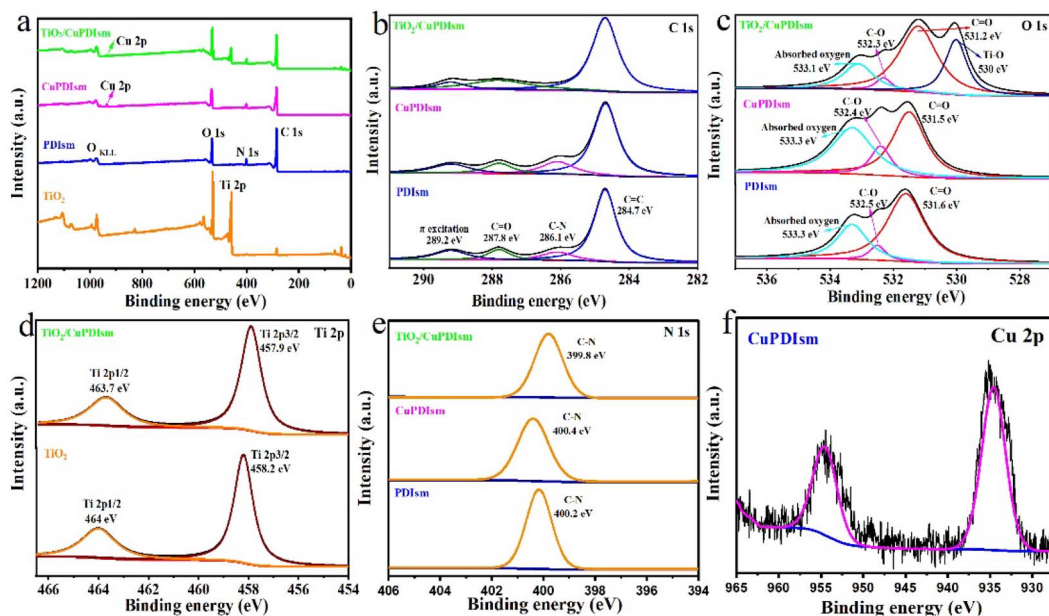


Fig. 3 XPS spectra of PDIsm, CuPDIsm and $\text{TiO}_2/\text{CuPDIsm}$ samples. Survey scan (a) and fine scan energy spectra of C 1s (b), O 1s (c), Ti 2p (d), N 1s (e) and Cu 2p (f).

interface.⁴⁵ For N 1s pattern in Fig. 3e, a single high-intensity peak presents at 400.2 eV in PDIsm, corresponding to C–N bond. A small shift of this peak is detected in CuPDIsm (400.4 eV) and $\text{TiO}_2/\text{CuPDIsm}$ (399.8 eV), indicating that the chemical environment of N 1s changed after PDIsm was doped with Cu and coupled with TiO_2 .⁴⁸ The Cu 2p spectrum (Fig. 3f) exhibits that two major peaks at 934.6 and 954.6 eV are ascribed to Cu 2p_{3/2} and Cu 2p_{1/2} valence states, respectively, confirming the presence of Cu as Cu^{2+} oxidation state⁴⁹ in CuPDIsm.

3.2 Photoelectric properties

The UV-vis absorption spectra of the samples were measured to explore the light-absorption ability (Fig. 5a). TiO_2 exhibits high absorption intensity in the UV region but low absorption ability in the visible region. In contrast, PDIsm has a weaker

absorption of UV light, but strong response in the visible region of 400–700 nm. There are two absorption bands at 498 and 550 nm in the PDIsm, corresponding to the electronic transition in the isolated molecule and the π -electron delocalization between PDI chromophores, respectively.⁷ It is noteworthy that the introduction of Cu into PDIsm greatly increases the absorption ability and causes the red-shift of absorption edge. The loading of TiO_2 nanoparticles onto PDIsm surface is also advantageous to increasing the light absorption of PDI self-assembly. So both Cu-doped and TiO_2 coupling would enhance the photocatalytic performance of PDIsm under visible light irradiation. The band gap (E_g) of the as-prepared samples were estimated by Tauc's plot. As shown in Fig. 5b, the E_g of TiO_2 is estimated to be 3.1 eV, so TiO_2 is hardly photoexcited by visible light. The E_g of CuPDIsm and PDIsm are 1.91 and

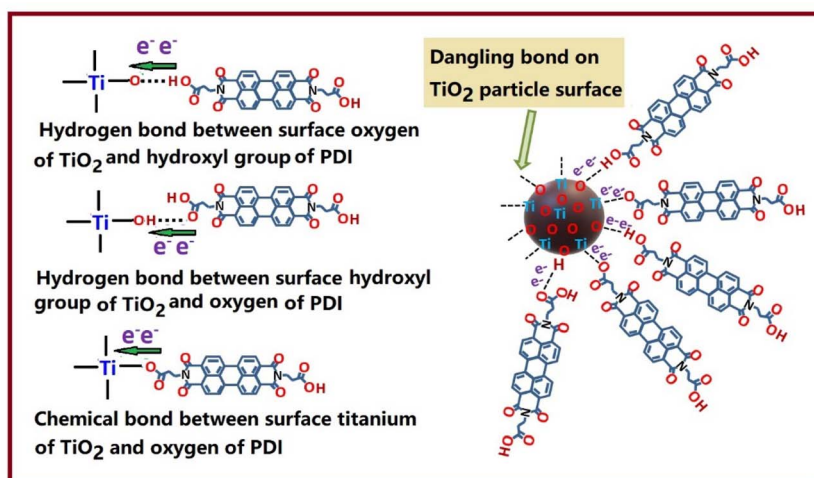


Fig. 4 Schematic illustration of binding interaction and charge transfer at the interface between PDIsm and TiO_2 .



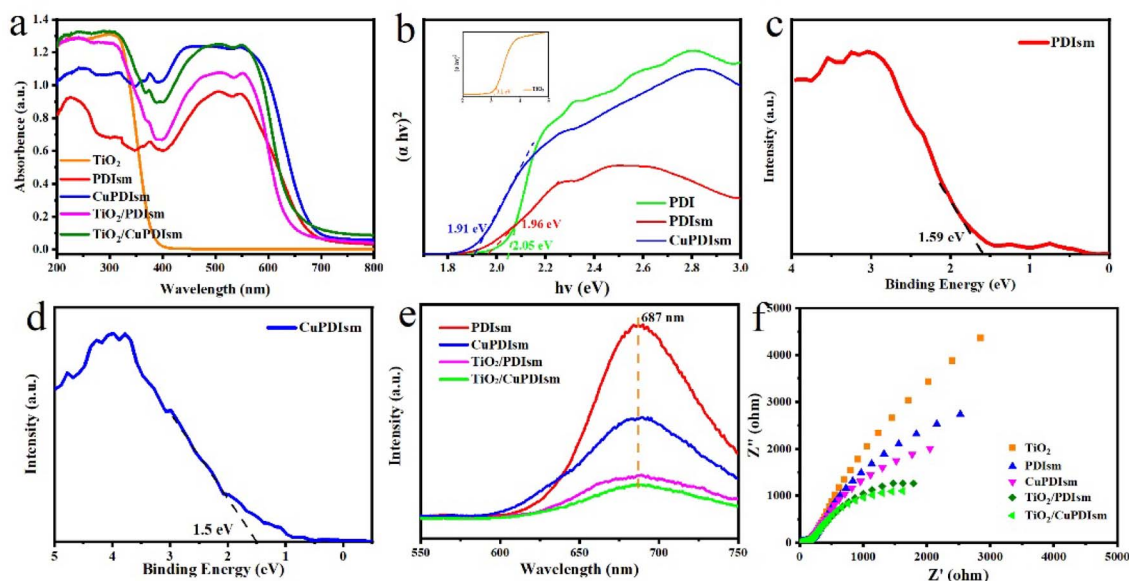


Fig. 5 UV-vis diffuse reflectance spectra of the samples (a) and the corresponding Tauc's plots (b), valence band potential diagrams of PDIsM (c) and CuPDIsM (d), photoluminescence profiles (e) and EIS profiles (f) of the samples.

1.96 eV, respectively, indicating their visible light photocatalytic activity. The valence band (VB) is determined as 1.59 eV for PDIsM and 1.5 eV for CuPDIsM by using XPS measurement (Fig. 5c and d). Thus, the conduction band (CB) of samples can be calculated according to the following equation:

$$E_{CB} = E_{VB} - E_g \quad (1)$$

the E_{CB} of PDIsM and CuPDIsM is determined as -0.37 and -0.41 eV, respectively.

Though PDIsM has good absorption of visible light and easily excited by visible light, its photocatalytic performance is still poor due to the high charge recombination.^{7,14,15} Photoluminescence (PL) spectroscopy was used to measure the separation efficiency of photogenerated carriers (Fig. 5e). Compared with PDIsM, CuPDIsM has low luminescence intensity, proving its low charge recombination. This indicates that the metal-ligand interaction between adjacent PDI molecules is more beneficial to the electron transfer than hydrogen bond link. Additionally, the mesoporous structure and high specific surface areas of CuPDIsM help to accelerate the migration of photo-generated electrons from the inside to surface as discussed above. $\text{TiO}_2/\text{CuPDIsM}$ shows the lowest luminescence intensity among all the samples, which is attributed to the more efficient transfer of photogenerated charge through the hydrogen bond and $\text{C}=\text{O}-\text{Ti}$ bond formed at the heterojunction interface.⁵⁰ The transfer efficiency of photogenerated electron was further investigated by the electrochemical impedance (EIS) spectra (Fig. 5f). The charge transfer resistance decreases in order: $\text{TiO}_2 > \text{PDIsM} > \text{CuPDIsM} > \text{TiO}_2/\text{PDIsM} > \text{TiO}_2/\text{CuPDIsM}$, further identifying that the conductivity is enhanced and electron transfer rate is accelerated when PDIsM is doped with copper and coupled with TiO_2 . So, it can be concluded that the incorporation of Cu and TiO_2 with PDIsM not only expands the photo-absorption but also

enhances the carrier separation efficiency, which were favorable for photocatalytic reaction.

3.3 Photocatalytic activity

TC, MO and MB were chosen as the target pollutants to investigate the photocatalytic activity of the samples. For anionic TC and MO organics (Fig. 6a and b), the photodegradation ratios of the samples increase in order: $\text{TiO}_2 < \text{PDIsM} < \text{CuPDIsM} < \text{TiO}_2/\text{PDIsM} < \text{TiO}_2/\text{CuPDIsM}$, so both Cu-doping and TiO_2 coupling can effectively increase the photocatalytic activity of PDIsM. $\text{TiO}_2/\text{CuPDIsM}$ composites show the highest degradation ratios, reaching the maximum values of 89.87% toward TC and 68.65% toward MO. The excellent photodegradation efficiency of $\text{TiO}_2/\text{CuPDIsM}$ is attributed to the noticeable photocharge transfer, high visible light absorption and large specific surface area as discussed above.

For cationic MB (Fig. 6c), $\text{TiO}_2/\text{CuPDIsM}$ also shows excellent photodegradation ratio, reaching the maximum value of 97.26%. It is notable that the MB degradation curves of PDIsM and CuPDIsM almost overlap and the total MB adsorption finishes in the dark adsorption stage, indicating the strong adsorption ability of these two samples. To understand the adsorption capacity of PDIsM toward organic pollutants, the adsorption experiment was conducted using 20 mg of adsorbent in 40 mL of 300 mg L^{-1} MB for a duration of 9 h. In Fig. S5,[†] PDIsM shows much better adsorption capacity (528.7 mg g^{-1}) toward cationic MB, while poorer adsorption toward anionic TC and MO. The remarkable adsorption capacity toward cationic MB is attributed to the strong electrostatic attraction because PDIsM and CuPDIsM are negative charged with the ionization of abundant carboxyl groups in the aqueous solution. The high negative charge are identified by the zeta potential (Fig. S6[†]) of PDIsM (-85.15 mV) and CuPDIsM (-70.99 mV), so PDIsM and CuPDIsM have better



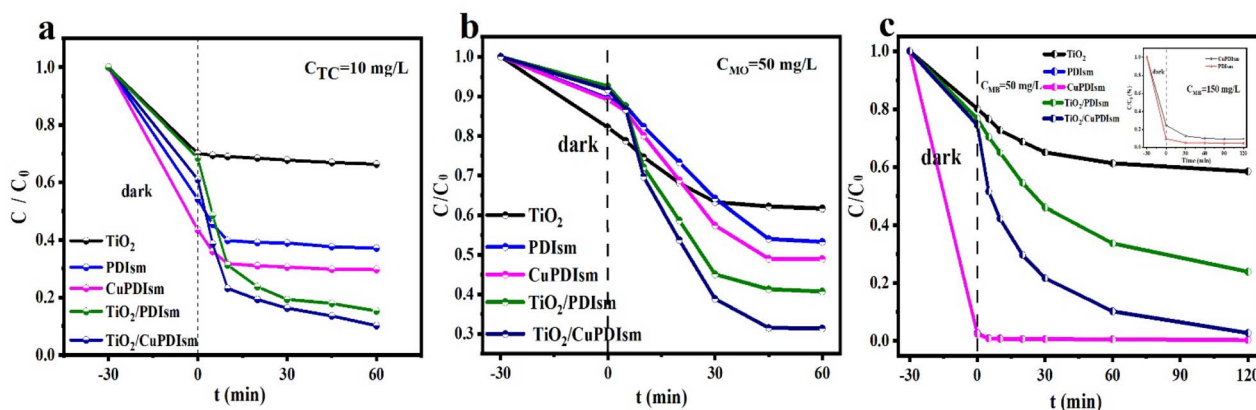


Fig. 6 The photodegradation ratios of the samples toward TC (a), MO (b) and MB (c).

adsorption ability toward cationic MB. However, it is difficult for anionic TC and MO to be adsorbed onto PDIsM and CuPDIsM surface due to the electrostatic repulsion. The excellent surface adsorption capacity is beneficial to enhancing the photo-performance of PDIsM and CuPDIsM based composites, explaining the better degradation ratio of PDIsM based composites toward MB than toward MO and TC.

The apparent rate constants of MO and MB were fitted by the pseudo-first-order equation (Fig. S7a and b†). CuPDIsM and $TiO_2/CuPDIsM$ show higher K_{MO} than PDIsM and $TiO_2/PDIsM$, respectively, further proving that the photocatalytic activity of PDIsM can be effectively improved by Cu-doping and TiO_2 -coupling. In addition, the cycling tests of $TiO_2/CuPDIsM$ show that the degradation ratio of TC decreases slightly with the increase of cycle times and remains 71.45% after five cycles (Fig. S8a†). The XRD patterns (Fig. S8b†) show that the crystal structure of $TiO_2/CuPDIsM$ hardly changes after the cycling experiment, indicating the good structural stability and favorable reusability of $TiO_2/CuPDIsM$.

3.4 Mechanism of photocatalytic activity

Radical trap experiment of $TiO_2/CuPDIsM$ composites was performed to explore the photocatalytic mechanism. The isopropanol (IPA), benzoquinone (BQ) and ethylenediaminetetraacetic acid disodium (EDTA) were used as scavengers of $\cdot OH$, $\cdot O_2^-$ and h^+ species, respectively. As shown in Fig. 7a, the addition of IPA to the reaction system results in very limited change in the photodegradation of TC, MB and MO, indicating that $\cdot OH$ plays a negligible role in the photodegradation reaction. The introduction of EDTA leads to a more significant decrease of the degradation efficiency than BQ addition, suggesting that the h^+ is identified as the main active species and $\cdot O_2^-$ is the secondary active species. The ESR spin-trap experiment shows that the characteristic peaks corresponding to the DMPO/ $\cdot O_2^-$ adducts were detected (Fig. 7b) under visible light irradiation while no signals in dark, further implying that the $\cdot O_2^-$ radicals are produced and involved in the photodegradation process. This indicates that the photoinduced electrons in $TiO_2/CuPDIsM$ can be captured by dissolved oxygen to generate $\cdot O_2^-$ radicals.^{51,52} However, after 5, 10 and 20 min of

light, the signal of DMPO- $\cdot O_2^-$ was significantly enhanced. This indicated that the photoinduced electrons in $TiO_2/CuPDIsM$ can be captured by O_2 to generate $\cdot O_2^-$ radicals.⁵¹

The presence of the mainly generated species can also be theoretically inferred from valence band and conduction band of CuPDIsM. According to the obtained E_g (1.91 eV), E_{CB} (−0.41 eV) and E_{VB} (1.5 eV) of CuPDIsM as discussed above, $\cdot O_2^-$ species can be generated because the E_{CB} of CuPDIsM was more negative than the reduction potential of $O_2/\cdot O_2^-$ (−0.33 eV) and the photo-induced e^- can be captured by dissolved oxygen to form $\cdot O_2^-$.^{50,51} But very limited amount of $\cdot OH$ species is generated since the E_{VB} of CuPDIsM is less positive than the oxidation potential of $OH^-/\cdot OH$ (1.99 eV) and $H_2O/\cdot OH$ (2.37 eV).⁵³ So, under visible light irradiation, PDIsM can be photo-excited and facilitates $\cdot O_2^-$ and h^+ active species for the degradation of organic pollutants.

Though PDIsM can be theoretically photoexcited, it shows low photodegradation performance due to the low transport efficiency of photo-generated electrons, which depends on the migration path along the π - π long-range transport pathway with the hydrogen bond link between the carboxylic side chains. When Cu is doped into PDIsM, the hydrogen bond link is replaced by the stronger metal-ligand interaction between adjacent side chains and hence the electron transfer is greatly accelerated. Additionally, the mesoporous structure of CuPDIsM system can shortens the migration distance of electrons from inside to surface and so enhances the spatial charge separation. Moreover, when TiO_2 is coupled with CuPDIsM, surface hydrogen bond interaction ($-C=O-H$)²² and electronic coupling interaction ($C=O-Ti$)^{13,23} form at the heterojunction interface, which provide the fast-channel for the transfer of electrons from CuPDIsM to the reactive (101) surface and conduction band of TiO_2 .^{13,22,23} This substantial electron transfer from CuPDIsM to TiO_2 has been identified by the shift of Ti 2p peaks to low binding energy position, as shown in Fig. 7c. The photocatalytic mechanism of $TiO_2/CuPDIsM$ composite is shown schematically in Fig. 7d. Therefore, we corroborate that the highly efficient photocatalysis of CuPDIsM/ TiO_2 are ascribed to simultaneous introduction of Cu and TiO_2 with the great enhancement of charge separation.



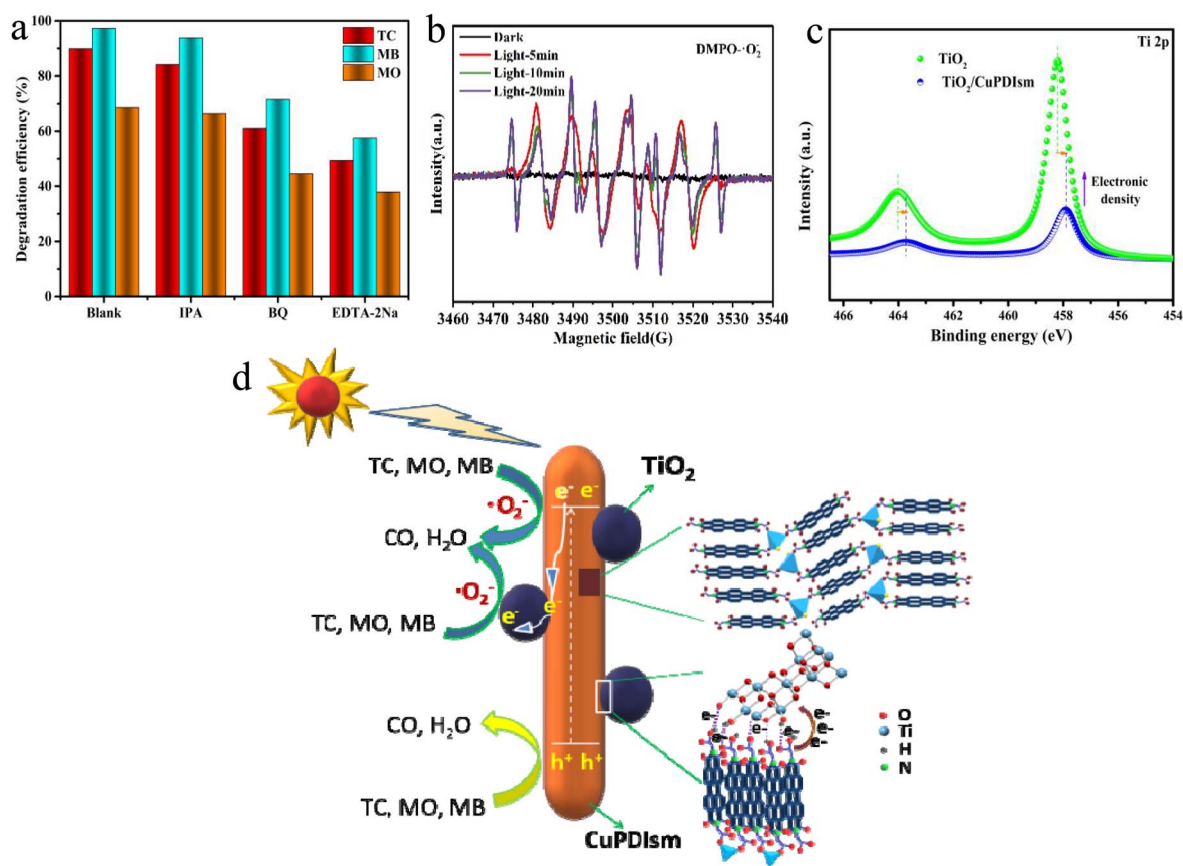


Fig. 7 Photodegradation of TiO₂/CuPDIsM with the addition of scavengers (a), ESR spectra of TiO₂/CuPDIsM in aqueous dispersion for DMPO-[•]O₂⁻ (b), high-resolution XPS of Ti 2p for TiO₂ and TiO₂/CuPDIsM (c), the photocatalytic mechanism of TiO₂/CuPDIsM composite (d).

4. Conclusion

We firstly prepared a novel CuPDIsM supramolecular by simple self-assembly of PDI molecules in copper salt solution and then designed the TiO₂/CuPDIsM system with the growth of TiO₂ on CuPDIsM by hydrothermal method. CuPDIsM with mesoporous structure is constructed by face-to-face organization of molecular planes through H-type π - π stacking and edge-to-edge connects of molecular side-chains through metal-ligand interaction. TiO₂ is closely immobilized on CuPDIsM surface through -C=O-H and C=O-Ti links at the heterojunction interface. Both Cu-doping and TiO₂ coupling could forcefully not only increase the adsorption intensity of visible light but also accelerate the spatial carrier separation, and hence greatly promote the photodegradation performance under visible-light irradiation. TiO₂/CuPDIsM composites construct an efficient electron transport system and show remarkable photodegradation efficiency as comparison with pure TiO₂ and PDIsM, with the degradation ratios of up to 89.87%, 68.65% and 97.26% toward tetracycline, methyl orange and methylene blue, respectively. This work demonstrates a bright application prospect of inorganic-organic composite catalysts for environmental purification territory.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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