Unprecedented Nonphotomediated Hole (h\textsuperscript{+}) Oxidation System Constructed from Defective Carbon Nanotubes and Superoxides

Junhui Wang, Jiaxing Yu, Qi Fu, Huangsheng Yang, Qing Tong, Zhengping Hao, and Gangfeng Ouyang*

ABSTRACT: Holes (h\textsuperscript{+}) on heterogeneous photocatalysts could act as important oxidative species or precursors for reactive oxygen species (ROS). However, due to the ultrafast recombination of photoinduced electrons and holes, a majority of carriers are consumed prior to surface reactions. Herein, we report an unprecedented non-photomediated hole oxidation system constructed from carbon nanotubes (CNTs) and superoxides. This system exhibited high catalytic activity for the degradation of organic pollutants, which outperforms the classical oxidation processes in the remediation of actual wastewater and is comparable to that of the best single cobalt atom catalyst. Theoretical and experimental results reveal that the intrinsic defects with unpaired spins on CNTs served as adsorptive sites to activate superoxides. This is the first report on exploring the oxidation properties of nonphotomediated hole carriers on heterogeneous catalysts, which will be of broad interest for researchers in environmental remediation, chemical synthesis, and biological fields.

INTRODUCTION

Photoinduced holes (h\textsuperscript{+}) generated on heterogeneous catalysts could serve as critical active oxidative species in splendid applications, including green energy exploration, chemical synthesis, and environmental technology.

As the core of photocatalytic activities, the electron–hole separation, transfer, and recombination often take place in an ultrasmall time scale from tens of femtoseconds to hundreds of picoseconds (Scheme 1a),

which result in the low utilization of photoexcited electrons and holes. For decades, tremendous efforts have been made to increase the separation efficiency of photoexcited electron–hole pairs, via facet/defect engineering or heterojunction construction on the catalysts.

Even so, still a majority of carriers are consumed by charge recombination prior to surface reactions.

Considering this, we wonder whether it is possible, through other facile strategies, to trap the electrons and create long-lasting hole carriers with high oxidative potential on the surface of catalysts. Actually, ascribed to the superior electronic conductivity of carbon nanotubes (CNTs), the chemical p-doping by electron-acceptors (e.g., O\textsubscript{2}, Br\textsubscript{2}) could create hole carriers on CNTs, due to electron transfer from the carbon π states to dopant molecules.

Most importantly, the hole carriers could exist for at least 5 min during the in situ Raman measurements.

This feature has offered CNTs exciting opportunities in various electronic/chemical applications. However, the oxidative properties of these hole carriers have never been explored.

Inspired by this observation, we presume that the adsorption of oxidants with strong electron-accepting capability on CNTs might fabricate holes with high oxidative capacity and long lifetime (Scheme 1b).

To increase the adsorption ability of oxidants on CNTs and electron transfer efficiency, more active adsorptive sites should be created. Previous theoretical calculations indicated that the enhancement of the charge or/and spin density of carbon materials could promote catalytic performance due to the increased adsorption affinity of oxidant molecules.

Recently, the intrinsic defect engineering has been regarded as an effective strategy, including the introduction of zigzag/armchair edges, vacancies, or topological defects (e.g., pentagons, heptagons, and Stone–Wales).

The defect sites can significantly tailor the surface electronic structures and act as charge-carrier acceptors, which might facilitate oxidant activation by the transfer of the electrons to the adsorbed oxidant molecules.

In this work, for the first time, we construct a non-photomediated hole oxidation system via a facile chemical strategy, utilizing defective CNTs and strong electron-accepting superoxides. Persulfate (PS) with a high redox potential (E\textsubscript{0}(S\textsubscript{2}O\textsubscript{8}\textsuperscript{2−}/SO\textsubscript{4}\textsuperscript{2−}) = 1.96 V\textsubscript{NHE}) was chosen as the model superoxide in this study. This system exhibited striking catalytic performances for the degradation of organic pollutants (e.g., bisphenol A, BPA), comparable to that of the reported single cobalt atom catalyst. Theoretical and
Experimental results verified that the intrinsic defects possess better adsorption affinities for PS and serve as the active sites. Raman, solid UV−vis, electron paramagnetic resonance (EPR), and scavenger experiments confirmed the generation of holes induced by the adsorbed PS on the defect sites and that the holes act as the critical active species for the rapid and selective degradation of BPA in actual wastewater. Intriguingly, the oxidative potentials of the holes could be tuned by proper defect construction on CNTs.

### RESULTS AND DISCUSSION

**Catalytic Performance of the Defective CNT/PS System.** To explore the possible effects of intrinsic defects on PS adsorption and activation, a series of CNTs with different levels of defect exposure were prepared via a facile acid etching and thermal reduction strategy as illustrated in Figure 1. This strategy could regulate the defect contents of CNTs along with maintaining other structural properties almost unchanged. In the acid etching procedure, as shown in TEM images (Figure S1) and reported in ref27, strong acids could etch the graphene’s surface, leading to a higher concentration of edges exposed, and covalently add to its edges and basal planes. And in the thermal reduction procedure, nearly all oxygen/nitrogen-containing functionalities can be desorbed from the CNT lattice by 1000 °C, leading to the formation of zigzag/armchair edges, single vacancy, and pentagon defects.18,27 Moreover, in CNTs, the Stone−Wales (5775) defect is presumed to be predominant with a relatively low formation energy of 3.5 eV.27 Therefore, combining the experimental procedure with the intrinsic properties of CNTs, we believe that in the defective CNTs prepared, there mainly existed zigzag edge, armchair edge, pentagon, Stone−Wales (5775), and single vacancy.

In an attempt to further clarify the effect of intrinsic defects on PS adsorption and activation, density functional theory (DFT) calculations were carried out on six CNT theoretical models (Figure S2). Figure 2 shows the local adsorption configurations and adsorption free energies of PS on pristine CNT surface and different kinds of defects on CNT. C gray, O red, S yellow.

Figure 1. Preparation route and model of defective carbon nanotubes, scale bars: 20 nm.

Figure 2. DFT calculations of adsorption affinities of PS on defective CNTs. Local adsorption configurations and adsorption free energies of PS on pristine CNT surface and different kinds of defects on CNT. C gray, O red, S yellow.
the pristine surface and consequently can serve as potential active adsorption sites for PS.

Limited by the available characterization techniques, quantification of different defects in defective carbon materials is still a great challenge. Therefore, in this study, all the defects in the defective CNTs prepared were treated as a whole and characterized by Raman spectroscopy which is an effective tool to characterize the defects in CNTs. As shown in the Raman spectra (Figure 3a), defect levels (the ratio \( I_D/I_G \) (D, defect; G, ground)) increased gradually with the acid treatment time. Nitrogen sorption analysis and X-ray photoelectron spectroscopy (XPS) characterization indicated no significant variation of other physical properties (textural structures and heteroatom contents) among the CNTs (Figures S3 and S4 and Table S1). As expected, the catalytic activity for BPA degradation shown in Figure 3b was remarkably enhanced by constructing defects on the CNTs. For instance, in the ROCNT-6/PS system, only 5 min was required for nearly total degradation of BPA. The BPA degradation intermediates were also analyzed and shown in Figure S5. To give a quantitative comparison of the catalytic activities, the BPA degradation kinetics were fitted by the pseudo-first-order model (Figure S6) and the obtained apparent rate constants (k) were listed in Table S1. Further, we correlated the rate constants \( k_{\text{BPA}} \) to \( I_D/I_G \) and found that \( k_{\text{BPA}} \) presents a positive linear relationship with \( I_D/I_G \) and the correlation coefficient \( R^2 \) is as high as 0.97 (Figure 3c). This result indicates that the defects make a crucial contribution to the catalytic activities of CNTs for BPA oxidation. Further investigations were subjected to the catalytic consumptions of PS by the CNTs (Figure S7 and Table S1). Quite similar to the observations above, the PS consumption rate constants \( k_{\text{PS}} \) increased with \( I_D/I_G \) and the calculated \( R^2 \) reached 0.96, suggesting that it is the defects that act as the active sites for PS adsorption and activation. Furthermore, XPS analyses excluded the possibility of heteroatoms (nitrogen dopants or oxygen-containing functional groups) as the catalytic active sites (Figures S8 and S9). In addition, it is worthwhile to mention that the optimal activity of ROCNT-6, with a \( k \)-value of 2.72 \( \mu \text{mol/g/s} \) (fitted by a modified kinetic model), is higher than the majority of photo/Fenton-like catalysts recently reported (Table S2) and comparable to that of the single cobalt atom catalyst (3.65 \( \mu \text{mol/g/s} \)).

Intrinsically, where do the catalytic activities of defects come from? Generally, carbon atoms with larger spin densities are most likely to serve as the active sites. To prove this assumption, we performed EPR measurements for different CNT catalysts. As shown in Figure 3d, the EPR spectrum of G-CNT has the narrowest line width (24 G), and those of ROCNTs become broader with the increment of defect levels. The signals are not observable for ROCNT-5 and ROCNT-6, due to the broadening of the signals. The \( \pi \)-electron radical at the defects of CNTs is delocalized to some extent and exhibits fast spin–lattice relaxation through interaction with the adjacent \( \pi \)-electron system, leading to a broad line width in EPR. We also found that the line width \( \Delta H \) showed a positive correlation with \( I_D/I_G \) (\( R^2 = 0.98 \)) (Figure 3e). To further confirm whether the spin states are important in
catalysis, OCNT-6 (before the thermal reduction treatment)
with intensive carboxyl and hydroxyl groups attached on
the defects was measured. Indeed, the EPR peak is quite narrow
(6.1 G, Figure 3f), and this is associated with its poor catalytic
activity (2.5% BPA removal in 5 min) compared with
ROCNT-6 (93% BPA removal in 5 min) (Figure S10).
Furthermore, higher spin densities were proved to locate at the
zigzag edge by DFT calculation (Figure 3g). Clearly, both the
experimental and theoretical results reveal the vital role of
unpaired spins at the intrinsic defects in the high catalytic
activities of defective CNTs.

Mechanism Investigation. The interaction between the
active sites on CNTs and PS molecules was further investigated
by Raman, solid UV−vis, and XPS analyses. As shown in
Figure 4a, after the adsorption of PS, the characteristic D and G bands at 1345 and 1577 cm⁻¹ of ROCNT-6 upshifted by 3 cm⁻¹, indicating that PS serves as an electron acceptor attracting electrons from the carbon π states, consequently creating hole carriers on ROCNT-6.¹² Such electronic interaction was also identified by solid UV−vis spectroscopy (Figure 4b). The peak at 210 nm represents the π−π* transition of aromatic C−C bonds on ROCNT-6. While after PS adsorption, the absorbance peak downshifted to 206 nm, implying the decrease of electron density on CNT, due to the electron-withdrawing capability of PS.¹² XPS analyses further confirmed that ROCNT-6 could be oxidized by PS if treated for a long time, with oxygen content increased (Figure S11). These results clearly prove the assumption that PS adsorption could cause hole injection into CNTs.

Based on the above observations, quenching experiments
were further conducted to identify the active species generated
in the CNT/PS system. The commonly used hole scavengers
in photocatalysis, triethanolamine (TEA) and ethylenediamine
tetraacetic acid disodium (EDTA-2Na), were applied. Not
surprisingly, as shown in Figure 4c, TEA and EDTA-2Na both
inhibited the BPA degradation effectively, and an increasing
amount of EDTA-2Na could induce a higher inhibition effect.
Radical quenching experiments and EPR spin-trapping
measurements revealed the absence of radicals (SO₄⁻ or
•OH) and singlet oxygen (¹O₂) during the reaction, which
are common active species in traditional oxidation systems,
such as photo/Fenton-like reactions (Figure S12). Further-
more, catalytic activities of several commercial MWCNTs were
investigated. We observed a positive linear correlation between
k₁ and k₂PS (R² = 0.91) rather than the adsorption amount of
BPA on CNTs (R² = 0.43) (Figure 4d), indicating that the
generation of holes induced by PS is the prerequisite for BPA
oxidation. If more electrons are extracted by PS, more holes
will be generated on CNTs, and higher BPA removal e-
fi
fficiency will eventually be achieved. All of the above observations
con
fi
rm that holes induced by PS on CNTs were the active
species for BPA oxidation.

The reaction process could be further uncovered from the
PS consumption under different conditions as illustrated in
Figure S13. With only catalyst, the concentration of PS was
reduced by 33% in 20 min. In contrast, the presence of
electron donors (BPA or EDTA-2Na) led to noticeable
decomposition of PS, and the same amount of EDTA-2Na
could induce a higher PS consumption rate than BPA, mainly
due to its higher electron-donating capability. These results implied that the electrons from BPA or EDTA-2Na injected to the holes on CNTs could then be transferred to PS leading to its rapid decomposition.

The reusability of ROCNT-6 was also investigated as shown in Figure S14. The used catalyst exhibited reduced activity, and after thermal treatment the catalytic reactivity could be totally recovered. Deactivation was mainly due to the side reaction—oxidation of ROCNT by PS—and the oxygen functionalities could be removed by thermal treatment resulting in the recovery of catalytic activity.

Based on the previous discussions and results, a possible catalytic pathway in the CNT/PS system for BPA degradation is proposed (Figure 5). PS is first adsorbed at the defect sites of CNTs (State I) and then extracts unpaired electrons followed by the generation of holes (State II). In the meantime, preadsorbed BPA molecules inject electrons into the holes and are oxidized simultaneously. However, if hole scavengers with higher reducibility also adsorb on the CNTs, they are prone to transfer electrons in prior to the BPA molecules, leading to the decrease of BPA degradation rate. Meanwhile, at the defect site, PS is decomposed into two SO$_4^{2-}$ ions, and CNT was oxidized resulting in its deactivation (State III). The active sites could be regenerated by removing the oxygen functional groups through thermal treatment in an inert atmosphere.

Figure 5. Proposed mechanism of the CNT/PS system. Schematic pathway of defective CNT/PS system for BPA degradation.

Figure 6. Redox potential investigation of the CNT/PS system. (a) Schematic illustration of the energy diagram for CNTs with PS adsorption on different defect sites obtained from DFT calculations. (b) Catalytic degradation of BPA in the actual water body (Pearl River in China). Reaction conditions: [BPA] = 20 mg/L, [PS] = [PMS] = 1.5 mmol/L, [ROCNT-6] = [TiO$_2$] = [Fe$^{2+}$] = 0.1 g/L, T = 298 K.
The potentials of the holes generated on CNTs. For the pristine CNT, the work function is determined as 4.76 eV, in accordance with those reported in the literature. When PS adsorbed on the defects of CNT, the work functions increased to 5.66–6.14 eV (shown in Figure 6a), further proving the charge transfer from CNT to PS. The oxidation potentials of the PS-induced holes reflected by work functions are in the range of 1.12 to 1.74 V, indicating that the CNT/PS system possesses moderate oxidation capability. Moreover, from the theoretical calculations, it could be found that the holes generated by PS adsorbed on vacancy and Stone–Wales (5775) defects have more positive oxidation potentials than those on pentagon and zigzag/armchair edges, which indicates that vacancy and Stone–Wales (5775) defects might have better catalytic performances, and the oxidation potentials of this system could be tuned by proper defect construction. The moderate oxidation capability was further confirmed experimentally by the selective degradation of different organic compounds exhibited by the ROCNT-6/PS system (Figure S16). And in the actual water body (detailed water quality parameters shown in Table S3), the catalytic performance of ROCNT-6/PS was scarcely depressed (100% of BPA removal).
in 10 min) (Figure 6b). In comparison, only limited degradation of BPA was obtained by the free radicals produced in the classical Fe$^{3+}$/peroxymonsulfate (PMS) and TiO$_2$/UV systems. These exciting results highlight the great efficiency of this novel selective oxidation system for the elimination of target toxic organic contaminants in actual wastewater.

**Device Fabrication for Water Treatment in a Continuous Flow System.** Due to the rapid degradation of organic compounds in the ROCNT-6/PS system, we further tried to explore its practical application. To avoid the secondary pollution and difficulty of reusing the highly dispersed catalyst suspensions, CNT membranes were fabricated in situ simply via filtration of ROCNT-6/water dispersions on a PVDF substrate under gas pressure in a membrane reactor (Figure 7a–b). The SEM images in Figure 7c–e show the random stacking of carbon nanotubes in the membrane with a thickness of about 120 μm and pore size of 200–800 nm. The abundant nanochannels not only provide confined voids for efficient catalytic reaction but also allow fast permeation of the treated water.

After the in situ fabrication of CNT membrane, water mixed with BPA and PS was directly added into the membrane reactor, and the following continuous permeance experiments were carried out (Figure 7a, f–g). The removal of BPA in this system maintained 100% over 10 h at a high average permeance of 2850 L/m$^2$/h/bar (Figure 7h), which is much superior than the best reported MoS$_2$ membrane (90% BPA removal in 6 h and permeance of 154 L/m$^2$/h/bar).$^{30}$ Waste water containing only BPA was also applied, and leakage of BPA after only 30 min could be observed, implying that the adsorption of BPA by CNTs made negligible contribution to the high removal efficiency. Moreover, the membrane could also maintain its excellent catalytic performance in actual water. We believe that the high catalytic efficiency and nonleakage of any toxic metal ions render this CNT-membrane/PS configuration an ideal oxidation system to selectively remove low-concentration of organic pollutants in actual contaminated water in a continuous flow mode.

In summary, by utilizing defective carbon nanotubes and persulfate, we have fabricated a novel hole oxidation system under dark conditions. This system exhibited striking catalytic performance for the degradation of BPA, and theoretical and experimental results revealed that the intrinsic defects with unpair spins were the active sites for the adsorption and activation of PS. Holes induced by PS at the defects of CNTs were proved to act as the critical active species with moderate and tunable oxidation potentials, rendering the CNT/PS system to have highly efficient and selective elimination toward target pollutants in the actual water body, which could outperform the classical oxidation processes. Furthermore, the optimal CNT catalyst could be fabricated in situ into membranes for water treatment in continuous flow. This is the first time to explore the oxidation properties of nonphotomediatesd hole carriers on heterogeneous catalyst, and the proposed novel oxidation process could offer interesting possibilities for the nonradical selective oxidation in water treatment, chemical synthesis, and biologic applications.

### Experimental Procedures

**Chemicals and Materials.** The following reagents were used as received: G-MWCNT (XF NANO, outer diameter: 10–20 nm, length: 5–30 μm), MWCNT (Aladdin, 95%, inner diameter: 3–5 μm), NH$_2$-MWCNT (Aladdin, 95%, inner diameter: 3–5 μm, outer diameter: 8–15 nm, length: 50 μm), OH-MWCNT (J&K, 95%, inner diameter: 5–10 nm, outer diameter: 10–20 nm, length: 10–30 μm), COOH-MWCNT (J&K, 95%, inner diameter: 3–5 nm, outer diameter: 8–15 μm, length: 50 μm), CON-MWCNT (J&K, 95%, inner diameter: 3–5 nm, outer diameter: 8–15 μm, length: 50 μm), potassium peroxysulfate (Innochem), bisphenol A (Sigma-Aldrich), carbamazepine (Sigma-Aldrich), benzoic acid (Sigma-Aldrich), phenol (Sigma-Aldrich), 4-chlorophenol (Sigma-Aldrich), methanol (Sigma-Aldrich), potassium iodide (Innochem). Ultrapure deionized water (>18 MΩ·cm), produced with a Millipore system, was used for the preparation of all experimental solutions. All chemicals were of reagent grade and were used without further purification or treatment.

**Preparation of MWCNTs with Variable Defect Contents.** 0.5 g G-CNT was added into a round-bottom flask, followed by slow addition of 120 mL of 68% HNO$_3$ or 68% HNO$_3$–98% H$_2$SO$_4$ (v/v = 1:3). Then the mixture was magnetically stirred for different periods at 120 °C with refluxing. After cooling down to room temperature, the final mixture was dropped slowly into 500 mL of deionized water under stirring. Then the mixture was filtered and washed with deionized water until the pH reached ~7 and was centrifuged at 11000 rpm for 10 min to obtain the solid. Finally, the obtained solid was dried at 120 °C for 24 h and annealed at 1000 °C for 3 h with a heating rate of 3 °C/min under argon atmosphere. The samples treated by HNO$_3$ for 3 h, 6 h, and 15 h were denoted as ROCNT-1, ROCNT-2, and ROCNT-3, and those treated by couple mixing of HNO$_3$ and H$_2$SO$_4$ for 0.5 h, 1 h, and 1.5 h were denoted as ROCNT-4, ROCNT-5, and ROCNT-6, respectively.

**In Situ Fabrication of ROCNT-6 Membrane.** 50 mg ROCNT-6 was dispersed in 10 mL Milli-Q water and sonicated for 0.5 h. The ROCNT-6 membrane was then prepared by directly filtering the resultant CNT solution on a PVDF substrate (Millipore) with a pore size of 0.22 μm in the membrane reactor with gas pressure of 0.2 bar. The membrane could be obtained in 30 min.

**Characterization.** Raman spectra were recorded on a Renishaw InVia spectrometer with a model 100 Ramascope optical fiber instrument. X-ray photoelectron spectroscopy (XPS) analysis was conducted on an ESCALAB 250 spectrometer (Thermo Fisher Scientific Corporation, USA) with Al Kα radiation as the exciting source (150 W). The binding energies of the recorded XPS spectra were corrected according to the C 1 s line at 284.8 eV. After subtracting the Shirley-type background, the core-level spectra were deconvoluted into their components with mixed Gaussian–Lorentzian (20:80) shape lines using the CasaXPS software.

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Catalytic Activity Measurements. The entire experiments were performed in 50 mL glass reactors at room temperature. The pH value of the reaction system was not further adjusted and to avoid any misleading conclusions, buffers were not used. The reaction solution was prepared by suspending CNTs (100 mg/L) in the solution of the target organic compound (0.09 mM for all the pollutants). After reaching the adsorption equilibrium, the reaction was initiated by adding an aliquot of a prepared solution of PS (150 mM). Samples were withdrawn using a 1 mL syringe at certain intervals during the 20 min reaction, and the solid was removed by filtration (0.22 μm PTFE filter).

The concentration of PS was measured by a spectrophotometric method using potassium iodide, which was modified from the method suggested by Liang et al.\textsuperscript{30} 0.8 mL sample solution was withdrawn and added into a prepared 3 mL KI water solution (containing 0.015 g of NaHCO₃ and 0.3 g of KI). The resulting solutions were hand shaken and allowed to equilibrate for 15 min for spectrophotometric measurement at 352 nm.

The concentration of the target organic compounds (i.e., bisphenol A, carbamazepine, phenol, benzoic acid, 4-chlorophenol) was analyzed using a HPLC (Shimadzu LC-20AD). Separation was performed on a Poroshell 120 EC-C18 column (4.6 × 100 mm, 2.7 μm, Agilent Technology, USA) using a mobile phase consisting of a binary mixture of water and methanol at a flow rate of 0.6 mL/min. The degradation intermediates of BPA were identified by direct immersion SPME (solid-phase microextraction) coupled with GC-MS (Agilent 6890-5975 B) equipped with a HP-5 capillary column, using a PA (polyamide) commercial SPEM fiber.

Density Functional Theory (DFT) Calculations. All the calculations are performed in the framework of the density functional theory with the projector augmented plane-wave method, as implemented in the Vienna ab initio simulation package.\textsuperscript{40} The generalized gradient approximation proposed by Perdew, Burke, and Ernzerhof is selected for the exchange-correlation potential.\textsuperscript{41} The cutoff energy for the plane wave is set to 400 eV. The energy criterion is set to 10⁻⁵ eV in the iterative solution of the Kohn–Sham equation. The Brillouin zone integration is performed at the Gamma point. All the structures are relaxed until the residual forces on the atoms have declined to less than 0.05 eV/Å. In this work, single wall carbon nanotubes (SWCNTs) are taken as an example to consider the catalytic effect for persulfate adsorption and activation as shown in Figure S1. The simulated carbon nanotube is with a size of a × b × c = 10 × 10 × 9.838 Å and built in a cubic simulation cell with lattice content of 30 Å. The vacuum layer is more than 15 Å along three directions to avoid unphysical interaction between the nanotube and the periodic image because of the periodic boundary condition. To investigate the adsorption affinities of PS molecules on different sites of SWCNTs, the adsorption energy is studied by DFT calculations, defined as

\[ E_{\text{ads}} = E_{\text{total}} - E_{\text{substrate}} - E_{\text{molecule}} \]

where \( E_{\text{total}} \), \( E_{\text{substrate}} \), and \( E_{\text{molecule}} \) denote the total energy of substrate with adsorbate, substrate, and free molecule, respectively.

In all experiments performed, no unexpected or unusually high safety hazards were encountered.

## ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscentsci.0c01600.

- HRTEM images of G-CNT, ROCNT-3, and ROCNT-6; favorite adsorption configuration of PS on pristine surface and different kinds of defects on CNT; N₂ adsorption–desorption isotherms of CNT catalysts; XPS spectra of CNT catalysts; GC-MS spectra of the BPA degradation intermediates by the ROCNT-6/PS system; kinetic analysis of BPA degradation on different CNT catalysts; PS consumption on different CNTs and the kinetic analysis of PS consumption according to the pseudo-first-order model; high-resolution XPS O 1s and N 1s spectra of the CNT catalysts; contents of different O and N species on the CNT catalysts analyzed from XPS spectra; catalytic degradation of BPA by OCNT-6; XPS spectra and contents of O species of ROCNT-6 and ROCNT-6; variation of \( \text{SO}_4^{2-} \) and \( \text{SO}_3^{2-} \) ions during the reaction of ROCNT-6/ROCNT-3 with PS solution for 1 h; pH variation during the reaction process under different conditions; EPR spectra obtained by spin trapping with DMPO and TEMP in the presence of PS and ROCNT-6; degradation of BPA by the ROCNT-6/PS system with and without methanol; PS consumption under different conditions; reusability investigation of ROCNT-6; XPS 1Os spectra of used catalyst and regenerated catalyst after 900 °C treatment; catalytic degradation of BPA on ROCNT-6 with other different superoxides adsorbed; catalytic performance of ROCNT-6/PS system for treating different pollutants; physicochemical parameters of the CNTs; comparison of recently reported photo/Fenton-like catalysts in BPA removal; and water quality parameters of Pearl River.

## AUTHOR INFORMATION

### Corresponding Author

**Gangfeng Ouyang**—MOE Key Laboratory of Bioinorganic and Synthetic Chemistry, School of Chemistry, Sun Yat-Sen University, Guangzhou, Guangdong 510275, China; Chemistry College, Center of Advanced Analysis and Gene Sequencing, Zhengzhou University, Zhengzhou 450001, China; orcid.org/0000-0002-0797-6036; Phone: +86 020 84110845; Email: cesoygf@mail.sysu.edu.cn; Fax: +86 020 84110845/0953

### Authors

**Junhui Wang**—MOE Key Laboratory of Bioinorganic and Synthetic Chemistry, School of Chemistry, Sun Yat-Sen University, Guangzhou, Guangdong 510275, China; orcid.org/0000-0001-9022-7417

**Jiaxing Yu**—School of Materials Science and Engineering, Sun Yat-Sen University, Guangzhou, Guangdong 510275, China

**Qi Fu**—MOE Key Laboratory of Bioinorganic and Synthetic Chemistry, School of Chemistry, Sun Yat-Sen University, Guangzhou, Guangdong 510275, China

**Huangsheng Yang**—MOE Key Laboratory of Bioinorganic and Synthetic Chemistry, School of Chemistry, Sun Yat-Sen University, Guangzhou, Guangdong 510275, China
Qing Tong — Jiangsu Key Laboratory of Vehicle Emissions Control, Center of Modern Analysis, Nanjing University, Nanjing 210093, China
Zhengping Hao — National Engineering Laboratory for VOCs Pollution Control Material & Technology, University of Chinese Academy of Sciences, Beijing 101408, China

Complete contact information is available at: https://pubs.acs.org/10.1021/acscentsci.0c01600

Author Contributions
J.W. and O.G. conceived the project. J.W., J.Y., Q.F., H.Y. and Q.T. designed, analyzed, and discussed the experimental results. J.Y. and Q.F. carried out the catalytic performance tests. J.W. and H.Y. performed the measurements. Z.H. and O.G. gave suggestions on the experiments and drafted the manuscript.

Notes
The authors declare no competing financial interest.

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