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## Highly selective generation of singlet oxygen from dioxygen with atomically dispersed catalysts†

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Singlet oxygen ( ${}^1\text{O}_2$ ) as an excited electronic state of  $\text{O}_2$  plays a significant role in ubiquitous oxidative processes from enzymatic oxidative metabolism to industrial catalytic oxidation. Generally,  ${}^1\text{O}_2$  can be produced through thermal reactions or the photosensitization process; however, highly selective generation of  ${}^1\text{O}_2$  from  $\text{O}_2$  without photosensitization has never been reported. Here, we find that single-atom catalysts (SACs) with atomically dispersed  $\text{M}\text{N}_4$  sites on hollow N-doped carbon ( $\text{M}_1/\text{HNC}$  SACs,  $\text{M} = \text{Fe, Co, Cu, Ni}$ ) can selectively activate  $\text{O}_2$  into  ${}^1\text{O}_2$  without photosensitization, of which the  $\text{Fe}_1/\text{HNC}$  SAC shows an ultrahigh single-site kinetic value of  $3.30 \times 10^{10} \text{ min}^{-1} \text{ mol}^{-1}$ , representing top-level catalytic activity among known catalysts. Theoretical calculations suggest that different charge transfer from  $\text{M}\text{N}_4$  sites to chemisorbed  $\text{O}_2$  leads to the spin-flip process and spin reduction of  $\text{O}_2$  with different degrees. The superior capacity for highly selective  ${}^1\text{O}_2$  generation enables the  $\text{Fe}_1/\text{HNC}$  SAC as an efficient non-radiative therapeutic agent for *in vivo* inhibition of tumor cell proliferation.

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## Introduction

Dioxygen ( $\text{O}_2$ ) occupies a critical position in a great variety of oxidation reactions involved in both chemical reactions and biological processes.<sup>1–6</sup> In aerobic biology,  $\text{O}_2$ -related oxidation reactions are achieved by a series of evolutionary metalloenzymes such as cytochrome P450 through the interaction with  $\text{O}_2$ , resulting in the formation of reactive oxygen species (ROS) such as singlet oxygen ( ${}^1\text{O}_2$ ), hydroxyl radical ( ${}^{\cdot}\text{OH}$ ) and superoxide anion ( $\text{O}_2^-$ ) or reactive metal- $\text{O}_2$  intermediates including metal superoxo, (hydro)peroxo, and metal-oxo species.<sup>7–11</sup> Among the active species,  ${}^1\text{O}_2$  with an unoccupied  $\pi^*$  orbital has strong electrophilicity and unique reactivity and selectivity to serve as a synthetic reagent, attracting enormous interest from both fundamental studies and practical application fields.<sup>12–16</sup> To this end, increasing attention has been drawn

to the development of efficient approaches to highly selective generation of  ${}^1\text{O}_2$ .<sup>17–22</sup>

Conventional approaches to  ${}^1\text{O}_2$  production mainly include thermal processes using enzymatic or chemical reactions and photosensitization of  $\text{O}_2$  with rationally designed photosensitizers.<sup>23–26</sup> However, the production of  ${}^1\text{O}_2$  through these approaches is always accompanied by other ROS generation. In addition, thermal processes often require harsh conditions such as rigorous pH and particular solvents.<sup>27,28</sup> Despite the simplicity and controllability of the photosensitization route, it faces inherent limitations like poor selectivity of  ${}^1\text{O}_2$ , photo-bleaching of sensitizers, and difficulty in large-scale production.<sup>29–31</sup> Therefore, development of new approaches to highly selective production of  ${}^1\text{O}_2$  is of great significance not only in the investigation of  ${}^1\text{O}_2$ -related biological and physical processes but also in the development of new materials and biological tools with  ${}^1\text{O}_2$ .

Recently, single-atom catalysts (SACs) have shown considerable potential in industrial chemical processing and photo/electrochemical energy conversion with high catalytic activity and unique selectivity due to their excellent properties like maximum atom utilization efficiency, well-defined active centers, and tunable coordination environment.<sup>32–34</sup> Among the developed SACs, metal-nitrogen ( $\text{M}\text{N}_x$ ) sites embedded in carbon skeletons represent a significant series of SACs with excellent catalytic activity toward oxygen-related reactions including the oxygen reduction reaction and advanced oxidation processes, benefitting from precise tuning of electronic structures of active sites.<sup>35–38</sup> Thus,  $\text{M}\text{N}_x$  SACs provide great possibilities and opportunities in the design and fabrication of

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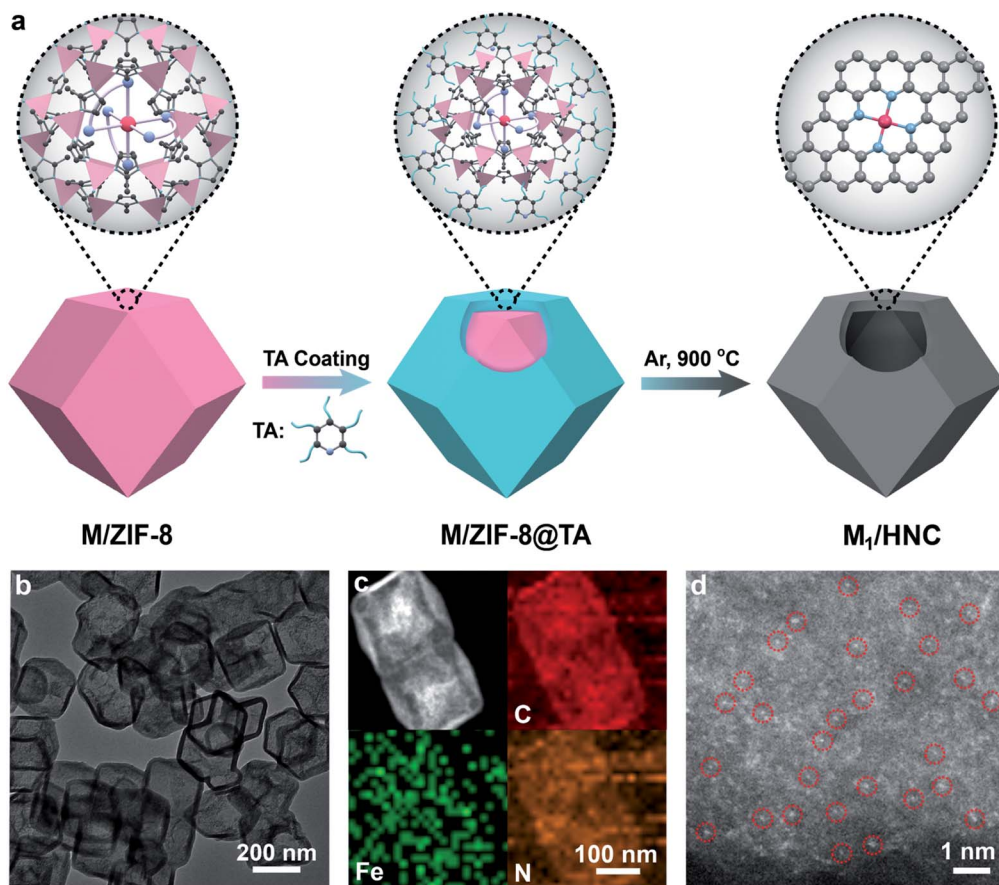
selective catalysts for  $^1\text{O}_2$  production. However, despite some investigations on the capacity of  $\text{MN}_x$  SACs toward oxygen activation with non-selective ROS production, the atomic engineering of  $\text{MN}_x$  SACs is still needed to optimize the metal–O<sub>2</sub> interaction to achieve selective  $^1\text{O}_2$  generation.

Here, we demonstrate a new approach to highly selective generation of  $^1\text{O}_2$  from O<sub>2</sub> without photosensitization with single transition metal atoms anchored on hollow N-doped carbon as atomically dispersed catalysts (M<sub>1</sub>/HNC SACs, M = Fe, Co, Cu, Ni). The  $^1\text{O}_2$  generation efficiency is highly related to the metal centers, following the sequence of Fe<sub>1</sub>/HNC > Co<sub>1</sub>/HNC > Cu<sub>1</sub>/HNC > Ni<sub>1</sub>/HNC. Among the catalysts examined, the Fe<sub>1</sub>/HNC SAC with a single atomic motif of FeN<sub>4</sub> coordination shows the best kinetic value of 0.140 min<sup>-1</sup>, exceeding those of Co<sub>1</sub>/HNC (0.033 min<sup>-1</sup>), Cu<sub>1</sub>/HNC (0.019 min<sup>-1</sup>) and Ni<sub>1</sub>/HNC (0.016 min<sup>-1</sup>), and the single-site kinetic value of the Fe<sub>1</sub>/HNC SAC reaches up to as high as  $3.30 \times 10^{10}$  min<sup>-1</sup> mol<sup>-1</sup>, representing top-level catalytic activity among known catalysts. Density functional theory (DFT) calculations demonstrate that the selective  $^1\text{O}_2$  generation capacity originates from significant charge transfer from MN<sub>4</sub> sites to chemisorbed O<sub>2</sub>, leading to the spin-flip process and spin reduction of O<sub>2</sub> with the lowest value of 0.25 for O<sub>2</sub> adsorbed on the FeN<sub>4</sub> site. Based on the

efficient production of  $^1\text{O}_2$  enabled by the Fe<sub>1</sub>/HNC SAC, we develop a non-radiative therapeutic platform for *in vivo* inhibition of tumor cell proliferation.

## Results and discussion

To synthesize M<sub>1</sub>/HNC SACs, metal acetylacetonate encapsulated zeolitic imidazolate framework-8 (M/ZIF-8) was first synthesized as the raw material to form core–shell composites by tannic acid (TA) coating. The core–shell precursors were then pyrolyzed at 900 °C under an Ar atmosphere to obtain M<sub>1</sub>/HNC catalysts (Fig. 1a). Hollow N-doped carbon (HNC) was prepared with pure ZIF-8 without the encapsulation of metal acetylacetone as the control catalyst. X-ray diffraction (XRD) patterns of the catalysts show peaks for (002) and (101) planes of graphitic carbon, located at around 24° and 43°, respectively, and no observable peaks for metal-related nanoparticles were found (Fig. S1†), indicating the absence of metal nanoparticles in M<sub>1</sub>/HNC catalysts.<sup>39</sup> Raman spectra of the M<sub>1</sub>/HNC SACs and HNC reveal similar intensity ratios of the D band to the G band (Fig. S2†), suggesting that the catalysts possess a similar structure of the defective carbon skeleton. In addition, the nitrogen sorption isotherm analysis of M<sub>1</sub>/HNC SACs and HNC (Fig. S3†)



**Fig. 1** Synthesis of M<sub>1</sub>/HNC SACs and structural characterization of the Fe<sub>1</sub>/HNC SAC. (a) Schematic depiction of the route to the fabrication of M<sub>1</sub>/HNC SACs. (b) TEM image of Fe<sub>1</sub>/HNC. (c) HAADF-STEM image and the corresponding elemental mapping images of Fe<sub>1</sub>/HNC (C, red; Fe, green; N, orange). (d) Aberration-corrected HAADF-STEM image of Fe<sub>1</sub>/HNC showing atomically dispersed Fe atoms as bright dots highlighted with red circles.



gave a similar Brunner–Emmet–Teller (BET) surface area at around  $700 \text{ cm}^2 \text{ g}^{-1}$ , further indicating the structural similarity of the prepared catalysts. Transmission electron microscopy (TEM) and high-angle annular dark-field scanning TEM (HAADF-STEM) images show that  $\text{Fe}_1/\text{HNC}$  has a hollow polyhedral morphology with an average size of 200 nm and thickness of 10 nm and no metal nanoparticles were observed (Fig. 1b). The elemental mapping images show a homogeneous distribution of Fe, N, and C elements over the whole domain of the  $\text{Fe}_1/\text{HNC}$  SAC (Fig. 1c). Furthermore, the aberration-corrected HAADF-STEM image was obtained to confirm single atomic Fe in the  $\text{Fe}_1/\text{HNC}$  SAC. As shown in Fig. 1d, the bright dots highlighted with red circles clearly demonstrate that the atomically dispersed Fe atoms exist over N-doped carbon. Similarly, the morphology of other three  $\text{M}_1/\text{HNC}$  SACs and the atomic dispersion were confirmed by using TEM and HAADF-STEM images, all showing a hollow polyhedral structure and atomically dispersed metal atoms (Fig. S4–S6†).

The atomic electronic structure and coordination configuration of metal species in  $\text{M}_1/\text{HNC}$  SACs were explored by X-ray absorption near-edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) characterization experiments. Fig. 2a shows the XANES spectra of  $\text{Fe}_1/\text{HNC}$  at the Fe  $K$ -edge in comparison with those of references including Fe foil and  $\text{Fe}_2\text{O}_3$ . The peak position of  $\text{Fe}_1/\text{HNC}$  is situated between those of  $\text{Fe}_2\text{O}_3$  and Fe foil, indicating that the valence state of the Fe atom is between 0 and +3. The Fourier-transform EXAFS spectrum of  $\text{Fe}_1/\text{HNC}$  shows only one peak at about 1.5 Å, ascribed to the first coordination shell of Fe–N, and no Fe–Fe

peak like that of Fe foil at around 2.2 Å was observed (Fig. 2b), suggesting the inexistence of metal-related nanoparticles or clusters in  $\text{Fe}_1/\text{HNC}$ . In order to further investigate the atomic distribution of Fe in  $\text{Fe}_1/\text{HNC}$ , wavelet transform (WT) analysis of Fe  $K$ -edge EXAFS was carried out. As illustrated in Fig. 2c, only one intensity maximum at approximately 4.2 Å<sup>−1</sup> related to Fe–N coordination was observed from the WT contour plot of  $\text{Fe}_1/\text{HNC}$ , which is different from the contour WT plots of Fe foil and  $\text{Fe}_2\text{O}_3$  with the intensity maximum corresponding to Fe–Fe contribution, elucidating that no Fe–Fe bond is present in  $\text{Fe}_1/\text{HNC}$ , but isolated Fe atoms exist. Moreover, the coordination configuration and structural parameters of Fe atoms were obtained from EXAFS fitting curves (Fig. 2d and S7†). As listed in Table S1,† the coordination number of Fe atoms and the bond length of Fe–N are 4.2 and 2.02 Å, respectively. From the above results, the atomic structure configuration of  $\text{Fe}_1/\text{HNC}$  was constructed (Fig. 2e). The local structure of other three  $\text{M}_1/\text{HNC}$  SACs was also confirmed with XANES and EXAFS spectra (Fig. S8–S10†), all possessing similar atomic configuration with  $\text{Fe}_1/\text{HNC}$ .

In order to systematically evaluate the oxidizing capacity of  $\text{M}_1/\text{HNC}$  SACs, the oxidation of colorless 3,3',5,5'-tetramethylbenzidine (TMB) to blue oxidized TMB (ox-TMB) with the characteristic adsorption at *ca.* 652 nm was used as the catalytic model reaction.<sup>40–43</sup> Considering that the pyrolysis temperature has significant effect on the coordination microenvironment of SACs,<sup>44,45</sup> the TMB oxidation activities of the catalysts prepared at different temperatures were explored. As shown in Fig. 3a, the catalyst obtained at 900 °C (denoted as  $\text{Fe}_1/\text{HNC}$ -900) exhibits

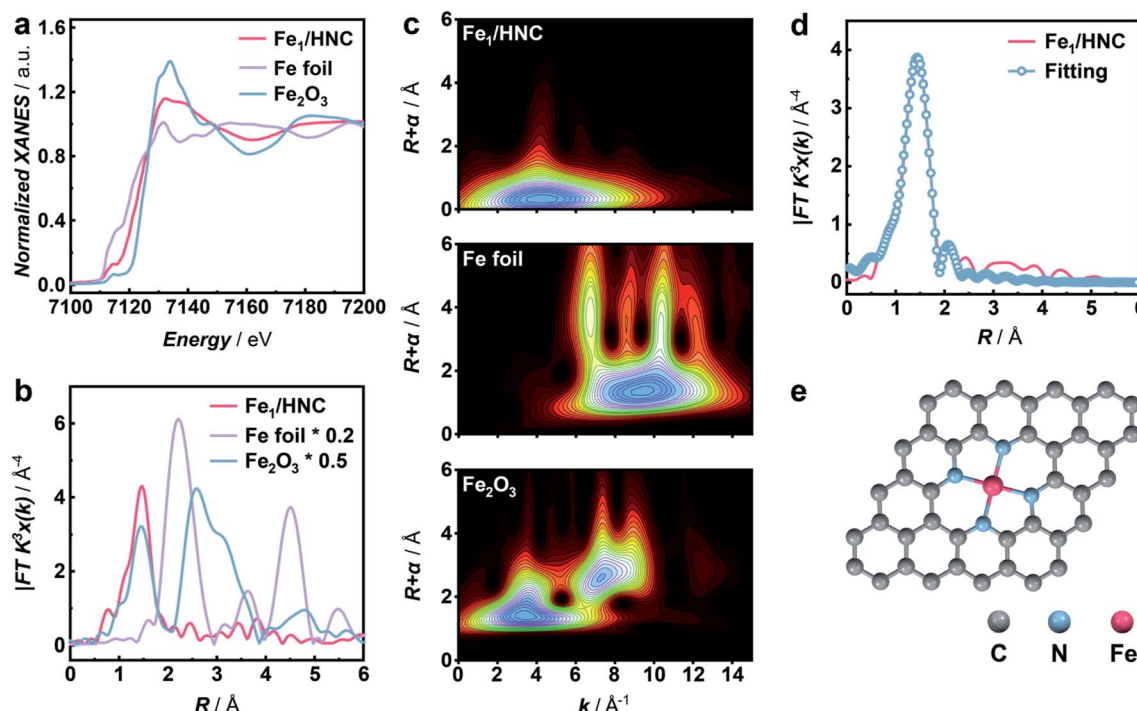
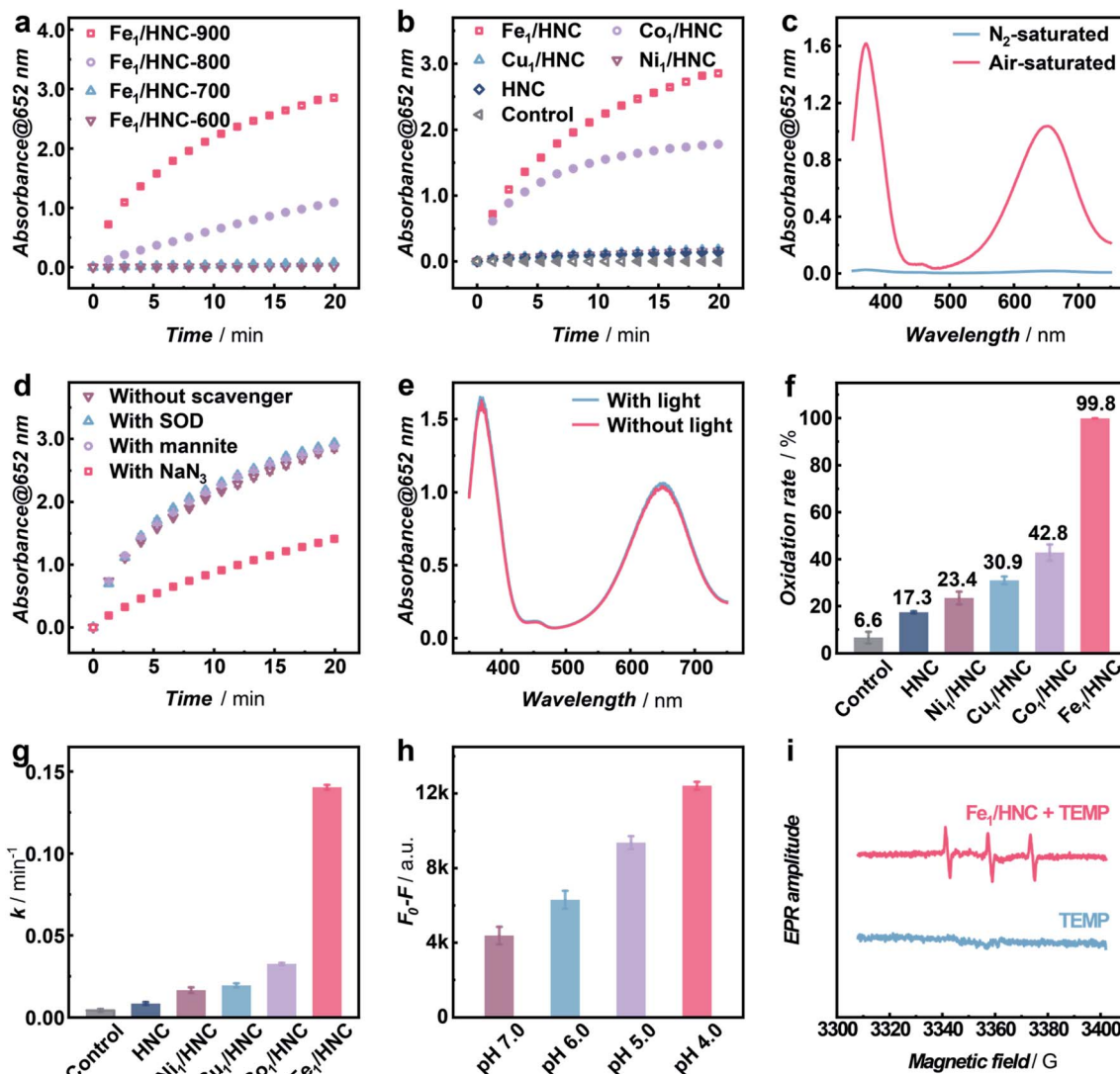


Fig. 2 XAFS analysis of the  $\text{Fe}_1/\text{HNC}$  SAC. (a) Normalized XANES spectra and (b) corresponding Fourier-transform EXAFS spectra at the Fe  $K$ -edge of  $\text{Fe}_1/\text{HNC}$  (red line), Fe foil (purple line), and  $\text{Fe}_2\text{O}_3$  (blue line). (c) Wavelet transforms of the Fe  $K$ -edge EXAFS spectra for  $\text{Fe}_1/\text{HNC}$ , Fe foil, and  $\text{Fe}_2\text{O}_3$ . (d) Fourier-transform EXAFS spectrum and the corresponding fitting curve of  $\text{Fe}_1/\text{HNC}$  in  $R$ -space. (e) Schematic structure of  $\text{Fe}_1/\text{HNC}$ .





**Fig. 3** Selective generation of  $^1\text{O}_2$  with  $\text{M}_1/\text{HNC}$  SACs. Time-dependent absorbance changes of ox-TMB monitored at 652 nm (a) with  $5 \mu\text{g mL}^{-1}$   $\text{M}_1/\text{HNC}$  synthesized at different temperatures or (b) with  $5 \mu\text{g mL}^{-1}$   $\text{M}_1/\text{HNC}$  and HNC in the Britton–Robinson (B–R) buffer (pH 4.0). (c) UV-vis absorption spectra of ox-TMB catalyzed by  $5 \mu\text{g mL}^{-1}$   $\text{Fe}_1/\text{HNC}$  at 2 min in the  $\text{N}_2$ -saturated (blue line) and air-saturated (red line) B–R buffer (pH 4.0). (d) Time-dependent absorbance changes of ox-TMB monitored at 652 nm catalyzed by  $5 \mu\text{g mL}^{-1}$   $\text{Fe}_1/\text{HNC}$  in the B–R buffer (pH 4.0) without scavengers (brown line) or with SOD (blue line), mannite (purple line), and  $\text{NaN}_3$  (red line). (e) UV-vis absorption spectra of ox-TMB catalyzed by  $5 \mu\text{g mL}^{-1}$   $\text{Fe}_1/\text{HNC}$  in 2 min without (red line) or with (blue line) light in the B–R buffer (pH 4.0). (f) Oxidation rate of ABDA with  $\text{M}_1/\text{HNC}$  and HNC in the B–R buffer (pH 4.0). (g)  $k$  value of the ABDA oxidation reaction with  $5 \mu\text{g mL}^{-1}$   $\text{M}_1/\text{HNC}$  and HNC in the B–R buffer (pH 4.0). (h) Decrease in fluorescence intensity ( $\lambda_{\text{ex}}/\lambda_{\text{em}} = 380/433 \text{ nm}$ ) of ABDA with  $5 \mu\text{g mL}^{-1}$   $\text{Fe}_1/\text{HNC}$  in 5 min in the B–R buffer at different pH values. (i) TEMPO ESR signals in the absence (blue line) and presence of  $20 \mu\text{g mL}^{-1}$   $\text{Fe}_1/\text{HNC}$  in the B–R buffer (pH 4.0).

the highest catalytic activity towards TMB oxidation, and  $900^\circ\text{C}$  was thus chosen to prepare other SACs. We next compared the catalytic activity of  $\text{M}_1/\text{HNC}$  SACs with different transition metal centers toward TMB oxidation. The results displayed in Fig. 3b and S11<sup>†</sup> show that the activity trends follow the sequence of  $\text{Fe}_1/\text{HNC} > \text{Co}_1/\text{HNC} > \text{Cu}_1/\text{HNC} > \text{Ni}_1/\text{HNC} > \text{HNC}$ , indicating that the single metal atom center plays a vital role in the oxidation reaction. Among the catalysts investigated,  $\text{Cu}_1/\text{HNC}$  and  $\text{Ni}_1/\text{HNC}$  SACs exhibit quite low catalytic activity towards TMB oxidation like HNC. Comparatively, both  $\text{Fe}_1/\text{HNC}$  and  $\text{Co}_1/\text{HNC}$  SACs show a high catalytic oxidation rate, far surpassing those of  $\text{Cu}_1/\text{HNC}$  and  $\text{Ni}_1/\text{HNC}$  SACs. The

significant difference in catalytic performance of  $\text{M}_1/\text{HNC}$  SACs may arise from the binding strength of different  $\text{MN}_4$  centers with O species.<sup>46–48</sup>

To verify whether  $\text{O}_2$  is the only substrate for the oxidation reaction,  $\text{Fe}_1/\text{HNC}$  was selected as the representative catalyst to evaluate the TMB oxidation reaction under a  $\text{N}_2$  atmosphere. As shown in Fig. 3c, the absorbance intensity of ox-TMB undergoes a sharp decrease in the  $\text{N}_2$ -saturated B–R buffer compared with that under air-saturated conditions, indicating the indispensable role of  $\text{O}_2$  in the oxidation of TMB catalyzed by the  $\text{Fe}_1/\text{HNC}$  SAC. In addition, as displayed in Fig. S12–S15,<sup>†</sup> the catalytic

TMB oxidation reaction with  $M_1$ /HNC (Fe<sub>1</sub>/HNC and Co<sub>1</sub>/HNC as examples) is dose- and pH-dependent.

It is generally known that the catalytic oxidation reactions with O<sub>2</sub> are always accompanied with the generation of ROS like ·OH, O<sub>2</sub><sup>−</sup> and <sup>1</sup>O<sub>2</sub>, thus, it is vital to identify the exact species produced during the process of the oxidation reaction by  $M_1$ /HNC SACs. For this purpose, mannite, superoxide dismutase (SOD), and NaN<sub>3</sub> were chosen as the ROS scavengers for ·OH, O<sub>2</sub><sup>−</sup>, and <sup>1</sup>O<sub>2</sub>, respectively.<sup>49</sup> Notably, only NaN<sub>3</sub> exhibits efficient inhibition of TMB oxidation catalyzed by the Fe<sub>1</sub>/HNC SAC, while the others display no obvious effects on the oxidation reaction (Fig. 3d), suggesting that the Fe<sub>1</sub>/HNC SAC selectively activates O<sub>2</sub> into <sup>1</sup>O<sub>2</sub>. In consideration of photosensitization usually implemented to produce <sup>1</sup>O<sub>2</sub>, the comparison of the Fe<sub>1</sub>/HNC-catalyzed TMB oxidation reaction under normal light conditions and in a dark environment was performed. As shown in Fig. 3e, a negligible difference in characteristic absorbance of ox-TMB was observed, indicating

the striking generation of <sup>1</sup>O<sub>2</sub> without photosensitization. Collectively, highly selective generation of <sup>1</sup>O<sub>2</sub> from O<sub>2</sub> can be achieved by using the Fe<sub>1</sub>/HNC SAC without the assistance of the externally applied stimulus.

To further validate the selective production of <sup>1</sup>O<sub>2</sub> from O<sub>2</sub> enabled by  $M_1$ /HNC SACs, an <sup>1</sup>O<sub>2</sub>-specific probe 9,10-anthracenediyl-bis(methylene) dimalonic acid (ABDA) was adopted to selectively recognize and react with <sup>1</sup>O<sub>2</sub>.<sup>50,51</sup> ABDA is a fluorescent molecule with  $\lambda_{\text{ex}}$  and  $\lambda_{\text{em}}$  at *ca.* 380 nm and 433 nm, respectively, which turns to be a non-fluorescent endoperoxide product upon selective oxidation by <sup>1</sup>O<sub>2</sub> *via* a [4 + 2]-cycloaddition. With ABDA as the probe, we evaluated the catalytic efficiency of  $M_1$ /HNC SACs in generating <sup>1</sup>O<sub>2</sub> from O<sub>2</sub>. As can be seen in Fig. 3f and S16,<sup>†</sup> the activity towards ABDA oxidation follows the rate sequence of Fe<sub>1</sub>/HNC (99.8%) > Co<sub>1</sub>/HNC (42.8%) > Cu<sub>1</sub>/HNC (30.9%) > Ni<sub>1</sub>/HNC (23.4%) > HNC (17.3%), which is consistent with the trends in TMB oxidation. Notably, Fe<sub>1</sub>/HNC displays the highest catalytic activity toward

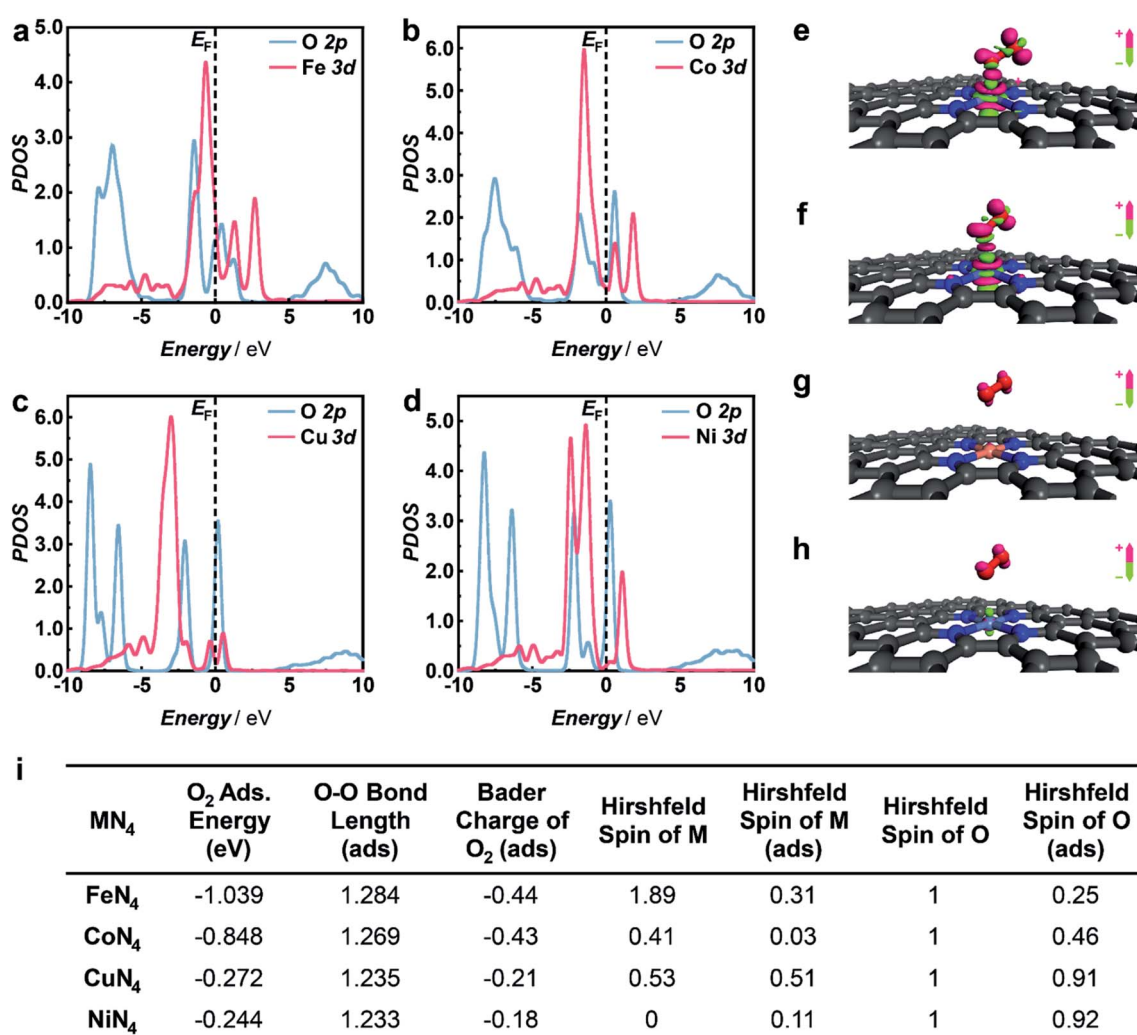


Fig. 4 DFT calculations. PDOS of the M 3d (red line) and O 2p (blue line) orbitals of (a) FeN<sub>4</sub>, (b) CoN<sub>4</sub>, (c) CuN<sub>4</sub> and (d) NiN<sub>4</sub> sites adsorbed with O<sub>2</sub>. Calculated charge density differences of (e) FeN<sub>4</sub>, (f) CoN<sub>4</sub>, (g) CuN<sub>4</sub> and (h) NiN<sub>4</sub> centers adsorbed with O<sub>2</sub>. (i) Comparison of the DFT results for constructed MN<sub>4</sub> structures.



ABDA oxidation with a rate of nearly 100%, indicative of more efficient  $^1\text{O}_2$  generation catalyzed by  $\text{Fe}_1/\text{HNC}$  in comparison with other SACs. In addition, the kinetic profiles of ABDA oxidation were examined to assess  $^1\text{O}_2$  generation capacities of  $\text{M}_1/\text{HNC}$  SACs by calculating the oxidation rate of ABDA with pseudo-first-order approximation. As shown in Fig. S17,<sup>†</sup> an exponential decrease in the fluorescence intensity of ABDA against the reaction time was observed and the pseudo-first-order kinetic constants ( $k$  values) were obtained from the linear relationship. The results shown in Fig. 3g reveal that  $\text{Fe}_1/\text{HNC}$  catalyzes the generation of  $^1\text{O}_2$  for ABDA oxidation with a  $k$  value of  $0.140\text{ min}^{-1}$ , considerably higher than that of  $\text{Co}_1/\text{HNC}$  ( $0.033\text{ min}^{-1}$ ),  $\text{Cu}_1/\text{HNC}$  ( $0.019\text{ min}^{-1}$ ) and  $\text{Ni}_1/\text{HNC}$  ( $0.016\text{ min}^{-1}$ ), further highlighting the critical role of different single metal atom coordination. The single-site kinetic constant ( $k_{\text{single-site}}$  value) of the  $\text{Fe}_1/\text{HNC}$  SAC was further evaluated to be as high as  $3.30 \times 10^{10}\text{ min}^{-1}\text{ mol}^{-1}$  due to the high catalytic efficiency with a low metal content of 0.17 wt% determined *via* inductively coupled plasma optical emission spectrometry (ICP-OES). Besides, the  $\text{Fe}_1/\text{HNC}$ -enabled selective generation of  $^1\text{O}_2$  was investigated with different concentrations of the catalyst and in reaction media with different pH values (Fig. 3h and S18<sup>†</sup>). As shown in Fig. S19,<sup>†</sup> the ABDA oxidation reaction shows a dose-dependent behavior. In addition, the generation of  $^1\text{O}_2$  for ABDA oxidation is peculiarly prone to occur under acidic conditions.

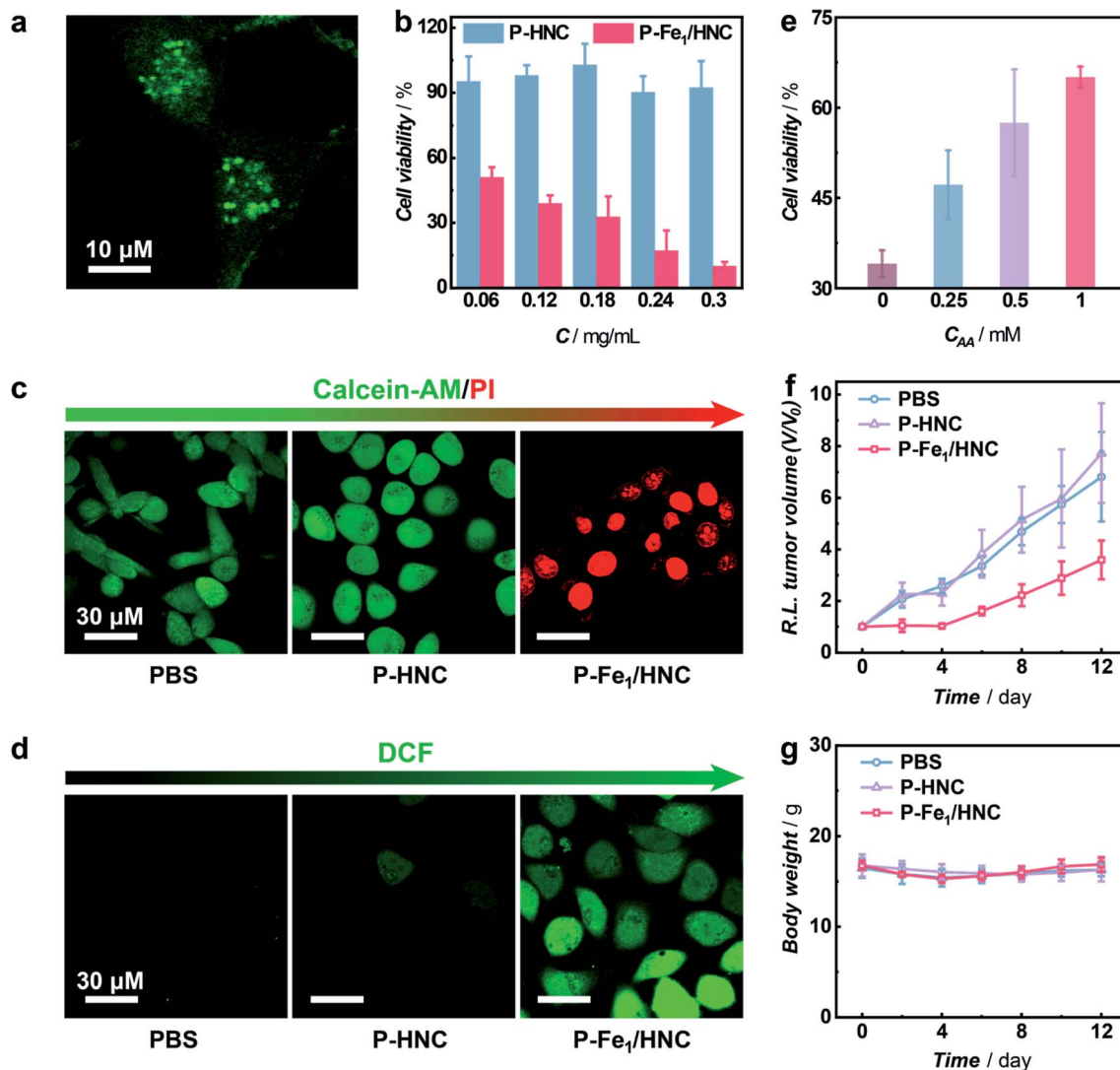
To further confirm the selective generation of  $^1\text{O}_2$  enabled by  $\text{Fe}_1/\text{HNC}$ , colorimetric and fluorescent probes for other ROSs were utilized, thereinto,  $\rho$ -phthalic acid (PTA) and iodonitrotetrazolium chloride (INT) were selected to detect  $\cdot\text{OH}$  and  $\text{O}_2^-$ , respectively. PTA, as a widely used  $\cdot\text{OH}$  probe, can be selectively hydroxylated upon reaction with  $\cdot\text{OH}$  to form 2-hydroxy terephthalic acid with strong fluorescence at  $\lambda_{\text{ex}}/\lambda_{\text{em}} = 315/400\text{ nm}$ .<sup>52</sup> INT is a specific probe for  $\text{O}_2^-$ , which can be reduced by  $\text{O}_2^-$  to generate the colored formazan product with the maximum absorption at 510 nm. As shown in Fig. S20,<sup>†</sup> no obvious changes in the fluorescence intensity of PTA or absorbance of INT were observed in the oxidation reaction, demonstrating negligible generation of  $\cdot\text{OH}$  and  $\text{O}_2^-$  but highly selective production of  $^1\text{O}_2$  during the  $\text{O}_2$  activation process. Furthermore, electron spin resonance (ESR) characterization using 2,2,6,6-tetramethylpiperidine (TEMP) as the  $^1\text{O}_2$  trapping agent and 5-*tert*-butoxycarbonyl-5-methyl-1-pyrroline *N*-oxide (BMPO) as the trapping agent for  $\cdot\text{OH}$  and  $\text{O}_2^-$  were also conducted to give more direct evidence of the produced ROS.<sup>53,54</sup> Fig. 3i shows the typical 1 : 1 : 1 triplet ESR signal for 2,2,6,6-tetramethylpiperidine-*N*-oxyl (TEMPO) formed due to the  $^1\text{O}_2$  generation in the presence of the  $\text{Fe}_1/\text{HNC}$  SAC. However, there are no observable signals when BMPO was used as the trapping agent (Fig. S21<sup>†</sup>), indicating that negligible  $\cdot\text{OH}$  and  $\text{O}_2^-$  are produced in the process of  $\text{Fe}_1/\text{HNC}$ -catalyzed  $\text{O}_2$  activation.

To gain in-depth insight into the mechanism of selective  $^1\text{O}_2$  generation enabled by  $\text{M}_1/\text{HNC}$  SACs, DFT calculations were performed to explore the electronic structure of the active sites and their interactions with  $\text{O}_2$ .<sup>55-58</sup> In accordance with the atomic structural analysis, the models of  $\text{M}_1/\text{HNC}$  SACs adsorbed with  $\text{O}_2$  were constructed and optimized. The adsorption

energy of  $\text{O}_2$  on the  $\text{FeN}_4$  site in the  $\text{Fe}_1/\text{HNC}$  SAC was calculated to be  $-1.039\text{ eV}$ , which is less than those of other  $\text{M}_1/\text{HNC}$  SACs (Fig. 4i), illustrating the strong interaction between the  $\text{FeN}_4$  site and  $\text{O}_2$ . In order to further elucidate the electronic interactions between  $\text{MN}_4$  sites and absorbed  $\text{O}_2$ , the projected density of states (PDOS) for  $\text{M}$  3d and  $\text{O}$  2p was calculated. As shown in Fig. 4a-d, more hybridization between the PDOS for Fe 3d orbitals and O 2p orbitals can be evidently observed in contrast with those for  $\text{CuN}_4$  or  $\text{NiN}_4$  centers. Meanwhile, the Fe 3d orbitals shift toward the Fermi level upon  $\text{O}_2$  adsorption (Fig. S22-S25<sup>†</sup>), further demonstrating the occurrence of  $\text{O}_2$  activation on  $\text{FeN}_4$ .<sup>59,60</sup> In addition, the Bader charge analysis and charge density difference analysis (Fig. 4e-h) reveal distinct charge transfer with 0.44 e from the  $\text{FeN}_4$  site to  $\text{O}_2$ , leading to the elongation of the O–O bond with a length from 1.22 Å in free  $\text{O}_2$  to 1.284 Å in  $\text{O}_2$  adsorbed on the  $\text{FeN}_4$  site (Fig. 4i), which is longer than those on other  $\text{MN}_4$  sites. The longest O–O bond of  $\text{O}_2$  upon adsorption on the  $\text{FeN}_4$  site manifests the strongest interaction between the  $\text{FeN}_4$  site and  $\text{O}_2$ . The transferred electrons will occupy the half-filled antibonding  $\pi^*$  orbitals of  $\text{O}_2$  and give rise to the reduction of the Hirshfeld spin value of  $\text{O}_2$  from 1 in free  $\text{O}_2$  to 0.25, suggesting that a significant spin-flip process occurs for chemisorbed  $\text{O}_2$  on the  $\text{FeN}_4$  site. The spin value of  $\text{O}_2$  upon chemisorption on the  $\text{FeN}_4$  site (0.25) is close to that of  $^1\text{O}_2$  (0) and relatively lower than those on other  $\text{MN}_4$  sites, elucidating the superior capacity of the  $\text{FeN}_4$  site toward the generation of  $^1\text{O}_2$ . Given the above, the charge transfer from the  $\text{MN}_4$  site to adsorbed  $\text{O}_2$  contributes to different spin-flip degrees of  $\text{O}_2$ , thus bringing about a discrepancy in their capacities toward selective  $^1\text{O}_2$  generation.

Having demonstrated the high efficiency of selective  $^1\text{O}_2$  generation without photosensitization from molecular  $\text{O}_2$  enabled by the  $\text{Fe}_1/\text{HNC}$  SAC, we then employed the  $\text{Fe}_1/\text{HNC}$  SAC as a therapeutic agent for suppressing tumor cell growth. To improve the dispersibility and biocompatibility of  $\text{Fe}_1/\text{HNC}$ , an amphipathic molecule DSPE-PEG2000 was modified on the surface of  $\text{Fe}_1/\text{HNC}$  to form PEGylated  $\text{Fe}_1/\text{HNC}$  (denoted as P- $\text{Fe}_1/\text{HNC}$ ) through the hydrophobic interaction.<sup>61</sup> The cellular uptake of P- $\text{Fe}_1/\text{HNC}$  was visualized by confocal laser scanning microscopy (CLSM) imaging with fluorescein isothiocyanate (FITC)-loaded P- $\text{Fe}_1/\text{HNC}$ . As displayed in Fig. 5a and S26,<sup>†</sup> the bright green fluorescence and the black dots in cytosol clearly indicate the efficient internalization of P- $\text{Fe}_1/\text{HNC}$  in HeLa cells. Next, the inhibition effect of P- $\text{Fe}_1/\text{HNC}$  on cell proliferation was systematically explored by the cell viability test with a standard MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium] assay. The results show that the treatment of HeLa cells with PEGylated HNC (P-HNC) without single Fe atomic centers exhibits a negligible effect on the proliferation of HeLa cells with cell viability exceeding 90% even at a concentration of P-HNC as high as  $0.3\text{ mg mL}^{-1}$  (Fig. 5b), indicating the excellent biocompatibility and low cytotoxicity of P-HNC. In striking contrast, an obvious decrease in cell viability of HeLa cells to less than 50% upon the treatment of P- $\text{Fe}_1/\text{HNC}$  with a low concentration of  $0.06\text{ mg mL}^{-1}$  was observed and the cell viability decreases with increasing P- $\text{Fe}_1/\text{HNC}$  concentration,





**Fig. 5** *In vitro* and *in vivo* inhibition of tumor growth with the Fe<sub>1</sub>/HNC SAC. (a) CLSM fluorescence image of HeLa cells treated with FITC-loaded P-Fe<sub>1</sub>/HNC for 6 h. (b) Cell viability of HeLa cells after 12 h-treatment of P-HNC or P-Fe<sub>1</sub>/HNC at the concentrations of 0.06–0.3 mg mL<sup>-1</sup>. CLSM fluorescence images of HeLa cells stained with Calcein-AM and (c) PI or (d) DCFH-DA after the 12 h-treatment with PBS, P-HNC and P-Fe<sub>1</sub>/HNC. (e) Cell viability of HeLa cells pre-incubated with AA (0.25 mM, 0.5 mM, and 1 mM) after 12 h-treatment with 0.1 mg mL<sup>-1</sup> P-Fe<sub>1</sub>/HNC. *In vivo* (f) tumor proliferation and (g) body weight curves of mice treated with the intravenous injection of PBS (blue line), P-HNC (purple line), or P-Fe<sub>1</sub>/HNC (red line).

providing direct evidence for the antiproliferation effect on HeLa cells.

In the meantime, the cytotoxicity of P-Fe<sub>1</sub>/HNC was further investigated by co-staining HeLa cells with calcein-AM (green fluorescent dye for viable cell staining) and propidium iodide (PI, red fluorescent dye for dead cell staining) and visualized with CLSM imaging.<sup>62</sup> As exhibited in Fig. 5c and S27,† in comparison with the group treated with PBS or P-HNC showing bright green fluorescence, cancer cells treated with P-Fe<sub>1</sub>/HNC show strong red fluorescence, indicating the high cytotoxicity against HeLa cells of Fe<sub>1</sub>/HNC SACs. According to the results demonstrated above, such capacity of killing cancer cells was deemed to originate from the intracellular oxidative stress induced by Fe<sub>1</sub>/HNC SAC-enabled selective production of <sup>1</sup>O<sub>2</sub>, which was distinctly confirmed with a cell-permeable ROS-

sensitive probe (2',7'-dichlorofluorescein diacetate, DCFH-DA).<sup>63,64</sup> Obviously, cancer cells treated with P-Fe<sub>1</sub>/HNC show the brightest green fluorescence of DCF (Fig. 5d and S28†), indicative of most <sup>1</sup>O<sub>2</sub> generation enabled by Fe<sub>1</sub>/HNC. In order to further demonstrate the cytotoxicity against HeLa cells of the Fe<sub>1</sub>/HNC SAC from the oxidative damage induced by the generation of <sup>1</sup>O<sub>2</sub>, ascorbic acid (AA) as an antioxidant was used to pre-incubate HeLa cells.<sup>61</sup> As shown in Fig. 5e, the pre-incubation of AA can efficiently reduce the cytotoxicity of the Fe<sub>1</sub>/HNC SAC through the elimination of generated <sup>1</sup>O<sub>2</sub> with a concentration-dependent behavior.

The antitumor effect of the Fe<sub>1</sub>/HNC SAC *in vivo* was also investigated with HeLa tumor-bearing mice as a model. When the tumor volume reached about 100 mm<sup>3</sup>, the mice were randomly divided into three groups and intravenously injected

with PBS, P-HNC, and P-Fe<sub>1</sub>/HNC, respectively. As shown in Fig. 5f, the mice treated with P-HNC exhibit a similar tumor growth tendency with that of PBS, indicating that HNC had no obvious therapeutic effect on suppressing tumor growth. In contrast, the group treated with P-Fe<sub>1</sub>/HNC displays a relatively higher tumor inhibition effect with the reduction of the tumor size to 46.5% in 12 days compared with the control group, which was attributed to the high capacity of the Fe<sub>1</sub>/HNC SAC for <sup>1</sup>O<sub>2</sub> generation within the tumor site. Moreover, photographs of the excised tumors further confirmed the good anti-tumor effect of Fe<sub>1</sub>/HNC (Fig. S29†). In addition, no obvious changes of the body weight of mice were observed during the experimental period upon administration of different catalysts, indicating the favorable biocompatibility of the Fe<sub>1</sub>/HNC SAC (Fig. 5g). Further blood biochemical analysis also implied the good biosafety of these catalysts (Fig. S30†).

## Conclusions

In summary, we have demonstrated that atomically dispersed MN<sub>4</sub> sites on hollow N-doped carbon can be used for the selective generation of strong oxidizing <sup>1</sup>O<sub>2</sub> from O<sub>2</sub> without photosensitization. The as-prepared M<sub>1</sub>/HNC SACs show different <sup>1</sup>O<sub>2</sub> generation efficiencies in the sequence of Fe<sub>1</sub>/HNC > Co<sub>1</sub>/HNC > Cu<sub>1</sub>/HNC > Ni<sub>1</sub>/HNC. Among the developed catalysts, the Fe<sub>1</sub>/HNC SAC shows the best kinetic value of 0.14 min<sup>-1</sup> and a single-site kinetic value of  $3.30 \times 10^{10}$  min<sup>-1</sup> mol<sup>-1</sup>, originating from the spin-flip process and spin reduction of O<sub>2</sub> to 0.25 induced *via* significant charge transfer with 0.44 e from the FeN<sub>4</sub> site to O<sub>2</sub>. More importantly, the Fe<sub>1</sub>/HNC SAC with superior <sup>1</sup>O<sub>2</sub> generation capacity has been successfully utilized as an efficient non-radiative therapeutic agent for inhibiting tumor cell proliferation *in vitro* and *in vivo*. We believe that this finding would provide a facile method for the selective production of <sup>1</sup>O<sub>2</sub>, opening up a new avenue for the development of functional SACs in biomedical applications.

## Data availability

All experimental and computational data is available in the ESI.†

## Author contributions

L. M., W. M. and J. M. conceived the idea for the project. W. M. and J. M. designed the experiments and conducted material synthesis, structural characterization, performance test and data analysis. C. H. carried out the DFT calculations. L. S. contributed to the animal experiments. J. L. and M. W. helped in the cell experiments. P. Y. contributed to the discussion of the catalytic part. W. M. drafted the manuscript, and L. M. finalized the manuscript. All authors discussed and commented on the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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