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1. Introduction

Carbocatalytic ozonation toward advanced water purification

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Ya Liu,^{ab} Chunmao Chen,^a Xiaoguang Duan, ^{bb} Shaobin Wang^{bb} and Yuxian Wang^{**}

Carbon-based catalytic ozonation is state-of-the-art technology with high oxidation capabilities for wastewater remediation, taking advantage of the synergies of direct ozone oxidation and the generated reactive oxygen species (ROS). Replacing metal-based materials with active and robust carbonaceous catalysts in catalytic ozonation will lead to outperformed catalytic performance, minimized operational cost, and no secondary contamination. Additionally, the variety of allotropes and maneuverable surface chemistry of carbons facilitate structural and surface engineering, which enables the regulation of reactivity, stability and reaction pathways. This review summarizes the recent development of structural control and surface modification of carbocatalysts and their applications in catalytic ozonation. The structure-performance relations and mechanisms are elucidated by a novel model based on the interaction intensity between reactants and carbon surface. Meanwhile, influences of the water matrix parameters on the catalytic system are unveiled. Finally, we provide directions to the rational design of reaction-oriented carbocatalysts, the methodology for mechanistic explorations, and the implementation of ozone-based AOPs in real wastewater treatment.

Advanced oxidation processes (AOPs) exploit reactive oxygen species (ROS), such as hydroxyl radicals (HO[•], $E^0 = 2.7$ V, NHE) and sulfate radicals (SO₄^{•–}, $E^0 = 2.5$ –3.1 V, NHE) generated from diverse peroxides and have emerged as powerful

"State Key Laboratory of Heavy Oil Processing, State Key Laboratory of Petroleum Pollution Control, China University of Petroleum-Beijing, Beijing 102249, China. E-mail: yuxian.wang@cup.edu.cn

^bSchool of Chemical Engineering and Advanced Materials, The University of Adelaide, Adelaide, SA 5005, Australia



Yuxian Wang received his B.Eng. with first-class honor and PhD from Curtin University in 2013 and 2016, respectively. He then joined the College of Chemical and Environmental Engineering, China University of Petroleumthe Beijing as associate professor. His current research interests include seeking feasible applications of nanomaterialbased advanced oxidation processes (AOPs) in environ-

mental remediation and facilitating mechanistic insights into carbocatalysis with advanced protocols. techniques in wastewater decontamination.^{1,2} Among AOPs, ozonation treatments relying on the high oxidation potential of O_3 ($E^0 = 2.05$ V, NHE) can directly destruct unsaturated organics via the Criegee reaction and outperform other AOPs in terms of low energy demand and facile implementation at a full-scale.^{3,4} However, the selectivity and low solubility of O₃ in the aqueous medium limited the mineralization efficiency of the ozonation process.5 Catalytic ozonation treatments achieve improved reaction kinetics and mineralization efficiency compared with ozonation treatment. The heterogeneous catalysts in the ozonated solutions accelerate O3 activation into ROS, which possess greater oxidation capacities and are less selective, thus facilitating the interactions of O₃ molecules and organic contaminants.6 Additionally, the tunable oxidation regimes in catalytic ozonation improve the adaptability to complicated water matrixes. Various metal-based heterogeneous catalysts have been developed for O3 activation, yet the secondary pollution arising from metal leaching becomes the bottleneck in practical applications.⁷

The past two decades witness a rapid development of carbons to replace conventional metal-based systems in environmental remediations on account of fast reaction kinetics and environmentally benign nature of carbocatalysis.⁸ As a result, various carbonaceous materials, including bulky carbons (*e.g.*, activated carbon, biochar and carbon fiber) and novel structured nanocarbons (*e.g.*, carbon nanotubes, graphene-based materials and three-dimensional porous nanocarbons), have been applied in heterogeneous catalytic

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ozonation (Fig. 1).⁹ A broad range of allotropes with different dimensions and porous structures facilitate the in-depth investigation of the mechanism of dimensional effects on catalytic performance. Additionally, the facilely-tuned surface chemistry together with the large surface area of carbonaceous materials provide an advanced platform to build and maneuver surface engineered active sites such as defects, surface functionalities, heteroatom dopants, and anchored metal species.^{10,11} Hence, the engineered carbonaceous materials with optimized physiochemical and electronic properties stimulate O_3 activation to induce either radical or nonradical-based reaction pathways to destroy the recalcitrant organics.

The innovative carbocatalysis in AOPs significantly expedites the development of new carbocatalysts and excavation of the underlying mechanisms. Till now, several reviews have been reported to summarize the development of carbocatalysis in wastewater decontamination,9,12-14 while few focused on the application of carbonaceous materials in catalytic ozonation. Recently, two reviews dedicated to the mechanistic investigations on nanocarbon-based catalytic ozonation.15,16 To the best of our knowledge, there is still lack of a review that comprehensively summarizes the recent advances in carbon-based catalytic ozonation treatments and the synthesis/modification strategies for advanced design of carbocatalysts with desirable activity, selectivity, and stability. In addition, the effects of water matrix parameters such as solution pH and background organic/inorganic substances on the performance of carbonbased catalytic ozonation treatments are rarely mentioned.

In this review, apart from showcasing the roadmap of catalytic ozonation treatments using different carbon allotropes, we

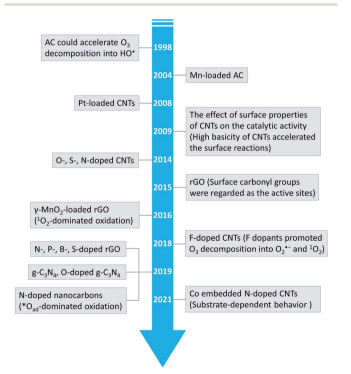


Fig. 1 Timeline of the important development of carbon-based catalysts in catalytic ozonation.

will discuss the catalytic origin and establish the surface engineering strategies for rational design and fast screen of the highly efficient carbocatalysts. For better elucidation of the mechanistic insights into ROS generation and differentiation of reaction pathways, a novel concept based on the intensity of interactions between O_3 and catalysts surface will be proposed. The environmental implications of solution pH, ionic strength, and other background substances such as inorganic anions and natural organic matters (NOMs) on the efficiency of the carbonbased catalytic ozonation treatments will be analyzed. Furthermore, the challenges and future research opportunities for the elaborate design of reaction-oriented carbocatalysts and systematically unravelling the mechanistic insights into carbonbased catalytic ozonation will be envisaged.

2. Synthesis and surface engineering protocols for surface active sites

In general, pristine carbon catalysts are catalytically inactive and their catalytic performances are lower than those of metalbased homogeneous and heterogeneous catalysts. For example, pristine carbon nanotubes (CNTs) obtained by the chemical vapor deposition (CVD) method generally possess a low density of functional groups with C-H terminations on the surface and poor dispersibility in aqueous solution.^{11,17} Surface chemistry and pore structure are key properties of activated carbon (AC) that directly influence their performances in various applications.18 Thus, surface modification and functionalization are conducive to promoting their catalytic performances in the development of highly effective carbon-based materials. Diverse modification methods have been summarized in several reviews,^{8,17,19,20} in which all modification principles are dedicated to creation/tailoring of catalytically active sites and electronic structures of the carbocatalysts. To rationally design and synthesize catalysts with improved activity, selectivity, and stability for a target application, it is crucial to unravel the catalytic origin and intrinsic active sites that govern the catalytic activity.

2.1 Carbon dimensions

Dimensional effects have a great impact on the catalytic activities of catalysts, especially for carbon materials (Fig. 2a). The pristine graphene network with free-flowing π electrons is beneficial to the adsorption and activation of reactants and the electron transfer process. In the catalytic ozonation processes, the conjugated π system of the graphitic carbon framework facilitates the adsorption and activation of O3 molecules.21 The difference in the electron density between carbocatalysts and O₃/organics is of great significance to the strength of chargetransfer interactions.²² Generally speaking, strong activation/ binding is more likely to occur when charge density difference is significant.^{23,24} Moreover, the highly delocalized π system endows sp² carbon with a faster electron transfer rate, which helps the redox reactions for O3 activation and organics oxidation.²⁵ The delocalized π electrons were also reported to be closely correlative to superoxide radical $(O_2^{,-})$ formation in

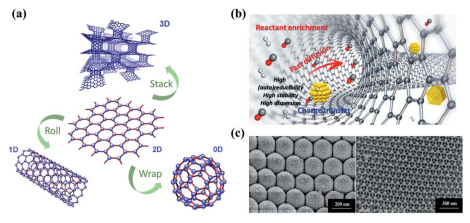


Fig. 2 (a) Construction of different dimensional carbon materials based on graphene. Elements are adapted with permission from ref. 32. Copyright 2012, American Chemical Society; adapted with permission from ref. 33. Copyright 2012, Elsevier. (b) Nanoconfinement effects in carbon nanotubes. Reprinted with permission from ref. 28. Copyright 2020, American Chemical Society. (c) HRTEM images of 3DOM structures. Reprinted with permission from ref. 34. Copyright 2015, Wiley-VCH.

catalytic O₃ decomposition.²⁶ On the other hand, the carbon framework is able to function as a charge-transfer mediator that induces the direct charge transport from the co-adsorbed organics (the electron donor) to oxidants (the electron acceptor) *via* the surface region of the carbocatalyst to achieve contaminant destruction.²⁷ In addition, aromatic organics tend to be adsorbed or activated on the basal plane due to the π - π interactions or electrostatic interactions, followed by an electrophilic attack by ROS.¹⁶ It is worth mentioning that the π orbitals are perpendicular to the graphene layer in graphene, while in CNTs, the curvature structures induce a bent π system and the redistributed electron density from the inner to the outer surface, thereby affecting the catalytic activity.²⁸

Engineering three-dimensional (3D) carbon catalysts with mesoporous and macroporous structures is a promising

strategy to improve the catalytic activity by shortening the mass transport/diffusion length, increasing specific surface area (SSA) and the number of exposed active sites, and further accelerating the reaction kinetics. Nanoconfinement effects are also expected in the porous structure, which facilitate both mass and charge transfer (Fig. 2b). 3D hierarchical carbon structures further enhance electron mobility and facilitate the adsorption of reactants in water media.¹⁶ In the presence of metal catalysts at a high pyrolysis temperature, 3D hybrid nanostructures can be synthesized by intercalating onedimensional (1D) wirelike nanocarbons between the stacked two-dimensional (2D) sheets (Fig. 2a).²⁹ The 3D ordered macroporous (3DOM) carbon architecture with an increased number of active sites and enlarged SSA also demonstrates a favorable application in environmental catalysis (Fig. 2c).^{30,31}

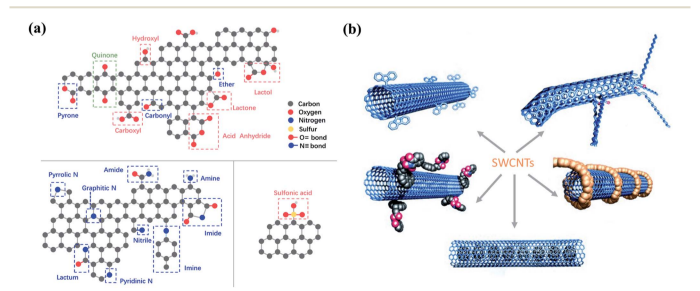


Fig. 3 (a) Possible oxygen-, nitrogen- and sulfur-containing functional groups on carbon surface. Reproduced with permission from ref. 49. Copyright 2013, Royal Society of Chemistry. (b) Functionalization strategies for CNTs. Adapted with permission from ref. 50. Copyright 2002, Wiley-VCH.

2.2 Surface functionalization

The surface chemistry such as surface charge and the electronic properties of the carbon materials is largely dependent on the surface functionalities, especially the surface oxygen functionalities, which have been proposed as the active sites in various carbon-based catalysts (Fig. 3a). Carbonyl groups (-C=O) with high basicity and nucleophilicity are catalytically active. It was found that reduced graphene oxide (rGO)/g-C₃N₄ with a higher content of carbonyl groups exhibited better catalytic performance toward O₃ activation.^{35,36} The electron-rich carbonyl groups with lone-pair electrons can act as Lewis basic sites to activate O₃ by an electron-transfer process. Similarly, carbonyl groups are critical active sites in persulfate activation. It was reported that carbonyl groups located at the boundaries showed a high affinity to interact with peroxymonosulfate (PMS), leading to the significant elongation of the peroxide O-O bond in PMS.37,38 Carboxyl groups (-COOH) can act as Lewis basic sites to facilitate O₃ activation due to its similar properties with -C=O in catalytic ozonation systems. Qu et al. reported that carboxyl functionalized CNTs (CNTs-COOH) possessed a better activity compared with hydroxyl functionalized CNTs (CNTs-OH), because the high charge density of -COOH was beneficial for the direct oxidation of pollutants by O₃.³⁹ Moreover, the deprotonated carboxyl groups reacted with O₃ to form electrophilic oxygen species, which served as radical promoters to produce HO'.40,41 Hydroxyl groups (-OH) as electron-donating groups would reinforce the electron density of the conjugated π system to facilitate the electron-transfer process. Ren and coworkers found that, as compared with carbonyl and carboxyl groups, hydroxyl groups on the CNTs surface were more conducive to persulfate adsorption and electron migration in the peroxodisulfate (PDS)/CNTs system.⁴² In catalytic ozonation, O₃ will electrophilically attack the surface -OH groups to generate ROS (HO' and O2'-).43 Furthermore, -OH groups might be oxidized into -C=O groups by O_3 , and the redox cycle of carbonyl-hydroxyl groups is beneficial to O3 activation.44

Tailoring the type and amount of the surface functionalities is a practical step to enhance the catalytic performance (Fig. 3b). Thus, surface engineering is a crucial way to manipulate the surface functionalities, which can be categorized into surface oxidation and reduction modifications.45 Surface oxidative modification generally employs oxidants (e.g., HNO₃, KMnO₄, H₂SO₄, H₂O₂, and O₃) to introduce a large number of oxygencontaining groups (e.g., -COOH, -OH, -NO₂, -SO₃H, etc.) on the carbon surface, thereby increasing the acidic property. During the surface reduction modification, carbon materials are treated at a high temperature in a reducing atmosphere or are reduced in alkaline solutions, leading to the formation of basic surface functionalities (e.g., -C=O, -NH₂, etc.). Noteworthly, the carbon structure can be entirely damaged by HNO₃ at its boiling point.⁴⁶ For surface-oxidized multi-walled CNTs (MWCNTs), the increased oxygen-containing functionalities (-C=O) on MWCNTs facilitated the adsorption of reaction intermediates and thus accelerated the reaction via tuning the electronic structures of the adjacent carbon atoms.⁴⁷ Graphene modified with amine groups (-NH₂) demonstrated better catalytic activity compared with unmodified graphene for persulfates activation.⁴⁸ Nevertheless, excessive surface functionalities are not desirable.38 Overabundant surface oxygen groups hindered the catalytic activity due to increased electrostatic repulsions and stereo-hindrance effect.42 Moreover, the extra surface oxygen moieties decrease the reductive degree of the carbon lattice for charge transfer and occupy the defective sites or edges, which are also crucial active sites for oxidant activation. Therefore, fabricating carbon materials with optimized oxygen contents to enhance the catalytic performance is highly imperative for future studies. In addition, unveiling the roles of the individual oxygen-containing groups on carbocatalysts is challenging since it is hard to regulate single type of surface functional groups without influencing others.

2.3 Defects engineering

Defect engineering has been demonstrated as an effective strategy to tailor the electronic properties of carbon catalysts. Nanocarbons are inherently produced with edges and topological defects (Fig. 4a). These defective sites with unpaired

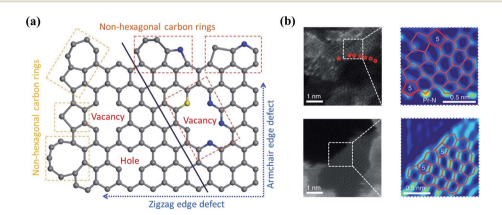


Fig. 4 (a) Defect-engineered nanocarbon. Adapted with permission from ref. 61. Copyright 2021, Springer. (b) The high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images of carbon defects. Reprinted with permission from ref. 62. Copyright 2019, Nature.

electrons disturb the homogeneity of carbon skeleton and tune the electronic structures of basal planes (Fig. 4b).⁵¹ The catalytic activity of edges originates from the presence of dangling bonds, which are high-energy sites and possess high reactivity toward the reactants to promote the radical chain oxidations. The carbon atoms at the edging sites obtain unpaired electrons in a "localized state", which strongly interact with the small molecules to form an attached complex.⁵² It was revealed that the electron transfer at graphene edges was much faster than that on the honeycomb basal plane.⁵³ With the merits, edge sites have been reported to promote O₃ adsorption and subsequent activation.^{25,52,54,55} On the other hand, topological defects (*e.g.*, vacancies and non-hexagonal carbon rings) can break the conjugated π network and redistribute the electrons.⁵³

Defects in a carbon framework can be further created either by downsizing carbons or by removal of the template materials. A novel catalyst for oxygen reduction reaction (ORR) based on graphene quantum dots (GQDs) supported on graphene nanoribbons (GNRs) was developed through a one-step reduction of methylbenzene and hexabromobenzene by Na.56 Such a treatment resulted in abundant surface and edge defects on the GQD/GNR surfaces or at their interface, which served as active sites to promote charge transfer. During the synthesis of Ndoped graphene, both edge and topological defects can be produced by high-temperature pyrolysis for decomposition of the N-rich precursors.57 Meanwhile, topological defects can be created through the direct carbonization of carbon/nitrogen precursors on templates of MgO/Zn-metal-organic frameworks (MOFs).58 Additionally, microwave reduction is an efficient method to synthesize rGO with abundant structural

defects, attributed to the microwave absorbing property of graphene oxide (GO) and the severe and uniform heating ambience provided by microwave irradiation.⁵⁹ Nevertheless, these carbocatalysts were fabricated under high temperatures or using template precursors, in which an accurate control of the generated defects was difficult. Plasma technology for etching the surface of nanocarbons has been proposed as an effective strategy to prepare defect-rich electrocatalysts in a controllable manner by altering treatment time and plasma power.⁶⁰ It is worth mentioning that, although nanocarbons with high-defect levels obtained good reactivities in AOPs, they usually show poor stabilities, because the unsaturated dangling bond and long-paired electrons on defects would induce the electrophilic attack of the oxidants. Moreover, defects might also be changed or recombined under the working conditions.60 Hence, manipulating the defective level is required to balance the trade-off between reactivity and stability.

2.4 Heteroatom doping

Introduction of heteroatoms (*e.g.*, B, N, F, P or S) to the carbon skeleton results in charge/spin redistribution and structural disorder because of the differences in electronegativities and atomic radius (Fig. 5a).^{10,63,64} As a result, the heteroatom dopants and the C atoms adjacent to the dopants with variations in electronic properties are often regarded as the active sites favoring the interaction with O_3 .⁶⁵ N-dopants alter the charge distribution and the density of states (DoS) of the doping area, owing to the great difference in electronegativity between N and C (3.07 *vs.* 2.55); N dopants also endow the

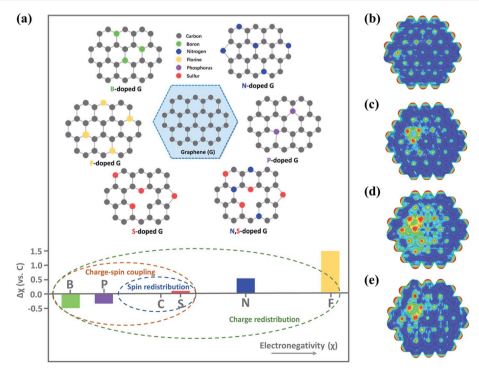


Fig. 5 (a) Illustration on heteroatom-doped carbon materials. Reproduced with permission from ref. 64. Copyright 2018, Wiley-VCH. Reprinted with permission from ref. 16. Copyright 2020, American Chemical Society. Electrostatic potential mapping from charge density matrix for (b) S-rGO, (c) N-rGO, (d) S, N-rGO, and (e) S, S, N-rGO. Reprinted with permission from ref. 67. Copyright 2015, Wiley-VCH.

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adjacent C atoms with a highly positive charge density, facilitating the adsorption and activation of the reactants. Moreover, the intrinsic chemical and electronic coupling induced by Ndoping synergistically promoted proton adsorption and the reduction kinetics.66 Meanwhile, heteroatom doping results in the redistribution of spin density.⁶⁴ Although the similar electronegativities of S to C weakened the intramolecular charge transfer, the higher spin density of S-doped graphene than that of pristine graphene (0.39 vs. 0) favored adsorption of reactants.⁶⁷ Co-doping of the heteroatoms can further improve carbocatalysis due to the synergy between different dopants. The performances of binary (N, P-carbon), ternary (N, P, Fcarbon) and quaternary (S, N, B, P-carbon) doped carbon catalysts have well outperformed the single doping counterparts.⁶⁸⁻⁷⁰ In a N, F co-doped system, F dopants tailored the Ndoping-induced charge transfer and decreased the work function, which accelerated charge transfer for redox reactions.⁷¹ Meanwhile, in N, S co-doped graphene, S atoms could activate the C atoms adjacent to N, which was advantageous for the interaction with PMS to lower the energy barrier for ROS generation.^{67,72} Additionally, the synergistic effect arising from co-doping dramatically enlarged the positively charged regions. Nevertheless, overloading of the S dopants restrained the synergistic effect due to the redistribution of electrons/spins and reduced active regions (Fig. 5b-e).67 Therefore, the doping species, level and configuration need to be carefully controlled.

Typically, heteroatom doping could be achieved by two strategies, direct/*in situ* synthesis and post-treatment.^{11,73} Direct/*in situ* synthesis strategy starts from molecular carbon and dopant precursors; the formation of graphitic carbon framework and heteroatom doping occur concurrently at the elevated temperature by CVD, solvothermal, segregation growth, or arc-discharge approaches.⁷³ Homogeneous doping could be attained by precisely controlling the direct/*in situ*

synthesis parameters. The post-treatment method is to pyrolyze the derived carbon materials with heteroatom-rich resources by a hydrothermal treatment, pyrolysis, ball milling method or plasma treatment.

Multiple factors can affect the catalytic performances of heteroatom-doped carbon catalysts. First, the dopant types and the bonding configurations determine the catalytic activity. N doping introduces graphitic, pyridinic, and pyrrolic Ns in the carbon framework, among which the electron-donating graphitic N is more favorable for O3 adsorption. According to density functional theory (DFT) simulation, graphitic N showed a superior catalytic activity to pyridinic and pyrrolic Ns.74 Altering the annealing temperature and ambience could manipulate the types and compositions of the N-dopants because of the different thermal stability of N species.75,76 Second, the content of heteroatom dopants can also influence doping performance. Although increasing the N content in Ndoped CNTs was reported to enhance the electrocatalytic activity,77 the excessive doping amount might hinder the catalytic activity because of the resulted unfavorable electronic structure and poor structural robustness.78 Therefore, it is significant to optimize the doping content by tuning the amount of heteroatom-containing precursors, substrate structures and treatment parameters. Third, catalytic activity may vary with the dopant location even with the same doping element and configuration. It was suggested that N doping at graphene edges demonstrated the best catalytic activity because the edges obtained a much faster electron-migration rate than the basal plane in graphene.^{74,79} Employing the heteroatomcontaining precursors with well-controlled locations of heteroatom atoms is a promising strategy to control the location of the heteroatom dopants and to establish structure-property relationships in specific applications.¹⁹ Noteworthily, the type, content, and location of dopants in heteroatom-doped carbon

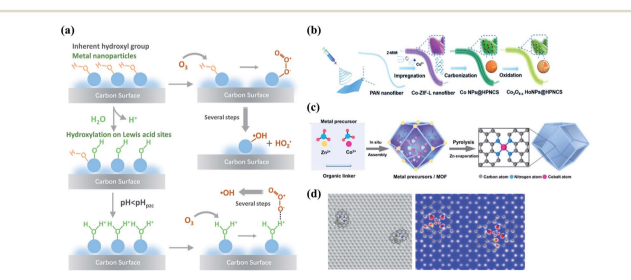


Fig. 6 (a) Schematic illustration of the interactions between O_3 and the surface hydroxyl groups on the metal sites of carbon materials. Reproduced with permission from ref. 16. Copyright 2020, American Chemical Society. (b) Preparation process of Co nanoparticles embedded carbon nanoflakes. Reprinted with permission from ref. 106. Copyright 2019, Wiley-VCH. (c) The formation procedure of Co-SA. Reprinted with permission from ref. 107. Copyright 2021, Wiley-VCH. (d) Simulated HRTEM and STM images of TM-N_x architecture in atomically disperse S|NiN_x species embedded in porous carbon (PC) nanosheets. Reprinted with permission from ref. 108. Copyright 2019, Nature.

catalysts are difficult to be precisely manipulated owing to the complexity of heteroatom dopants and the variety in the surface properties. Novel doping strategies need to be developed, and combined experimental and theoretical approaches will help unveil the chemical nature of the dopants.

2.5 Compositing and doping with metals

Compositing with metal species is another appealing strategy to enhance the performance of carbocatalysis because of the interfacial coupling and charge transfer between the dissimilar counterparts. The defects, heteroatom dopants, and surface oxygen functional groups on the carbon surface provide unique sites for anchoring the metal species owing to the strong binding strength and accelerated electron transfer. Metal species with electron-donating d-orbital electrons can improve electron conductivity and electron-transfer capacity of carbon materials by regulating their electronic properties such as spin density and work functions.^{80–82} Therefore, the strong couplings between the metal species and carbon support significantly change the interface charge density and the coordination environment of active sites, ultimately regulating the catalytic activity.

The redox couples of metal cations (Me^{n+}/Me^{n+1}) facilitate O₃ decomposition into ROS via creating the electron-transfer environment during catalytic ozonation.83 Moreover, the surface of metal oxides is commonly covered by hydroxyl groups, including the inherent hydroxyl groups and the hydroxyl groups arising from the surface hydroxylation process, which act as the Lewis acid sites to accelerate O3 decomposition or induce a substitution reaction with O3 to form surfaceadsorbed active species (Fig. 6a).^{16,84,85} Oxygen vacancies (OVs) are typical defects in metal oxides which contain unsaturated coordination sites and unpaired electrons.86 OVs can enhance the mobility of the surface oxygen and thus facilitate electron transfer in catalytic reactions. Electronic properties of metal/ carbon composites can be regulated with the existence of OVs. The abundant OVs within the surface in NiCo2O4-loaded hollow carbon spheres (HCS) increased DoS near the Fermi level and decreased work function and also optimized the valence state of active sites, resulting in a remarkable enhancement in ORR activity.87 Meanwhile, OVs of TiO2 resulted in inter-bandgap states and a lower flat band potential in a CNT/TiO₂ hybrid, promoting the photocatalytic activity.88 Recently, the generation of high-valent metal species has been successfully verified in various kinds of AOPs.⁸⁹⁻⁹¹ In the Fe(II)/O₃ system, the involvement of high-valent iron-oxo species (Fe(IV)-oxo) formed from the reaction of Fe(II) with O_3 was identified by ¹⁸O isotopelabeling technique.92 The generated high-valent metal-oxo species are highly reactive and capable of oxidizing the organic contaminants.

Impregnation has been widely employed for the synthesis of metal-loaded carbon catalysts, whereas the loading rate should be below the threshold of 30 wt% to attain the well-distributed metal particles.⁹³ To achieve both high metal loading and optimal particle size distribution, MOFs can be utilized as templates.^{94,95} Compared with metal-loaded carbon composites, encapsulation

of metal clusters/nanoparticles within carbon layers by forming a core@shell structure successfully avoid undesirable metal leaching (Fig. 6b). Meanwhile, the encapsulated metal species will modulate the electronic state of outer-sphere graphitic layers via an electron tunnelling effect at the carbon-metal interfaces and enhance electron-transfer capability.96,97 Additionally, the size of anchored transition metals (TM) can be downsized to an atomic level to form single-atom catalysts (SAC), which can be synthesized by a variety of approaches, such as CVD, atomic layer deposition, pyrolysis, wet chemical method, and electron/ion irradiation (Fig. 6c).98,99 Extensive attention has been given to single-atom transition metal (TM) species coordinated N-doped nanocarbons (TM $-N_x$ -C) because of their high catalytic activities stemmed from the unique TM-N coordination (Fig. 6d).¹⁰⁰⁻¹⁰² Such TM-N coordinations have been regarded as the principal active centers, because the hybridizations between the d-orbitals of TM and the 2p-orbitals of nitrogen increased the electron densities near the Fermi level and the work function of carbon basal plane.¹⁶ The inexpensive precursors, such as inorganic salts, carbon, and nitrogen sources, have been extensively employed for the preparation of TM-N_x-C catalysts.¹⁰³⁻¹⁰⁵

3. Catalytic ozonation with different carbon allotropes

3.1 Bulky carbons

By virtue of its high surface area, large porosity, plentiful surface functionalities and low costs, activated carbon (AC) has been widely used to remove micropollutants from water. Combined AC with ozonation was found to be an attractive method for wastewater decontamination owing to the synergistic effect between adsorption and catalytic oxidation (summarized in Table 1). The rich porosity and maneuverable surface functionalities expedite the interactions of both organic pollutants and O₃ on the surface of AC. Moreover, it was reported that AC or carbon black could act as the initiator for radical chain reactions to accelerate O₃ decomposition into HO^{.109,110} In ACinitiated catalytic ozonation, variations in solution pH and surface chemistry of AC will exert a profound influence on the catalytic performance.

Solution pH can be a critical factor affecting the ozonation activation pathways, yet the mechanism is still controversial. Beltrán et al. applied AC in catalytic ozonation of oxalic acid (OA) at acid pH and proved that catalytic degradation of OA at pH 2.5 could be ascribed to a HO'-based radical pathway occurring in the bulk solution.111 In another study, O3 mainly adsorbed on the surface of AC without generating free radicals at acidic pH to near neutral pH (pH 2 to 6), while it tended to react with the surface hydroxyl groups on the AC to produce HO' in bulk solution when solution pH > 6.110 Faria et al. reported that surface oxidation was the dominant degradation route of OA at pH 3 since the addition of HO' scavenger resulted in a marginal inhibition on the destruction rate of OA.¹¹² At alkaline pH, both surface oxidations and bulk reactions relying on HO' were responsible for OA destruction. Additionally, the greater degradation efficiency of OA was achieved at acidic pH

Table 1 Application of c	Application of carbonaceous materials in catalytic ozonation for the abatement of pollutants	on for the abatement of pollutants			
Catalysts	Synthesis/modification methods	Target pollutants	Active sites	ROS	Ref.
AC	Commercially purchased	Oxalic acid, gallic acid, 1,3,6- naphthalenetrisulphonic acid, diethvl phthalate	Delocalized-π system, surface acid/basic groups	HO', activated surface-O ₃ complex	40, 111, 117 and 160
GAC	Commercially purchased	<i>P</i> -Nitrophenol, chemical industrial wastewater	Surface basic groups	.0Н	113-115
ACHNO ₃	Modification with HNO ₃	Oxalic acid, oxamic acid, benzenesulfonic acid, sulfanilic acid	Surface basic groups	.OH	112 and 161
AC-NO ₂ AC-NH ₃	Modification with HNO ₃ / ammonia	Oxalic acid	Surface basic group (–NH ₂ , pyrone groups), acid oxygenated surface groups	.0Н	118
AC-N ₂ AC-H ₂ AC-NH,	Thermal treatment under $N_2/H_2/NH_3$ atmosphere	Oxalic acid	Sitrogen-containing groups, basal planes, basic oxygen-containing oronnes	.0Н	120
AC-NH ₃ AC-NH ₄) ₂ CO ₃ AC-urea	Modification with ammonia/ ammonium carbonate/urea	Naphthalenesulphonic acid	Nitrogen-containing groups, basic oxygen-containing groups	.0Н	119
Biochar	Pyrolysis of pistachio hull	Reactive red 198	Surface functional groups	HO', activated surface-O ₃ complex	125
Biochar Biochar	Pyrolysis of sludge Pyrolysis of sludge, treatment with HCl/NaOH	Phenol, oxalic acid Petroleum refinery wastewater	Surface carbonyl groups Functional C groups, Si–O structures, metallic oxides	02	124 and 126 122
CNTS	Commercially purchased	<i>P</i> -Chlorobenzoic acid, oxalic acid	Graphene basal plane, surface oxygen groups, defects	.0Н	21 and 55
MWCNTs	Pretreatment with O ₃	Oxalic acid	Surface basic groups	HO', surface- oxygenated radicals	129
CNTs-COOH CNT-N ₂ CNT-H ₂	Pretreatment with HNO_3/H_2SO_4 Pretreatment with HNO_3 followed by thermal treatment under N_2/H_2 atmosphere	Indigo Oxalic acid	Surface carboxylic groups Surface basic groups	.0Н	39 130
MWCNT-HNO ₃ MWCNT-H ₂ O ₂ MWCNT-O ₂	Pretreatment with HNO_3/H_2O_2 , thermal treatment in the gas phase with 5% O_2 in N_2	Oxalic acid, oxamic acid	Surface basic groups	HO', surface- oxygenated radicals	131
N-doped CNTs	Ball milling of CNTs with urea or melamine followed by thermal annealing	Oxalic acid	Doped N functionalities	I	136
N/S-doped CNTs	Liquid-phase oxidation with HNO ₃ /H ₂ SO ₄	Oxalic acid, phenol	Doped N/S functionalities	.0н	135
F-doped CNTs	Mixing of CNTs with hydrofluoric acid	Oxalic acid	Doped F, surface basic groups, delocalized-π system	$0_2^{-1}, {}^10_2$	138
GO	Modified Hummer's method, commercially purchased	5-Chloro-2-methyl-4- isothiazolin-3-one, <i>N</i> , <i>N</i> - diethyl- <i>m</i> -toluamide	Surface oxygen groups	.0Н	139, 140 and 148

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Table 1 (Contd.)					
Catalysts	Synthesis/modification methods	Target pollutants	Active sites	ROS	Ref.
rGO	Thermal reduction of GO under N, atmosphere	P-Hydroxybenzoic acid	Surface carbonyl groups	$0_2^{}, {}^10_2$	35
rGO	Thermal reduction/chemical reduction/hvdrothermal method	Oxalic acid, 4-nitrophenol	Defect sites	HO', 0_2 ^{'-'} , 1O_2	142
rGO N-doped rGO	Microwave reduction method, mixing of GO with ammonium nitrate followed by microwave irradiation	4-Nitrophenol	Edging structures, dangling bonds, doped N	но', 0 ₂ ' , ¹ 0 ₂	59
N-doped rGO	Ball milling of GO with urea or melamine followed by thermal annealing	Oxalic acid	Doped N functionalities	.0Н	145
N/P-doped rGO	Mixing of GO with ammonium nitrate/ammonium phosphate monobasic followed by thermal annealing	Sulfamethoxazole	Defective sites, dopants	.он	146
N/P/B/S-doped rGO	Mixing of GO with melamine/ H ₃ PO ₄ /H ₃ BO ₃ /mercaptoacetic acid followed by thermal annealing/hydrothermal treatment	Benzotriazole, <i>p</i> - chlorobenzoic acid	Surface hydroxyl groups, the free electron in heteroatoms, delocalized π electron	HO', 0 ₂ ^{, 1} 0 ₂	25
Layered N-doped nanocarbon	Pyrolysis of β -cyclodextrin and melamine under N ₂ atmosphere	Oxalic acid	Pyridinic N and C atoms adjacent to graphitic N	HO', *O _{ad}	74
Nanocarbon with built-in nanotubes	Pyrolysis of <i>a</i> -cyclodextrin and cobalt nitrate hexahydrate under Ar atmosphere	Oxalic acid	Defective sites	HO', *O _{ad}	159
N-doped hollow sphere carbon	Pyrolysis of polydopamine using a template-etching method	Ketoprofen	Doped N functionalities	HO', electron transfer mechanism	158
g - C_3N_4	Pyrolysis of melamine/urea	4-Chlorobenzoic acid, benzotriazole	N vacancies	но', о ₂ '	36
O-doped $g-C_3N_4$	Pyrolysis of melamine followed by hydrogen peroxide hydrothermal treatment	Atrazine	Surface oxygen groups, N vacancies	HO', O ₂ '-	155
O-doped g-C ₃ N ₄	Pyrolysis of melamine and oxalic acid	<i>P</i> -Chlorobenzoic acid, atrazine	Surface carbonyl groups	.0Н	154 and 156

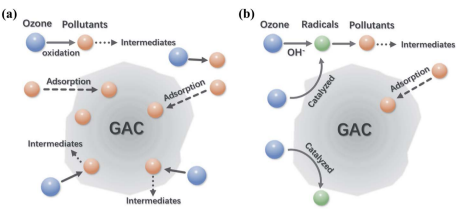


Fig. 7 Schematic illustrations of the main reactions on GAC surface under (a) acidic conditions and (b) basic conditions. Reproduced with permission from ref. 113. Copyright 2008, American Chemical Society.

than at basic pH, though a higher pH promoted the decomposition of O_3 into more reactive species.

Variations in solution pH not only affect the decomposition rate of O_3 , but also alter the surface charge of AC as well as the dissociation of organic molecules in aqueous solutions. Surface of AC is positively charged when solution pH is lower than pH at the point of zero charge (pH_{pzc}) of AC, and its surface becomes negatively charged *vice versa*. Meanwhile, the organic molecules could dissociate to anionic species at a solution pH higher than its pK_a. Hence, the resulted electrostatic forces (attraction or repulsion) between the surface of AC and the organic molecules at different solution pHs affect their interactions, which influence the catalytic performance and oxidation pathways.¹¹² The principal reactions on AC at different pH conditions are schematically illustrated in Fig. 7a and b.¹¹³

The activity of AC can also be correlated with the types and contents of surface functionalities. Increasing the amounts of acidic surface oxygen functionalities with electron-withdrawing ability can be accounted for the activity loss of AC after the catalytic ozonation reaction.114 The increased amount of acidic functionalities reduces the affinity to the electrophilic interaction with O₃ molecules and prevents O₃ decomposition in solution. Acid or alkali surface modifications are effective in optimizing the catalytic activity of AC. Faria and co-workers modified AC by nitric acid treatment (ACHNO₃), and the increased surface acidity by introducing carboxyl groups on AC surface led to the decrease in catalytic ozonation activity.¹¹² The delocalized π electron system of basic groups on AC surface favored OH⁻ adsorption and acted as catalytic centers to catalyze HO' generation.110,115,116 Also, pyrone-like and chromenelike surface oxygen functionalities react actively with O₃ via Criegee mechanism to yield hydrogen peroxide (H_2O_2) , which reacts with the AC surface or with O3 itself for HO' generation.113,114,117 It was also observed that the adsorption capacity of basic AC was superior to more acidic ones because of the increased electronic attraction resulted from the higher pH_{pzc} of alkaline AC.¹¹² In addition, surface basic oxygen functionalities facilitate the occurrence of surface-based oxidations. However, the generated HO' around the high basicity active

sites was difficult to diffuse into the bulk solution owing to its short lifetime and rapid reaction with surface-adsorbed organics. Additionally, the surface basic oxygen functionalities on AC tended to dissociate into cationic species with positive charges, which impeded the electrophilic attack of O3 for HO' generation. Thus, the basicity of AC may indirectly weaken HO'dominated oxidation in bulk solution.¹¹⁸ Despite of its electronwithdrawing character, the deprotonated acid oxygenated surface groups can enhance the formation of O₂^{•-} and O₃^{•-} as a consequence of its reaction with O₃, which function as radical initiators to generate HO' through the radical chain reactions.40 Surface nitrogen-containing functionalities also influenced the catalytic activity of AC. ACs with various types and concentrations of nitrogenated functional groups were prepared, and urea-treated AC with enriched pyrrole groups and low concentrations of pyridine and pyridone groups demonstrated the best catalytic ozonation activity among ACs treated by different nitrogenating agents.¹¹⁹ The pyrrole groups extensively increased the electronic density of the graphitic layers in AC, which served as the catalytic active sites and promoted O₃ decomposition into HO¹²⁰ On the contrary, the π -deficient pyridine groups diminished the electronic density of AC basal plane. This is in line with another study of Cao's group, where the pyrrole groups mainly accelerated HO' generation in the bulk solution but the pyridine groups had a detrimental effect on the surface reaction.120

Biochar is a type of bulky carbon prepared from the pyrolysis of biomass or solid wastes, which possesses the merits such as abundant precursors with low costs, bulk level preparation and manipulatable surface chemistry. Thus, biochar as a highly modifiable green platform has drawn widespread attention in environmental remediation.¹²¹⁻¹²⁴ The properties of biochar are highly dependent on different factors such as feedstock type and pyrolysis temperature. Moussavi and Khosravi converted pistachio hull into biochar *via* thermal pyrolysis (500 °C for 2 h).¹²⁵ The pistachio hull-derived biochar had a strongly basic surface (pH_{pzc} = 11.5) and a macroporous structure in favor of adsorption of O₃ and dye molecules. Therefore, a higher decolorization efficiency of reactive red 198 was observed in the catalytic ozonation process with pistachio hull-derived biochar than that with AC. Pyrolysis temperature has certain effects on the morphological, physicochemical, and catalytic capacity of biochar. For instance, Zhang et al. produced biochar using raw sludge obtained from coking wastewater at different pyrolysis temperatures (300, 500, 700, 900 °C) for catalytic ozonation of phenol.¹²⁶ Raising the pyrolysis temperature enhanced the graphitization level of the biochar and stabilized the carbon structure and the inherently integrated metal components. Substantial reduction in total organic carbon (TOC) releasing and metal ions (e.g., Co²⁺, Fe³⁺, Ni²⁺, Cu²⁺ and Mn²⁺) leaching were observed when pyrolysis temperature was 900 °C. In addition, high-temperature pyrolysis also favored the formation of a porous structure with an increased surface area by promoting the evaporation of the volatile matters in biochar feedstock. The increasing amount of oxygen-bonded carbon groups formed at elevated pyrolysis temperature endows the biochar with the ability to remove contaminants, since the functionality will facilitate the adsorption of O₃ within the surface region and participate in the electron-transfer process.

3.2 Nanocarbons

Although bulky carbonaceous materials are cost-effective and facile to prepare, their microporous structures limit the diffusion of both reactants and products in catalytic reactions. Thus, bulky carbons usually demonstrate poor activity and reusability in catalytic ozonation. The mixed mesoporous or microporous structures in nanocarbons such as carbon nanotubes (CNTs), graphene allotropes, carbon nitride (g-C₃N₄), and 3D hierarchical carbons shorten the length of mass diffusion, facilitate the exposure of active sites, and accelerate the reaction kinetics. Additionally, the surface chemistry of nanocarbons can be easily regulated. The introduced defects or heteroatoms will further improve the catalytic activity. Herein, nanocarbons with well-defined structures and chemistry are promising platforms to gain insights into catalytic ozonation as well as other carbonbased AOPs. Table 1 summarizes carbonaceous materials in catalytic ozonation for the removal of organics.

3.2.1 Carbon nanotubes. CNTs belongs to the 1Dstructured allotrope of nanocarbons, which can be regarded as a tubular structure of rolled graphene layers. The mesoporous structure enables CNTs with a large external specific surface area which facilitates the dispersion of active components and promotes the interactions with reactants. Surfaceengineered CNTs have been widely used as heterogeneous catalysts for ozonation in aqueous solution to degrade organic pollutants.55,127,128 The highly graphitic degree of CNTs with limited structural topological defects favors grafting oxygen functional groups. Surface functionalization and heteroatom doping are two main strategies to modify the surface chemistry of CNTs, thereby enhancing the catalytic activity. The role of surface functional groups in the catalytic activity has been studied by Ma's group.129,130 Compared with untreated MWCNTs, the catalytic activity of MWCNTs was significantly reduced after O3 oxidation pretreatment, which was ascribed to the remarkable decrease of surface basic groups and the

increase of acidic groups, especially for carboxyl acid groups.129 N2 or H2 heat treatment of CNTs further increased the pHpzc value and afforded higher catalytic activity in OA degradation.130 It was found that higher pH_{pzc} of CNTs resulted in a greater rate constant for OA removal. Compared with N₂ heat treatment, H₂ heat treatment had a more significant effect in eliminating acidic groups because of the reducibility of H2. Gonçalves et al. also reported similar results that low acidity of MWCNTs promoted the catalytic ozonation of aliphatic acids.131 In these scenarios, surface basic functionalities stimulated the evolution of O₃ or H₂O₂ via Criegee reactions into active species, such as HO'. Additionally, CNTs with a negative surface charge density were favorable to OA adsorption, which accelerated the surface reactions.129,130 Confirmed by the quenching tests with the presence of tert-butanol (TBA), a typical HO' scavenger, only minor inhibition on the activity of heat-treated CNTs was observed compared with CNTs. Therefore, similar to AC, CNTs with a higher basic character were more conducive to catalytic ozonation of OA through surface reaction.

The effects of oxidized surface of CNTs on the catalytic activity are debatable, which can be favorable,55 unfavorable129,130 or marginal.131,132 One obstacle in the clarification of this argument is that it is difficult to alter the surface chemistry without changing the morphology or dispersity of CNTs after a prolonged exposure in the highly oxidative environment during catalytic ozonation processes; additionally, it is challenging to directly correlate the catalytic activity with one or more surface properties.^{133,134} To distinguish the influence of CNTs surface chemistry and CNTs aggregation on catalytic activity, Oulton et al. investigated the effects of MWCNTs surface properties and suspension stability on HO' production in a catalytic ozonation system by functionalizing MWCNTs with HNO3.55 It was found that the oxidized MWCNTs possessed a higher rate of HO' formation compared with the pristine MWCNTs. Dynamic light scattering and sedimentation studies indicated that oxidized MWCNTs exhibited a lower degree of aggregation than pristine MWCNTs, and the better dispersion favored O3 decomposition and HO' formation. Moreover, surface oxygen functionalities (e.g., -COOH, -C=O, and -OH) resulting from HNO3-pretreatment were also responsible for HO' formation.

Tailoring the surface chemistry and the electrochemical properties of CNTs by substituting carbon atoms with metalfree heteroatoms (e.g., N, S, and F) can be an alternative strategy to promote catalytic activity. Restivo et al. reported that the introduction of N and S species led to improved catalytic efficiency of CNTs in catalytic ozonation.135 Soares et al. further prepared N-doped CNTs by a simple ball-milling method.¹³⁶ The elevated catalytic activity and the high stability were observed for as-prepared N-CNTs, which may be primarily attributed to the pyridine N groups. However, compared with non-doped ball-milled CNTs, no significant improvement in the mineralization degree of the organic contaminants was observed for Ndoped CNTs.136,137 It was suggested that ball milling led to the breakage of nanotubes and the decrease in the entanglement, thus resulting in the increase in surface area and more exposure of active sites; however, N-doping decreased the surface area. F

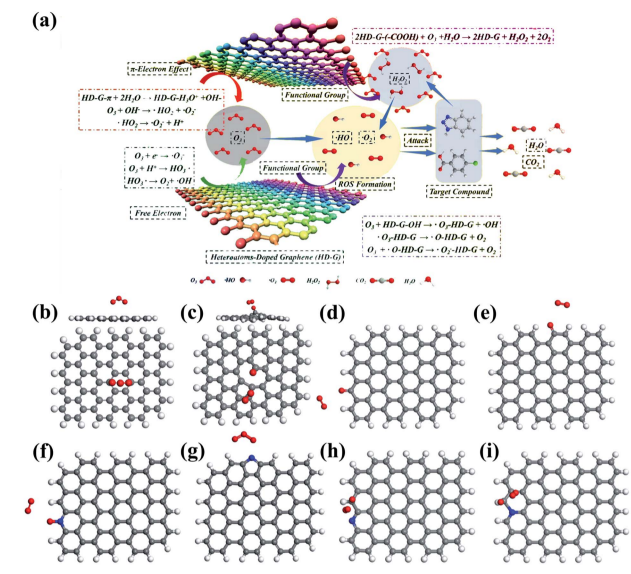


Fig. 8 (a) Mechanism of catalytic ozonation by heteroatom-doped graphene. Adapted with permission from ref. 25. Copyright 2019, American Chemical Society. DFT simulations of O_3 adsorption on the (b) basal plane, (c) vacancy, (d) zigzag edge, and (e) armchair edge of graphene. Reprinted with permission from ref. 142. Copyright 2018, Elsevier. DFT simulations of O_3 adsorption on (f) pyridinic N, (g) pyrrolic N, C with high charge density adjacent to (h) pyridinic N, and (i) edge graphitic N. Reprinted with permission from ref. 74. Copyright 2019, Elsevier.

atoms are the most electronegative elements with an extraordinary electron-withdrawing ability. Wang and co-workers found that F-doped CNTs obtained a significantly improved catalytic ozonation activity in OA destruction than the undoped CNTs, and outperformed N-CNTs and several metal oxides (*e.g.*, ZnO, Al₂O₃, Fe₂O₃, and MnO₂).¹³⁸ The covalent C–F bonds in F-CNTs formed in sp³ hybridization maintained the integrity of the delocalized π networks, which were beneficial to O₃ activation *via* an electron-transfer process.

3.2.2 Graphene and derivatives. Graphene is a promising 2D material consisting of honeycomb-like sp² carbon layers. Graphene-based materials have attracted intensive research attention due to their unique chemical and electronic properties. Different from CNTs, topological defects can be introduced during the synthesis or post-treatment processes. The high

surface area of graphene facilitates the adsorption of reactants and the exposure of active sites for O₃ to decompose into ROS.

GO¹³⁹⁻¹⁴¹ and rGO^{35,36,59,142} have been widely employed to activate O₃ for the degradation of various contaminants. rGO displayed much higher catalytic performance than GO ascribed to the higher defective level and a proper amount of surface oxygen functionalities. By the purification of the graphite from the anode of waste lithium-ion battery (LIB), Wang *et al.* synthesized defects-rich rGO (LIB-rGO).¹⁴² The as-prepared LIBrGO outperformed rGO obtained from commercial graphite powder with a mediocre defective level in catalytic ozonation for 4-nitrophenol degradation. DFT simulations validated that the peroxide bond in O₃ would be broken to generate surfaceadsorbed atomic oxygen (*O_{ad}) and free peroxide species (*O₂ free) when O₃ molecules were placed on vacancies, zigzag, and armchair edges of rGO, giving rise to the adsorption energies of -3.88, -5.69, and -4.64 eV, respectively (Fig. 8c-e). In contrast, the peroxide bond in O₃ only slightly prolonged when placed above the graphene basal plane, resulting in adsorption energy of -0.32 eV (Fig. 8b). The more negative adsorption energy of O_3 on defective sites than that on the basal plane demonstrated the vital role of structural defects, especially the edging sites.¹⁴² Additionally, O₃ interacting with different oxygen functional groups (-OH, -C=O, -COOH, and C-O-C) of graphene were also performed by DFT calculations. Among different configurations, -OH groups on the zigzag edge of graphene exhibited the highest affinity toward O₃ as it obtained the most negative adsorption energy with O_3 (-1.00 eV). Moreover, the longest peroxide bond (l_{O-O}) in O₃ (1.360 Å) also indicated the greatest possibility for O₃ decomposition on surface hydroxyl groups.¹⁴² However, as discussed above, the number of surface oxygen functionalities should be manipulated at a proper level, otherwise, the overabundant surface oxygen functionalities, such as the ones on GO, can impede the catalytic activity by affecting the electron-transfer process and inhibiting the adsorption of both O₃ and organics due to the steric effect.

Doping of metal-free heteroatoms in the graphene structure can further improve the catalytic activity and stability by

regulating the chemical and electronic properties of rGO.143,144 N-rGO was prepared by a simple ball milling method using urea (rGO-U) or melamine (rGO-M) as nitrogen precursors.145 The higher SSA (102 vs. 47 m² g⁻¹) and a greater amount of doped N (9.3% vs. 7.5%) endowed rGO-M with higher catalytic activity than rGO-U. Furthermore, metal-free doping also created a large number of defective sites.146 Song and co-workers prepared N-, P-, B-, and S-doped rGO and the catalytic ozonation activity for wastewater decontamination was in the order of N-rGO > P-rGO > B-rGO > rGO > S-rGO.²⁵ They suggested that the unpaired electron/free electrons and delocalized π electrons from defective sites in heteroatom-doped rGO participated in the electrontransfer process, in which unpaired electrons were easily captured by O_3 to generate HO' and the delocalized π electrons expedited O2^{•-} formation. Furthermore, surface -OH and -COOH groups also acted as the active centers for ROS evolution in this study (Fig. 8a). DFT simulations revealed insights into the activity improvement induced by heteroatom doping.74 It was found that N-doping redistributed the charge density of the graphene basal plane and enabled the pyridinic and graphitic N-doping sites as well as C atoms adjacent to graphitic N to be electron-rich zones, which acted as active sites for catalyzing O₃ dissociation (Fig. 8f-i). The chemisorbed O₃ molecules on the

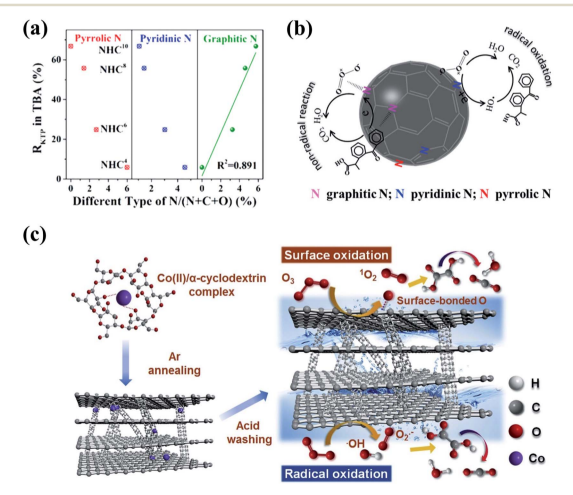


Fig. 9 (a) Relationships between the removal rates of ketoprofen and the content of different N species of NHCs. (b) Mechanism of ketoprofen degradation by catalytic ozonation on NHCs. Reprinted with permission from ref. 158. Copyright 2019, American Chemical Society. (c) Schematic diagram of catalytic ozonation of OA with CPGs. Reprinted with permission from ref. 159. Copyright 2020, Elsevier.

active sites tended to dissociate into ${}^{*}O_{ad}$ and ${}^{*}O_{2}$ free based on *in situ* Raman spectrum and DFT simulations. With a high oxidation potential of 2.43 V, ${}^{*}O_{ad}$ could directly attack the adsorbed OA molecules *via* nonradical oxidation, and it was able to react with the ambient H₂O to generate surface-adsorbed HO^{*}.

Despite of their high catalytic activities, graphene-based materials in catalytic ozonation reactions often demonstrated poor stability and became passivated after the long-term operation. In Song's study, the increase of TOC was observed in both 4-chlorobenzoic acid and benzotriazole degradation using GO as the catalyst.³⁶ The soluble GO would react with O₃ and ROS during catalytic ozonation, leading to the destruction of its structure and the change in surface chemistry. O3 would attack GO first by oxidation and follow by decomposition. In the oxidation stage, oxygen-containing groups were generated on the surface of GO, and the decomposition stage would give rise to the decay of carbon framework and release of CO₂/CO.147 Du et al. reported that O₃ was only capable of oxidizing functionalized aromatic rings in GO; the C-O-C/C-OH groups could be oxidized to C=O and O-C=O by O₃, which could lead to the opening of the aromatic ring.148 The combination of HO' and O3 could oxidize both functionalized and unfunctionalized aromatic carbon rings. Previous studies on applications of GO in other oxidative systems, such as UV/H2O2,149 UV/PS,149 and photo-Fenton process¹⁵⁰ also reported the similar observations, indicating that GO was not an appropriate candidate for AOPsbased technologies. Though rGO exhibited better structural robustness than GO in catalytic ozonation,36 it would still experience gradual deactivation with the increase of cyclic numbers,35,54,142 which was mainly ascribed to the change of surface chemistry and the blockage of active sites by the degradation intermediates. Herein, the stability and recyclability of graphene-based materials should be evaluated for applications in AOPs, and the treatment time needs to be optimized to avoid the increase of TOC caused by graphene decomposition.

3.2.3 Other nano-scale carbon-based materials. Apart from CNTs and graphene-based catalysts, graphitic carbon nitride (g- C_3N_4) with a 2D layered structure composed of intralayer heptazine units with sp²-hybridized C and N atoms has been employed in catalytic ozonation. Pristine g-C₃N₄ usually obtains insufficient activity for the degradation of oxalic acid,151,152 bisphenol A,151 and p-hydroxybenzoic acid.153 Modification of g-C₃N₄ with metal-free heteroatoms will disrupt the sp²-hybridized carbon network and induce new physicochemical and electronic properties for photocatalytic ozonation.12,154 Yuan et al. found that oxygen functionalized $g-C_3N_4$ (O@g-C₃N₄) composite could enhance the removal efficiency of atrazine in catalytic ozonation systems.155 The introduction of O atoms with lone electron pairs effectively regulated the electronic properties of surrounding atoms and facilitated their interactions with O_3 to generate HO' and O_2 .⁻ through radical chain reactions. Furthermore, compared with direct oxidation, pre-doping O atoms into g-C₃N₄ followed by HNO₃ oxidation generated extra attachment sites at which oxygen-containing groups could be anchored.156 The extensively grafted carboxyl groups at an apex angle in the heptazine unit of g-C₃N₄ were the principal active sites. Meanwhile, the construction of N vacancies on the g-C₃N₄ framework also enhanced the catalytic ozonation activity. As reported by Song et al., delocalized electrons generated by N vacancies and carbonyl groups in g-C3N4 facilitated the decomposition of O₃ into HO' for 4-chlorobenzoic acid and benzotriazole degradation as well as bromate elimination.³⁶ However, the overabundant hydrogen bonds might hinder electron transfer and thus impact catalytic activity. It should be noted that the heptazine unit of g-C₃N₄ would be attacked by HO' into nitrates, while g-C₃N₄ could maintain structural stability under the attack of O2.⁻ and O3.¹⁵⁷ Due to the competitive reactions between organic pollutants and g-C₃N₄ for HO', the decomposition of g-C₃N₄ could be completely or partially impeded in the presence of organic pollutants with small molecular weights. On the contrary, in the case of pollutants with larger molecular weights, both g-C₃N₄ and pollutants were simultaneously oxidized by HO'.

3D nanocarbons with porous structures and large SSA are conducive to the maximum exposure of active sites and the mass transfer in heterogeneous catalytic processes. Ma's group prepared in situ N-doped hollow sphere carbons (NHCs) via the facile pyrolysis of polydopamine using a template-etching method, and principally figured out the specific role of three N species including pyrrolic, pyridinic, and graphitic Ns in catalytic ozonation (Fig. 9a).¹⁵⁸ The extraordinary catalytic ozonation activity of NHCs for ketoprofen destruction was attributed to the synergy between HO' radical-based oxidation and a nonradical intra-electron-transfer pathway. Pyrrolic and pyridinic Ns at edges and structural vacancies functioned as the "radical-generation" region to initiate HO'-based oxidation, while the graphitic N served as the "electron-mobility" region to accelerate the electron transfer from the adsorbed organics to the activated O3 via a nonradical oxidation pathway (Fig. 9b).158 Metal-free 3D mesoporous graphitic carbons with built-in nanotubes (CPGs) were synthesized via a facile one-pot copyrolysis method using cost-efficient precursors (Fig. 9c).159 The CPGs showed superior catalytic ozonation activities in OA destruction compared with CNTs, rGO, and LaMnO₃ perovskite catalysts, which could be ascribed to the nanoconfinement effect brought by the presence of few-layer (5-10) built-in nanotubes within the bulk structure of graphitic carbons.

3.3 Carbon-based nanocomposites

Heterogeneous metal-based catalysts show desirable activities, but generally suffer from poor stability due to metal leaching and agglomeration. Anchoring metal species on supports can significantly improve the stability and dispersion of metal species, thereby suppressing the metal leaching and facilitating the exposure of active metal sites (Fig. 10a). Moreover, the electron conductivity and electron-transfer capacity can be enhanced due to the synergistic effect between the metal species and the supports, which further contributes to the catalytic activity. Carbonaceous materials have been exploited as versatile catalyst supports because of their adjustable

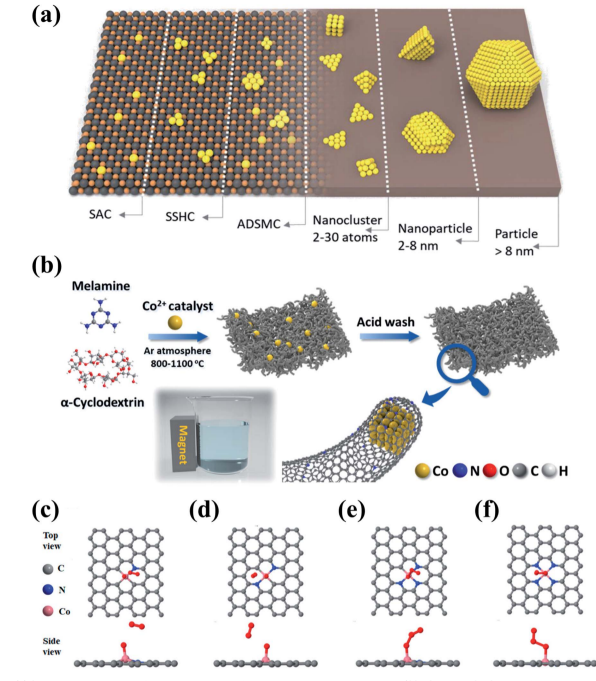


Fig. 10 (a) Schematic illustration of the various types of metal–carbon nanocomposites. (SSHC and ADSMC are defined as single site heterogeneous catalyst and atomically dispersed supported metal catalyst, respectively.) Reprinted with permission from ref. 195. Copyright 2019, Wiley-VCH. (b) Schematic illustration of the synthesis route for Co–N@CNTs. DFT simulations for mimicking the adsorption of O_3 onto (c) Co– N_1 , (d) Co– N_2 , (e) Co– N_3 , and (f) Co– N_4 . Reprinted with permission from ref. 96. Copyright 2021, American Chemical Society.

dimensional architecture, SSA, and surface chemistry which facilitate the anchoring and dispersion of metal species.

Transition metals have been recognized as the catalytic active sites in ozone activation owing to their multivalence states originated from their 3d orbitals, which accelerate the electron-transfer process *via* redox reactions. Different transition metals have been applied to modify carbon materials, such as Mn, Fe, Co, and Zn, as well as their oxides (summarized in Table 2). Ma *et al.* anchored manganese oxides (MnO_x) on

granular active carbon (GAC) to enhance the ozonation of nitrobenzene, and higher catalytic treatment efficiency was observed at low pH (2.74–3.52) than that at high pH (6.72–9.61) because the low pH condition benefited the adsorption of nitrobenzene on the catalyst.¹⁶² AC derived from sewage sludge (SBAC) was also used to load Mn.^{163,164} The organic degradation pathways were different in these studies which were either attributed to HO⁻based degradation or the surface electron transfer among the loaded metal oxides and the adsorbed

Table 2 Application of carbon-based nanocomposites in catalytic ozonation for the abatement of pollutants

Catalysts	Synthesis/modification methods	Target pollutants	Active sites	ROS	Ref.
0	d				
Supported mangan MnO _x /GAC	Impregnation method	Nitrobenzene, oxalic acid	Mn species	HO', surface adsorbed ROS	162 and 163
MnO ₂ /biochar	Impregnation method	Piggery bio-treatment effluents	Surface hydroxyl groups	но.	196
MnO _x /biochar	Impregnation method	Atrazine	Lewis acid sites	HO', O ₂ '-	197
MnO _x /MWCNT	Impregnation method	Ciprofloxacin	Mn^{n+}/Mn^{n+2}	HO.	198
MnO ₂ -NH ₂ /GO	Hydrothermal method	Cephalexin	Mn^{3+}/Mn^{4+}	02	188
z-MnO ₂ /rGO	Hydrothermal method	Bisphenol-A	_	HO.	165
β-MnO ₂ /rGO	Hydrothermal method	Dichloroacetic acid	_	но.	166
γ-MnO ₂ /rGO	Hydrothermal method	4-Nitrophenol	Mn(III)/Mn(IV)	$O_2^{*-}, {}^1O_2$	83
Supported iron oxi	des				
Fe/AC	Dipping method, commercially purchased	Dibutyl phthalate, formate	Fe species	но.	169 and 172
FeO _x /biochar	Impregnation method	Atrazine	Lewis acid sites	HO', O ₂ '-	197
Fe ⁰ /CNTs	Electrophoresis deposition followed by calcinations	Methylene blue	Fe(III)/Fe(IV)	но.	199
Fe ₃ O ₄ /MWCNTs Fe ₃ O ₄ /GO	Hydrothermal method Precipitation method	Bisphenol A <i>p</i> -Chlorobenzoic acid, ibuprofen	Surface oxygen groups —	но. Но.	173 141
Supported cobalt o	vides				
Co/AC	Impregnation method	Nitrobenzene	Co species	но.	200
Co–N@CNTs	Two-step carbonization method	Oxalic acid, phenol	Two pyridinic N- coordinated Co (Co–N ₂)	*O _{ad} , ¹ O ₂	96
Supported zinc oxid	des				
GAC@ZnO Zn/CNTs	Deep-coating method Thermal solvent treatment process using zinc power, MWCNTs and 40% polyethylene glycol followed by pyrolysis under N ₂ atmosphere	Metronidazole 4-Chloro-3-methyl phenol	Zn species Zinc oxide (hydroxide)	НО [•] НО [•] , О₂ ^{•−}	201 174
ZVZ/g-C ₃ N ₄	Pyrolysis of zero-valent zinc (ZVZ) and $g-C_3N_4$	Atrazine	Surface hydroxyl groups, ZVZ/ZnO	HO', O ₂ ' ⁻ , ¹ O ₂	175
ZnO/g-C ₃ N ₄	Mixing and calcination method	Atrazine	Surface hydroxyl groups, Zn species	HO', $O_2'^-$, ${}^{1}O_2$	94
Supported copper of					
CuO/AC	Impregnation method	Heavy oil refinery wastewater	Cu species	но'	202
CuO/g-C ₃ N ₄	Reflux condensation method	Oxalic acid	Oxygen vacancies, surface hydroxyl groups	HO', O ₂ ' ⁻	203
Supported platinur					
Pt/CNT	Incipient wetness impregnation method	Oxalic acid	Pt_{red}/Pt_{ox}	но	176
Supported cerium			2.		
Ce/AC	Precipitation method	Textile dyes, dimethyl phthalate, <i>p-</i> toluenesulfonic acid	Ce ³⁺ species	но.	177, 178 and 180

Table 2 (Contd.)

Catalysts	Synthesis/modification methods	Target pollutants	Active sites	ROS	Ref.
CeO ₂ /AC	Impregnation method	Oxalic acid, aniline	Ce ³⁺ species	но.	179 and 181
CeO ₂ @AC	Urea-mediated hydrothermal method	Oxalic acid	Oxygen vacancies	но'	204
CeO ₂ /AC, CeO ₂ / MWCNT	Impregnation method	Erythromycin	Ce ³⁺ species	HO', surface- oxygenated radicals	182
CeO ₂ /OCNT	Pretreatment of CNTs with HNO ₃ , and hydrothermal treatment of OCNT and $Ce(NO_3)_3 \cdot 6H_2O$ followed by calcinations	Phenol	Surface hydroxyl groups, Ce ³⁺ /Ce ⁴⁺	но.	183
CeO ₂ /rGO	Hydrothermal method	<i>N,N</i> -Diethyl- <i>m</i> - toluamide	Ce^{3+}/Ce^{4+}	но.	205
Ce/g-C ₃ N ₄	Pyrolysis of Ce(NO ₃)₃·6H₂O and g- C₃N₄	Oxalic acid	Ce(m), surface hydroxyl groups	но'	96
Supported bimetall	ic oxides				
Fe ₂ O ₃ -CeO ₂ /AC	Impregnation method	Sulfamethoxazole	Metal oxides	HO'	206
MnO ₂ -Co ₃ O ₄ /AC	Impregnation– precipitation method	Incineration leachate	_	но.	192
Fe-Ni/AC	Impregnation method	2,4- Dichlorophenoxyacetic acid	Iron–nickel	но.	207
Fe-Mn/GAC	Impregnation– desiccation method	Methyl orange	Metal species	но'	208
MnFe ₂ O ₄ /rGO	Electrospinning method	Di- <i>n</i> -butyl phthalate	Surface hydroxyl groups, Mn ²⁺ /Mn ³⁺	но.	193 and 194
LaCoO ₃ /g-C ₃ N ₄	Pyrolysis of $LaCoO_3$ and $g-C_3N_4$	Benzotriazole	N vacancies, $-C-O-Co$ bonds, Co^{2+}/Co^{3+}	HO', O ₂ '-, ¹ O ₂	97

organics and O₃.^{163,164} However, the multivalence of Mn stimulated the electron-transfer in both processes. Great efforts have also been put into the hybridization of rGO and MnO₂ for catalytic ozonation.^{83,165,166} 2D γ -MnO₂/rGO nanohybrids (MnO₂/rGO) were fabricated and used for catalytic ozonation by Wang and co-workers.⁸³ The redox reactions between Mn³⁺ and Mn⁴⁺ *via* single electron transfer acted as the active sites for the catalytic reaction. The electron transfer process was further promoted by the tight interaction between MnO₂ and rGO.

Mn/CNTs composites were developed to intensify catalytic ozonation by uniting the merits of both homogeneous and heterogeneous reactions.¹⁶⁷ The reversible desorption and adsorption of Mn^{2+} on the surface of CNTs could be regulated by the spontaneous variations of solution pH during pollutant mineralization. At the first stage, organic acids as the degradation intermediates would gradually accumulate and lower the solution pH. CNTs surface was positively charged when solution pH < pH_{pzc} of CNTs, leading to the desorption of Mn^{2+} by the electrostatic repelling forces. The dissolved Mn^{2+} in the solution allowed homogeneously catalytic ozonation. With the mineralization of organic acid intermediates, solution pH gradually increased and Mn^{2+} re-adsorbed on the electronegative CNTs surface when solution pH was higher than the pH_{pzc} of the

catalyst, thereby achieving the recovery of Mn^{2+} . However, during this process, Mn^{2+} was inevitably oxidized into MnO_2 during catalytic ozonation and resulted in the decrease of catalytic efficiency.

Iron-based catalysts have been widely employed because of their comparatively low price and high catalytic performance. In combination with Fe-loaded AC for catalytic ozonation, the treatment efficiency for various refractory pollutants such as crystal violet,168 dibutyl phthalate,169 and organic contaminants in heavy oil refining wastewater¹⁷⁰ and biologically pretreated coal gasification wastewater171 were greatly enhanced. In Yuan's study, an iron oxide-impregnated AC catalyst (JBX) was only effective to activate O3 for formate/oxalate degradation at pH 3, while the alkaline environment (pH 7.3 and 8.5) inhibited the catalytic activity.¹⁷² As revealed, the positively charged protonated iron oxide surface sites at acidic pH obtained a higher affinity toward O₃ than the deprotonated sites at basic pH. Fe₃O₄/MWCNTs samples with the similar surface area but different surface oxygen functional groups were also fabricated to investigate the effect of oxygen-containing groups on catalytic ozonation activity.173 Fe3O4/MWCNTs with the maximum amount of surface oxygen-containing groups exhibited the highest adsorption capability and catalytic activity, because

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surface oxygen functionalities could combine with organics *via* hydrogen bonding and induce O₃ decomposition.

The built-in electric field at the Zn-CNTs interface facilitated O_3 adsorption and subsequent conversion into H_2O_2 , which further evolved into HO' for organics degradation.174 Additionally, the formed zinc oxide and zinc hydroxide with small particle sizes and large SSA could also promote the decomposition of O₃ to yield HO'. The introduction of zero-valent zinc (ZVZ) modified the electronic structure of pristine g-C₃N₄, and the redox couple of ZnO/ZVZ was favorable to the electrontransfer process.175 Loading ZnO on g-C3N4 also induced a higher density of defective sites.94 Meanwhile, the formed Zn-N bonds between ZnO and g-C₃N₄ adjusted the electron density of the nanocomposites near the Fermi level and optimized the work function for promoting the generation of photoinduced electrons. Platinum (Pt) has also been loaded on the carbonaceous supports as the ozonation catalysts.176 CNTs supported Pt catalyst (Pt/CNTs) was found to be more active than Pt/AC in the ozonation of OA, because of the higher accessibility of Pt particles locating on the outer surface of CNTs.¹⁷⁶ It was reported that the redox couple of Pt was decisive for HO' generation.

As a typical rare earth oxide, cerium oxide (CeO_2) is more active than other rare earth oxides due to its special structure of 4f orbit and high oxygen mobility.177 A coupling effect of AC and CeO₂ was reported to enhance the catalytic ozonation efficiency in the treatment of textile effluents,178 pharmaceutical wastewater¹⁷⁷ as well as the removal of individual or mixed micropollutants.^{179,180} The free electrons on the basal plane of AC were beneficial for the formation of Ce³⁺ species and thus promoted the redox reactions involving the cerium redox pair (Ce^{3+} / Ce⁴⁺).^{178,181} The catalytic activity and selectivity of Ce/AC depend on the cerium precursors (Ce3+ or Ce4+). For oxalic acid destruction, the catalysts obtained from the "Ce4+" precursor possessed the highest mineralization efficiency, whereas aniline mineralization was favored by the catalysts prepared from the "Ce³⁺" precursor.¹⁷⁹ Gonçalves et al. compared the catalytic performance between CeO2/MWCNTs and CeO2/AC for the ozonation of erythromycin (ERY).182 The synergic effect between carbon materials and CeO₂ was more evident in CeO₂/ MWCNTs because of the higher dispersion of CeO2 on MWCNTs than that on AC. Additionally, ERY with large molecular weight tended to adsorb and react on the mesoporous surface of CeO2/MWCNTs rather than on the microporous surface of CeO₂/AC. The dispersion of CeO₂ on CNTs can be regulated by chemical treatment to increase the surface functionalities of CNTs. The well-dispersed CeO2 was beneficial to the exposure of active sites and decrease the mass-transfer resistance.¹⁸³ The abundant π - π electrons on the surfacefunctionalized CNTs facilitated the mobility of electron and thus accelerated the redox cycle of Ce^{3+}/Ce^{4+} for O₃ activation.

The bonding strength between metal species and carbon supports is a decisive factor influencing the activity and stability of carbon-supported catalysts. Previous studies indicated that the presence of strong chemical bonds between graphene and active components could lead to the improvement of electrochemical performance as well as cyclic stability of constituents.^{184,185} Therefore, reinforcing the interface bonding force can be a viable strategy to improve both the activity and the long-term stability of carbon–metal nanocomposites. The introduction of surface functionalities on the carbon supports provides strong binding sites for metal species and helps to stabilizing the structure of catalysts.^{186,187} Given that the hybrid structure of MnO₂–GO was not stable enough because of the weak physical adsorption and electrostatic interaction between MnO₂ and GO, Xu *et al.* synthesized a novel structured catalyst *via* amino-functionalizing the hybrids of MnO₂ and GO (MnO₂– NH₂–GO).¹⁸⁸ The amino-functionalization endowed MnO₂– NH₂–GO higher stability than MnO₂–GO *via* strengthening the covalent bonding between MnO₂ and GO to prevent their decoupling. The formed strong covalent bonds bridged by NH₂ would accelerate the charge transfer.

In addition to the surface functionalization, the introduction of heteroatoms into the carbon lattice alters the electronic and adsorption properties in the local region and increases the reactivity of adjacent carbon atoms, which possess efficient localized grounds for bonding with metal species.189,190 Taking advantage of Co₃O₄ and N modification, the N-doped graphene (NG) supported Co₃O₄ nanocrystals (Co₃O₄/NG) showed outstanding catalytic activity and stability compared with its sole counterparts.191 Co3O4/NG obtained a smaller particle size and better dispersion of Co₃O₄ compared with Co₃O₄/G because of NH₃-Co coordination. The increased surface π electronic density and the basic active centers (e.g., pyridinic-N and pyrrolic-N) brought by N dopants promoted electron transfer between the π orbitals of graphene and the d orbitals of Co atoms, leading to the enhanced phenol adsorption and O₃ activation. Recently, Wang et al. reported the remarkable catalytic activity and chemical stability of N-doped carbon nanotubes with encapsulated Co (Co-N@CNTs).96 The embedded Co nanoparticles endowed Co-N@CNTs with magneticity, which was favorable for magnetic separation and recycling (Fig. 10b). DFT calculations suggested that pyridinic N-Co manifested more negative adsorption energy than other types of N-Co coordination, and Co coordinated with two pyridinic N atoms (Co-N₂) demonstrated the greatest affinity toward O₃ adsorption to induce surface-based nonradical oxidation (Fig. 10c-f). Meanwhile, the high DoS of Co-N2 at the Fermi level facilitated the electron transfer from Co-N2 to O3 and the redox cycle at the active sites.

Additionally, the activity and durability of carbocatalysts can also be improved through bimetallic loading. Li and co-workers used $MnO_2-Co_3O_4/AC$ in catalytic ozonation and observed the increased biodegradability (BOD₅/COD) of incineration leachate containing abundantly refractory humic-like and fulvic-like components, compared with AC loaded with single metal.¹⁹² The relationship between the concentration of surface hydroxyl sites on rGO-MnFe₂O₄ and the removal rate of di-*n*butyl phthalate was in positive linearity.^{193,194} The hydroxyl groups on the surface of rGO-MnFe₂O₄ hybrids bonded with MnFe₂O₄ and C atoms to form chemisorbed oxygen groups (Mn-O-H, C-O-H, or C-OOH), which acted as the active sites for catalytic ozonation.¹⁹⁴ In a LaCoO₃/g-C₃N₄ activated O₃ system, the charge transfer between electron-rich centers

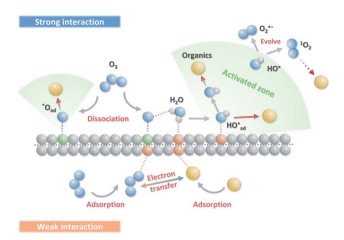


Fig. 11 Schematic illustration of the interactions of O_3 on carbon surface for ROS generation.

created by -C-O-Co or nitrogen vacancies and electron-poor centers promoted HO' generation.⁹⁷

4. Catalytic mechanisms in carbondriven ozonation

4.1 Correlations of O₃@carbon interaction intensity and reaction pathways

Carbon-based ozonation processes typically involve free radicals (such as HO' and HO' /O2'-) and nonradical (surfaceadsorbed activated species and singlet oxygen (¹O₂)) oxidation pathways.^{15,16} The adsorption of O₃ on the catalysts surface is a vital process for O3 activation and the subsequent ROS formation. The intensity of interactions between the active sites and the reactants (O_3 and organics) governs the oxidation pathways (Fig. 11). Defects and active sites with high electron density could induce strong interactions with O₃, resulting in the dissociation of O₃ into surface-adsorbed atomic oxygen (* O_{ad}) and a free peroxide species (* $O_{2 \text{ free}}$) (eqn (1)). * O_{ad} with a high oxidation potential of 2.43 V will elevate the surface potential, which facilitates the activation of the ambient water molecules to generate HO' and triggers the subsequent radical chain reactions (Fig. 11).74,96,142 *Oad also favors the hydrogen abstraction reaction with surrounding water molecules or H⁺, and such a process is governed by the solution pH. The asformed *O_{ad}-H might further activate O₃ molecules for HO' formation. In situ characterizations with the isotopic substitution of ${\rm ^{16}O_3}$ or ${\rm H_2}{\rm ^{16}O}$ by ${\rm ^{17}O_3}$ or ${\rm H_2}{\rm ^{17}O^{209,210}}$ and the DFT simulations on thermodynamics feasibility of the reactions between *O_{ad} and H₂O via calculating the free energies of intermediates and the energy barriers of transition states can be performed in the future studies to unveil the origin of HO'.^{211,212} The HO'-based ozonation obtains fast reaction rates toward almost all kinds of organics because of the high oxidizing capability and nonselectivity of HO'. Nevertheless, the short half-life of HO' ($t_{1/2} \le 1 \ \mu s$) limits the effective mass transfer and long-range attack, making HO' only effective around the surface region of catalysts.²¹³ Additionally, *O_{ad} can induce nonradical oxidation. The high oxidation potential makes it capable of mineralizing recalcitrant organics such as aliphatic acids. The strong interactions between the active sites and O_3 could form a positively charged zone on catalyst surface. Even if organics cannot adsorb on the surface, organics within this activated zone can be captured and oxidized (Fig. 11).

During the radical chain reactions, HO'_2/O_2 ⁻ and ${}^{1}O_2$ can be produced as side products (eqn (2)–(7)). With a mild oxidation potential of 0.89–1.7 V, HO'_2/O_2 ⁻ favors nucleophilic attack of certain organics (*e.g.*, phenolics) *via* nucleophilic substitution. Singlet oxygen, as one of nonradical ROS, also possesses a mild oxidation potential (0.81 V). The reaction rates of ${}^{1}O_2$ with organic compounds range from 10⁴ to 10⁷ M⁻¹ s⁻¹, which are much inferior to those with free radicals.²

$$O_3 \xrightarrow{active sites} *O_{ad} + *O_2 free$$
 (1)

$$O_3 + OH^- \rightarrow HO_2^{\cdot} + O_2^{\cdot -}, \ k = 70 \ M^{-1} \ s^{-1}$$
 (2)

$$O_3 + HO' \rightarrow O_2 + HO'_2 \tag{3}$$

$$\mathrm{HO}_{2}^{\bullet} \leftrightarrow \mathrm{O}_{2}^{\bullet-} + \mathrm{H}^{+}, \ \mathrm{p}K_{\mathrm{a}} = 4.8 \tag{4}$$

$$*O_2 \xrightarrow{\text{electron transfer 1}} O_2$$
 (5)

$$\mathbf{O_2}^{\cdot -} + \mathbf{HO_2}^{\cdot} + \mathbf{H}^+ \rightarrow \mathbf{H_2O_2} + {}^1\mathbf{O_2}$$
(6)

$$O_2^{\bullet-} + HO^{\bullet} \rightarrow {}^1O_2 + OH^-$$
(7)

Surface active sites on carbon-based catalysts with moderate electron density would lead to weak interactions with O3, which is conducive to forming a surface-O3 complex. The activated complex would potentially elevate the redox potential of the carbon framework, making it capable of oxidizing the adsorbed organics via an intramolecular electron-transfer process (Fig. 11).^{158,214,215} In this process, the carbon substrate acts both as a catalyst and an electron tunnel for electron transport. A similar activated surface-oxidant complex was reported in persulfate/nanocarbon systems, where the formed carbon-persulfate complex demonstrated a mild oxidation potential (<1.2 V) with particular selectivity toward electron-rich organics.216,217 Although PDS obtains a similar oxidation potential to O₃ (2.01 vs. 2.05 V), the more complicated molecular structure of PDS dissipates the energy transferred from the adsorption site to the O-O bond and hinders its further activation to a higher-energy state. Moreover, O₃ presents a higher oxidation potential than PMS (2.05 vs. 1.75 V). Therefore, a greater oxidation ability of the activated surface-O₃ complex is expected than that of activated persulfates (PDS and PMS) complex, which has been proved by higher efficiency of aliphatic acids mineralization.218,219

4.2 Analytical approaches for ROS identification

Tracing the occurrence of ROS and confirming their contributions during catalytic ozonation processes are indispensable for mechanistic studies. Due to the diversity of ROS generated from O_3 activation, it is difficult to precisely evaluate the species, quantity, and roles of the reactive species by a single strategy.

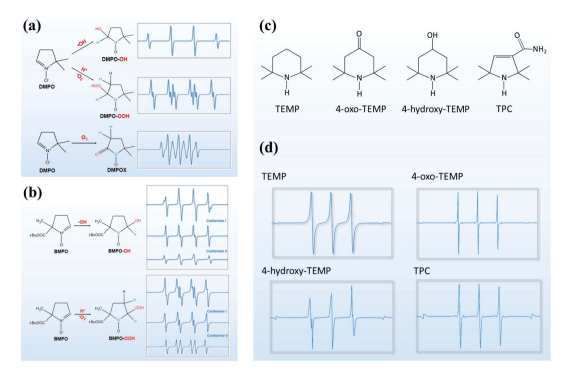


Fig. 12 (a) Structures and simulated EPR signals of DMPO–HO[•] and DMPO–HOO[•] adducts formed by trapping HO[•] and O_2^{--} with DMPO and DMPOX formed by oxidation of DMPO with O_3 . (b) Structures and EPR signals of BMPO–HO[•] and BMPO–HOO[•] adducts formed by trapping HO[•] and O_2^{--} with BMPO. Adapted with permission from ref. 16. Copyright 2020, American Chemical Society. (c) Chemical structures of four sterically hindered amines. (d) EPR signals of nitroxide radicals formed by trapping ${}^{1}O_2$ with four sterically hindered amines. Reproduced with permission from ref. 230. Copyright 2017, Royal Society of Chemistry. Reproduced with permission from ref. 231. Copyright 2011, Royal Society of Chemistry.

Therefore, multi-strategies would be encouraged to obtain an in-depth understanding of their generation and contributions.

4.2.1 Electron spin resonance/paramagnetic resonance. Electron spin/paramagnetic resonance (ESR/EPR) spectroscopy provides an efficient way to identify the types of radicals based on the characteristic signals of the spin adducts from the reactions between ROS and their corresponding spin trapping agents. 5,5-Dimethylpyrroline-N-oxide (DMPO) and 5-tertbutoxycarbonyl-5-methyl-1-pyrroline-N-oxide (BMPO) are the most prevailing spin trapping agents for HO' and O2'-, respectively, due to their good solubility and stability (Fig. 12a and b). The signal of DMPO-HO' adducts (peak intensity ratio: 1 : 2 : 2 : 1, hyperfine splitting constants: $a_{\rm N} = a_{\rm H}^{\beta} = 14.9$ G) can be easily discriminated by EPR, while the identification of HO'_2/O_2 ·- is challenging in catalytic ozonation. This is because the oxidation potential of $HO_2^{\cdot}/O_2^{\cdot-}$ is relatively weak and their production often accompany by other ROS such as HO' and ${}^{1}O_{2}$. 35,220 Additionally, the reaction rate of DMPO with O_{2} . is extremely lower than that with HO' (2–170 M^{-1} s $^{-1}$ vs. 1.9–4.3 \times $10^9 \text{ M}^{-1} \text{ s}^{-1}$),²²¹ and the characteristic signals for DMPO-HOO' adducts ($a_{\rm N} = 14.2 \text{ G}, a_{\rm H}{}^{\beta} = 11.4 \text{ G}, \text{ and } a_{\rm H}^{\gamma 1} = 1.2 \text{ G}$) and DMPO-HO' adducts normally overlap. In order to identify the DMPO-HOO' adducts, adopting absolute ethanol as the reaction medium instead of ultrapure water can prolong the half-life time $(t_{1/2})$ of $O_2^{\cdot-}$ and scavenge HO' in the meantime.³⁵ However, carbon-centered radicals, as the byproducts of the quenching reaction, can also be captured by DMPO, which

interfere with the distinguish of $O_2^{\cdot-.^{222}}$ Spin-fittings can thus be performed for the further differentiation. Furthermore, DMPO-HOO' adducts are unstable ($t_{1/2} = 45$ s) and tend to convert into DMPO-HO' adducts.²²² In some cases, the characteristic signal of 5,5-dimethyl-2-oxo-pyrroline-1-oxyl (DMPOX) with the peak intensity ratio of 1:2:1:2:1:2:1 ($a_N = 7.3 \pm$ 0.1 G, $a_H = 3.9 \pm 0.1$ G) is observed, which is attributed to the oxidation of DMPO by O₃ or surface-activated species (Fig. 12a).²²³ DMPO-HO' can also be oxidized into DMPOX *via* electron extraction by HO'.²²⁴ Compared with DMPO, BMPO is a better candidate for O₂⁻⁻ spin trapping. The higher stability and longer half-life of BMPO-HOO' adducts ($t_{1/2} = 23$ min) facilitate the identification of HO'₂/O₂⁻⁻ against HO'.²²⁵

Singlet oxygen can be detected by EPR in aqueous solution by four different sterically hindered amines as trapping reagents, including 2,2,6,6-tetramethylpiperidine (TEMP), 2,2,6,6tetramethyl-4-piperidone (4-oxo-TEMP), 2,2,6,6-tetramethyl-4piperidinol (4-hydroxy-TEMP) and 2,2,5,5-tetramethyl-3pyrroline-3-carboxamide (TPC) (Fig. 12c), which can be oxidized by ¹O₂ to generate nitroxide radicals.²²⁶ The EPR signals of nitroxide radicals generated from each amine demonstrate distinctive triplet peaks (Fig. 12d). Interestingly, 4hydroxy-TEMP can be oxidized into two different EPR signals: 4hydroxy-2,2,6,6-tetramethylpiperidine-N-oxyl (TEMPOL) and 4oxo-2,2,6,6-tetramethyl-1-piperidinyloxy (4-oxo-TEMPO). Among the four sterically hindered amines, 4-hydroxy-TEMP and TPC have a proper sensitivity and a wide detectable range for the yield of ${}^{1}O_{2}$.²²⁶ It is noteworthy that other reactive species (*e.g.*, HO', HO'_2/O_2'⁻) can also oxidize TEMP to produce similar 3-line characterization signals as ${}^{1}O_{2}$.^{221,227-229} Introducing radical scavengers to minimize the effects of other radicals would cause the competitive reactions and further mislead the results. Additionally, TEMP-based spin trapping agents are alkaline and might induce the base activation of O₃. Therefore, the dosage of TEMP and solution pH should be precisely controlled.

4.2.2 Selective ROS scavenging tests. Quenching tests are based on the differences in the reactivity and rates of the reactions between the scavengers and the active species, which can be performed to differentiate the generated ROS and identify their contributions in removing target pollutants. Considering the co-existence of various oxidants, the ideal scavengers should have high selectivity toward the target ROS; otherwise the competitive reactions between the target ROS and other oxidative species (e.g., O₃ and other ROS) would mislead the differentiation of dominant ROS. For instance, on account of the high oxidation potential of HO', it can react rapidly with most of the quenchers, such as tert-butanol (TBA), para-benzoquinone (p-BQ), furfuryl alcohol (FFA), and sodium azide (NaN₃). Thus, the corresponding scavengers are required to exclude the contribution of HO'. When TBA is employed as the HO' quenching agent ($k = 6 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$), it is only effective for scavenging HO' in the bulk solution rather than the surface-adsorbed HO',232 because of the low affinity of TBA toward the catalyst surface.^{233,234} Meanwhile, fine bubbles might be produced with the presence of TBA which would enhance the mass transfer.²³⁵ Alternatively, methanol is capable of quenching HO' both in bulk solution and on the catalyst surface, as well as preventing the generation of fine bubbles. However, quenching tests using radical scavengers are hard to distinguish the weak-oxidative ROS from high-oxidative ROS because of their differences in reactivities. Furthermore, the organics or reductive agents might attain fast reaction rates with O3, resulting in competitive

consumption. p-BQ can react rapidly with HO₂/O₂^{'-}, HO' and O_3 , $(9.8 \times 10^8, 1.2 \times 10^9 \text{ and } 2.5 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$, respectively).²³⁶ Meanwhile, the low solubility of *p*-BQ hinder its contact with O_2 ·-. NaN₃ is a strong reductant which would only quench 1O_2 $(2 \times 10^9 \text{ M}^{-1} \text{ s}^{-1})$, but react with O₃ with a mediocre constant $(2.5 \times 10^{6} \text{ M}^{-1} \text{ s}^{-1})$.²³⁷ Moreover, N₃⁻ is a strong alkaline agent, which would influence solution pH and cause base activation of O₃. Taking advantage of the kinetic solvent isotope effect (KSIE), the contribution of ${}^{1}O_{2}$ can be better evaluated via using $D_{2}O$ instead of H₂O as the solvent, because the lifetime and steadystate concentration of ¹O₂ would increase in D₂O owing to the lower reaction rate of ¹O₂ in D₂O than that in H₂O ($k_d(H_2O) =$ $2.4-3.2 \times 10^5 \text{ s}^{-1}$; $k_{\rm d}({\rm D}_2{\rm O}) = 1.4-2.3 \times 10^4 \text{ s}^{-1}$).²³⁸⁻²⁴⁰ The KSIE effect using D₂O as the reaction medium can also be employed in EPR technique for ¹O₂ identification, and augmented signal intensity is expected for a ¹O₂-based system. Nevertheless, similar effects can also be caused by other ROS because of their reduced reaction rates in D2O prolonged the life time.241 Therefore, careful attention should be paid to the explanation of the experimental results. Potassium iodine (KI) and dimethyl sulfoxide (DMSO) have been used as scavengers for surfaceadsorbed ROS in other AOPs systems.96,242 However, KI can react with O_3 (1 \times 10⁸ M⁻¹ s⁻¹) and other oxidative species with high reaction rates, leading to the depletion of both the quenchers and dissolved O3. DMSO obtains an inferior reaction rate with O_3 (0.4162 M⁻¹ s⁻¹), whereas it can fast react with both HO' and surface-adsorbed ROS. Hence, the type and concentration of the scavengers should be carefully selected, in view of the undesired competition reactions with ozone and other ROS as well as the impact on solution properties. In a complex system involving various types of ROS, the scavenger might become unspecific to the desired ROS; thus comparing the reaction rates between scavengers and different oxidants is essential to determine the contribution of the specific ROS. In addition, the combination of scavengers with higher selectivity

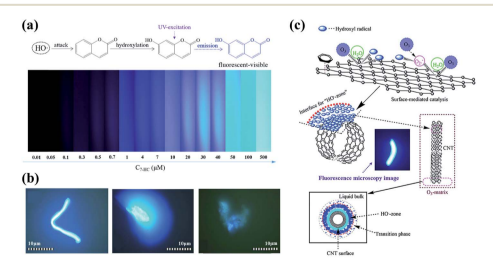


Fig. 13 (a) Colorimetric references of 7-hydroxycoumarin solutions with the changes to the 7-hydroxycoumarin concentration for FMI. (b) Fluorescence microscopy images of the produced HO⁺ during the catalytic ozonation on the surface of the carbonaceous materials. (c) Schematic illustration for the generation of interphase "HO⁺ zone" on the surface of CNTs. Reprinted with permission from ref. 259. Copyright 2017, Elsevier.

is necessary to comprehensively evaluate the contribution of each ROS during pollutants degradation.

4.2.3 Molecular probes. Chemical probes can be employed for ROS identification via detecting the byproducts formed from the reactions between target ROS and chemical probes. In addition, the amount of ROS can be semi-quantified by measuring the degradation reaction constants of chemical probes and the concentration of the generated byproducts. Probes such as p-chlorobenzoic acid,243-245 benzoic acid,246,247 salicylic acid,248,249 N,N-dimethyl-p-nitrosoaniline,250 dimethyl sulfoxide,^{251,252} terephthalic acid^{253,254} and 1,5-diphenylcarbohydrazide255 can be employed for HO' detection. For example, HO' can convert the non-fluorescent salicylic acid to the fluorescent 2,3-dihydroxybenzoic acid.248,249 HO' will react with DMSO to generate formaldehyde (HCHO). The concentration of the produced HCHO could be measured by 2,4-dinitrophenylhydrazine (DNPH) derivatization method, in which hydrazone (HCHO-DNPH) is formed and its concentration can be facilely quantified by chromatography analysis.²⁵² Guo et al. employed *p*-chlorobenzoic acid (*p*-CBA) and tetrachloromethane (CCl_4) as the probes for HO' and O2'- in ozonation processes, respectively.244,245 Based on the measured ROS exposures, the evolution of HO' and O2' concentrations were simulated using chemical kinetic models.245 Moreover, nitro blue tetrazolium (NBT) is also efficient to estimate the concentration of O_2 . based upon the absorbance intensity of the corresponding byproduct at 560 nm.²⁵⁶ For ¹O₂ identification, diphenylanthracene (DPA) has been considered as the preferably chemical probe besides furfuryl alcohol (FFA). Formation of the characterized dibenzanthracene peroxides (DAPO₂) from DAP indicates the direct singlet oxygenation.²⁵⁷ Also, chemical probes can be utilized to locate the area where ROS would be generated. As a probe for HO', coumarin could be oxidized by HO' via hydroxylation, forming the stable compound 7-hydroxvcoumarin, which is photoluminescence with the UV characteristic absorption peak at 456 nm.258 Using fluorescence microscopy image (FMI), the region that HO' produced on CNTs was visualized (Fig. 13a-c).²⁵⁹ However, careful consideration is required when interpreting the results of chemical probes, because O₃ has a strong oxidation capability and might directly oxidize the added chemical probes during the reaction. Furthermore, the added probes might not be "chemically selective" to the target ROS. In persulfate-based AOPs, methyl phenyl sulfoxide (PMSO) is used as a chemical probe to distinct the high-valent metal species from HO' or SO₄.⁻, because PMSO can be oxidized by high-valent metal species to methyl phenyl sulfone (PMSO₂) via an oxygen-transfer reaction, while the reactions of PMSO with HO' and SO4⁻⁻ lead to the formation of hydroxylated products and biphenyl compounds, respectively.^{89,212} Inspired by this method, it is promising to seek for certain chemical probes that obtain different reactivities toward ROS to yield characteristic products, which can be differentiated by chromatography or spectrum analysis.

4.2.4 Intermediates-directed prediction of oxidation pathways. With different oxidation potentials and electronic properties, the generated ROS obtain quite dissimilar mechanisms for organics oxidation, giving rise to featured degradation

intermediates and pathways. Therefore, the contribution of a certain ROS can be determined. HO' with a high oxidation potential tends to attack the aromatic ring to generate polyhydroxylated products; this is usually the very first step during the oxidation.260,261 Additionally, HO' also favors the attack of phenolics by hydrogen abstraction and hydroxylation reactions, resulting in the formation of hydroquinone and catechol, which might be further converted to p-benzoquinone and o-benzoquinone as a result of the extraction of hydrogen from the O-H bond.^{262,263} Similarly, O2^{•-} also favors the hydrogen abstraction reactions but the process will produce carbon-based radicals rather than hydroquinones in O2. -- based oxidation. 264,265 Subsequently, the formed carbon-based radicals can further react with O₂ to generate peroxy intermediates.²⁶⁶ In a ¹O₂driven process, the unsaturated organic compounds generally suffer from electrophilic attack and electron abstraction.267,268 Meanwhile, ¹O₂ tends to attack phenolic compounds to form corresponding phenolic radicals, which further evolves into quinone-like and ring-opening products.²⁶⁹ So far, few studies have been focused on differentiating the reaction pathways and intermediates of specific pollutants degraded by different ROS in catalytic ozonation. Excavating the relationships between the characteristic intermediates and ROS can be an indirect way to verify the role(s) of potential ROS during ozonation.

4.3 Selectivity of the ROS toward organic pollutants

The free and surface-confined HO' can non-selectively destroy most of organic contaminants at a high reaction rate (10⁸ to 10¹⁰ M^{-1} s⁻¹) because of its strong oxidizing capability, while it might not be the dominant ROS responsible for the abatement of some certain electrophilic organic compounds such as CCl₄ $(k_{\rm HO} < 2 \times 10^6 \text{ M}^{-1} \text{ s}^{-1})$, perfluorooctanesulfonic acid (PFOS, $k_{\rm HO} < 10^4 \text{ M}^{-1} \text{ s}^{-1}$), and perfluorooctanoic acid (PFOA, $k_{\rm HO} <$ $10^4 \text{ M}^{-1} \text{ s}^{-1}$).^{270,271} O₂.⁻ with a mild oxidation potential of 0.89-1.7 V can oxidize less recalcitrant organics. Particularly, O_2 . exhibits higher reaction rates with some halogenated hydrocarbons such as CCl₄, PFOS, and PFOA ($k_{O_2} = 10^7 - 10^8 \text{ M}^{-1} \text{ s}^{-1}$) than HO' due to its strong nucleophilicity.244,245 Additionally, the halogenated hydrocarbons are recalcitrant for O_3 attack (k_{O_3}) $< 0.1 \text{ M}^{-1} \text{ s}^{-1}$), which have been used as the chemical probes for quantifying the concentrations of HO' and HO' /O2'- generated in catalytic ozonation systems.244,245 Nonradical-based oxidations in catalytic ozonation reactions also demonstrated a substrate-dependent behavior.96,142 The surface-confined nonradical species (including activated surface-O3 complex and $*O_{ad}$) with high oxidation potentials tend to attack aliphatic compounds with saturated carbon bonds rather than organics with unsaturated carbon bonds. Similar to O₂^{•-}, ¹O₂ obtains a moderate oxidation capacity ($E^0 = 0.81$ V, NHE), which exhibits high selectivity in the decomposition of organics with unsaturated carbon bonds and electron-rich functionalities (e.g., phenols, sulfides, and anilines) via electrophilic addition and electron abstraction, but not reactive to saturated alcohols (e.g., methanol, ethanol and tertiary butanol).272 Some studies revealed that O2^{•-} or ¹O2 played a predominant role in the degradation of phenolic compounds during the catalytic

ozonation processes, even though the existence of HO' was confirmed by EPR spectra and quenching tests.^{142,220,273} It is worth noting that, although studies have reported that O_2^{-} or 1O_2 were primarily responsible for the TOC abatement, O_2^{-} and 1O_2 with mild oxidation potentials are substrate-dependent and cannot completely mineralize saturated compounds.^{274–276} Therefore, these conclusions are questionable and require a careful reevaluation. In addition, it is also quite challenging to justify the contribution of $O_2^{-/1}O_2$ in the carbon-catalyzed ozonation system since the formation of $O_2^{-/1}O_2$ is often accompanied by other ROS.

5. Impacts of water matrix

5.1 Solution pH

Solution pH is a crucial factor in catalytic ozonation processes, affecting the surface properties of catalysts, the nature of organic contaminants, and the generation and transformation of ROS, thereby influencing the catalytic performances. The surface charge of the catalyst can be affected by solution pH. The catalyst surface will be negatively charged when the solution pH is greater than pH_{pzc} of the catalysts, which favors the electrophilic adsorption of O₃. In contrast, the interaction of O₃ with the positively charged surface is weak. Additionally, the charge and molecular states of organics are closely related to their pK_a values, which are also affected by solution pH. The states of organics will subsequently influence the electrostatic interactions with the surface of the catalysts and the affinities toward ROS. The opposite charge between the catalysts surface and the organics will induce electrostatic attraction and facilitate surface reactions, 277, 278 while the deprotonated organics are more likely to be electrophilically attacked by ROS.279 Additionally, the electrochemical parameters of organics, such as ionization potential, half-wave potential, and Hammett constant, will vary with solution pH and closely relate with the redox capacity of the oxidation system.^{23,159,280,281} For example, the half-wave potential of phenol declined with the increased pH, leading to a higher degradation rate. However, when solution pH was further elevated beyond the pK_a of phenol, the phenolate would form and demonstrate strong recalcitrance for degradation.²¹⁶ The variations in surface interactions and affinities toward ROS at different pHs might result in different reaction mechanisms. Altering solution pH might change the adsorption capacity of organics and alter the reaction pathway from a nonradical pathway to radical-dominated oxidation.¹¹²

Solution pH can also influence the conversion of reactive species and their redox potentials. O_3 molecules can be activated by alkaline solution into HO[•] ($k = 70 \text{ M}^{-1} \text{ s}^{-1}$) (eqn (2)).²⁸² However, deprotonation of HO[•] takes place at pH > 11.9, and the formed conjugate base radical (O^{•-}) is less reactive than HO[•] and thus reduces the treatment efficiency (eqn (8)).²²¹ For O₂^{•-}, it can be oxidized with HO₂[•] in acidic solutions to generate ¹O₂ (eqn (6)), while HO₂^{•-} will deprotonate into O₂^{•-} when solution pH is over 4.8, thereby changing from electrophilic attack to nucleophilic attack (eqn (4)).⁶ The reduction potentials of ROS are pH-dependent, because the concentrations of H⁺/OH⁻ affect the activity of the generated ROS and their conjugated reductive

pairs.²⁸³ Generally, an increased H⁺ concentration demonstrates an improved activity of ROS, resulting in an increase in reductive potential as the solution pH decreases.^{221,283}

$$\text{HO}^{\cdot} + \text{OH}^{-} \rightarrow \text{O}^{\cdot-} + \text{H}_2\text{O}, \, \text{p}K_a = 11.9$$
 (8)

5.2 Inorganic anions

Inorganic anions such as Cl⁻, CO₃²⁻, HCO₃⁻, and SO₄²⁻ ubiquitously exist in practical wastewater and natural water systems, which can potentially affect O3 activation even at low concentrations. Inorganic anions are commonly considered as radical scavengers to consume ROS and generate less reactive radicals (eqn (9)-(14)), resulting in the decrease in the overall degradation efficiency.272,284 The scavenging capacity can be affected by solution pH. For instance, Cl⁻ can scavenge HO' to produce Cl'. However, the scavenging effect was only effective within an acidic pH range and was limited under neutral conditions, because the intermediate of the radical chain reaction (ClOH^{•-}) would rapidly revert to HO' rather than forming Cl' (eqn (9) and (10)).²⁸⁵ A similar effect was observed when HCO₃⁻ was involved. The elevated solution pH would deprotonate HCO₃⁻ into CO₃²⁻, resulting in an increased scavenging effect because of the higher reaction rate of HO[•] with CO_3^{2-} (3.9 × 10⁸ M⁻¹ s⁻¹) than that with HCO_3^- (8.5 \times 10⁶ M⁻¹ s⁻¹) (eqn (11) and (12)).^{232,286} CO_3^{2-} also increases the stability of O_3 by removing the dominant chain carrier radicals, such as O3, , 287 while SO_4^{2-} and NO_3^{-} showed negligible effects on O_3 decomposition.²⁸⁸ The presence of inorganic anions might also result in the improved selectivity of oxidative systems, because the generated reactive halogen species or CO3. are more selective than HO' and favor the attack of electron-rich organics.2,289

Cl⁻ + HO[•] ↔ ClOH^{•-},
$$k = 4.3 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$$
 (9)

ClOH^{•−} + H⁺
$$\leftrightarrow$$
 Cl[•] + H₂O, $k = 2.1 \times 10^{10} \text{ M}^{-1} \text{ s}^{-1}$ (10)

$$\text{CO}_3^{2-} + \text{HO}^{\bullet} \to \text{CO}_3^{\bullet-} + \text{OH}^-, k = 3.9 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$$
 (11)

$$\text{HCO}_3^- + \text{HO}^\bullet \to \text{CO}_3^{\bullet-} + \text{H}_2\text{O}, k = 8.5 \times 10^6 \text{ M}^{-1} \text{ s}^{-1}$$
 (12)

$$\text{HCO}_{3}^{-} + \text{O}_{2}^{\cdot -} \rightarrow \text{CO}_{3}^{\cdot -} + \text{HO}_{2}^{-}, k = 1.5 \times 10^{6} \text{ M}^{-1} \text{ s}^{-1}$$
 (13)

$$PO_4^{3-} + HO^{\bullet} \rightarrow PO_4^{\bullet 2-} + OH^{-}, k = 1.1 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$$
 (14)

The effect of inorganic anions on catalytic ozonation processes also depends on their concentrations. It was reported that the Cl⁻ concentration can influence the scavenging products and thus affect the catalytic ozonation efficiency.^{35,159} At low concentrations, Cl⁻ would quench HO[•] ($E^0 = 2.7$ V, NHE) to form the less reactive Cl[•] ($E^0 = 2.4$ V, NHE). In contrast, a high concentration of Cl⁻ favored the transformation of Cl⁻ into reactive chlorine species (Cl₂ and HClO), which could couple with HO[•] for organics degradation. However, opposite conclusions were derived in another study.²⁹⁰ The discrepancies in these studies might be explained by the combined effects of ionic strength, which influences the solubility and mass transfer of O₃, the reaction rates of ROS, and the interactions of O₃ and organics on catalyst surface. It was reported that the solubility and mass transfer of O₃ decrease as ionic strength grows, especially when Cl^- , CO_3^{2-} and PO_4^{3-} are present, while SO_4^{2-} gives rise to an insignificant effect.²⁹¹ Moreover, positive effects of ionic strength on reaction rates were observed in various reaction systems.^{292–294} Studies suggested that the reaction rates between oxidants and organics raised with the increase in ionic strength.²⁹⁵ The ionic strength created by different ions might also influence reaction rates. The reaction rate between O3 and Cl⁻ improved with the increase in the ionic strength caused by NaClO₄ or KNO₃, whereas simply increasing the concentration of Cl- resulted in the decrease of the corresponding reaction rate.296 Additionally, increasing the ionic strength reduces the zeta potential of catalyst surface via compressing the thickness of the electric double layer and therefore affects the weak interactions (via electrostatic bonding) of O3 and organics on catalyst surface, such as the activated surface-O3 complex.297 However, marginal influence occurs to strong interactions, *i.e.*, the *O_{ad}-induced oxidation model. Therefore, variations in ionic strength can be utilized to distinguish the different interaction models of O₃ on the catalyst surface. Ionic strength also affects the adsorption capacity of the organics on catalysts. As ionic strength increases, the adsorption capacity decreases when the electrostatic forces between the catalyst surface and organics are attractive.298

Inorganic anions may compete on the surface-active sites of catalysts with O₃ or organics, thus inhibiting the oxidation performance. The positively charged catalysts can provide suitable adsorption sites for inorganic anions. PO₄³⁻ as a strong Lewis base shows the strong inhibitory effect by occupying surface Lewis acid sites and impeding O3 activation.299,300 Notably, the induced alkalinity with the presence of CO_3^{2-} and PO_4^{3-} as basic anions would promote the base activation of O_3 and affect the surface charge of the catalyst.287,290 Therefore, special attention should be paid to the variation in solution pH with the presence of basic anions. In the most of current studies, individual effects of inorganic anions on the activation of O₃ have been well evaluated. Nevertheless, considering the complexity in compositions of the practical wastewater, the synergistic effects of various inorganic anions cannot be overlooked.

5.3 Natural organic matters

NOMs are pervasive constituents in natural water, and are known to decrease the efficiency of AOPs-based treatments by competing with organic contaminants for HO[•] consumption $(1.6-3.3 \times 10^8 \text{ M}^{-1} \text{ s}^{-1})$.³⁰¹ NOMs with electron-rich sites could also deplete the dissolved O₃ *via* direct O₃ attack, thus lowering the utilization efficiency of O₃.³⁰² Furthermore, NOMs containing carboxyl and phenolic hydroxyl groups would adsorb on the surface of carbonaceous materials and block the active sites, giving rise to inhibited interactions between the carbocatalysts and O₃/organics.³⁰³ For HO[•]-dominant oxidation pathway in catalytic ozonation, the inhibitory effects would continue until most of NOMs are mineralized because of the non-selectivity of HO[•].³⁰⁴ The nonradical pathways based on the surface oxidations are highly selective to the aliphatic acids with small molecular weight; therefore, NOMs may have a less impact on nonradical-based oxidation.¹⁵⁹ Nevertheless, debates on the effect of NOMs on catalytic ozonation still exist. Some studies reported that NOMs could act as initiators and promotors for radical chain reactions of O_3 decomposition to HO', which enhanced the degradation rate.^{305,306} Specific moieties, such as aromatic and aliphatic unsaturated constituents in NOMs, could directly activate O_3 and accelerate the radical chain reactions to generate more HO'.³⁰⁷ Moreover, the presence of humic acid (HA), a major constituent of NOMs, could introduce oxygen functional groups on Fe₃O₄/MWCNTs and thus accelerate HO' production.¹⁷³

6. Conclusions and future perspectives

In summary, this review presents the current application of carbon-based catalysts for O₃ activation and degradation of organic pollutants. The engineering principles for regulating the active sites and electronic structures of carbocatalysts have been discussed, which will provide directions and general guidelines for developing high-performance carbocatalysts. This review also puts forward new insights into ROS generation and the reaction pathways based on the interaction intensity model between O3 and catalyst surface. In addition, the effect of the water matrix factors, including pH, inorganic anions and NOMs, on catalytic ozonation processes are highlighted. Based on our review, future directions to material design, ROS identification and mechanism regulation aiming at enhancing the application potential of carbon-based catalytic ozonation in real wastewaters are proposed as follows:

(1) Encapsulating metal species beneath graphitic carbons (metal@C) has been proven as an effective strategy to increase both the reactivity and stability of the composites. Additionally, the magneticity endowed by the encapsulated metal species facilitates the separation of the nano-scaled catalysts. Apart from the dimensional effects, surface properties of the graphitic carbons and the types of the encapsulated metals are critical factors governing the electron-transfer ability and the catalytic activity of the nanocomposites. Therefore, it is promising to design metal@C catalysts with various numbers of coated graphitic shells, defective levels, and heteroatom dopants and manage the types and exposed crystal facets of the encapsulated metal species. The electron hybridizing, distribution, and transfer process at the metal-carbon heterojunctions can be evaluated by the DFT simulations which guide the theoretical design and screening the metal@C catalysts.

(2) In the current stage, intensive research attentions have been paid to identifying the types of ROS and elucidating ROS evolution mechanisms. However, the interactions of reactants on the active sites of the carbocatalysts are normally transient and thus hinder the unravelling of the in-depth insights. Future mechanistic studies require combined theoretical and experimental methodologies. The development of *in situ*, cryo-, or time-resolved characterization techniques with the isotopic substitutions will help unravel the interactions of the O_3 and/or ambient molecules on catalyst surface and differentiate the types of generated ROS. Additionally, evaluating the existence and effects of other ROS apart from HO' in catalytic ozonation is challenging because of the high oxidation capacity and non-selectivity of HO'. Therefore, it is crucial to develop probes or sensors that selectively react with targeted ROS, especially those with mild oxidation potentials.

(3) Water matrix parameters such as pH, inorganic anions and NOMs can significantly affect the interactions of O₃ and organics on the catalyst surface and the reactivities of the generated ROS. HO'-based radical pathway demonstrates a highly oxidative and nonselective behavior, yet the treatment efficiency would be decreased in the presence of inorganics anions and NOMs as radical scavengers. Nonradical pathways arising from the weak interactions between O₃ and catalyst surface is vulnerable to be influenced by solution pH and ionic strength. Nonradical oxidation induced by strong interactions are typically recalcitrant toward the ionic strength and NOMs, while solution pH might influence the catalytic performance by affecting the surface charges. Therefore, developing reactionoriented carbonaceous materials that can precisely regulate the reaction pathways based on the characters of different types of wastewater is highly desirable. DFT calculations of the interactions and electron properties at the atomic level can be employed to guide the design of on-demand active sites on carbocatalysts to trigger specific reaction pathways for different application scenarios.

Conflicts of interest

There are no conflicts to declare.

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