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Magnetic recyclable CoFe₂O₄@PPy prepared by *in situ* Fenton oxidization polymerization with advanced photo-Fenton performance†

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Here we present a magnetic recyclable photo-Fenton catalyst $CoFe_2O_4$ @PPy with uniform morphology and excellent dispersibility prepared *via* simple *in situ* Fenton oxidization polymerization. The $CoFe_2O_4$ core provides good magnetic recyclability for the catalysts as well as the ion source for catalyzed decomposition of H_2O_2 in PPy coating. The optimal catalytic effect can be obtained by adjusting the ratio of $CoFe_2O_4$ and PPy. Methylene blue, Methyl orange and Rhodamine B (RhB) employed as model pollutants certificated that the catalyst exhibits a wide range of photodegradability. The decoloration rates reach nearly 100% in the photodegradation of 10 mg L^{-1} RhB after 2 h visible-light irradiation and only low toxicity small molecules are detected by LC-MS. Moreover, the catalytic activity remains after 5 cycles with decoloration rates up to 90%. The degradation measurement in the presence of scavengers of reactive species reveals that the positive holes (h⁺) and hydroxyl radical (·OH) are the main reactive oxygen species in the $CoFe_2O_4$ @PPy system. The performance enhancement may be attributed to the combination of improved Fenton activity by coordinated Fe^{2^+} and PPy redox pairs and photo-catalytic activity by broaden adsorption and photo-generated charge separation.

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Introduction

Nowadays, organic pigments and dyes are increasingly popular because of the booming chemical industry and the pursuit of better life. The discharge of waste water containing such organic pigments and dyes brings about enormous threats to our environment and health. As an advanced oxidation process, the Fenton reaction has advantages of complete degradation and mineralization of organic pollutants by in situ generated highly oxidative free hydroxyl radicals. However, the traditional Fenton reaction encounters problems such as low pH conditions, sluggish Fe³⁺ reduction kinetics and secondary pollutants caused by mass production of slurry containing Fe³⁺. Therefore, heterogeneous Fenton systems $^{1-3}$ in which Fe^{2+} of the catalyst was replaced by $Fe(\pi)$ active sites from solid iron metals,4 alloys,5-7 oxides7-9 and other compounds are developed for better recyclability. Nevertheless, the rapid re-generation of Fe(II) active site from Fe(III) on the surface of heterogeneous Fenton catalysts remains challenges. Photo-Fenton 10-12 in which the solar

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energy is utilized to drive the reduction is a promising route to conquer the sluggish kinetic. After the excitation of semi-conductors with photo-catalytic activity, photo-generated electrons on conductive band play a role of reductant for Fe(III) while the photo-generated holes play a role of additional oxidant for organic pollutant.

Transition mixed metal oxide, CoFe₂O₄, with spinel structure13 is regarded as one of effective heterogeneous photo-Fenton catalysts along with magnetic recyclability14,15 as well as unique electrical properties,16 physicochemical stability,17 abundant resources, and eco-friendliness.¹⁸ However, poor electron transfer activity and dispersibility severely limit the catalytic activity of the ferrite.19 Therefore, it is of particular importance to transform the steric hindrance and electrical conductivity of the nanomaterial by changing the surface of the nanomaterial and appropriate doping, thereby obtaining stable nanoparticles and facilitating electron conduction.20 Coating ferrite with inorganic layers e.g. silica, 10,21 inhibits the agglomeration of magnets, thus improves its dispersibility. Modifying ferrite with metal nanoparticles acting as electron acceptors such as Ag,22 Au,23 Pd,24 Cu25 etc. on the surface enhances the separation of photo-generated electron-hole pairs. Carbon nanotubes, 26 graphene, 27,28 C3N4 (ref. 29 and 30) or other organic semiconductors decorated on ferrite, on the one hand, can act as a collector and transmitter of excited electrons because of its conjugated large Π bond, on the other hand, can improve the light absorption, thus enhance the

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photo-catalytic activity. In recent years, thanks to the high carrier mobility and chemical stability, conductive polymers such as polythiophene (PTH),³¹ polyaniline (PANI),^{32,33} polypyrrole (PPy),^{34,35} *etc.* are used to modify the ferrite, giving the composites good electron transport capability in photocatalysis process.³⁶ The synergistic effect is revealed by the combination of the organic phase and the inorganic phase, as improved photo-catalytic performance compared to each

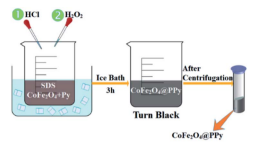
single component is exhibited due to better electron transfer

ability through multiple redox pairs. So far, PPy has been confirmed to effectively enhance the photocatalytic performance of semiconductor photo-catalysts,³⁷ such as BiOI@PPy, 38 TiO2@PPy, 39 Fe3O4@PPy, 40 etc. Moreover, Yu et al.41 found that PPy showed better enzyme-mimic activity than that of horseradish peroxidase (HRP) in catalyzing the decomposition of hydrogen peroxide. Yang et al.42 also tried to combine Fe₃O₄ with PPy to prepare a peroxide nanozyme with magnetic properties and excellent conductivity. It is deduced that the enzyme-mimic activity similar with Fenton catalytic ability may be related to its porphyrin-like structure. Accordingly, combination of PPy and ferrites may provide novel catalysts with enhanced performance in photo-Fenton degradation of organic pollutants due to their improved dispersibility, recyclability, light-absorption, and electron transfer. However, so far as we know, there are few reports on the application of such kind of catalysts.

Herein, we present a facile preparation of an effective photo-Fenton catalyst with good recyclability and reusability. The PPy coated CoFe₂O₄ microspheres (CoFe₂O₄@PPy) were prepared via Fenton oxidation induced in situ polymerization coating method. In brief, in situ generated Fe³⁺ and Co²⁺ by dissolution of CoFe₂O₄ in acidic solution were used to catalyze the decomposition of H_2O_2 for the oxidation polymerization of PPy. The coating of PPy on the surface of CoFe₂O₄ produces a ferromagnetic CoFe₂O₄@PPy with uniform morphology, good dispersibility and recyclability. The catalyst shows enhanced photo-Fenton activity in the removal of model water pollutants RhB under neutral condition with 95% decoloration within 30 min irradiation in the presence of 45 mM H₂O₂. The catalytic activity remains 90% of decoloration rate after 5 cycles of degradation, revealing good reusability. The degradation of the two dyes, methylene blue and methyl orange, demonstrates the lower selectivity of catalysts.

Results and discussion

The schematic illustration of preparation procedure and principle is shown in Scheme 1. By adding hydrochloric acid to the reaction system, an acidic environment is created and partial of $CoFe_2O_4$ is dissolved. The released Fe^{3+} was reduced to Fe^{2+} by pyrrole and the polymerization is initiated. As soon as the addition of hydrogen peroxide, it reacted with the resulted Fe^{2+} to generate a hydroxyl radical in Fenton mechanism, which act as oxidant together with Fe^{3+} to oxidize the pyrrole to form a polypyrrole shell on the surface of $CoFe_2O_4$. The presence of small amount of surfactant SDS in the solution is expected to



Scheme 1 Schematic illustration for the synthesis procedure of $CoFe_2O_4@PPy$.

prevent the agglomeration between polypyrrole coated ferrite microspheres. Therefore, a well dispersed, spherical core–shell structure with uniform particle size distribution was prepared successfully.

Characterization of the catalyst

The amount of Py used for coating is varied as 2 mmol, 4 mmol, 6 mmol and 8 mmol. Accordingly, the obtained CoFe₂O₄@PPy were named as S1, S2, S3 and S4. The morphologies of synthesized catalysts S1, S2 and S3 characterized by SEM and TEM were shown in Fig. 1a-c. It can be seen from Fig. 1 that the prepared CoFe₂O₄@PPy exhibits spherical morphology with diameter of 200 nm on average and excellent dispersibility. With the increase of PPy content, the size of CoFe₂O₄@PPy microspheres hardly increases but their surface becomes increasingly rougher than that of pure CoFe2O4 (Fig. S1†), revealing that the surface of CoFe₂O₄ is coated with a very thin layer of PPy. However, there's no obvious PPy layer can be resolved from TEM images as shown in Fig. 1 when the PPy dosage no more than 6 mmol, implying the PPy layer on S1, S2 and S3 is too thin. Legible PPy coating layer is observed for S4 when the dosage of Py is up to 8 mmol. From the TEM as shown in the inset of Fig. 2a, a boundary between PPy and CoFe₂O₄ is appeared vividly. The magnified TEM image of S4 as shown in Fig. 2b indicates that the surface of CoFe₂O₄ microspheres was covered with an amorphous layer of about 10 nm thickness. To get a deep insight on the structure of S4, high resolution TEM and selected area electron diffraction (SAED) measurements were performed. The SAED pattern as shown in Fig. 2c shows a series of discontinuous diffraction rings with inter-planar distance of 0.299, 0.254, 0.174, 0.166 and 0.150 nm which are corresponding to the distance between crystal planes of {220},

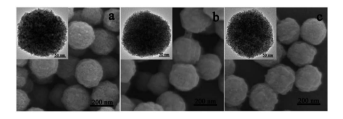


Fig. 1 SEM images of S1 (a); S2 (b); S3 (c). Insets are TEM images corresponding to the SEM, respectively.

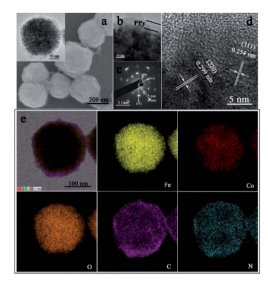


Fig. 2 The SEM image of S4 (a); the TEM image of S4 (b); the SAED pattern of S4 (c); the HRTEM image of S4 (d); the TEM-EDS image of S4 (e)

{311}, {422}, {511} and {440}, revealing a polycrystalline with cubic lattice structure of $CoFe_2O_4$ microspheres. From HRTEM image of Fig. 2d, the inter-planar distance is calculated to be 0.299 and 0.254 nm, corresponding to (220) and (311) plane of $CoFe_2O_4$ respectively. Furthermore, the EDS mapping of a single microsphere of S4 as shown in Fig. 4e clearly signifies that $CoFe_2O_4$ @PPy contains five elements of Co, Fe, O, C and N, among which C and N elements evenly distributed on the

surface of $CoFe_2O_4$, demonstrating that PPy is successfully coated on $CoFe_2O_4$ microspheres.

The XRD patterns of original CoFe₂O₄, PPy and CoFe₂O₄@PPy loaded with different content of PPy are depicted in Fig. 3a. The diffraction peaks at 2θ values of 30.084° , 35.437° , 43.058° , 53.445°, 56.973°, 62.585°, and 74.009° correspond to the standard peak positions of spinel CoFe₂O₄ structure (PDF no. 22-1086). However, the pattern of PPy has only one broaden diffraction peak with a 2θ value of 26° due to the amorphous structure of PPy polymer. The patterns of CoFe₂O₄@PPy have no obvious PPy diffraction peak but strong diffraction signal of CoFe₂O₄, which is attributed to the strong signal of CoFe₂O₄ tending to overwhelm the weak carbon peak. Thermogravimetric analysis (TGA) in air was also carried out for further confirming the existence of PPv and roughly estimating its amount. Due to the weight loss caused by the volatilization of adsorbed water is observed at around 100 °C, the thermo-gravimetric analysis is started from 100 °C. The temperature was raised from 100 °C to 800 °C for the pyrolysis of PPy. 43 As shown in Fig. 3b, PPy is almost completely oxidized at 650 °C and the weight loss is near 100%, whereas the curve for CoFe₂O₄ only shows a slight weight loss (\sim 5%) at 600 °C. The weight loss for the composites consecutively increases with the increase of the PPy content. From the weight loss of composites, the content of PPy in S1, S2, S3 and S4 is estimated to be 40, 23, 16 and 11 wt%. The Raman spectra of CoFe₂O₄@PPy and bare PPy, CoFe₂O₄ are shown in Fig. 3c. There are two distinct broad peaks ranging from 1200 cm⁻¹ to 1800 cm⁻¹ in CoFe₂O₄@PPy and pure PPy. The peak at 1350 cm⁻¹ is assigned to the D band (C-N stretching vibration), which mainly due to the presence of structural defects

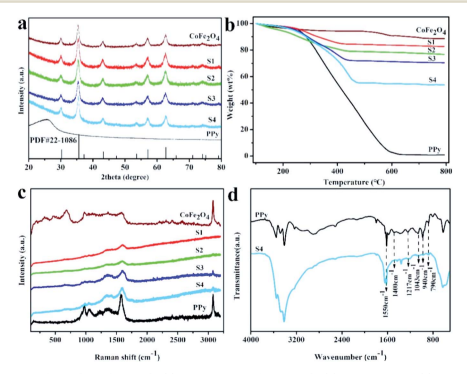


Fig. 3 Powder XRD patterns of PPy, CoFe₂O_{4,} and S4 (a); FT-IR spectra of PPy and S4 (b); Raman spectra (c) and TGA patterns (d) of PPy, CoFe₂O_{4,} and S4.

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and marginal unsaturated carbon, while the characteristic peak at 1580 cm⁻¹ is attributed to the G band (C=C stretching vibration), representing the in-plane stretching vibration of carbon atom sp2 hybrid.34,44 It can be seen from Fig. 3c that unloaded CoFe₂O₄ has no obvious G band or D band, whereas there is a significant double peaks after PPy loading, indicating the existence of PPy in the composites. In addition to the existing D band and G band, two peaks at 1045 and 977 cm⁻¹ are observed because of the existence of PPy with the polaron (NH⁺) and bipolaron structures (N⁺). It is worth noting that after the addition of CoFe₂O₄ to the PPy matrix, the characteristic absorption peak of PPy at 1579 cm⁻¹ was blue-shifted to 1592 cm⁻¹. This transfer may be owing to the interaction between CoFe₂O₄ and PPy backbones. The pure polypyrrole has a sharp G band and a flat D band, while for CoFe₂O₄@PPv, the relative intensity of the D band and the G band decreases, manifesting that the degree of disorder of PPy increases due to the introduction of CoFe₂O₄ nanoparticle.⁴⁵ FT-IR measurement was also performed. As shown in Fig. 3d, the absorption peak at 1550 cm⁻¹ is attributed to the anti-symmetric stretching of pyrrole ring due to the stretching vibration of the C=C bond; the absorption peaks observed at 1400 and 1043 cm⁻¹ correspond to the N-H and C-H deformation vibrations; and the peak at 1217 cm⁻¹ is derived from C-N⁺ stretching, 46,47 moreover 940 cm⁻¹ and 790 cm⁻¹ are the C-C bond vibration and the outof-plane vibration of C-H bond,41 respectively.

The specific surface area and pore size distribution of the samples can be determined by isothermal nitrogen adsorptiondesorption curves. The results as shown in Fig. 4a indicate that the curves of CoFe₂O₄, PPy, CoFe₂O₄@PPy own hysteresis loop consistent with the IV type adsorption-desorption behavior. The BET surface areas of pure CoFe₂O₄, PPy, CoFe₂O₄@PPy calculated by isotherms are 90.48, 8.65, 69.86 cm 2 g $^{-1}$, respectively. The coating of PPy reduces the specific surface area of CoFe₂O₄@PPy, which may be attributed to the filling or covering of the micropores of CoFe₂O₄. CoFe₂O₄ and CoFe₂O₄@PPy have a slight capillary condensation step in the relative pressure range of 0.44-0.87, indicating that the material has a large mesopores. 48 In addition, the hysteresis loop in the curve shows high permeability between the mesopores. Fig. 4b shows that the pore size is mainly distributed at 1-20 nm, which suggests that CoFe₂O₄ and CoFe₂-O₄@PPy are mainly mesoporous structures, giving them physical adsorption capacity.

The valence state and element ratio in CoFe₂O₄@PPy and CoFe₂O₄ were analyzed by XPS as shown in Fig. 5. From the full XPS spectra shown in Fig. 5a, the peaks of elements C, O, Fe, Co were observed at nearly 280, 530, 720 and 780 eV successively. Apart from those observed for CoFe₂O₄, the signal for CoFe₂-O₄@PPy emerged at approximately 400 eV is ascribed to nitrogen element in coated PPy layer. The magnification spectra of signals assigned to specific element C, N, O, Co and Fe were shown in Fig. 5b-f. C 1s XPS spectra in Fig. 5b was fitted by four component peaks at 284.4, 285, 286, and 288.5 eV, corresponding to C-C, C-H, C-N, C=O groups⁴⁹ respectively. The main peak at 399.8 eV in the deconvoluted N 1s XPS spectrum (Fig. 5c) is assigned to neutral amine nitrogen(-NH-),50,51 whereas the small peak at 401.6 eV could be originated from iron-nitrogen related bond (Fe-N type) which is regarded as catalytic active Fe-N sites. The O 1s spectra as shown in Fig. 5d can be divided into several peaks at around 530.36, 531.73 and 533.1 eV, which can be assigned to the surface adsorbed hydroxyl oxygen, water and crystal lattice oxygen O²⁻ (ref. 16) respectively. For CoFe₂O₄@PPy, the peak locating at 530.36 eV is weakened due to the cover of PPy layer. Notably, an additional peak at 531.1 eV appeared, probably due to the formation of Fe-O-C bond between PPy and Fe2+, which is beneficial for the rapid transfer of electrons during the photo-Fenton reaction.⁵² The PPy coating also reduced the relative strength of Fe and Co. From Fig. 5e, the three characteristic peaks with binding energies of 780.05 \pm 0.2 eV, 781.89 \pm 0.2 eV and 785.9 eV are ascribed to Co $2p_{3/2}$. The binding energies of 780.05 ± 0.2 eV and 781.89 \pm 0.2 eV are $ext{Co}^{2+}$ ions at octahedral and tetrahedral positions,⁵³ respectively, confirming the chemical state as Co(II). The signal at 785.9 eV is characterized to be the satellite peak of Co 2p_{3/2} main line. In the Fe 2p spectrum of CoFe₂O₄ (Fig. 5f), signals appear at 711.10 and 724.7 eV with the separation of \sim 13 eV are ascribed as Fe 2p_{3/2} and Fe 2p_{1/2}, respectively, indicating the presence of Fe³⁺. Furthermore, a satellite peak appears at the binding energy of 720.14 eV, which is a typical feature of Fe³⁺. However, the two significant peak positions of CoFe₂O₄@PPy shifted to a low binding energy by 1.35 eV with the separation of \sim 6 eV,⁵⁴ demonstrating the existence of Fe²⁺. The appearance of Fe²⁺ may be due to the reduction of Fe³⁺ by Py during the in situ growth of Py and then Fe2+ strongly coordinated with the nitrogen atom in PPy,35 controlled growth of PPy

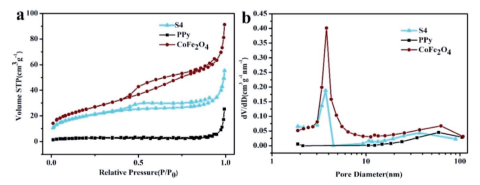


Fig. 4 Nitrogen absorption-desorption isotherms (a) and the pore size distribution curve (b) of pure CoFe₂O₄, PPy and S4 composite.

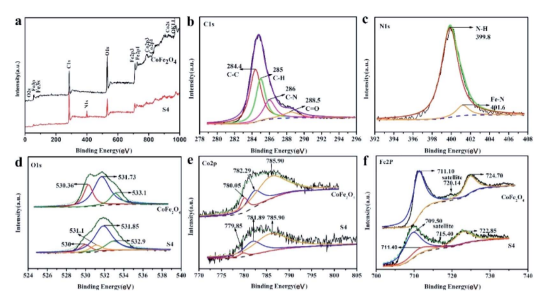


Fig. 5 XPS spectra of the pure $CoFe_2O_4$ and S4. Survey of the samples (a); C 1s narrow scan (b); N 1s narrow scan (c); O 1s narrow scan (d); Co 2p narrow scan (e) and Fe 2p narrow scan (f).

into the surface of CoFe₂O₄. Accordingly, Fe²⁺ is embedded in the interior of PPy which is consistent with the result of EDS.

The catalytic activity and reusability of catalysts in Fenton degradation of RhB

The abilities of different samples to degrade organic substance under dark Fenton, photocatalysis, photo-Fenton were detected using RhB as a model pollutant, and the results were shown as Fig. 6a–c. It can be seen from Fig. 6a that the adsorption of RhB is significantly increased with the incorporation of PPy, which may be attributed to the strong interaction between RhB and PPy. Compared with $CoFe_2O_4$ and PPy, PPy modified $CoFe_2O_4$ showed obvious dark Fenton activity when H_2O_2 was added in, as the concentration of RhB decreased gradually because of the Fenton-like degradation. The improved catalytic performance of $CoFe_2O_4$ @PPy and PPy may be attributed to the presence of Fe^{2+}

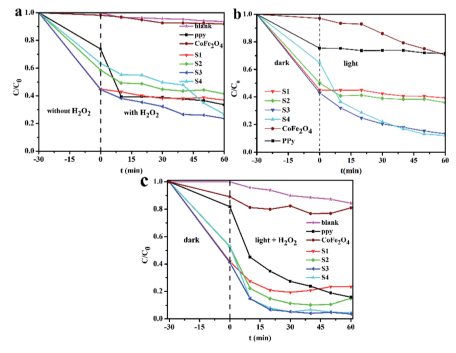


Fig. 6 Rhodamine degradation ability under dark Fenton for all samples (a); the photocatalytic degradation of RhB over all samples under visible light irradiation (b), the photo-Fenton degradation of RhB solution with the presence of H_2O_2 under visible light irradiation (c) ($C_{RhB} = 5 \text{ mg L}^{-1}$, $C_{H_2O_2} = 45 \text{ mM}$, pH = 7).

coordinated in PPy, which may be dissolved and released into the solution to catalyze the decomposition of H₂O₂, similar with that reported in the enzyme-mimic behavior. It is worth mentioning that PPy decolorized RhB by 50% in the first 10 minutes of dark Fenton. This may be due to the large amount of Fe²⁺ anchored into PPy during synthesis process. With the addition of H2O2 during the dark Fenton process, it is consumed in a short time. Subsequently, the ability of PPy to degrade RhB tends to be gentle, while other composites still maintain a stable activity. Among them, the dark Fenton ability of S1 and S2 is poor, which may be due to the relatively small amount of PPy coating. Although S3 has a higher degree of decoloration at 60 minutes compared with S4, we believe this is mainly due to the strong adsorption effect during dark process. Comparing the dark Fenton activity, S4 has a better catalytic ability, and the photocatalysis is consistent with this result. Fig. 6b shows that the photocatalytic ability of S4 is better than that of S3. This strongly indicates that S4 has improved electron-hole transport ability and more oxidizing substances are generated. The photo-Fenton degradation behavior shows significantly better performance than that of dark Fenton and photocatalysis shown as Fig. 6c. Similarly, the activity of CoFe₂O₄@PPy increases with the increase of polypyrrole

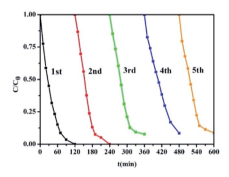


Fig. 7 Cycling runs for the photodegradation of RhB in the presence of S4 under visible light irradiation. ($C_{RhB}=10~mg~L^{-1}$, $C_{H_2O_2}=45~mM$, pH = 7).

content. Since S4 is the composite with the best synergy effect, we chose S4 as the main research object in subsequent experiments. In order to prove the non-selectivity of catalytic degradation, we also selected methyl orange and methylene blue as substances to be degraded, which showed outstanding photodegradation effect (Fig. S2–S4†).

The reusability of $CoFe_2O_4$ @PPy was investigated by cycling test. To reduce the influence of RhB adsorption in the degradation, the RhB concentration is increased to 10 mg L⁻¹ and the time of each cycle is set as 2 h. As shown in Fig. 7, the degradation efficiencies of RhB in these five cycles are 99.7%, 99.5%, 92.1%, 91.4% and 91.2%, respectively. Although the degradation efficiency was slightly reduced, the overall decolorization effect was still significant, demonstrating that the stability of the system is still brilliant. The slight decrease of decolorization rate may be owing to the adsorption of small molecules on the surface of $CoFe_2O_4$ @PPy after decomposition of RhB molecule, occupying the catalytic position on the surface active site. We compare this work with others, as shown in Table 1, which demonstrates that $CoFe_2O_4$ @PPy has better photocatalytic ability.

Degradation pathway and species of RhB

The photo-Fenton degradation mechanism of RhB may be that the RhB molecule is eventually oxidized. ⁵⁵ As the reaction progressed, the RhB solution decolored from rose red to a colorless chromophore. However, the decolorization of RhB does not mean complete mineralization, and the intermediate products produced during the degradation process may also be contaminated and toxic, and cannot achieve the purpose of our initial preparation of the catalyst. Therefore, during the experiment, we chose to scan a wavelength range of 200–650 nm. Fig. 8a shows that the peak at 553 nm is decreasing, and the peak at 259 nm, 310 nm, and 320 nm gradually disappears, indicating that the aromatic chain, N-position ethyl group and conjugated structure in the RhB molecule are gradually destroyed during the reaction resulting in small organic molecules. ⁵⁵ Due to the destruction of the C=N and C=O

Table 1 The comparison of this work with others	Table 1	The comparison	of this work	with others
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Catalyst	Dye	Reaction conditions	Degradation	Reusability	Ref.
$CoFe_2O_4@PPy$	RhB	[Dye] = 5 mg L; $[H_2O_2]$ = 45 mM; pH = 7; catalyst = 0.01 g	99.5% (30 min)	5 th (90%)	This work
Fe ₃ O ₄ /P(MAA-MBAA)-PPy/Au/void/TiO ₂	RhB	[Dye] = 5 mg L; pH = 3; catalyst = 4 mg	100% (90 min)	6 th (70.2%)	62
$Ag@CoFe_2O_4/Fe_2O_3$	RhB	[Dye] = 5 mg L; $[H_2O_2]$ = 225 mM; pH = 7; catalyst = 2 cm × 2 cm square	99.15% (1 h)		63
Reduced graphene oxide-CoFe ₂ O ₄	Malachite green	[Dye] = 5×10^{-6} M; [H ₂ O ₂] = 112.5 mM; pH = 7; catalyst = 0.05 g	100% (30 min)		64
rGO/PB/PPy	RhB	[Dye] = 10 mg L; $[H_2O_2]$ = 22.5 mM; pH = 6.18; catalyst = 0.01 g	95.2% (30 min)	4 th (91.5%)	65
$rGS/Fe_xO_y/NCL$	RhB	[Dye] = 10 mg L; $[H_2O_2]$ = 180 mM; pH = 7; catalyst = 0.03 g	100% (150 min)	5 th (94.6%)	66
PPy NF/Zn–Fe LDH	Safranin	[Dye] = 5 mg L; catalyst = 0.02 g; pH = 7	88% (120 min)	6 th (58%)	67

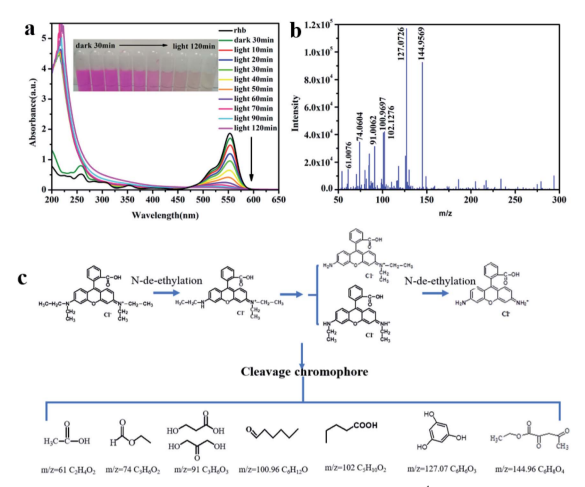


Fig. 8 The photocatalytic degradation of RhB over S4 under visible light irradiation ($C_{RhB} = 10 \text{ mg L}^{-1}$, $C_{H_2O_2} = 45 \text{ mM}$, pH = 7) (a). LC/MS spectra of end products of the photo-Fenton degradation of RhB (b), the degradation path of possible intermediates (c).

conjugated structures, the characteristic absorption peak at 553 nm gradually decreases.⁵⁶ The disappearance of the absorption peak at 259 nm indicates that the structure of the benzene ring has been destroyed. However, the peak at 225 nm has been rising. It is speculated that the conjugated structure of the RhB molecule is gradually destroyed during the reaction, and other non-polluting small molecules may be formed by deethylation at the N-position. In order to gain a deeper understanding of the colorless products after degradation, we conducted a LC/MS test. After two hours of visible light irradiation, the final colorless product of Rhodamine B degradation was used for LC-MS testing to analyze the components. From Fig. 8b, the final products were mainly small molecule compounds, which means RhB has been completely decomposed into low-toxic or even non-toxic substances. Based on the results of LC-MS, the structures of major intermediates can be inferred as Fig. 8c. The degradation process mainly involves: (i) at the beginning, the peaks differ exactly by 28 mass units successively, indicating sequential N-de-ethylation of RhB. (ii) The intermediate is further broken down into several compounds by the cleavage of the azo (C-N) bond. Fragmented compounds have no chromophore, resulting in colorless. A ring-opening reaction occurs, decomposing the intermediate

into small molecules. Further mineralization into $\rm H_2O$ and $\rm CO_2$ eventually.

The mechanism in performance enhancement

To deep understand the mechanism in performance enhancement, PL tests and quench experiments were conducted. The room temperature PL spectrum was used to study the separation and recombination rate of photoexcited charges of charge carriers. As shown in Fig. 9a, the PL spectrum of CoFe₂O₄ shows only one emission peak at 427 nm. Whereas, that of CoFe₂-O₄@PPy shows two emission peaks at 427 nm and 620 nm, which are attributed to the band PL and the exciton PL respectively.⁵⁷ Compared to the PL of CoFe₂O₄, the lower intensity of band PL at 427 nm in PL of CoFe₂O₄@PPy indicates that the load PPy contributes to the separation of the photogenerated charges, resulting less charge recombination. The incorporation of PPy also introduces in a broad emission peak range from 500 to 750 nm, which can be attributed to the oxygen-related defects (such as oxygen gaps).58 This observation reveals that the coating of PPy results defect-rich surface and oxygen vacancy of CoFe₂O₄, which are also evidenced by the XPS results. Quenching experiments were carried out to investigate the active species of photocatalytic properties. In the photoPaper

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Fig. 9 PL spectra of CoFe₂O₄ and S4 (a); trapping experiment of active species during the photocatalytic degradation of RhB over S4 under visible light irradiation (b) ($C_{\text{RhB}} = 10 \text{ mg L}^{-1}$, $C_{\text{H}_2\text{O}_2} = 45 \text{ mM}$, pH = 7).

Fenton process, superoxide radicals from oxygen reduction, photo-generated holes, and hydroxyl radicals from H₂O₂ decomposition and water oxidation are responsible for the degradation of organic pollutants. Therefore, EDTA as the electron donor, ethanol (EA) containing α -H as the free radical scavenger and benzoquinone (BQ) as the superoxide radical quencher were added into the photo-Fenton system individually. As shown in Fig. 9b, compared with the control experiment without any quencher, the RhB decolorization rate decreased by 57.74%, 69.25% and 76.16% after adding EA, EDTA and BQ respectively, revealing a complicate contribution of decolorization. According to the results of the quenching test, we speculated the mechanism of performance improvement. CoFe₂O₄ and PPy will generate photo-induced electrons and holes under light irradiation⁵⁹ (reaction (1)). Under light irradiation, the photo-generated electrons of CoFe₂O₄ move to the PPy shell and react with oxygen attached to the surface to generate Oo-(reaction (2)), which contributed to the degradation of organic matter. Simultaneously, Fe²⁺ located between the core and shell can form a Fenton reaction system with H₂O₂ to form Fe³⁺(reaction (3)). Hydrogen peroxide can be captured by Fe³⁺, generating Fe²⁺ and performing Fe²⁺/Fe³⁺ cycles (reaction (4)–(6)), which makes recycling and stability possible. Moreover, ·OH generated in this process is the most important active species according to the trapping experiment. The accumulated h⁺ can oxide the organic substrates directly (reaction (7)).

$$PPy + CoFe_2O_4 + h\nu \rightarrow e_{CB}^- + h_{VB}^+$$
 (1)

$$O_2 + e_{CB}^- \rightarrow O_2^- \tag{2}$$

$$Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + \cdot OH + OH^-$$
 (3)

$$Fe^{3+} + H_2O_2 \rightarrow Fe^{2+} + \cdot OOH + H^+$$
 (4)

$$Fe^{2+} + \cdot OOH \rightarrow Fe^{3+} + \cdot OOH^{-}$$
 (5)

$$Fe^{3+} + e_{CB}^{-} \to Fe^{2+}$$
 (6)

$$\cdot$$
OH + \cdot O₂⁻ + h⁺ + RhB \rightarrow degraded products (7)

Generally, the larger specific surface area can provide more active sites for photodegradation and promote photocatalytic reaction. However, the relatively poor catalytic performance of

CoFe₂O₄ reveals that the factor response for photo-Fenton activity is far beyond the specific surface area. The quench experiments also indicate that in CoFe2O4@PPy catalyzed photo-Fenton, the contributor on degradation is more than hydroxyl radicals. Therefore, the improvement of degradation performance may be attributed to several factors. Firstly, there is an appropriate amount of Co²⁺, Fe²⁺ and Fe³⁺ embedded in the interior of PPy during the polymerization, which may accelerate the decomposition of H2O2 through Fenton way, which may mainly contribute to the formation of hydroxyl radicals. Secondly, oxygen vacancy and lattice defect may be introduced into the CoFe2O4 during the PPy coating. The existence of oxygen vacancy enhances the adsorption of substrates, provides more active sites, and promotes the photo-generated charge separation. 61 Thirdly, as an organic semiconductor, PPy not only promotes the light harvesting, but also accelerates electron transfer in the re-generation of active site for Fenton reaction through redox pairs. In summary, the improvement on degradation activity of photo-Fenton can be attributed to the combination of improved Fenton oxidation and additional photo-catalyzed degradation provided by coated PPy layer on CoFe₂O₄ microspheres.

Experimental

Materials and reagents

 ${\rm CoSO_4\cdot 7H_2O}$, ${\rm FeCl_3\cdot 6H_2O}$, ethylene glycol (EG), pyrrole (Py), rhodamine B (RhB), hydrogen peroxide (${\rm H_2O_2}$, 30%), hydrochloric acid (HCl) were obtained from Aladdin (China). Sodium dodecyl sulfate (SDS), sodium acetate anhydrous (NaAc, AR), polyethylene glycol (PEG), ethylenediaminetetraacetic acid (EDTA), methanol (MA) and benzoquinone (BQ) were purchased from Macklin (China). All chemicals were used without further purification.

Preparation of CoFe₂O₄@PPy

CoFe₂O₄ was prepared according to the previous work, ^{43,68} and the details are shown in the ESI.† The CoFe₂O₄@PPy was synthesized *via in situ* Fenton oxidation polymerization of pyrrole. Typically, 50 mg CoFe₂O₄, 100 mg SDS and various amounts of pyrroles were mixed with 40 mL water under magnetic stirring to form a homogeneous brown dispersion. To the dispersion cooled under ice bath, 280 μ L HCl and 2 mL 30% H₂O₂ were successively dropped in, and vigorous stirring was continued for other 3 hours. The product was collected by centrifugation and repeated washed, then vacuum dried at 65 °C for 12 h. For comparison, PPy nanoparticles were also prepared according to the previous report. In this synthesis procedure, the molar ratios of CoFe₂O₄/PPy were adjusted at 1:2,1:4,1:6 and 1:8, and the corresponding CoFe₂O₄@PPy samples were named Sx (x = 1–4).

Characterization

The morphology and composition of samples were observed by scanning electron microscope (SEM, SU-70) equipped with EDS and Transmission electron microscope (TEM) (JEM-21200F).

The magnetic properties of $CoFe_2O_4$ and $CoFe_2O_4$ @PPy composites were measured in a vibrating sample magnetometer (Quantum Design Corporation, USA) with a maximum applied field of ± 2 T. X-ray diffraction (XRD) curves were recorded with an X-ray diffractometer (Smart Lab) using Cu K α radiation (λ = 1.5418 Å). The chemical structure of samples was characterized by FT-IR spectrometer.

(Nicolet 6700) and FT Raman Microscope (RENIDHAW invia). The chemical valence state of samples was analyzed by XPS (PHI5000 Versa Probe II). TGA was performed to understand the content of PPy on a thermal analyzer (TGA 55) at a heating rate of 10°C min⁻¹ from ambient to 800 °C in air. The nitrogen adsorption desorption isotherms were measured at 77 K using a Micromeritics ASAP 2460 system, and the surface area and pore size distribution were calculated using the Brunauer-Emmett-Teller (BET) and Barrett-Joyner-Halenda (BJH) method, respectively. The UV-Vis absorption spectrum (UV-Vis) was characterized with a UV-Vis spectrometer (PerkinElmer Lambda 950). Liquid chromatography tandem mass spectrometry (LCMS) with OTRAP6500+ was employed to determine the final product of rhodamine decolorization. Photoluminescence (PL) emission spectra was performed on a Renishaw in via spectrometer with 285 nm UV light as the excitation source.

Degradation of RhB by heterogeneous catalyst

The photo Fenton activity was assessed by the degradation of rhodamine B (Rh B) under visible light irradiation with a 300 W xenon lamp with a 420 nm cut-off filter. Typically, 10 mg sample was dispersed in 50 mL 10 mg L⁻¹ of RhB in a baker with cycling water cooling and magnetic stirring. After 30 min for adsorption in dark, 200 µL of 30% H₂O₂ was added in and light on. Solution was taken out at preset intervals and centrifuged prior to UVvisible spectrometry analysis. The dark Fenton activity was assessed similarly except lighting off. The catalyst after 1 cycle of 2 h degradation test was washed with ethanol and dried thoroughly, then used for the other cycle of degradation to evaluate its reusability. Photo degradation in the presence of ethylene diamine tetra-acetic acid (EDTA), methanol (MA) and benzoquinone (BQ) as a quencher for holes, hydroxyl radicals and superoxide radicals respectively, was also carried out to study the active species during degradation.

Conclusion

In summary, CoFe₂O₄@PPy prepared *via* a simple *in situ* Fenton oxidization polymerization coating was applied for the photo-Fenton degradation of organic pollutants. The CoFe₂O₄ core in PPy coating provides outstanding magnetic recyclability (Fig. S5†) for the catalysts as well as provides the ion source for catalyzed decomposition of H₂O₂. The prepared CoFe₂O₄@PPy exhibited improved performance in Fenton and photo-Fenton degradation of RhB, which was superior to those of pure CoFe₂O₄ and PPy. The LC-MS of RhB degradation residues indicates that only low-toxic species are reserved. The possible performance enhancement mechanism is investigated *via* degradation measurement in the presence of scavenger of

reactive species and PL spectra of catalysts. Photodegradation experiments of methyl orange and methylene blue confirm the wide applicability of the catalyst. It is deduced to be attributed to the synergistic effects of the following factors: firstly, the rapid Fenton oxidation brings about massive defects in PPy layer, thereby strongly improves its adsorption to RhB; secondly, the *in situ* dissolved Fe³⁺ is also reduced to Fe²⁺ and embedded into the PPy layer through N–Fe coordination thus improves the activity in Fenton reaction; thirdly, the presence of PPy coating layer prevents the agglomeration of magnetic particles; the last but the most important, the PPy layer on $CoFe_2O_4$ enhances the absorption of light, inhibits the combination of photo-generated charge carriers and promotes the electron transfer for H_2O_2 decomposition.

Conflicts of interest

There are no conflicts to declare

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