EES Catalysis



REVIEW ARTICLE

View Article Online



Cite this: EES Catal., 2023, **1**, 103

Received 18th December 2022 Accepted 23rd January 2023

DOI: 10.1039/d2ey00107a

rsc.li/eescatalysis

Evolution of singlet oxygen in peroxymonosulfate activation: a review

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Given that sulfate-radical-based advanced oxidation processes (SR-AOPs) have attracted great attention in recent years, PMS activation has been reckoned to be an alternative method to the Fenton process in environmental remediation. Hence, in this review, we deliberately retrospect and study the recent progress using singlet oxygen (102) in SR-AOPs since 2016. First, the fundamental principles and means of characterization of ¹O₂ are carefully presented. Several categories of activators, such as metal-free catalysts (e.g., carbon tubes and graphene), metal-based (i.e., the elements Co and Mo), and other metallic-based catalysts (i.e., the elements Fe, Mn, and noble metals), are then specifically proposed. Under these circumstances, the mechanisms of ${}^{1}O_{2}$ formation via direct electron transfer, selfdecomposition, superoxide radical mediation, oxygen vacancy, perxoymonosulfate radical recombination, etc., have been deliberately summarized in sections. Notably, the research gaps and perspectives for 102 in environmental remediation have been critically put forward. Hopefully, this review can offer detailed and theoretical guidance for researchers participating in the study of ¹O₂ in SR-AOPs.

Broader context

As peroxymonosulfate (PMS) activation has received great attention in recent years, researchers have revealed that not only sulfate radicals (SO₄•-) and hydroxyl radicals (*OH) are involved the removal of organic contaminants (OCs); singlet oxygen (102) also plays a leading role under some circumstances. Due to the complexity of 102 formation during sulfate-radical-based advanced oxidation processes (AOPs), and the fact that topic has been less-discussed, it is imperative to discuss the formation mechanism of ${}^{1}O_{2}$ intensively. In this review, we summarize several formation mechanisms based on various nanomaterials and point out the research gaps and perspectives. It is expected that this review will provide some guidance for the future development of ¹O₂ in AOPs.

Introduction

Facing the thriving and prosperous human society are emerging, recalcitrant, and persistent organic contaminants (OCs), which create potential and subtle risks to water and food security, as well as human health. Typically, these OCs are widespread in wastewater; thus, feasible and efficient wastewater treatment technology is of paramount importance to be

$$M^{n+} + HSO_5^- \rightarrow M^{n+1} + SO_4^{\bullet-} + OH^-$$
 (1)

$$M^{n+} + S_2 O_8^- \rightarrow M^{n+1} + SO_4^{\bullet -} + SO_4^{2-}$$
 (2)

Moreover, in recent years, SR-AOPs have become more convincing and better recognized by researchers than the conventional Fenton process due to the greater stability of the oxidants in solid powder form, facile transportation of common chemicals used in the process, and a more broadly adaptable pH region (pH 2.0-8.0). In contrast, the hydroxyl-radical-dominated Fenton

developed. Advanced oxidation processes (AOPs) are known to be promising choices to remove these adverse pollutants via the production of efficient reactive oxidizing species (ROSs), including hydroxyl radicals (${}^{\bullet}OH$) and sulfate radicals ($SO_4^{\bullet-}$). Among the AOPs, the sulfate-radical-based AOPs (SR-AOPs) have attracted great attention. They are primarily driven by peroxymonosulfate (PMS, KHSO₅) and persulfate (PS, K₂S₂O₈) activation.^{3,4} Specific illustrations of PMS and PS activation to generate SO4 • and •OH radicals are provided in eqn (1) and (2).

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process is usually effective under only acidic conditions. However, SR-AOPs would be hampered by the aquatic environment, such as inorganic anions (e.g., HCO₃⁻, Cl⁻, and H₂PO₄⁻) and natural organic matter (NOM) in bodies of water.⁵⁻⁷ Nevertheless, more research has indicated that singlet oxygen (1O2) is emerging as a predominant ROS in SR-AOPs. The formation of ¹O₂ occurs mainly through a non-radical pathway, whereas the formation of SO₄ and OH radicals occurs via a radical pathway. 8-12 In comparison to $SO_4^{\bullet-}$ and $^{\bullet}OH$ radicals, singlet oxygen possesses advantages: (1) higher selectivity toward electron-rich organic complexes owing to its electrophilic attributes, 13,14 so as to preferentially degrade pharmaceuticals and endocrine-disrupting compounds (EDCs) in the presence of inorganic ions and organic compounds 13,14 and disinfect toxic pathogens (e.g., E. coli and the MS-2 bacteriophage); 15,16 (2) higher resistance to frequently-used free radical scavengers, such as methanol and tert-butanol. Principally, the generation of ¹O₂ initially involves a photosensitization effect using photo-induced energy transfer to molecular oxygen (O2); additionally, superoxide radicals $(O_2^{\bullet-})$ often act as intermediates of the singlet oxygen. Alternatively, activating PMS is considered to be a feasible and green chemical process for ¹O₂ production.

Many correlative studies regarding the production of ¹O₂ in PMS activation have been reported. It has been summarized that $SO_4^{\bullet-}$ and $^{\bullet}OH$ radicals can be obtained by the activation of PMS using ultraviolet (UV) light, heat, and transition metals, 17,18 etc., whereas singlet oxygen is primarily generated by metal-free catalysts (e.g., carbonaceous materials). 19-21 Considering the above information, the interest in PMS activation has increased dramatically, as can be evidently deduced from the number of publications since 2011 depicted in Fig. 1 (from the Scopus database). Specifically, the number of publications concerning PMS activation has grown continuously, and the number of publications concerning singlet oxygen in PMS activation has doubled since 2016. However, it can be observed that before 2016, most research focused on the generation of SO₄ • and •OH radicals, while comprehensive analysis of ¹O₂ formation during PMS activation has rarely been reported. Additionally, current

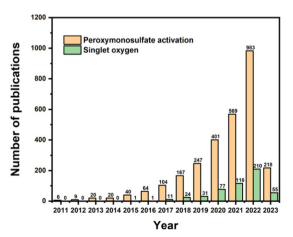


Fig. 1 Number of papers published on peroxymonosulfate activation and singlet oxygen in PMS activation in the Scopus database

studies rarely summarize the recent progress in ¹O₂ formation in PMS activation systems. Thus, after a brief introduction of ¹O₂, this review critically discusses the performance of various types of catalysts and proposes a deep insight into the possible mechanism. Finally, the research gaps and future perspectives are indicated in the last section. This review will provide more theoretical support for future research progress in PMS activation for wastewater treatment.

Singlet oxygen in PMS activation

Characters of singlet oxygen

Compared to the $SO_4^{\bullet-}$ ($E_0 = 2.5-3.1 \text{ V}$) and ${}^{\bullet}OH$ ($E_0 = 2.8 \text{ V}$) radicals, singlet oxygen, as a mild ROS, possesses a lower redox potential $(E_0 = 2.2 \text{ V})^{22}$ Consequently, many scientists have found that singlet oxygen could be resistant to background water matrices, such as NOM and inorganic ions, but its mineralization rates for total organic carbon (TOC) and chemical oxygen demand (COD) are not as good as those of SO₄• and •OH radicals. For the generation of ¹O₂, several approaches have been summarized: (1) photosensitized excitation of oxygen;^{23,24} (2) PDS/PMS activation;^{8,25} (3) periodate;¹³ and (4) reaction of H₂O₂ with NaClO.²⁶ Due to its specific properties, singlet oxygen can be applied in tumor therapy, 27 organic contaminant degradation, 25,28 and pathogenic bacteria inactivation.29

In light of its potential for environmental remediation, the use of ¹O₂ as a crucial ROS during AOPs and its controversial formation mechanism have been extensively studied. Generally, singlet oxygen can be qualitatively identified using chemical scavenger tests and electron paramagnetic resonance (EPR). Sodium azide (NaN₃),^{30,31} furfuryl alcohol (FFA),^{13,32} and tryptophan^{33,34} are commonly adopted as scavengers in chemical probing tests; additionally, spin-trapping agents, such as TEMP, can be used to capture 1O2, forming TEMPO with an intense 1:1:1 signal. Moreover, several investigations have showcased the fact that ¹O₂, as an excited ROS, prefers to approach electronrich compounds with unsaturated C=C bonds and amine and sulfide groups, endowing it with the capability to degrade pharmaceutical and personal care products (PPCPs). 5,20,34,35

Metal-free catalysts

Traditionally, the transition metals Fe, Co, and Mn and their oxides have been considered effective activators for the PMS activation process. However, in recent years, metal-free materials have gathered an intensive impetus due to their metal-free nature, sufficient surface area, good biocompatibility, superior stability toward acidic and alkaline conditions, and adjustable electronic and physicochemical characteristics. 36,37 More importantly, metal-free catalysts can overcome the drawback of metallic secondary contamination in AOPs. Since Duan et al. discovered that a series of carbonaceous materials, including graphene,38 carbon nanotubes,39 and nanodiamond,40 can act as PMS and/or peroxydisulfate (PDS) activators, this research area has been extensively investigated, and has inspired other

researchers involved in this state-of-the-art technology to treat wastewater. In this section, we will talk about the evolution of ¹O₂ over carbonaceous nanomaterials, for instance, carbon tubes (CNTs), graphene, metal-nitrogen-carbon (M-N-C), single-atom nanomaterials (SACs),41 etc. Assuming that in a non-radical reaction system, metal-free catalysts usually serve as an intermediary agent for electron transfer between PMS and organic contaminants, the catalysts contain: (1) delocalized π -electrons; ⁴² (2) defects and vacancies; ⁴³ (3) heteroatoms bonded to carbon;44 and (4) pyran-like oxygen functional groups and C=O bond.45

Carbon tubes

Unlike metallic catalysts, which induce PMS activation relying on SO₄ • and •OH radicals, carbonaceous nanomaterials follow three pathways: (1) the radical pathway producing $SO_4^{\bullet-}$ and OH; (2) the non-radical pathway involving complicated complexes, ¹O₂, and/or direct PMS oxidation; and (3) a combination of radical and non-radical pathways. For example, researchers have also found that organics could mediate PMS activation to produce ¹O₂; for instance, Zhou et al. ⁴⁶ proposed that the established PMS/benzoquinone (BQ) system demonstrated that BQ could efficiently activate PMS for the degradation of sulfamethoxazole (SMX). In this case, ¹O₂ played a leading role, and neither *OH nor SO4*- were involved. A possible catalytic mechanism was proposed involving the formation of an intermediate between PMS and BQ, and subsequently the decomposition of this intermediate into ¹O₂ (Fig. 1a). This mechanism is also referred to in previous works. 43,47,48

Zhou et al.49 further employed ten types of phenols as activators in PMS activation systems, including methyl phenol, phenol, methoxy phenols, and dihydroxybenzenes. Consequently, it was found that phenols can effectively activate PMS to generate ¹O₂ under an alkaline environment (pH 8.5 and 10), with quinone intermediates playing an important role (Fig. 2b). Afterward, Qi et al.50 found that sodium hydroxide itself could act as an activator in PMS activation, and ¹O₂ stemmed from superoxide radicals (O2 •), in accordance with eqn (3) and (4). Therefore, in this part, we will systematically discuss the mechanism of the evolution of ¹O₂ based on carbon-tube-related nanomaterials.

$$2O_2^{\bullet -} + 2H^+ \rightarrow {}^1O_2 + H_2O_2$$
 (3)

$$O_2^{\bullet^-} + {}^{\bullet}OH \rightarrow {}^{1}O_2 + OH^-$$
 (4)

In particular, the PMS activation efficiency of CNTs is correlated with oxygen functionalities (i.e., ketonic groups), exposed edge sites, vacancies and heterogeneous atoms doped into the carbon matrix.⁵¹ For example, Yun et al.⁵² degraded furfuryl alcohol (FFA) using a CNT/PMS system and proposed singlet oxygenation and mediated electron transfer as plausible nonradical mechanisms. In addition, CNT/PDS systems are complicated; they yielded both $SO_4^{\bullet-}$ and 1O_2 as the main ROS catalyzed by quinone groups (i.e., CNT-C=O).53 However, some researchers have pointed out that ketonic groups that are initially on a surface or externally introduced onto

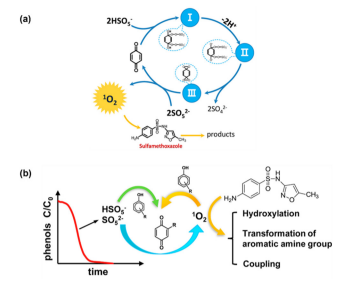


Fig. 2 (a) The proposed catalytic mechanism in the BQ/PMS system; (b) possible mechanism for several types of phenolic compounds as activators in PMS activation. Copied with permission from ref. 46. Copyright 2018, Elsevier.

carbonaceous materials can induce the formation of 1O2 from PDS without interference of the pH value. 53,54 The presence of C=O bonds in carbon-based catalysts is also reported to enhance the efficiency of PMS activation via a non-radical pathway. 55,56 For instance, in 2018, Gao et al. 57 fabricated oxygen-doped graphitic carbon nitride (O-CN) and asserted that electron-deficient C atoms were responsible for generating ${}^{1}O_{2}$. Afterward, Gao et al. continued to dispel the mist surrounding the generation pathway of ${}^{1}O_{2}$ involving PMS oxidation over the electron-deficient carbon atoms neighboring graphitic N atoms in nitrogen-doped carbon nanosheets (NCN-900).58 Distinct from the above-reported mechanism, in their recent work, Sun et al.⁵⁹ reported ¹O₂ originating from self-decomposition. Table 1 lists some representative works for the evolution of ¹O₂ over carbon tubes, including pure CNTs, modified CNTs, and metallic-atom- and/or metal-oxide-doped CNTs. Correspondingly, Guan et al.60 utilized CNTs for the degradation of bromophenols (BrPs) via the nonradical activation of PMS, in which multiple ROS (i.e., SO₄•-, •OH, and ¹O₂) and PMS-CNT* complexes were the active sites in the PMS/CNT system, unlike in the case of PDS, which involved only nonradical ROS (i.e., reactive PDS-CNT complexes). Interestingly, Yun et al. 52 drew the same conclusion that the CNTs/PMS system relies on a major radical route (SO₄ • - and •OH) via electron (from organics) mediation and a minor non-radical route (i.e., ¹O₂). Heterogenous atom doping is another methodology to manipulate the radical generation pathway. In 2015, Duan et al. 61 for the first time discovered that both radical (i.e., SO₄•- and •OH) and non-radical pathways (i.e., graphitic N) contribute to phenol degradation on N-doped CNTs with PMS activation, which has been intensively studied.

Further, Sun et al.65 prepared N/S co-doped ordered mesoporous carbon (NS-CMK-3) as an activator in PMS activation,

Table 1 Performance and mechanism for PMS oxidation of pollutants in typical heterogenous systems

| Catalyst | Target/pollutant | Degradation rate | Radicals | Involved sites | Ref. |
|---|--------------------------------------|--|--|--|------|
| Polyimide-modified carbon nanotubes (PI/CNTs) | Acid Orange 7 (AO7) | 98.9% within 15 min | ¹ O ₂ | N and C=O groups | 62 |
| xFe-N-C | Phenol | 97% within 10 min | $^{1}O_{2}$ | FeN _r sites | 63 |
| Co/NCNT | Rhodamine B (RhB) | 98% within 7 min | $SO_4^{\bullet-}$ and 1O_2 | Co species, pyridinic nitrogen, sp ² -hybrid carbon, ketonic groups | 64 |
| N/S co-doped ordered mesoporous carbon (NS-CMK-3) | Acetaminophen (ACT) | Nearly 100% within 30 min | ¹ O ₂ and catalyst-PMS* | Graphite-N and thiophene-S | 65 |
| Carbon nanotube-magnesium oxide composite (CNTs/MgO) | Rhodamine B (RhB) | 100% of RhB was degraded in 20 min | $^{1}O_{2}$ | Mg–C and C–O bond, surface-active groups, and oxygen vacancies | 66 |
| Carbon nanotubes (CNTs) | Bromophenols (BrPs) | Nearly 100% within 30 min | SO ₄ • and OH, 1O ₂ | PMS-CNT* complexes | 62 |
| FeN ₄ -doped carbon nanotubes | Acid orange 7 (AO7) | 100% within 5 min | $^{1}O_{2}$ | Spin states in FeN ₄ | 67 |
| Fe ₃ C@BN-C | Doxycycline hydrochloride (DOX-H) | 91.9% within 120 min | $^{1}O_{2}$ | Fe ₃ C nanoparticles, oxygen functional groups, pyridinic N, pyrrolic N, graphite N, and B, N dopants | 68 |
| Nitrogen-doped carbon nanosheets (NCN-900) | Bisphenol A | Complete degradation of bisphenol A within 2 min | $^{1}O_{2}$ | Electron-deficient carbon atoms neighboring graphitic N | 21 |
| Carbon nanotube (CNT) | Furfuryl alcohol (FFA) | 100% within 60 min | ¹ O ₂ and electron | Electron transfer from organics to PMS | 52 |

which followed a nonradical mechanism. Fig. 3a illustrates the possible mechanism, which is attributed to the synergy of N/S co-doping (i.e., graphite-N and thiophene-S, Fig. 3c and d) contributing to the highly efficient catalytic performance of NS-CMK-3 compared to that of pristine and single N-doped CMK-3 (Fig. 3b). The scavenger tests and EPR spectra (Fig. 3e and f) indicated that NS-CMK-3 triggers two nonradicaldominated oxidations (i.e., singlet oxygen and surface-confined

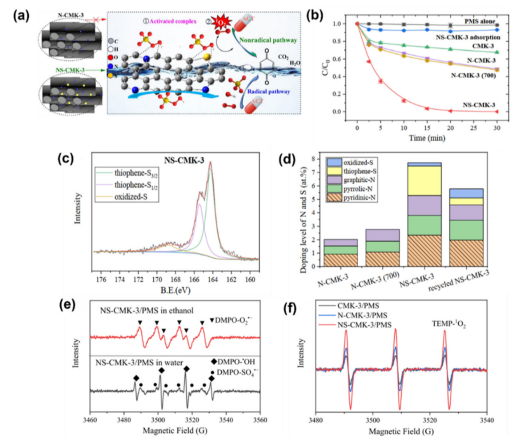


Fig. 3 (a) Brief illustration of the PMS activation mechanism for N/S co-doped ordered mesoporous carbon (NS-CMK-3). (b) Catalytic performance of acetaminophen (ACT) degradation in various oxidation systems. (c) XPS of S_{2p} for NS-CMK-3, and (d) N and S doping levels of multiple materials. (e) EPR spectra of $O_2^{\bullet-}$, ${}^{\bullet}OH$, and $SO_4^{\bullet-}$ and (f) ${}^{1}O_2$ in the NS-CMK-3/PMS system. Copied with permission from ref. 65. Copyright 2021, Elsevier.

activated PMS). Moreover, Li et al.69 used boron (B)-doped carbon tubes for PMS activation, revealing the intrinsic theory toward the catalytic oxidation of pollutants. The involved ROS, such as $SO_4^{\bullet-}$, ${}^{\bullet}OH$, $O_2^{\bullet-}$ and ${}^{1}O_2$, were generated after activation by the active sites; additionally, a B-MWNT-PMS* complex was confirmed as an active site via a non-radical pathway and electron transfer mechanism. Although heteroatom doping is well recognized as an efficient protocol because of its capability to modulate the electronic properties of the sp²-hybridized carbon matrix and provide abundant active sites for catalysis,⁷⁰ the catalytic efficiency of metal-free catalysts is usually impeded by the non-radical process. Regarding this, one promising strategy is to embed transition metal-nitrogen coordination centers into a porous carbon structure via structural engineering.⁷¹ In particular, the nitrogen-coordinated iron (FeN_x) moiety has aroused particular attraction. 72 Thus, for PMS activation, Li et al.73 proposed the use of FeN4-doped carbon nanotubes, which were derived from MIL-101, to degrade acid orange 7 (AO7) in the presence of PMS. The ¹O₂ stemmed from the selfdecomposition of the PMS molecules. In addition, the Fe-N moiety is the active site for the formation of oxygen intermediates. The N atoms in the FeN4 moiety draw electrons from neighboring C atoms, facilitating the production of ¹O₂. Additionally, Du et al. 74 prepared atomically dispersed Fe-N_r site doped N-CNTs derived from ZIF-8; the as-prepared catalyst was used to activate PDS for chloramphenicol (CAP) degradation. Again, a singlet oxygen-dominated process was found in this work, revealing the role of the single-atom site in singlet oxygen evolution (i.e., atomically dispersed Fe-N_x and graphitic N) and offering a new approach for selective removal of trace organic pollutants in complex water matrices.

Graphene

Graphene has emerged as a single-atom-thick sheet composed of sp²-hybridized carbon, 75 and has received great attention in both the scientific and engineering fields since the first fabrication of this "dream material" through the mechanical exfoliation method in 2004. Due to its high thermal conductivity, good electrical conductivity, abundant specific surface area (SSA), environmentally friendly compatibility and low production cost, graphene has been widely applied in the optical, electronic, biological, and catalytic fields. 76-78 In particular, since the discovery of the metal-free activation of persulfates (PMS and PDS) by graphene, 38 the mists surrounding carbon catalysis in AOPs have gradually been dispelled. Generally, the PMS and/or PDS activation efficiency are strongly dependent on the inherent complexity of graphene or its derivatives, such as the graphitic degree, surface oxygen groups, and metallic or heterogenous doped atoms.

Hence, in this section, we briefly introduce the graphenebased nanomaterials in SR-AOPs, and systematically unravel their intrinsic nature in catalytic oxidation. Table 2 summarizes some representative materials, including N-doped graphene, single cobalt atoms embedded in N-doped graphene, N,Sdoped graphene, etc. Accordingly, it was reported that reduced graphene oxide (rGO) can stimulate PMS to evolve SO₄• for the degradation of phenolics and dyes; the zigzag edges and ketone (C=O) groups at the graphene boundaries are the active sites.³⁸ However, for PDS activation using graphene-like nanosheets (GNS), ¹O₂ is mainly produced from O₂•-, and the transfer of surface-confined electrons controls the major nonradical oxidation pathway.⁷⁹ Further, Wang et al.⁸⁰ found that in the nitrogen-doped graphene (NRGO)/PMS system, NRGO can act not only as a PMS activator but also as an electron transfer mediator. In addition, S and N co-doping exhibits an enhanced synergistic effect for catalysis compared with single doping. 81 This reveals that doping rGO at an optimized sulfur loading can effectively break the inertness of carbon systems, activate the sp²-hybridized carbon lattice and facilitate the electron transfer from covalent graphene sheets for PMS activation. A similar result was obtained by Sun et al.,82 in which N doping and additional S doping played pivotal roles in enhancing catalytic

Table 2 Performance and mechanism for PMS oxidation of pollutants in typical heterogenous systems

| Catalyst | Target/pollutant | Degradation rate | Radicals | Involved sites | Ref. |
|---|--------------------------|--|--|--|------|
| Graphitic N-rich graphene (GNG) | RhB | 100% RhB removal was achieved within 40 min at 45 °C | $^{1}\mathrm{O}_{2}$ | Graphitic N | 84 |
| N-doped graphene | Phenol | Nearly 100% within 20 min | $^{1}O_{2}$ | Surface functionalities | 85 |
| Cobalt single atoms embedded in nitrogen-doped graphene (SACo@NG) | Benzyl alcohol (BzOH) | Over 90% within 180 min | $^{1}O_{2}$, $SO_{4}^{\bullet-}$ and $^{\bullet}OH$ | Co ²⁺ and Co ³⁺ atoms, as well as the abundant surface N and nucleophilic C = O groups | 86 |
| FeCo@NC | BPA(bisphenol A) | Complete degradation within 4 min | $^{1}O_{2}$ | CoN ₄ site with a single Co atom | 87 |
| Nitrogen-doped graphene (NRGO) | Sulfamethoxazole (SMX) | 91.7% at 240 min | $^{1}O_{2}$ | Pyrrolic N sites | 80 |
| Nitrogen-doped graphene (N-IrGO) | Benzophenone-1 (BP-1) | 100% within 60 min | $^{1}O_{2}$ | Graphitic-like nitrogen | 88 |
| Nitrogen-sulfur co-doped indus- trial graphene (i-rGO-NS) | Methyl paraben (MP) | Complete degradation within 5 min | $^{1}O_{2}$ | Graphitic N | 82 |
| Cu-rGO LDH nanohybrid | Bisphenol A (BPA) | 100% within 40 min | $^{1}O_{2}$ | Electron transfer from PMS to Cu centers | 89 |
| Nitrogen and sulfur co-doped graphene (N, S-G) | Phenol | Nearly 100% within 6 min | $^{1}O_{2}$ | Metastable PMS@carbon complex | 90 |

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performance. Additionally, the nitrogen and boron-co-doped graphene developed by Chen et al. 83 shows a synergistic effect between pyridinic N and BC3 (B-C-C-pyridinic N), and subsequently has more positively charged carbon atoms as active sites for PMS activation.

As for single atomic nanomaterials, their high atom utilization efficiency has aroused tremendous interest in the PMS activation process. A Co-based SAC on nitrogen-doped graphene (SACo@NG) material synthesized by Li et al.91 was successfully utilized to selectively oxidize BzOH via PMS activation, and showed superior catalytic performance compared to its counterparts such as NG and CoNP@NG. In the reaction process, the single atomic Co clusters function as active sites via both radical and non-radical pathways, with the latter being preferred. This non-radical pathway is achieved by the adsorption of a BzOH/PMS complex on the surface of SACo@NG followed by electron transfer throughout the carbon matrix.

Furthermore, Li et al. 92 put forward single cobalt atoms anchored on porous N-doped graphene with dual function in PMS activation for the rapid degradation of bisphenol A (BPA). A nitrogen-doped graphene-coated FeCo bimetallic nanocage (FeCo@NC) was formed using nanospheres of a Prussian blue analog (FeCo PBA) in nitrogen. FESEM, TEM, and HR-TEM

images of FeCo-NC-2 (Fig. 4A, inset a-d) showed a large number of hollow graphene spheres, with a small amount of FeCo nanocrystals remaining after acid treatment. The energy dispersive X-ray spectrum (EDX) mapping images of FeCo-NC-2 (Fig. 4A, inset e) illustrate the distribution of N, Fe and Co species in the nanocrystals and the porous graphene shells. The dispersion of individual Co/Fe atoms anchored to porous nitrogen-doped graphene can be clearly observed by HAADF-STEM with aberration correction. As shown in Fig. 4A, inset f-i, the bright spots corresponding to heavy atoms (single Co/Fe atoms) are well dispersed throughout the graphene sphere. In this case, bisphenol A was selected as the target pollutant, and the PMS catalytic performance of the single-Co-atom catalyst was studied. Surprisingly, 100% BPA removal was achieved in 4 minutes using FeCo-NC-2. In addition, the first-order reaction rate constants of FeCo-NC-1 (1.18 min⁻¹), FeCo-NC-2 (1.25 min^{-1}) , and FeCo-NC-3 (0.47 min^{-1}) were found to match the relative content of cobalt catalysts (Fig. 4B, inset b). In contrast, the reaction rate constant did not correlate with the content of iron. However, the removal efficiency of BPA by Co-NC with the highest Co content (19.5 wt%) was lower than 73%, indicating that other factors besides the single cobalt atoms may be dominant factors in the Fenton-like catalytic

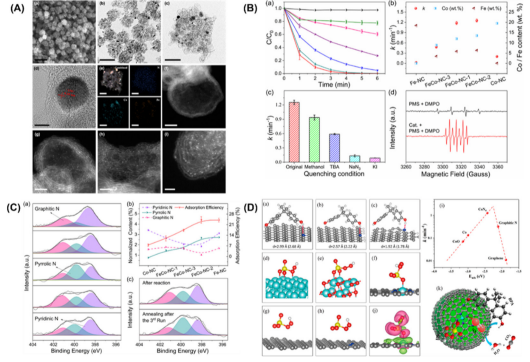


Fig. 4 (A) FESEM image (a), TEM images (b-c), HRTEM image (d), EDX mappings (e), and HAADF-STEM images of FeCo-NC-2 (f-i). (B) Catalytic performance in BPA degradation in different reaction systems by the leaching solution (a). (b) Reaction rate constants with the contents of Co and Fe using different catalysts. (c) Reaction rate comparison in different quenching conditions. (d) EPR spectra with/without FeCo-NC-2. (C) Determination of the adsorption sites. (a) XPS spectra of N 1s for Fe-NC, FeCo-NC-1, FeCo-NC-2, FeCo-NC-3, and Co-NC. (b) Relationships between BPA adsorption ability and normalized N species content. (c) XPS spectra of N 1s for FeCo-NC-2 after reaction (top) and regeneration by annealing in N2 (bottom). (D) Proposed possible mechanism of the PMS activation reaction. Adsorption configurations of BPA on (a) pyrrolic, (b) pyridinic, and (c) graphitic N-doped graphene. Optimized configurations of PMS adsorbed on (d) Co (0001), (e) CoO (100), (f) CoN4-graphene, (g) graphitic N-doped graphene, and (h) graphene, respectively. (i) Plot of the reaction rate against the adsorption energy of PMS on Co (0001), CoO (100), CoN₄-graphene, graphitic N-doped graphene, and graphene. (j) Charge density difference in CoN₄-graphene (ρ total $-\rho$ substrate $-\rho$ PMS). (k) Fenton-like reaction mechanism on single-Co-atom catalyst. Copied with permission from ref. 92. Copyright 2018, American Society Chemistry

reaction. The scavenger test and EPR spectrum for the determination of free radicals are shown in Fig. 4B (insert C and D), respectively. The results show that the reactive oxygen species (*OH, SO₄*-) exhibit little impact on the degradation efficiency of BPA, while NaN3 played a crucial role in the degradation of ¹O₂. Furthermore, KI, as a strong quenching agent of surfacebound free radicals, can quench the reaction, indicating that the reaction relying on ¹O₂ occurs throughout the whole surface catalytic process. EPR spectra (Fig. 4B, insert D) showed that OH can be produced by hydrolysis of the PMS molecules.

In addition, an apparent ¹O₂ signal appeared after FeCo-NC-2 was added to the system, confirming its dominant role in this case. XPS N 1S high-resolution spectra of the prepared catalysts were collected to quantify the type and content of the nitrogen dopant, as shown in Fig. 4C (inset a). Three peaks with binding energies of 398.4 eV, 399.8 eV and 401.1 eV were deconvoluted and attributed to pyridine, pyridine, and graphite N, respectively. The concentration of pyrrole N was observed to be correlated with the BPA adsorption efficiency (Fig. 4C, inset b), suggesting that pyrrole N may act as the adsorption site of BPA. To further confirm this hypothesis, the XPS N1s high-resolution spectra of the catalyst was investigated (Fig. 4Cinset (c)). It indicated that the adsorption of BPA and its corresponding intermediates on the pyrrole N position is the main cause of the gradual inactivation of the mono-Co catalyst. Finally, DFT calculations verified that the electron donor of BPA is the pyrrole N and the electron acceptor is the -OH group. Fig. 4D (inset a-c) shows the optimized configuration of BPA for the adsorption of pyrrole, pyridine and graphene N, where BPA has the optimal adsorption of pyrrole N, the shortest N-H distance (1.78 Å) and the highest adsorption energy (-0.31 eV). In addition, DFT calculations were performed to provide theoretical insight into the activation of PMS at the CoN4 sites containing single atoms of Co. Fig. 4D (inset d-h) reveals the complete relaxation atomic configurations of PMS on Co (0001), CoO (100), CoN₄, graphite N, and graphene. Using the PMS adsorption energy descriptor, the relationship with the reaction rate is shown as a volcano plot in Fig. 4D (inset i). Thus, the low-reactivity cobalt metal and cobalt oxide of PMS lead to poisoning of the active site due to their strong binding to PMS. On the other hand, the binding of PMS to graphite N and graphene is too weak to activate the PMS molecules effectively. In addition, from the charge density analysis (Fig. 4D, inset j), it can be seen that there is significant electron transfer between PMS and CoN₄, reflecting the chemisorption of PMS at the CoN4 site.

Transition metals

Transition metals with lower valence states, like that of the Fenton regent (i.e., Fe²⁺ and H₂O₂), can activate PMS/PDS molecules well according to eqn (1) and (5). Typical PMS and PDS activation by electron transfer and energy transfer are depicted in Fig. 5. In this section, we will discuss the utilization of transition metal ions in SR-AOPs.

$$M^{n+} + HSO_5^- \rightarrow M^{n+1} + SO_4^{\bullet-} + OH^-$$
 (1)

$$M^{n+} + S_2O_8^- \rightarrow M^{n+1} + SO_4^{\bullet-} + SO_4^{2-}$$
 (5)

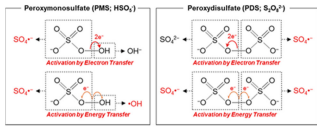


Fig. 5 Activation of PMS and PDS via electron- and energy-transferring processes. Copied with permission from ref. 91. Copyright 2020, American Chemical Society

Co-based activator

Dionysiou et al. 93 reported that Co²⁺ outperforms most reported transition metal ions, such as Fe²⁺, Cu²⁺, Ce³⁺, Mn²⁺, and Ni²⁺, for PMS activation; the intrinsic reason has been ascribed to its highest standard reduction potential among various metallic redox couples, as shown in Table 3. The basic mechanism for the activation of PMS by Co^{2+} ions is summarized in eqn (6)–(8). In this part, we will list some relevant examples, as comparative studies on the formation of singlet oxygen in elemental Co-based catalysts have progressed greatly.

$$CO^{2+} + HSO_5^- \rightarrow CO^{3+} + SO_4^{\bullet-} + OH^-$$
 (6)

$$CO^{3+} + HSO_5^- \rightarrow CO^{2+} + SO_5^{\bullet-} + H^+$$
 (7)

$$SO_4^{\bullet -} + H_2O \rightarrow {}^{\bullet}OH + SO_4^{2-} + H^+$$
 (8)

For instance, Chen et al. 94 employed magnetic Co-N-doped carbon hybrid catalysts (Co-NC-x) to treat recalcitrant organic pollutant effluent by using a facile cation exchange and selfreduction method to activate PMS. EPR spectroscopy and scavenger testing demonstrated that $SO_4^{\bullet-}$ and 1O_2 were responsible for the degradation of RhB in the Co-NC-850/PMS system; additionally, ¹O₂ contributed ca. 86.2% to RhB removal. Remarkably, the synergistic effect of Co⁰ nanoparticles (NPs) and NC on Co-NC-850 acted as active sites to trigger PMS activation, and the direct oxidation of O_2^{ullet-} by ${}^{ullet}OH$ played a crucial role for forming 1O2. Zhang et al. 35 proposed a CoOOH/ PMS system to degrade 2,4-dichlorophenol. Again, ¹O₂ played a leading role; however, the singlet oxygen was derived from the reaction between PMS molecules and water molecules with the formation of superoxide as an intermediate. Interestingly, Zeng et al. 95 utilized oxygen-vacancy-enriched cobalt aluminum hydroxide@hydroxysulfide (CoAl-LDH@CoSx) hollow flowers

Table 3 Reduction potentials of transition metallic couples

| Metallic redox couple | E_0 (V) | | |
|------------------------------------|-----------|--|--|
| Co ³⁺ /Co ²⁺ | 1.92 | | |
| Fe^{3+}/Fe^{2+} | 0.77 | | |
| Ce^{4+}/Ce^{3+} | 1.72 | | |
| Mn^{3+}/Mn^{2+} | 1.54 | | |
| Cu^{2+}/Cu^{+} | 0.15 | | |
| V^{4+}/V^{3+} | 0.34 | | |

for the degradation of sulfamethoxazole (SMX) in PMS activation; ¹O₂ was verified to be dominant ROS responsible for SMX degradation via quenching tests. Mechanism investigation revealed that the oxygen vacancies, redox cycles of Co(II)/ Co(III) and S²⁻ as well as sulfate species were responsible for singlet oxygen formation. Similar results were obtained by Dong et al.,95 who employed natural illite microsheets via PMS activation. Accordingly, abundant oxygen vacancies were created by the indistinct lattice boundaries. DFT calculations verified that the oxygen vacancies significantly reduced adsorption energy and accelerated electron transfer, further promoting PMS activation. In addition, the oxygen-vacancy-rich Co₃O₄/ illite exhibited superior catalytic efficiency in a real water matrix, which was ascribed to $SO_4^{\bullet-}$, $\bullet OH$ and 1O_2 (generated from oxygen vacancies, eqn (9)-(12)).

$$O_v + HSO_5^- \rightarrow O_2^{\bullet -} + HSO_4^- \tag{9}$$

$$O_2^{\bullet -} + H_2O \rightarrow {}^{\bullet}OOH + OH^-$$
 (10)

$$O_2^{\bullet -} + {}^{\bullet}OOH \rightarrow {}^{1}O_2 + HOO^{-}$$
 (11)

$$^{\bullet}$$
OOH + $^{\bullet}$ OOH \rightarrow 1 O₂ + H₂O₂ (12)

Dong et al.³³ reported CoP/N-g-C₃N₄ nanosheets in which the doping of graphitic N atoms could modulate the electronic properties of the g-C₃N₄ nanosheets. As previously mentioned, in the N-g-C₃N₄ nanosheets, the electron-deficient carbon atoms neighboring graphitic N took charge of PMS oxidation in generating ¹O₂. However, in the visible-light-driven CoP/N-g-C₃N₄/PMS system, the PMS molecules were preferentially adsorbed onto the electron-deficient carbons near the newly formed N=N bonds for PMS reduction (SO4 o and OH radicals), whereas the CoP nanoparticles were responsible for the PMS oxidation (i.e., ¹O₂ production) and PMS reduction at the same time. For the ROS, ¹O₂ plays a leading role, and is derived from $SO_5^{\bullet-}$ and $O_2^{\bullet-}$. The specifics are shown in Fig. 6.

In terms of single atomic materials, it has been clearly elaborated that the Co-N₄ site with a single Co atom serves as the active site with optimal binding energy for PMS activation. 91 However, Zhan and his group members 21 recently found that CoN2+2 could be recognized as the active site and



Fig. 6 The proposed mechanism in the CoP/N-g-C₃N₄ nanosheets catalyzed visible-light-driven PMS activation. Copied with permission from ref. 33. Copyright 2021, Elsevier.

singlet oxygen was the predominant ROS with almost 100% PMS conversion.

More specifically, this work by Zhan and coworkers focuses on the CoN₂₊₂ monoatomic active sites, which regulate PMS oxidation at the atomic level. Unlike in the case of the rupture of the O-O and/or O-H bonds in PMS, electrons are transferred from PMS to a single Co atom, forming ¹O₂. Co-SA materials were synthesized via self-assembly and pyrolysis (Fig. 7A, inset a). After the Zn atoms evaporated, the remaining Co atoms were successfully fixed on the N-C support. Co-SA presents a smooth and dodecagonal shape (Fig. 7A, inset b). No obvious aggregation of the finite atoms exists, and individual finite atoms (shown in white circles) are observed fixed throughout the architecture (Fig. 7A, inset c and d). Additionally, mapping of Co-SA revealed the uniform distribution of C, N species (Fig. 7A, inset e). X-ray absorption fine structure (XAFS) analysis suggested the presence of atomic structure of the catalyst. As can be seen from the XANES spectrum (Fig. 7B, inset a), the rising edge of Co-SA is located between that of Co foil and Co-NP, indicating that the average oxidation state of a single Co atom is between Co⁰ and Co³⁺. The EXAFS spectrum of Co-SA has a main peak at 1.41 Å, corresponding to the Co-N coordination

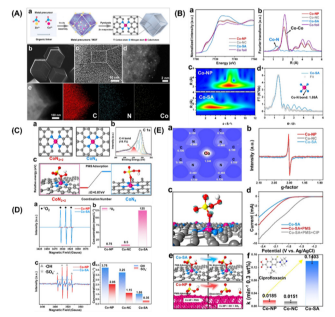


Fig. 7 (A) (a) Preparation steps of Co-SA. (b)-(e) SEM, TEM, HAADF-STEM and EDS mapping images of Co-SA. (B) (a) Co normalized K-edge XANES spectra and (B) K^3 -weighted Fourier transform spectra of Co foils and Co-based samples. (c) Wave transformations of Co-NP and Co-SA. (d) EXAFS fitting curve corresponding to Co-SA in R space. (C) Atomic structure of CoN₂₊₂ and CoN₄ active sites of Co-SA (a) and XPS spectrum of C of Co-SA (b). The free energy evolution of the active sites for PMS adsorption of CoN_{2+2} and CoN_4 (c). (D) EPR spectrum and quantitative determination of ¹O₂, •OH and SO₄•-. (E) Two-dimensional valence electron density map of the CoN_{2+2} site in Co-SA (a). EPR spectrum of catalysts (b). The electron density difference diagram (c) of Co-SA and PMS adsorbed at the CoN_{2+2} site. (d) Linear sweep voltammetry (LSV) curves of Co-SA. (e) Adsorption of PMS on Co-SA and Co-NP surfaces. (f) Degradation efficiency of CIP in different systems. Copied with permission from Ref. 21. Copyright 2021, Wiley.

EES Catalysis

(Fig. 7B, inset b). Unlike for the Co foil, Co-NP and Co-NC, there is no Co-Co coordination peak at 2.1 Å, indicating the atomic dispersion of Co on the carbon carrier. The wavelet transform (WT) diagram of Co-SA shows that the WT maximum value is 5.1 Å^{-1} , corresponding to Co-NP (Fig. 7B, inset c). In addition, the Fourier transform of the Co-SA K-Edge EXAFS spectrum can be fitted by CoN from the scattering path of the CoN₄ structure (Fig. 7B, inset d). Considering the influence of the carbon matrix CoN₄ active sites, studies were first computed focusing

on the CoN₂₊₂ (CoN₄ partially bridged at two adjacent graphite

edges) and CoN₄ (CoN₄ half embedded in a complete graphite

layer) sites (Fig. 7C, inset c). The predicted Co-N bond length

was 1.88 Å at CoN₂₊₂ and 1.85 Å at CoN₄. Based on EXAFS

analysis, the CoN bond length at the CoN2+2 site was predicted

to be close to 1.89 Å. In addition, a C-H bond was detected, which is consistent with the CoN₂₊₂ configuration (Fig. 7C,

In addition, the adsorption of PMS by the CoN₂₊₂ site is more advantageous in terms of energy than that by CoN₄ site (Fig. 7C, inset c). Therefore, it can be inferred from the above results that the configuration of the active Co-SA site is CoN₂₊₂. To identify the ROS generated during PMS activation, EPR was used to directly detect ROS. The intensity of the triplet signal of TEMP-¹O₂ in the Co-SA system is 37 times higher than that in the Co-NP system (Fig. 7D, inset a). In addition, the ${}^{1}O_{2}$ concentration was quantitatively determined in all three systems (Fig. 7D, inset b). The ratio of ${}^{1}O_{2}$ to produce ROS Co-SA accounted for 98.89%, and SO4 and OH radicals were generated in the Co-NP and Co-SA systems (Fig. 7D, inset c), which implies that the major ROS in the Co-NP systems were

SO₄• and OH (Fig. 7D, inset d). These results show that different reaction pathways and dominant ROS-catalysed activation of PMS on the CoN2+2 active sites. The simulated configuration of the charge properties of the CoN_{2+2} active sites was determined (Fig. 7E, inset a). The electronic properties of Co-NP and Co-SA were detected using EPR (Fig. 7E, inset b). The EPR signal of Co-NP was strengthened compared to that of Co-SA, revealing a higher concentration of unpaired electrons for Co-NP. Furthermore, the adsorption and reaction of PMS on the prototypical Co-NP and Co-SA surfaces were theoretically modelled, and the results revealed a difference in electron density between PMS and Co-SA (Fig. 7E, inset c). It was found that electrons transfer from PMS to Co single atoms, indicating that the Co single atoms were reduced during the PMS adsorption process. In the LSV analysis, the number of the Co-SA electrodes significantly increased the electron transfer process in the presence of PMS (Fig. 7E, inset d). These results demonstrate that combined Co-NP can be used as an electron donor to reduce the activation of the PMS, and that the body of the molecules tends to transfer electrons to the Co-SA. In the joint Co-NP system, the PMS molecules are adsorbed by the Co-NP system because of the strongly positively charged nanoparticles (Fig. 7E, inset e). The strong interaction between the positively charged Co atom and the negatively charged O atom results in spontaneous dissociation of the adsorbed PMS and the formation of SO₄•- and •OH.

Thus, the dissociation of PMS on the Co-NP surface is thermally favourable, with no energy barrier. In the Co-SA system, PMS is adsorbed on a single Co atom, and an O atom on the -SO4 side is bonded to the Co atom on the surface, indicating that the Co atom is the active adsorption site regardless of the morphology of Co. The O-O and O-H bonds were not cleaved. The interaction between H and N is too weak for the hydrogen atom to attach stably to the negatively charged N atom. After normalizing the rate constant to the Co concentration, the constants of Co-SA are 10.55 times and 6.27 times those of Co-NP and Co-NC, respectively (Fig. 7E, inset f). In general, Co-SA with CoN₂₊₂ active sites regulates ¹O₂ production during PMS activation. For comparison, Co-NP and Co-NC were synthesized to study the generation of 1O2 during PMS activation. In the activation using the Co-NP system, the main ROS are SO4. and OH radicals. PMS is directly decomposed into SO4 and OH on Co nanoparticles, which is mainly attributed to the strong reducibility of the Co adsorption sites. Unlike the traditional catalytic system with free radical generation, the Co-SA activation of PMS involves the oxidation of PMS to ${}^{1}O_{2}$. The spontaneous dissociation of PMS on the Co-SA surface cannot be induced by the charge density of N and the coordination of Co atoms in CoN2+2. However, electron transfer to the Co single atom occurs during the adsorption process of PMS, resulting in ¹O₂ formation in 98.89% of the generated ROS. The generated singlet oxygen showed effective degradation activity toward several organic pollutants over a wide pH range. This study proposes a strategy to control ¹O₂ generation through PMS activation.

Mo-based activators

Xing and his group members 96 proposed that MoS2 could act as a highly efficient cocatalyst in the Fenton process, mainly driven by the fast circulation of Fe³⁺/Fe²⁺ through the exposure of Mo4+ active sites. Relevant research using the element Mo has since progressed greatly. 97,98 Xing et al. 99 also found that MoS2 could act as a cocatalyst for rapid disinfection of Escherichia coli K-12 (E. coli) and S. aureus. Afterward, with the popularity of PMS activation, Sheng et al. 100 further incorporated MoS₂ as a cocatalyst in the Fe(II)/PMS system (arising from Fe³⁺ to Fe²⁺), accompanied by the yield of a high concentration of SO4 • and •OH radicals, which resulted in effective removal of 2,4,6-trichlorophenol. In addition, Zhou et al. 101 employed carbamazepine as a target to unravel the different mechanisms of the MoS₂/PMS system and the MoS₂/PDS system. Zhang et al. 102 observed that the excellent degradation of acetaminophen is due not only to the acceleration of Fe ion circulation by MoS₂ but also the formation of a strongly oxidative Mo(VI) peroxo-complex intermediate. Additionally, ¹O₂ takes charge of the overall process; the 1O2 is derived from the oxidation of MoS_2 , as presented in eqn (13) and (14).

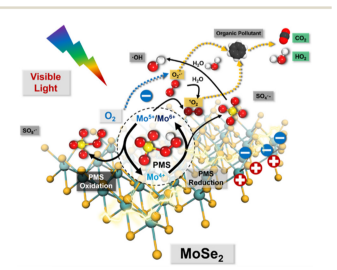
$$MoS_2 + 9HSO_5^- + 5OH^- \rightarrow MoO_4^{\ 2-} + 10H^+ + {}^1O_2$$
 (13)

$$MoOOH(O_2)_2^- + H_2O \rightarrow MoO_4^{2-} + {}^1O_2 + H^+$$
 (14)

Review

Accordingly, a series of Mo-based catalysts and/or cocatalysts were employed in AOPs (e.g., the Fenton process), such as Mo powder, 103,104 MoO₂, 105 two-dimensional (2D) MoS₂, 106 and three-dimensional (3D) MoS₂ sponge. 107 The relevant innovative mechanism for the Fenton process was proposed, which mostly revolves around the following two phenomena: (1) the oxidation process between Fe³⁺ and Mo⁴⁺; (2) the ¹O₂ stemming from the oxidation-reductive reaction between Mo^{6+} and $O_2^{\bullet-}$. However, the mechanism related to ${}^{1}O_{2}$ for SR-AOPs still lacks sufficient exploration. In this context, Dong et al. 33 tentatively synthesized bimetallic MoFe/TiO2 nanospheres for solar-lightdriven PMS activation. They pointed out that both photogenerated electrons and transition metallic redox couples (i.e., Mo⁶⁺/Mo⁴⁺, Fe³⁺/Fe²⁺ and Mo⁴⁺/Fe³⁺) were responsible for PMS activation. Most importantly, SO₄•-, •OH, and SO₅•participated in the transformation and generation of ¹O₂. Furthermore, Dong et al.⁵ used commercial MoSe₂ to replace MoS₂ to solve the problem of H₂S discharge in sewage. In this case, $O_2^{\bullet-}$ is dominant, and 1O_2 comes mainly from the transformation of $O_2^{\bullet -}$. The specific information is shown in Fig. 8 and eqn (15)-(24). If MoSe₂ were irradiated with visible light, carriers (i.e., electrons and holes) would be generated (eqn (15)), with photogenerated electrons reacting with oxygen to form $O_2^{\bullet-}$ (eqn (16)). Afterward, Mo^{4+} reacts with HSO_5^- to form $SO_4^{\bullet-}$, Mo^{5+} and Mo^{6+} (eqn (17)). After that, the $SO_4^{\bullet-}$ radical leads to the formation of *OH radicals, and excess HSO₅⁻ tends to produce hydrogen peroxide (eqn (19)), resulting in the reaction of *OH free radicals with H2O2 to produce $HO_2^{\bullet -}$ and $O_2^{\bullet -}$ (eqn (20) and (21)). Afterward, excess $O_2^{\bullet -}$ is converted into ${}^{1}O_{2}$. Of note, Mo $^{6+}$ can oxidize $O_{2}^{\bullet-}$ to ${}^{1}O_{2}$. 106 Accordingly, the ¹O₂ is mainly derived from the transformation of $O_2^{\bullet-}$ rather than stemming from the previously reported

$$MoSe_2 + h\nu \rightarrow e^- + h^+ \tag{15}$$



PMS self-decomposition or PMS oxidation. Ultimately, both

O₂• and ¹O₂ radicals contribute to the degradation of PPCPs.

Fig. 8 Proposed mechanisms for the visible-light-driven MoSe₂/PMS system. Copied with permission from ref. 5. Copyright 2021, Elsevier.

$$O_2 + e^- \rightarrow {}^{\bullet}O_2^- \tag{16}$$

$$Mo^{4+} + HSO_5^- \rightarrow Mo^{6+}/Mo^{5+} + SO_4^{\bullet-} + OH^-$$
 (17)

$$Mo^{6+}/Mo^{5+} + HSO_5^- \rightarrow Mo^{4+} + SO_5^{\bullet-} + H^+$$
 (18)

$$HSO_5^-(SO_5^-) + H_2O \rightarrow H_2O_2 + HSO_4^-(SO_4^{2-})$$
 (19)

$${}^{\bullet}OH + H_2O_2 \rightarrow HO_2{}^{\bullet -} + H_2O$$
 (20)

$$HO_2^- \to H^+ + {}^{\bullet}O_2^- \tag{21}$$

$$2^{\bullet}O_2^- + 2H_2O \rightarrow H_2O_2 + 2OH^- + {}^{1}O_2$$
 (22)

$$Mo^{6+} + {}^{\bullet}O_{2}^{-} \rightarrow Mo^{4+} + {}^{1}O_{2}$$
 (23)

$${}^{\bullet}\text{O}_2^{-}/{}^{1}\text{O}_2 + \text{CBZ} \rightarrow \text{intermediates} + \text{CO}_2 + \text{H}_2\text{O}$$
 (24)

Other metallic activators

In addition to materials based on the elements Co and Mo, iron-based catalysts are popular in AOPs. Generally, PMS/PDS activation using Fe involves the transformation from Fe²⁺ to Fe³⁺ and the generation of OH and SO₄. Notably, the concentration of Fe2+ has a crucial impact on PMS/PDS oxidation of organic pollutants. However, the relative progress involving its non-radical pathway is rarely reported. Interestingly, Li et al. 108 found that the Fe⁰/PMS system for the degradation of Rhodamine B (RhB) could be enhanced by the addition of Cu2+, and that the dominant ROSs in the process were ¹O₂ and O₂•-. Yang et al. ¹⁰⁹ also confirmed ¹O₂ and O₂•to be the dominant species, with OH also coexisting, in the Fe⁰-montmorillonite/PMS system.

In addition, Mn-based catalysts are considered to be effective PMS/PDS activators with the advantage that Mn is an Earthabundant element and less toxic than Co. 110 In terms of the non-radical pathway, Huang et al. 111 revealed that 1O2 was generated in the PMS/MnO2 system under acidic conditions. They also concluded that 1O2 was generated as the primary ROS through the direct oxidation of O2. by Mn(IV) and $O_2^{\bullet-}$ recombination, as well as the reaction between $O_2^{\bullet-}$ and metastable manganese intermediates under neutral conditions. Furthermore, metal oxides of perovskites (i.e., ABO₃ structure) and spinel (i.e., AB_2O_4 structure)¹¹² have attracted increasing interest, and the corresponding oxygen vacancies on the surface of the metal or metal oxide account for the generation of ¹O₂. Due to the facilitated oxygen adsorption on the surface and lattice oxygen vacancies, the rapid generation of O2. leads to rapid formation of 1O2, as a greater amount of oxygen vacancies is beneficial to the interfacial electron transfer in PMS activation. 113 In addition to the conventional metallic catalysts, Al₂O₃ and TiO₂ loaded with noble metals including Pt, Pd, Au, and Ag can mediate electron transfer from organic pollutants to PMS molecules, 114 and subsequently achieve the non-radical oxidation of PMS through ¹O₂. The degradation efficiency follows the order Pd > Pt \approx Au \gg Ag. Wang et al. 115 intensively investigated the catalytic mechanism of PMS. The dispersed Pd particles on the g-C₃N₄ (Pd/g-C₃N₄) surface could activate PMS, inducing the formation of ¹O₂ and O2. of Tor BPA degradation. It is worth noting that Pd⁰ is

Performance and mechanism for other metallic-based activators in PMS activation

| Catalyst | Target/pollutant | Degradation rate | Radicals | Involve sites | Ref. |
|----------------------------------|---------------------------------|------------------------|--|--|------|
| Fe-Mt-C-H ₂ | BPA (25 mg L ⁻¹) | 99.3% within 120 min | ¹ O ₂ and O ₂ • − | Fe ⁰ | 121 |
| Fe-N-C | BPA (25 mg L^{-1}) | 88% within 120 min | $^{1}O_{2}$ | Metal-N-C catalytic site | 119 |
| MnO_2 | Bisphenol A (BPA) | 94.5% within 10 min | $^{1}O_{2}^{^{2}}$ | Mn ⁴⁺ and Mn ³⁺ | 111 |
| CuO-CeO ₂ | Rhodamine B | A complete degradation | $^{1}O_{2}^{^{2}}$ | Cu ²⁺ /Cu ⁺ , oxygen vacancies, | 122 |
| | | within 60 min | | electron transfer | |
| $LaCo_{0.4}Cu_{0.6}O_3$ | Phenol | 100% within 12 min | $^{1}\mathrm{O}_{2}$ | Cu ²⁺ /Cu ⁺ , oxygen vacancies | 123 |
| LaNiO ₃ | Ofloxacin | 93% within 10 min | $^{1}O_{2}^{-}$ | Ni ³⁺ /Ni ²⁺ , oxygen vacancies | 124 |
| Pd/C ₃ N ₄ | BPA | 91% within 60 min | $^{1}O_{2}^{-}$ | Pd^0 | 114 |
| CuO nanosheet | AO7 in high salinity wastewater | 95.8% within 30 min | $^{1}O_{2}$ | Cu ⁺ /Cu ²⁺ and Fe ²⁺ /Fe ³⁺ | 125 |

converted into Pd(II) during the process, and ¹O₂ is produced via eqn (25). Table 4 summarizes some other relevant metalbased activators in PMS activation.

$$2Pd^{0}O \rightarrow 2Pd^{0} + {}^{1}O_{2}$$
 (25)

Factors impacting singlet oxygen in PMS activation

The broad existence of PPCPs, pesticides, and surfactants in the aquatic environment has driven the development of SR-AOPs. The direct and indirect discharge of PPCPs leads to their accumulation in bodies of water and is detrimental to human health and aquatic life. In principle, the treatment efficiency of PMS/PDS activation is influenced by the practical water matrix, including the temperature, pH value, and salinity.4 The general production of $SO_4^{\bullet-}$ and $^{\bullet}OH$ radicals can be impeded by several inorganic anions (e.g., Cl⁻, CO₃²⁻/HCO₃⁻, NO₃⁻ and SO₄²⁻) and by natural organic matter (i.e., fulvic acid and humic acid). In particular, Cl⁻ can efficiently scavenge SO₄•and OH in SR-AOPs, thereby producing $\text{Cl}_2^{\bullet-}$ and $\text{Cl}^{\bullet.116}$ Generally, the alkalinity of a body of water is tuned by carbonate (CO₃²⁻) and bicarbonate (HCO₃⁻) ions, whose concentrations are in the range of 50-200 mg L⁻¹ in real bodies of water. 117 It has been found that both CO₃²⁻ and HCO₃⁻ act as radical scavengers in AOPs. Furthermore, the radicals SO4 •and OH can be suppressed by NO₃-, SO₄²⁻ and phosphate ions, whereas, for ¹O₂, the inhibitory effect of high salinity is significantly weakened in the non-radical process. 117 The insignificant impact of anions on the ¹O₂-dominated PMS activation process is mainly attributed to the dissymmetry of PMS molecules, which facilitates the self-decomposition of PMS molecules under nucleophilic attack by the anions. Additionally, natural organic matter (NOM) is also a complicated factor in SR-AOPs. Dong et al.⁵⁻⁷ found that the degradation efficiency of PPCPs is not impeded even in the presence of coexisting NOM, which is even self-degraded during the process. Moreover, NOM in aquatic systems acts as a photosensitizer for ¹O₂ formation rather than a scavenger;118 therefore, the effect of NOM on the ¹O₂-dominated systems can be ignored. ^{119,120}

In addition, the solution pH is another crucial parameter depending on the characteristics of the PMS activators. For instance, in homogeneous activation systems, ¹O₂ can be directly generated via PMS activation under neutral (6.5 \pm 0.3) and alkaline conditions, as the surface hydroxyl groups

enhance the chemical binding with PMS molecules. In heterogeneous systems, metal ions can activate PMS utilizing ¹O₂ over an even broader pH range.93 Overall, the catalyst structure, singlet oxygenation, and electron transfer are crucial factors behind the generation of ¹O₂. ⁹³

Conclusions and perspectives

In summary, we have specifically reviewed the evolution of ¹O₂ in SR-AOPs. Since 2016, studies of 1O2 in SR-AOPs have attracted dramatic attention. Compared to $SO_4^{\bullet-}$ and $^{\bullet}OH$ radicals, ¹O₂ has advantages, such as its longer lifetime, mild conditions, resistance to inorganic anions (e.g., HCO₃⁻, Cl⁻, and H₂PO₄⁻), NOMs and free-radical scavengers (e.g., methanol and t-butanol), and ability to degrade electron-rich organic complexes. Due to the advantages of ¹O₂, the ¹O₂-dominated PMS system is a promising technique in wastewater treatment plants. However, some critical points and perspectives need to be addressed in the future.

- · As the degradation efficiency of PMS activation using carbonaceous nanomaterials strongly correlates with the oxygen functionality, the defect degree and the heterogeneous atoms doped in the carbon matrix, the development of advanced carbon-based materials is an effective route to improve SR-AOPs. However, the insufficient oxidation ability of ¹O₂ is an obstacle to the process, and hence, the emerging M-N-C materials are a reliable choice because of their extremely low metallic leaching. In addition, in the context of the formation of ${}^{1}O_{2}$, the mechanism of PMS-Carbon* is worthy of exploration owing to its complexity.
- Among transition-metal-based materials, single transition metal oxides have more oxygen vacancies and exhibit higher degradation efficiency. Additionally, composites of carbon materials and transition metals provide a new way of thinking in SR-AOPs owing to the synergistic effect between them when the carbonaceous materials act as a mediator.
- ¹O₂ is insensitive to background inorganic ions (e.g., Cl⁻, CO_3^{2-}/HCO_3^{-} , NO_3^{-} , PO_4^{3-} and SO_4^{2-}) and NOM, being more accessible to electron-rich compounds. Hence, ¹O₂-mediated PMS systems have great potential for the treatment of actual wastewater and the effluents of river water, as well as drinking water, regardless of the background water matrix.

• Most importantly, based on the theoretical foundation, it is more desirable to think more about how to select and design an appropriate material via computational attempts (e.g., machine learning or artificial intelligence) rather than directly conducting experiments in order to save labour and chemical expenses.

Author contributions

Review

The manuscript was written through the contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Key R&D Program of China (2022YFE0107900), National Natural Science Foundation of China (21972040), the Program of Introducing Talents of Discipline to Universities (B20031, B16017), Innovation Program of Shanghai Municipal Education Commission (2021-01-07-00-02-E00106), the Science and Technology Commission of Shanghai Municipality (20DZ2250400, 22230780200) and Fundamental Research Funds for the Central Universities.

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