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${\rm FeN_4}$ -doped carbon nanotubes derived from metal organic frameworks for effective degradation of organic dyes by peroxymonosulfate: Impacts of ${\rm FeN_4}$ spin states

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Meng Li ^{a,d,1}, Zilong Li ^{a,1}, Xiaolong Yu ^b, Yinlong Wu ^a, Cehui Mo ^a, Mi Luo ^c, Ligui Li ^{c,*}, Shaoqi Zhou ^d, Qiming Liu ^e, Nan Wang ^{a,*}, King Lun Yeung ^f, Shaowei Chen ^{e,*}

^a Guangdong Provincial Research Center for Environment Pollution Control and Remediation Materials, College of Life Science and Technology; Guangdong Provincial Key Laboratory of Optical Fiber Sensing and Communications, Siyuan laboratory, Guangzhou Key Laboratory of Vacuum Coating Technologies and New Energy Materials, Guangdong Provincial Engineering Technology Research Center of Vacuum Coating Technologies and New Energy Materials, Department of Physics, Jinan University, Guangzhou, Guangdong 510632, PR China

^b Guangdong Provincial Key Laboratory of Petrochemical Pollution Processes and Control, School of Environmental Science and Engineering, Guangdong University of Petrochemical Technology, Maoming, Guangdong 525000, PR China

^c Guangzhou Key Laboratory for Surface Chemistry of Energy Materials, New Energy Research Institute, School of Environment and Energy, South China University of Technology, Guangzhou 510006, PR China

^d College of Resources and Environmental Engineering, Guizhou University, 2708 Huaxi Road, Guiyang 550025, PR China

^e Department of Chemistry and Biochemistry, University of California, 1156 High Street, Santa Cruz, CA 95064, USA

^f Department of Chemical and Biological Engineering, The Hong Kong University of Science and Technology, Clear Water Bay, Kowloon, Hong Kong SAR, PR China

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ABSTRACT

Development of low-cost, high-performance catalysts for the effective degradation of organic dyes is of both fundamental and technological significance for environmental remediation. Herein, FeN₄-doped carbon nanotubes exhibit a remarkable catalytic activity towards the degradation of acid orange 7 (AO7) in the presence of peroxymonosulfate (PMS). This is accounted for by the abundant well-dispersed FeN₄ sites and appropriate adsorption energy. Radical quenching experiments and electron paramagnetic resonance measurements suggest that singlet oxygen is the main reactive oxygen species. ⁵⁷Fe Mossbauer spectroscopy measurements show that the FeN₄ moieties contain multiple spin states, and in conjunction with density functional theory calculations, confirm that FeN₄ with a low/medium spin state serves as the key sites for PMS activation and hence AO7 degradation. Results from this study highlight the critical role of spin states in FeN₄ in the generation of singlet oxygen for effective degradation of organic pollutants.

1. Introduction

The accumulation of hazardous and toxic contaminants, such as pesticides, antibiotics [1], dyes [2], and endocrine disruptors [3], in surface water and groundwater has caused severe environmental pollution issues. Various technologies, such as biological degradation [4], physical adsorption [5], membrane filtration [6], electro-Fenton [7], and advanced oxidation processes (AOPs) [8], have been explored to remove organic pollutants from the environment. Among these, AOPs have shown a high efficiency in removing organic contaminants from water due to the generation of highly reactive species [9]. In fact,

sulfate-based AOPs have received extensive attention in the degradation of organic pollutants, due to strong oxidation ability, wide pH range, low cost, and high stability [10,11]. In this process, sulfate radicals (SO_4^-) can be generated by the activation of peroxydisulfate (PDS) and peroxymonosulfate (PMS) by thermal control [12], transition metals [13], UV [14], microwave irradiation [15], etc, as manifested in eqs. (1, 2), and responsible for the oxidative degradation of organic pollutants [16]. Nevertheless, the application of such processes is generally limited due to the high cost and energy input, non-ideal activation performance, and release of metal ions that may lead to secondary pollution [17-20]. Consequently, development of metal-free heterogeneous catalysts for

* Corresponding authors.

¹ These authors contributed equally.

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E-mail addresses: esguili@scut.edu.cn (L. Li), nanwang@jnu.edu.cn (N. Wang), shaowei@ucsc.edu (S. Chen).

PMS activation is urgently needed.

$$S_2 O_8^{2-} + thermal/UV/microwave/carbon \rightarrow 2SO_4^{-}$$
(1)

$$S_2 O_8^{2-} + M^{n+} \to M^{(n+1)+} + S O_4^{2-} + S O_4^{\bullet-}$$
(2)

Over the past few years, metal-free carbon-based materials have been used as PMS activators for the degradation of organics pollutants [21], such as carbon nanotubes (CNT) [22], activated carbon (AC) [23], graphene oxide (GO) [24], and nanocarbons [25]. Yet, the catalytic performance has remained subpar as compared to those of the metalbased counterparts. Also, whereas a good removal efficiency can be achieved by the addition of a large amount of oxidants [26], the accumulation of excessive sulfate ions may lead to corrosion of the pipeline in the drinking water and sewer systems [27]. Moreover, the stability and reusability of carbon-based materials is usually poor, limiting their practical applications. Consequently, it is important to develop metalfree PMS catalysts with excellent high catalytic activity and reusability. Recently, numerous studies have been carried out to enhance the catalytic activity of carbon-based materials by the incorporations of heteroatom into the carbon frameworks, such as nitrogen, sulfur, phosphorus, and boron [28-30], wherein heteroatom-doping is recognized as an effective method by regulating the charge distribution of the sp²-hybridized carbon frameworks and providing abundant active sites for catalysis [31]. For example, nitrogen-doped single-walled carbon nanotubes can significantly improve the catalytic activity for phenol degradation by activating PMS up to 57 times that of pristine carbon nanotubes [32]. Nevertheless, the catalytic activity of metal-free catalysts is usually suppressed via a non-radical process. This can be mitigated by further structural engineering such that practical application is feasible.

One strategy is to embed transition metal-nitrogen coordination centers within a porous carbon scaffold [33]. The resulting metalnitrogen-carbon (M–N–C, M = Mn, Co, Fe, Ni, etc.) catalysts possess a high density of active sites and exhibit promising catalytic activity in various reaction processes [34-36]. Of these, the nitrogen-coordinated iron (FeN_x) moieties are of particular interest [37], which can impact the electronic structure of the catalyst surface and electrical conductivity. Both factors are conducive to PMS activation [38], which is believed to follow three reaction pathways, i.e., radicals, singlet oxygen, and direct electron transfer [39]. Of these, singlet oxygen (¹O₂) is known to exhibit high selectivity and mild redox ability (2.2 V) for the degradation of organic contaminants, in comparison with other radical species, such as •OH (2.7 V) and SO_4^{--} (2.5–3.1 V) [40].

These Fe-N-C catalysts are usually prepared by fixing Fe atoms within a porous support, e.g., graphene oxide (GO), activated carbon (AC), and covalent organic frameworks [41]. Yet the preparation generally entails a complex process, producing a large and uncontrollable structure [42]. Metal-organic frameworks (MOFs), a class of porous crystalline materials, represent an effective alternative. MOFs are constructed by the self-assembly of transition metals and organic ligands [43], and have been extensively explored for various applications, such as molecular sensing, catalysis, pollutant adsorption, and gas separation, due to their excellent conductivity, high catalytic activity, good absorbability, and large surface area [44]. In addition, MOFs can serve as self-sacrificial templates or precursors to synthesize carbon composites based on the unique structure of metal clusters and ligands [45]. For instance, Zn-based MOF (ZIF-8) with a cavity structure can incorporate Fe-containing salts for the preparation of FeN_x-doped carbon composites [46]. As the Zn species in ZIF-8 can be evaporated at elevated temperatures, a porous carbon skeleton is formed with a large surface area [39], where the Fe species are typically in form of FeNx and/or Fe nanoparticles (NPs) [47]. Note that the former is catalytically active whereas the latter inactive in PMS activation [48]. In most prior studies, Fe (NO₃)₃, FeSO₄ and FeCl₃ have been commonly used as the Fe precursors, where Fe(OH)3 is generally produced at high temperatures increasing

the amount of Fe NPs [49], and the number of active FeN_x species is in general limited by controlled pyrolysis of these Fe salt precursors [50].

In recent years, functional composites of M-N-C composites have been confirmed as an effective catalysts to activate persulfate [51-53], and the MN_x coordination moieties are found to be responsible for PMS activation and contaminant degradation [50,54]. For instance, Li et al. reported that high-performance Fenton-like Fe-N-C derived from Fedoped zeolitic imidazolate framework showed a significantly enhanced performance in the degradation of BPA, as compared to Fe nanoparticles-loaded N-doped carbon [50]. Note that the FeN_x moieties can regulate the electronic structures and electrical conductivity of the nanocomposite for the effective activation of PMS [47,55]. However, studies have been scarce focusing on the impact of the FeN_x spin state on the catalytic performance, which are known to influence the adsorption of reactants and key intermediates. This is the primary motivation of the present study.

Herein, FeN4-doped carbon nanotubes are derived from MIL-101 by ball milling and pyrolysis and exhibit an effective performance in the activation of PMS for the degradation of organic dves. The obtained catalysts retain the original morphology of ZIF-8 with a high surface area and abundant internal channels. In addition, due to the space confinement effect, FeN₄ sites are effectively isolated within the carbon framework, and no Fe NPs are formed. The resulting FeNC nanocomposites exhibit apparent catalytic activity in the activation of PMS and the degradation of a range of organic dyes, such as acid orange 7 (AO7), methyl red (MR), rhodamine B (RhB), acid orange G (AOG), and methyl orange (MG). Radical quenching experiments and electron paramagnetic resonance (EPR) spectroscopy measurements show that singlet oxygen is the reactive oxidative species (ROS) responsible for the catalytic activity. In conjunction with DFT calculations, medium- and low-spin FeN₄ is identified as the most active species that facilitate the adsorption and activation of PMS and ultimately the degradation of organic pollutants. Furthermore, the degradation mechanism and pathway are proposed based on the identification of key reaction intermediates by gas chromatography-mass spectrometry and Fukui index which suggests the atomic sites within the dye molecules for radical attacks.

2. Experimental section

2.1. Materials

Ferric chloride hexahydrate (FeCl₃·6H₂O, 99.5%), zinc nitrate hexahydrate (Zn(NO₃)₂·6H₂O, 99.5%), *p*-phthalic acid, dimethylformamide (DMF), ethanol (EtOH), methanol (MeOH), *tert*-butanol (TBA), 5,5-dimethyl-1-pyrroline N-oxide (DMPO), and 2,2,6,6-tetra-methyl-4-piperidone (TEMP) were purchased from Aladdin (Shanghai, China). Oxone (2KHSO₅•KHSO₄•K₂SO₄), L-histidine, *p*-benzoquinone, AO7, MR, RhB, AOG, and MG were obtained from Sigma-Aldrich (Shanghai, China). All chemicals in this study were used directly without further purification. Water was supplied with a Barnstead Water Purification System (18.2 MΩ cm).

2.2. Synthesis of catalysts

Synthesis of MIL-101

MIL-101 was prepared by following a literature procedure [56]. In brief, 1.71 g of FeCl₃·6H₂O and 0.41 g of *p*-phthalic acid were added into 30 mL of DMF under stirring for 30 min, and the resulting mixture was heated in a hydrothermal reactor at 110 °C for 20 h. The precipitate was rinsed several times with ethanol, DMF and deionized water, and dried in a vacuum oven at 70 °C overnight.

Synthesis of ZIF-8

ZIF-8 was prepared by using the traditional solution method [57]. Experimentally, 1.68 g of $Zn(NO_3)_2$ ·6H₂O was dispersed into 20 mL of MeOH, which was then combined with a 60 mL MeOH solution

containing 4 g of dimethylimidazole under magnetic stirring. The solution turned milky white, and was let to stand overnight at room temperature. The bottom product was collected by centrifugation, rinsed with distilled water and absolute ethanol, and then dried under vacuum at 70 $^{\circ}$ C overnight.

Synthesis of FeN4-doped carbon nanotubes

The two precursors of MIL-101 and ZIF-8 obtained above were mixed and ground by ball milling under the protection of a N₂ atmosphere at different mass ratios (1:15, 1:20, and 1:25), and the powders were placed in a tube furnace and heated at 900 °C under a N₂ atmosphere for 2 h, affording FeN₄-doped carbon nanotubes, which were denoted as FeNC-15, FeNC-20, and FeNC-25, respectively. The samples were then subject to a second heating treatment in an NH₃ atmosphere at 950 °C for half an hour to increase the amount of N doping. Preparation of Fe-Nx-fer catalyst, FeNC-fer uses the same preparation method as FeNC-20 but uses ferrocene or FeCl₃·H₂O instead of MIL-101 for the iron precursor.

2.3. Characterization

Transmission electron microscopic (TEM) measurements were conducted on a Tecnai G2-F20 equipped with an energy-dispersive X-ray spectroscopy (EDS) detector at the acceleration voltage of 100 kV. The TEM samples were prepared by dropcasting a catalyst dispersion onto a copper grid coated with a holy carbon film. The morphology and structure of the samples were studied by scanning electron microscopy (SEM, Gemini FE-SEM). Powder X-ray diffraction (XRD) patterns were recorded with a Rigaku-D/Max-IIIA diffractometer using Cu K_a radiation. X-ray photoelectron spectroscopy (XPS) experiments were carried out on a Thermo-ESCA-Lab 250 instrument with Al Ka radiation. The Brunauer-Emmett-Teller (BET) surface area was determined by using a Kubo-X1000 instrument at 77 K. Raman spectra were recorded on a RENISHAW inVia instrument with an Ar laser source of 488 nm in a macroscopic configuration. Inductively coupled plasma atomic emission spectrometric (ICP-AES) measurements were carried out with a Libertyax instrument. The X-ray absorption spectroscopys (XAS) measurements and standard contrast samples used were provided by Taiwan Light Source.

2.4. Catalytic degradation experiment

The catalytic degradation experiments were carried out in a 300 mL flat bottom breaker at ambient temperature at a stirring speed of 350 rpm. The degradation reaction was initiated by adding a calculated amount of PMS and FeNC catalysts into the dye solution. The solution pH was adjusted by NaOH (10 mM) and H₂SO₄ (10 mM). An adsorption equilibrium between the catalysts and target contaminants was achieved by stirring the mixture for 30 min. During the catalytic reaction, 1 mL of the sample solution was taken out at certain time points, and filtered with a 0.25 µm polytetrafluoroethylene (PTFE) Millipore membrane, before the optical absorbance was quantified. To test the durability and stability of the catalysts, the catalyst samples were collected by centrifugation from the reaction vessel, rinsed with deionized water, and used for repeated dye degradation. In the degradation process of AO7, the corresponding Total Organic Carbon (TOC) was measured by a TOC analyzer (Shimadzu, Kyoto, Japan) at certain time intervals. The reaction conditions included the dye concentration of 50 mg L⁻¹, catalyst concentration of 2 mg L⁻¹, PMS concentration of 0.5 mM, and solution pH = 7.

2.5. Analysis

The dye concentration was determined via the spectrophotometric method on a Shimadzu U2800 UV–vis spectrophotometer. For MR, RhB, AOG, MG, and AO7, their maximum adsorption wavelengths are 527, 553, 476, 464, and 484 nm, respectively. EPR measurements were

conducted on a Bruker A300 spectrometer with TEMP and DMPO as the spin-trapping agents. The degradation dynamics of AO7 was fitted with a pseudo-first-order kinetic equation, $lnC/C_0 = -kt$, where k was the reaction rate, t was the reaction time, and C and C_0 were the concentration of AO7 at time t and 0, respectively [58-60].

The degradation products of AO7 were determined by gas chromatography-mass spectrometry (GC–MS, 7890A-5975C, Agilent, USA) equipped with an HP-5MS capillary column (30.0 m × 250 µm × 0.25 µm, Shimadzu, Japan). High-purity helium was utilized as the carrier gas at the rate of 1 mL min⁻¹. The column temperature was initially set at 60 °C and held for 5 min, then raised to 280 °C at a rate of 10 °C min⁻¹ and held for 10 min, next to 260 °C, finally to 280 °C. The MS analysis was carried out in the EI mode (70 eV), and ion source and quadrupole temperatures were set at 200 and 150 °C, respectively. The scanning mass range was 35 to 500 *m/z*, and the corresponding degradation products were identified utilizing the MS database and NIST11 search as references.

2.6. Computational methods

The molecular orbital analysis was carried out by utilizing the Gaussian 09 program [61], and the optimized structural geometry with the minimum energy was conducted at the b3lyp/6–31 g* level. The lowest unoccupied molecular orbital (LUMO) and the highest occupied molecular orbital (HOMO) were obtained from the output files of Gaussian 09. The Fukui function values of electrophilic reaction (*f*) were calculated based on the charge distribution from Multiwfn 3.7 [62].

DFT calculations were performed with the CASTEP code of the Materials Studio package of Acclerys Inc. In all calculations, the generalized gradient approximation and the projector augments wave pseudopotentials with the exchange and correlation in the Perdew-Burke-Ernzerhof were employed. The plane-wave cutoff energy was set at 450 eV. The convergence of forces and energy on each atom during structure relaxation were set to 0.03 eV $Å^{-1}$ in force and 10^{-6} eV in energy, respectively. The Brillouin zone was sampled with a $3 \times 3 \times 1$ Monkhorst-Pack k-point grid. The optimized structure sizes for pyrrolic, graphitic, pyridinic and FeN₄ were 12.26 \times 12.26 \times 15 Å, and Fe₂O₃(001) and Fe(110) were 10.07 \times 10.07 \times 20 Å and 9.49 \times 9.49 \times 20 Å, respectively. The vacuum space along the z[1] direction was more than 15 Å, to avoid interplanar interactions. For geometric optimization of the slab models, the top two layers were allowed to relax. Van der Waals (vdW) interaction was taken into account at the DFT-D3 level as proposed by Grimme. The adsorption energy was defined as $E_{ads} = E_{total}$ – ($E_{substrates}$ + E_{PMS}), where E_{total} , $E_{substrates}$, and E_{PMS} denote the total energy of substrates with PMS, substrates, and PMS, respectively.

3. Results and discussion

3.1. Structural characterization

As shown in Fig. 1, the FeNC nanocomposites were prepared by ball milling of two precursors of ZIF-8 and MIL-101 followed by controlled pyrolysis of the mixture. ZIF-8 is a rhombic dodecahedron framework with a large specific surface area (crystal size 2 nm), and MIL-101 is a rhombic crystal framework containing unique monodisperse Fe centers and carboxylic acid bridge ligands (crystal particles about 30 nm in size, Figure S1). Their XRD patterns are consistent with those reported in the literature [63] (Figure S2). Ball milling helped create uniform mixing of the elements (i.e., C, N, Fe, etc.) in the sample, and pyrolysis facilitated the formation of a carbon skeleton with a tubular structure, which exhibited a large specific surface area, and contained abundant FeN₄ moieties. Furthermore, the second heat treatment in an ammonia atmosphere effectively increased the contents of Fe-N and C-N bonds, and the microporosity of the samples.

The crystalline structure of FeNC-20 was then characterized by XRD measurements. From Figure S2a, one can see that the FeNC samples all



Fig. 1. Schematic illustration of the preparation of FeN₄-doped nanotubes.

exhibited only two intense diffraction peaks at $2\theta = 26^{\circ}$, which can be indexed to the (002) lattice planes of graphite carbon (PDF#75–1621), indicating successful transformation of the MOF precursors to graphitized carbon. The additional diffraction peak at $2\theta = 44^{\circ}$ likely arose from metal nanoparticles of iron (PDF#89–7194) or Fe_xC (PDF#52–0512), which is consistent with the results of TEM measurements (Fig. 2). The pyrolytic product of MIL-101 exhibited similar XRD patterns, whereas only a carbon diffraction peak was observed for ZIF-8 after pyrolysis (Figure S3).

The structures of the obtained nanocomposites were first examined by TEM measurements. From Fig. 2a, 2b and 2d, FeNC-20 can be seen to exhibit a porous network of CNTs with a bamboo-like structure featuring a tube diameter of about 100 nm and shell thickness of ca. 6 nm. From the high-resolution TEM image in Fig. 2c, one can see that FeNC-20 displayed well-defined lattice fringes, with an interplanar spacing of ca. 0.40 nm that is consistent with the (002) crystal planes of graphitic carbon (card #75–1621). Similar structures were observed for FeNC-15 and FeNC-25 (Figure S4). Note that a small number of dark-contrast nanoparticles can also be identified in the TEM image of FeNC-20 (Fig. 2e); and elemental mapping analysis (Fig. 2f-2i) suggest that these are most likely Fe or Fe₃C nanoparticles, embedded within a N-doped carbon matrix. Additionally, isolated Fe can be seen to be distributed rather uniformly across the sample, along with C and N, suggesting the formation of FeNx moieties within the carbon nanotubes.

The nanotubular structure is also evident in SEM measurements (Figure S5). For the samples prepared at either higher or lower Fe feed (i. e., FeNC-15 and FeNC-25), the fraction of nanotubes was markedly lower, and the samples consisted mostly of irregular aggregates. This led to a reduced specific surface area and likely compromised the accessibility of the active sites and hence the catalytic activity (vide infra). In addition, the SEM image of ZIF-8, ZIF-8–900, MIL-101, and MIL-101–900 (Figure S6a-6d) confirm that ball milling can facilitate the



Fig. 2. (a-d) Representative TEM images and (e-i) EDS elemental maps of FeNC-20.

formation of the nanotubes. As is shown in Figure S7, the SEM images of FeNC-fer (uses ferrocene or $FeCl_3 \cdot H_2O$ as precursor) reveal the advantages of MIL-101 as an iron source for the formation of carbon nanotube structures.

From the nitrogen adsorption–desorption isotherms (Figure S8a-8b), FeNC-20 can be seen to show an obvious H₄ hysteresis loop, indicating the formation of micropores and mesopores in the sample (Figure S8b) [64], with a specific surface area of 926.3 m² g⁻¹, more than twice those of other samples in the series (inset to Figure S8a). The abundant micropores in FeNC-20 are anticipated to facilitate access to the active sites [28,65]; and mesopores are known to be advantageous for mass transfer of reactants and products (Figure S8c-8d).

Further structural insights were obtained in Raman spectroscopic measurements. From Figure S9, the three FeNC samples can be seen to display two vibrational bands at approximately 1350 and 1595 cm⁻¹, due to the structural defects in the graphitic structure (D band) and graphitic sp² C (G band), respectively [66], with an intensity ratio (I_D/I_G) of 1.196, 1.20, and 1.197 for FeNC-15, FeNC-20, and FeNC-25, respectively. This indicates a similar defective structure among the series of samples (slightly more defective with FeNC-20 than the other two).

In ⁵⁷Fe Mössbauer spectroscopy measurements (Fig. 3a-3c), FeNC-20 was found to contain two components, FeN₄ and Fe/Fe₃C, at different spin states: D₁ (Fe^{II}N₄, low spin), D₂ (Fe^{II}N₄, medium spin, like FePc), D₃ (N-Fe^{II}N₂₊₂, high spin) and α -iron or iron carbide [67]. Note that the quadrupole doublet of D₂–FeN₄ with an isomer shift (IS) of 0.27 mm s⁻¹ has been identified to be the main source of activity for heat-treated porphyrins [67,68]. The fact that only a sextet, but no obvious singlet, was observed in the Mössbauer spectra suggests that the content of the

superparamagnetic iron was relatively low in the catalyst. That is, the FeNC samples were composed primarily of FeN₄ moieties and only a small amount of α -Fe or iron carbide [67,69], as manifested in TEM measurements (Fig. 2).

Furthermore, in conjunction with ICP-AES measurements, the fractions of the Fe species in different spin states (Fig. 3d and Table S1) were quantitatively assessed. From Fig. 3e, one can see that FeNC-20 primarily contained low-spin FeN₄ (D₁, 0.641 at%) and medium-spin FeN₄ (D₂, 0.223 at%), which are markedly lower in FeNC-15 (0.218, 0.195 at %) and FeNC-25 (0.479 and 0.131 at%). In addition, in the latter two samples, there is a substantial amount of α -Fe or Fe_XC (0.321 and 0.501 at%), respectively.

The elemental composition and valence state of the samples were then evaluated by XPS measurements. From the survey spectra (Figure S10), the elements of C, N, O, and Fe can be readily identified at 284.8, 399.8, 531.5, 712.2 eV, respectively, consistent with the incorporation of Fe and N into the carbon matrix; and from the integrated peak areas, the elemental compositions were estimated and listed in Table S2. One can see that the series of FeNC samples exhibited rather consistent elemental compositions, and FeNC-20 contained a slightly higher N (7.10 at%) and Fe (0.84 at%) content than others. In the highresolution XPS spectra of the N 1 s electrons (Fig. 3f), deconvolution yields five peaks at 398.4, 399.7, 400.9, 402.0 and 405.1 eV, corresponding to the pyridinic, Fe-Nx, pyrrolic, graphitic and oxidized N (Fig. 3g, Table S3), respectively [28,70-73]. Figure S11a shows the C1 s spectra of the FeNC series, where sp² C represents the dominant species, consistent with the successful graphitization of the MOF precursors by pyrolysis. The Fe 2p spectra was shown in Figure S11b, where the peaks at 708.4, 710.3, 711.9, 714.6 and 718.1 eV were attributed to Fe-N, Fe



Fig. 3. 57 Fe Mössbauer transmission spectra of (a) FeNC-15, (b) FeNC-20, and (c) FeNC-25. (d) Schematic representation of Fe²⁺ in high- and low-spin state, (e) the relative atomic contents of different spin states FeN₄ in the three samples, (f) High-resolution XPS scans of the N 1 s electrons of FeNC-15, FeNC-20, and FeNC-25, and (g) Schematic illustration of the five nitrogen doping configurations in carbon.

(II), satellite of Fe(II), Fe(III), and satellite of Fe(III), respectively [74,75]. In addition, the Fe-N content of the three samples was estimated to be 0.92 at.%, 1.33 at.% and 0.84 at.% (Table S3), respectively. That is, the Fe-N content of the FeNC-20 sample is the highest among the series of samples. This is in good agreement with the results from Mössbauer spectroscopy measurements where FeNC-20 contained the highest concentration of low-/medium-spin species (Fig. 3a-3e).

The electronic structure and coordination configuration of the FeNx moiety in FeNC-20 was then examined by synchrotron-based X-ray absorption spectroscopy (XAS) measurements. Fig. 4a shows the normalized Fe K-edge X-ray absorption near-edge structure (XANES) spectra of FeNC-20 and reference samples. It can be seen that FeNC-20 and FePc displayed a similar absorption edge, white line, and post-edge oscillations, suggesting a rather consistent atomic structure and chemical states of the Fe atoms in the samples. In fact, in the k³-weighted Fourier transform (Fig. 4b) and wavelet transform (Fig. 4d-f and Figure S12), FeNC-20 exhibits a main peak at 1.58 Å in R space, which is close to that of FePc (1.57 Å). To clearly analyze the coordination structure of the FeNx active site, we first confirmed the range of fitting with the help of k-q comparison (Figure S13). Based on the fitting of the k^2 -weighted Fourier transform of the EXAFS data in Fig. 4c, the coordination number (N) of the Fe center was estimated to be about 6, and the bond length between the Fe atom and the four N atoms was ca. 2.0 Å (Table S4, with the misfit, R factor, of only 0.016%). Meanwhile, from the structural model in the inset to Fig. 4c, one can see that there is an additional oxygen atom adsorbed onto the central iron atom with a Fe-O bond length of 1.2 Å. The additional peak at 2.57 Å corresponds to the Fe-Fe bond, indicating that the presence of metallic Fe in the sample.

Interestingly, there is a clear variation of the adsorption energy of PMS onto the different Fe spin structures, as manifested in DFT calculations. Fig. 5a-5i shows the optimal configurations and differential surface charge density of PMS adsorption on $Fe^{II}N_4$ in three different spin structures (D_1 -Fe^{II} N_4 low spin, D_2 -Fe^{II} N_4 medium spin, D_3 -N-Fe^{II} N_4 high spin), in comparison to graphene, Fe₂O₃, Fe, pyridinic N, pyrrolic N, and graphitic N. One can see that medium-spin Fe^{II} N_4 exhibits the lowest adsorption energy, followed by low-spin Fe^{II} N_4 (Fig. 5j), implying that these two structures are the preferred adsorption sites for PMS. The fact that FeNC-20 possessed the highest contents of low- and medium-spin Fe species among the series of samples (Fig. 3e) suggests that it is

the optimal structure for the catalytic degradation reactions (an additional minor contribution may arise from pyrrolic N which is a close third in PMS adsorption, and FeNC-20 also contained a slightly higher pyrrolic N content than the others, as shown in Table S3).

3.2. Catalytic degradation of organic dyes

Remarkably, the FeNC nanocomposites exhibited apparent catalytic activity in the oxidative degradation of AO7 by PMS. As shown in Fig. 6a, AO7 degradation was negligible in PMS or FeNC-20 alone (the 13.9% loss of AO7 in the presence of FeNC-20 was due to adsorption). In contrast, when both FeNC-20 and PMS were present, 100% of AO7 was degraded within 5 min at the optimal pH of 7 (optimization of catalyst and PMS loadings, as well as solution pH can be found in Figure S14), suggestive of effective activation of PMS by FeNC-20 in the oxidative degradation of AO7. This performance is markedly better than those when ZIF-8 (39.6%) and MIL-101 (71.7%) was used instead for PMS activation. In addition, by fitting the AO7 degradation curves with the pseudo-first-order kinetic model, the reaction rate was estimated, which was the highest at 0.811 min^{-1} for FeNC-20, in comparison to 0.732 min^{-1} for FeNC-25, 0.403 min^{-1} for FeNC-15, 0.226 min^{-1} for MIL-101, and 0.154 \min^{-1} for ZIF-8 (Fig. 6a inset), confirming the effectiveness of the low- and medium-spin FeN4 moieties in activating PMS for AO7 degradation. The degradation performance of AO7 by FeNC-fer catalyst with PMS is still passable (Figure S15), which supported the above results.

Notably, the Fe-NC/PMS system was also active in the degradation of other organic dyes, such as MR, RhB, AOG, and MO (Fig. 6b), where the removal was all completed within 5 min. The corresponding reaction rate was estimated to be 0.291, 0.203, 1.726, and 1.094 min⁻¹ for MR, RhB, AOG, and MO, respectively (Fig. 6b inset). These results demonstrate that the Fe-N₄/PMS combination can be used as an effective catalyst for the degradation of a wide range of organic pollutants in wastewater. Notably, such performances were significantly higher than those of relevant catalysts reported recently in the literature (Table S5).

3.3. Analysis of reactive oxygen species and activation mechanisms

It has been known that AOPs generally involve reactive species, such



Fig. 4. (a) Fe K-edge XANES spectra and (b) Fourier transforms of the Fe K-edge EXAFS oscillations of FeNC-20 and reference samples (Fe foil, Fe0, Fe₂O₃ and FePc). (c) The corresponding EXAFS fitting result of FeNC-20 in R space (the inset is the coordination structure of FeN(O). (d, e, f) WT-EXAFS of FeNC-20, FePc and Fe foil.



Fig. 5. Optimized configurations (top panels) and differential surface charge density diagrams (bottom panels) of PMS adsorbed on (a) D_1 -Fe^{II}N₄, low spin, (b) D_2 -Fe^{II}N₄, medium spin, (c) D_3 -N-Fe^{II}N₄, high spin, (d) carbon, (e) Fe₂O₃, (f) Fe, (g) pyridinic N, (h) pyrrolic N, and (i) graphitic N. (j) Adsorption energy of PMS onto different carbon configurations. For the Fe-containing samples, the adsorption site is the Fe center, whereas for the metal-free samples, it is the C adjacent.

as $SO_4^{\bullet-}$, ${}^{\bullet}OH$, $O_2^{\bullet-}$, and nonradical oxidation pathways with ${}^{1}O_2$ [19,76]. To identify the relative contribution of these various reactive species in the FeN₄/PMS system, quenching experiments were conducted during the oxidation of AO7. Experimentally, TBA, EtOH, pbenzoquinone, and L-histidine were used as selective quenchers for [•]OH, $SO_4^{\bullet-}$, $O_2^{\bullet-}$, and 1O_2 , respectively [45]. As shown in Fig. 7a and inset, the addition of 0.1 and 0.5 M TBA into the solution decreased the degradation efficiency of AO7 only slightly from 100% to 96.9% and 91.2%, respectively, and the corresponding degradation reaction rate was reduced from 0.811 \min^{-1} to 0.604 and 0.557 \min^{-1} . Similarly, the catalytic degradation of AO7 was not significantly inhibited by the addition of 0.1 and 0.5 M EtOH or p-benzoquinone. These results demonstrated limited contributions from ${}^{\bullet}OH$, $SO_4^{\bullet-}$, and $O_2^{\bullet-}$ to AO7 degradation. In contrast, the AO7 degradation efficiency decreased markedly from 99.7% to 16.5% upon the addition of 0.1 M L-histidine, with an abrupt diminishment of the corresponding degradation rate from 0.811 to 0.046 min⁻¹, indicating the dominant contribution of ¹O₂ to the degradation of AO7 by the FeN₄/PMS system.

Consistent results were obtained in EPR measurements with DMPO and TEMP as the spin-trapping agents [49]. From Fig. 7b, the signals of DMPO-SO₄ and DMPO-OH were readily detected with the FeN₄/PMS system, confirming the formation of SO_4^- and •OH radicals. In addition, a sextet characteristic of DMPO-OOH was observed in Fig. 7c, due to the generation of O_2^- . Note that the signals of •OH, SO_4^- , and O_2^- were all relatively weak, suggesting that these radical species did not play a significant role in degrading AO7, in good agreement with results from the quenching experiments (Fig. 7a) [77,78]. By contrast, a strong triplet with an intensity ratio of 1:1:1, characteristic of TEMP-¹O₂, were observed upon the addition of TEMP (Fig. 7d), confirming the ready generation of ¹O₂ in the FeN₄/PMS system. Additionally, the signal intensity of the TEMP-¹O₂ adducts was obviously higher than that of MIL-101/PMS, signifying the important role of FeN₄ moieties in facilitating the generation of ${}^{1}O_{2}$. Taken together, these results suggest that the nonradical pathway (${}^{1}O_{2}$) played a dominant role in AO7 degradation, with minor contributions from the radical pathways involving ${}^{\bullet}OH$, SO_{4}^{-} , and O_{2}^{-} . Similar results have been obtained in carbon/PMS systems where ${}^{1}O_{2}$ was found to exhibit selective degradation towards organic pollutants with electron-donating groups [30,79].

Based on these results, the mechanism of PMS activation for AO7 degradation is shown in Fig. 8. Note that SO_4^{-} and •OH are typically generated in a series of reactions involving the loss of Fe by electron transfer between Fe⁰ and PMS (eqs. 3–6) [80], where the produced Fe³⁺ reacts with H₂O₂ (produced by oxygen reduction) to generate HO_2° , which is further decomposed into O_2° (eqs. 3–7). However, in the present study, the FeN₄ moieties were firmly embedded within the carbon skeletons, only a trace amount of Fe⁰ was released into the solution, hence producing a low concentration of the radicals. This most likely accounts for their minor contributions to PMS activation.

$$\operatorname{Fe}^{0}(s) + 2\operatorname{HSO}_{5}^{-}(l) \rightarrow \operatorname{Fe}^{2+}(1) + 2\operatorname{SO}_{4}^{\bullet-}(l) + 2OH^{-}(l)$$
 (3)

$$Fe^{2+}(l) + HSO_{5}^{-}(l) \rightarrow Fe^{3+}(l) + SO_{4}^{-}(l) + OH^{-}(l)$$
(4)

$$Fe^{0}(s) + O_{2}(g) + 2H^{+}(l) \rightarrow Fe^{2+}(l) + H_{2}O_{2}(l) \rightarrow Fe^{3+}(l) + OH^{-}(l) + \bullet OH(l)$$
(5)

$$\operatorname{Fe}^{3+}(l) + H_2O_2(l) \rightarrow \operatorname{Fe}^{2+}(l) + \operatorname{HO}_2(l) + H^+(l)$$
 (6)

$$HO_{2}(1) \rightarrow O_{2}^{-}(1) + H^{+}(1)$$
 (7)

$$HSO_{5}^{-}(l) + SO_{5}^{2-}(l) \rightarrow HSO_{4}^{-}(l) + SO_{4}^{2-}(l) + 1O_{2}(g)$$
(8)

$$Fe - N - C(s) + HSO_5^{-}(l) \rightarrow Fe - N - C \equiv HSO_5^{-}(s)$$
(9)



Fig. 6. Degradation properties of AO7 in different environmental conditions. (a) Degradation of AO7 by PMS with various catalysts. (b) Degradation of various organic dyes by FeN_{C} -20 and PMS. (c) Reusability of the FeNC-20 catalyst for the degradation of AO7. (d) Variation of total organic carbon (TOC) with time. Conditions: dye concentration 50 mg L⁻¹, catalyst concentration 2 mg L⁻¹.

 $3Fe - N - C \equiv HSO_{5}^{-}(s) + H_{2}O(l) \rightarrow 2^{1}O_{2}(g) + 3SO_{4}^{2-}(l) + 5H^{+}(l) + 3Fe - N - C(l)$

3.4. Degradation pathway

 $3Fe - N - C \equiv \bullet HSO_5^{-}(s) \rightarrow 3/2^1O_2(g) + 3SO_4^{2-}(l) + 3Fe - N - C(s)$ (11)

It has been shown that ${}^{1}O_{2}$ can be produced by the selfdecomposition of PMS at a rate constant of 0.2 M⁻¹ s⁻¹ (eq. (8)) [45]. However, the low catalytic activity of PMS alone (Fig. 6a) suggests that such a low yield of ${}^{1}O_{2}$ cannot be responsible for the markedly enhanced degradation performance observed with the FeNC composites. Note that production of ${}^{1}O_{2}$ can be readily facilitated by the formation of the unique active structure of Fe-N-C \equiv HSO₅⁻ (eq. 9). Here, the Fe-N moiety serves as the active site to produce oxygen intermediates, where the N atom in FeN₄ extracts electrons from adjacent C atoms, and the positively charged C atoms boost the production of ${}^{1}O_{2}$ (eqs. 9–10). The release of ${}^{1}O_{2}$ from the catalyst surface can also enhance the availability of ${}^{1}O_{2}$ by decreasing the migration distance of the active species [81]. This is consistent with the experimental observation that the degradation efficiency of AO7 was significantly enhanced in the FeN₄/PMS system, as compared to that in ZIF-8 or MIL-101/PMS (Fig. 6a). To explore the degradation pathways of AO7, the active sites of the AO7 molecule need to be identified. As ${}^{1}O_{2}$, an electrophilic oxidant, is the major ROS for degrading AO7 by the FeN₄/PMS catalyst (Fig. 7), it will most likely attack the electron-rich atomic sites in AO7 [48]. From the optimized structure of the AO7 molecule (Fig. 9a), the Fukui index (*f*) was calculated at varied atomic sites and listed in Table S6. It can be seen that 13C exhibited the highest *f* value of 0.057, and is therefore the most vulnerable site that may be attacked by ${}^{1}O_{2}$ in the AO7 molecule, leading to the cleavage of the C-N bond. Other atomic sites, such as 22C (0.054), 24C (0.047), 15C (0.046), 6C (0.045), 21C (0.044), and 29O (0.049), also exhibited a rather high *f* value and may serve as the attack sites, too.

(10)

In addition, the energies of the lowest unoccupied molecular orbital (LUMO) and highest occupied molecular orbital (HOMO) are calculated and shown in Fig. 9a. The red and green represent the electron-rich and -poor regions of the AO7 molecule. One can see that the HOMO mainly resides on the benzene, and electrons can be easily lost from the nitrogen atoms in AO7 molecule, leading to ring opening upon the attack of electrophilic species like ${}^{1}\text{O}_{2}$, ${}^{O_{2}^{--}}_{2}$, and h^{+} . Meanwhile, the carbon



Fig. 7. (a)Effects of different quenchers on PMS activation for AO7 degradation. Inset shows the corresponding degradation reaction rate. (b) EPR spectra of DMPO-SO₄, DMPO-OH, (c) DMPO-OOH, and (d) TEMP-¹O₂ adduct formed after 1 min in the FeN₄/PMS system. Conditions: $[AO7]_0 = 50 \text{ mg/L}$, $[FeNC-20]_0 = 2 \text{ mg/L}$, $[PMS]_0 = 0.5 \text{ mM/L}$, pH = 7.



Fig. 8. Proposed reaction mechanism of AO7 degradation by the FeN₄/PMS system.

atoms in AO7 can be attacked by nucleophilic species as good electron acceptors [59,82,83]. In conjunction with GC–MS studies where the reaction intermediates and products were identified (Figure S16), the

degradation pathway of AO7 is proposed in Fig. 9b, which starts with the breaking of the -C-N = bond between 13C and 12 N atoms due to the highest f value of 0.057 [84]. Meanwhile, C15 in aromatic rings was



Fig. 9. (a) Optimized structure of AO7 and energies of HOMO and LUMO; the possible degradation pathways of AO7 in FeN₄/PMS system.

replaced by oxygen due to the strong attacking effect of ${}^{1}O_{2}$, forming a new product of $C_{9}H_{7}O_{2}$ (Figure S16) which was unstable and easily oxidized into P2 (m/z-146 [M + H]⁺) due to the high redox potential of ${}^{1}O_{2}$ (Figure S17). In addition, based on the electrostatic potential (ESP) distribution on the $C_{9}H_{7}O_{2}$ molecule surface (Figure S18), it can be seen from that the region surrounding of 16O likely attracted ${}^{1}O_{2}$, and thus boosted the reaction. P1 is then further decomposed to P3 (m/z-108 [M + H]⁺), and P3 is mineralized to small-molecule carboxylic acids and eventually H₂O and CO₂. Meanwhile, P2 is transformed to P4 (m/z-120 [M + H]⁺) and P5 (m/z-148 [M + H]⁺), due to the high charge distribution of C15 (0.046). P4 and P5 are further degraded to P6 (m/z-120 [M + H]⁺) and P7 (m/z-194 [M + H]⁺) by ring opening of the heterocycle moiety. Further cleavage of the aromatic rings gave rise to the generation of carboxylic acids and finally to H₂O and CO₂.

3.5. Stability and reusability of FeNC catalysts

The stability and reusability of the FeN₄/PMS catalysts was then tested by repeating the operation of AO7 degradation under the same conditions for six times, where a removal rate of 97.1% can still be achieved in 5 min (Fig. 6c), indicating a robust structure of the FeN₄/PMS catalyst. In addition, the total organic carbon (TOC) of the solution was also measured to evaluate the mineralization efficiency of FeNC-20/PMS. From Fig. 6d, one can see that the final TOC removal efficiency was ca. 92%, much higher than those of N-doped graphene and kaolinite [30,85]. Such a high mineralization efficiency was likely due to the scavenging effect for reactive oxygen species.

Furthermore, the FeNC-20 catalyst after reaction was collected by filtration and some material characterizations were carried out. The XRD patterns of the materials before and after reaction have good coincidence (Figure S19a), which proves that the materials have not

undergone phase transformation. The ratio of micropores and specific surface area did not decrease significantly (Figure S19b-19c), indicating that the skeleton structure of the material was maintained well. In Figure S19d-19f, it is obvious that the tubular structure of FeNC-20 also remains good. The XPS survey spectra revealed the changes of FeNC-20 after the degradation reaction (Figure S20). The enhancement of O peak and the appearance of S peak indicated the presence of some sulfate impurities and the increase of oxygen content in the catalyst. The highresolution scan of the Fe 2p exhibits an increase in Fe³⁺ content (Figure S20b), which proves that Fe^{3+} is an important intermediate in the degradation of organic pollutants. This result verified the mechanism of PMS activation for AO7 degradation described above again. In addition, there is almost no change in C 1 s spectrum indicates the stability of the catalyst carbon skeleton (Figure S20c). It is worth mentioning that the relative content of FeNx (the active site) in N spectrum increases, which confirms the excellent stability of the material in catalytic degradation reaction.

4. Conclusions

In summary, carbon nanotubes embedded with FeN₄ moieties were successfully prepared by a ball milling and pyrolysis method and exhibited unique activity in the activation of PMS for effective degradation of a range of organic pollutants. Spectroscopic measurements suggest that the degradation reaction occurred primarily via the non-radical pathway involving ¹O₂ singlet oxygen, with minor contributions from the radical pathways involving **•**OH, SO_4^{--} , and O_2^{--} . DFT calculations revealed that the spin property of the FeN₄ moieties played a critical role in PMS adsorption and activation, and low- and medium-spin FeN₄ was the most active species for the generation of ¹O₂. Results from this study highlight the significance of carbon-supported

single atom catalysts in the degradation of organic pollutants, a key step towards environmental remediation.

Declaration of Competing 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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Appendix A. Supplementary data

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