A cocoon silk chemistry strategy to ultrathin N-doped carbon nanosheet with metal single-site catalysts

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Development of single-site catalysts supported by ultrathin two-dimensional (2D) porous matrix with ultrahigh surface area is highly desired but also challenging. Here we report a cocoon silk chemistry strategy to synthesize isolated metal single-site catalysts embedded in ultrathin 2D porous N-doped carbon nanosheets (M-ISA/CNS, M = Fe, Co, Ni). X-ray absorption fine structure analysis and spherical aberration correction electron microscopy demonstrate an atomic dispersion of metal atoms on N-doped carbon matrix. In particular, the Co-ISA/CNS exhibit ultrahigh specific surface area (2105 m$^2$ g$^{-1}$) and high activity for C-H bond activation in the direct catalytic oxidation of benzene to phenol with hydrogen peroxide at room temperature, while the Co species in the form of phthalocyanine and metal nanoparticle show a negligible activity. Density functional theory calculations discover that the generated O = Co = O center intermediates on the single Co sites are responsible for the high activity of benzene oxidation to phenol.
Single-site catalysts with isolated metal single-atomic sites supported by solid matrix are emerging as an exciting research frontier in catalysis. With atomically metal dispersion, such single-site catalysts display several distinct advantages in the maximized atom efficiency, strengthened selectivity towards targeted product, improved intrinsic activity, and facilitative recyclability. Indeed, single-site catalysts, including Au, Pt, Pd, Ru, Ir, Rh, Fe, Co, Ni, Mn, Mo, and W, are reported over past few years and exhibit interesting activity in CO oxidation, solar energy conversion, hydrogen production, hydrogenation, CO2 reduction, and oxygen reduction reaction. To date, several synthetic protocols have been reported for fabricating single-site catalysts including photochemical reduction, co-precipitation, atomic layer deposition (ALD), wet impregnation, and so on. Accordingly, the construction of support materials with rich anchoring sites and identical geometric structure is of fundamental importance for achieving desirable catalytic performance. Moreover, significantly enlarging surface area of support materials can expose the most active sites accessible to reactants, which has become considered as an effective route to boost the intrinsic activity of these heterogeneous supported single-site catalysts. Various zeolites, metal–organic frameworks (MOFs), and well-defined porous carbon materials have been employed as possible support for single-site catalysts. Therefore, establishing rational and general synthetic approaches to prepare single-site catalysts with controllable structure and ultrahigh surface area to take full advantage of their potential is highly desired but challenging.

The catalytic performances of single-site catalysts are highly dependent upon support materials. Considerable researches have demonstrated that support materials can not only firmly stabilize isolated single metal sites but often play a key role in controlling the local geometric and electronic structures of catalytic sites, leading to excellent retentivity of individual site state under catalytic conditions and superb catalytic activity. To date, the reported support materials generally consist of the oxides and carbon-based materials. N-doped porous carbon materials have been widely employed for (i) rich anchoring sites, (ii) favorable mass adsorption and transportation facilitated by controllable porous structure with high surface area, and (iii) good thermal stability under harsh catalytic conditions. Ultrathin 2D N-doped carbon nanosheets supported single-site catalysts stand for the ultimate low-end limit for active metal centers. Besides exposing the largest metal single-atomic sites on external surface, the ideal architecture can shorten diffusion pathway and provide sufficient interface area accessible to reactants, which has been well demonstrated in case of graphene supported single-site catalysts. Furthermore, the large planar area and atomic thickness of the such single-site catalysts make them a model platform to identify the accurate atomistic structure of active sites and the catalytic pathways in heterogeneous catalysis at molecular level.

The selective catalytic oxidation of aromatic inactive C–H bonds is one of the most active research topics but still remains a long-standing challenge. Phenol is the most important industrial chemical due to its broad application in the fields of phenolic resin, pharmaceuticals, agrochemicals, etc. However, current phenol production is still limited to the three-step cumene process, including propylation of benzene, autoxidation to cumene hydroperoxide, and Hock rearrangement. These reactions are conducted under elevated temperature, high pressure, and strongly acidic conditions, which result a low overall yield of phenol (<5%) and an equimolar amount of acetone byproduct. Thus, one-step direct catalytic oxidation of benzene to phenol with hydrogen peroxide over heterogeneous catalysts has remained a focal subject of the fundamental and applied research. Bal and co-workers have devoted great effort in such catalytic system based on novel heterogeneous catalysts including zeolite and CuCr2O4 spinel. Despite considerable achievements, the direct catalytic oxidation of benzene at room temperature is still very difficult due to the notoriously low reactivity of the aromatic C–H bonds.

Developing more efficient catalysts has been the most urgent task. Given that the achieved high catalytic performance in most single-site catalysts, we expect their preparation will provide exciting opportunity in direct catalytic oxidation of benzene to phenol.

Here we report a cocon silk chemistry strategy to prepare isolated metal single-atomic site catalysts embedded in ultrathin 2D porous N-doped carbon nanosheets (M-ISA/CNS; M = Fe, Co, Ni). Their atomistic structures were revealed by spherical aberration correction electron microscopy and X-ray absorption fine structure analysis. The as-synthesized Co-ISA/CNS exhibited a high nitrogen content (9.2 wt %) and large specific surface area (210 m2 g−1). As a proof of concept, the Co-ISA/CNS catalyst showed high activity for C–H bond activation with 68% conversion in the direct catalytic oxidation of benzene to phenol with hydrogen peroxide at room temperature. The density functional theory (DFT) calculations were further performed to investigate the reaction mechanisms of benzene oxidation over the Co-ISA/CNS catalyst. Furthermore, the synthetic strategy demonstrates efficient and general for preparing single-site catalysts, providing opportunities for single-site heterogeneous catalytic applications in atomic scale.

Results

Concept and procedure of cocon silk chemistry strategy. Our cocon silk chemistry synthetic strategy to Co-ISA/CNS consists of four steps: (i) cocoons degumming; (ii) silk fibroin regeneration; (iii) salts-assisted pyrolysis; and (iv) acid etching (Fig. 1a). This synthesis is based on the utilization of amino-group-rich chemistry and lamella-like sheet structure of silk fibroin to anchor single cobalt atom and simultaneously construct 2D morphology. Silk fibroin was first extracted from Bombyx mori cocoons through a modified degumming procedure. The degummed silk fibroin was then dissolved in a concentrated aqueous solution of the cobalt and zinc salt. Driven by self-assembly of the hydrophobic and hydrophilic blocks, the silk fibroin was subsequently restructured into regenerated silk fibroin (Supplementary Fig. 1). The traditional all-aqueous extraction could convert natural silk fiber into lamella-like layer structure with β-sheet crystal contents. The regenerated silk fibroin exhibits a geometrically sheet-like 2D morphology (Supplementary Fig. 2) with a rich chemistry and high nitrogen content of 20.47 wt% (Supplementary Fig. 3). Concurrently, a large amount of cobalt and zinc ions were intercalated into layer space or absorbed on layer surface of regenerated silk fibroin, and tightly anchored into the unique precursor framework by strong interaction with amino groups. The excess metal salts not only promote dissolution of natural silk but also provide a hypersaline conditions and act as effective activation agents or porogens in pyrolysis process. Following a pyrolysis at 900 °C, the layered biopolymer film was in situ converted into 2D porous N-doped carbon nanosheets, meanwhile the rich amino groups were degraded to N species inserting into carbon skeleton to prevent the aggregation and migration of previously incorpo- rated cobalt ions, resulting isolated single cobalt atoms. Finally, Co-ISA/CNS was obtained through acid leaching to remove soluble salts and generated cobalt nanoparticles.

The X-ray powder diffraction pattern (Supplementary Fig. 4a) of Co-ISA/CNS exhibits a broad graphic (002) peak.
Single atoms were circled in red for close observation. Electron microscopy (HAADF-STEM) image (Fig. 1e) shows several Co, N, and C are homogeneously distributed on the whole carbon matrix. Energy-dispersive X-ray spectroscopy (EDX) mapping analysis (Fig. 1d) suggests that no cobalt particles are present. High-angle annular dark-field (HAADF) image (Supplementary Fig. 4f) clearly reveals a transparent feature, which belongs to the fundamental characteristic in common of this type of ultrathin 2D graphene-like nanostructure. Figure 1b and Supplementary Fig. 4f clearly reveals a transparent feature, which belongs to the fundamental characteristic in common of this type of ultrathin 2D graphene-like nanostructure. Additionally, a continuous 2D porous network is clearly observed from the TEM images. The high-resolution TEM (HRTEM) image (Fig. 1c) shows the local atomic structure model of Co-ISA/CNS can be identified as shown by the inset in Fig. 2e, where one Co atom is coordinated by four N atoms (CoN4) configuration, which is evidently different from that of Co foil. The pre-edge peak at 7709.6 eV is caused by a 1s−4p3 transition, indexing the fingerprint of Co−X0 coordination (here X = N). As shown in Fig. 2b, the Co-ISA/CNS exhibits a main peak at 1.41 Å that is shorter than Co-O peak at 1.71 Å in CoO and 1.51 Å in Co3O4, and thus can be ascribed to the backscattering between Co and light atoms, corresponding to the Co−N/C first coordination shell. By contrast, the main peak for Co foil is observed at 2.20 Å, which can be ascribed to the metallic Co−Co coordination. The metallic Co−Co peak for the Co-ISA/CNS is negligible, and no other high shell peaks are detected. This result suggests that the Co atoms in Co-ISA/CNS are atomically dispersed. Additionally, the wavelet transforms (WT) of Co K-edge extended X-ray absorption fine structure (EXAFS) oscillations also well demonstrate the atomic dispersion of Co species in Co-ISA/CNS. As shown in Fig. 2c, the WT contour plots of Co-ISA/CNS present only one intensity maximum at 4.0 Å−1 that can be assigned to the Co−N/C coordination and no intensity maximum related to Co−Co coordination can be observed, compared with that of Co foil, CoO, and Co3O4 references. The coordination configuration of Co atom for Co-ISA/CNS was investigated by quantitative EXAFS curve fitting analysis (Fig. 2d, e). The best-fitting analyses clearly present a main peak originating from Co−N first shell coordination, which is evidently different from that of Co foil (Supplementary Fig. 8). The calculated coordination number of center Co atom is about 4 and the mean bond length of Co-ISA/CNS is 1.98 Å (Supplementary Table 2). So based on the experimental spectra results and the well-matched EXAFS fitting, the local atomic structure model of Co-ISA/CNS can be identified as shown by the inset in Fig. 2e, where one Co atom is coordinated by four N atoms (CoN4) confined in the carbon matrix. However, it is necessary to point out that the pyrolytic carbonaceous materials converted from metal-contained X-ray photoelectron spectroscopy (XPS) exhibits three obvious characteristic peaks of C, N, and O elements as well as a clear high-resolution Co 2p spectrum (Supplementary Fig. 5). The deconvoluted core level spectra of the N 1s region (Supplementary Fig. 5b) consists of two strong peaks at 400.6 and 398.4 eV, corresponding to pyrrolic-N and pyridinic-N, respectively, with N content of 9.2 wt %. Raman spectrum (Supplementary Fig. 6) presents two obvious signals at 1335 and 1593 cm−1, which are related to the D and G bands, respectively. The observed G-band suggests a typical sp2-hybridized carbon network, while the broad D-band with relatively high intensity reveals highly defective structure of Co-ISA/CNS. Additionally, the graphene-like Co-ISA/CNS exhibits high specific surface area with porous structure (Supplementary Fig. 7). The 2D porous nature of Co-ISA/CNS caused a large Brunauer–Emmett–Teller (BET) specific surface area of 2105 m2 g−1, which was comparable to the theoretical value of monolayer graphene (2630 m2 g−1). The dominant pore size distribution is calculated to be about 3.4 nm with a high total pore volume of 1.7 cm3 g−1. Supplementary Fig. 4c, d further display a highly flexible and gauze-like morphology of Co-ISA/CNS. Figure 1b and Supplementary Fig. 4f clearly reveals a transparent feature, which belongs to the fundamental characteristic in common of this type of ultrathin 2D graphene-like nanostructure. Additionally, a continuous 2D porous network is clearly observed from the TEM images. The high-resolution TEM (HRTEM) image (Fig. 1c) reveals some irregular fringes, corresponding few graphene layers. An amount of disordered pores with diameter of 3–4 nm can also be found, and no cobalt particles are present. Energy-dispersive X-ray spectroscopy (EDX) mapping analysis (Fig. 1d) suggests that Co, N, and C are homogeneously distributed on the whole carbon nanosheet. The Co content was determined to be about 0.6 wt % by inductively coupled plasma optical emission spectrometry (ICP-OES) analysis (Supplementary Table 1). The atomically dispersed single Co atoms can be clearly distinguished as the brighter spots on N-doped carbon nanosheet from aberration-corrected high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) image (Fig. 1e). Several Co single atoms were circled in red for close observation.
precursors may usually deliver various metal-nitrogen moieties (MNx, x = 0, 1, 2, 3, and 4) in carbon lattices caused by the disorder, edge, and in-plane atomic vacancy during the thermal treating process at high temperature. Furthermore, it is difficult to completely exclude the existence of other metal-nitrogen moieties based on the current experimental characterization technology. So, we carried out the DFT calculations on formation energies of various possible structures to examine the stability of CoN4 (x = 0, 1, 2, 3, and 4) and to access whether there are CoN3, CoN2, CoN, or CoC existing in the carbon matrix. The DFT calculations were conducted following a modified procedure based on that reported in Deng’s work41, and the detailed methods and results are summarized in Supplementary Fig. 9. After comparing the formation energies of different structures, it is evidently found that the CoN4 moiety is the most stable compound among MNx structures. Therefore, we propose that the CoN4 moieties are the most possible atomic structures for Co species and serve as the dominant active sites in Co-ISA/CNS catalyst.

By skillfully taking advantage of amino-group-rich chemistry and lamella-like sheet structure of silk fibroin, the Co-ISA/CNS catalysts are successfully synthesized for the first time. It is found that the regenerated silk fibroin precursor plays a crucial role in the formation of the sheet-like 2D structure. When replacing silk fibroin precursor by melamine, the identical treatments result into a hybrid of tube-like 1D carbon microstructure and Co nanoparticles (Supplementary Fig. 10), in which the Co nanoparticles are tightly encapsulated by multiple carbon layers and could survive in acid leaching process. However, for regenerated silk fibroin precursor, their lamella-like layer 2D backbone could be maintained in pyrolysis process. The reduced Co nanoparticles are anchored on surface of converted carbon layer and could be removed by acid leaching.

Furthermore, we also found that the cocoon silk chemistry strategy proves versatile in synthesis of Co single-site catalysts with tailorable central-metal atomic local structures. Our previously published researches have demonstrated that a certain amount of coordination bonds between N and metal atoms in pyrolytic carbon matrix can be broken up with pyrolysis temperature increasing60. Here, we synthesized different samples at much higher pyrolysis temperature. Microstructure characterizations and XAFS analysis (Supplementary Fig. 11, 12, 13, 14) suggest that both the samples exhibit a similar morphology of the
ultrathin 2D porous N-doped carbon nanosheets with atomically dispersed Co sites. The dominant reactive sites for Co single-site catalysts obtained at 950 and 1000 °C are identified as CoN3 and CoN2, designated as Co-ISA/CNS-3 and Co-ISA/CNS-2, respectively.

Additionally, the present cocoon silk chemistry strategy is also versatile in synthesis of porous carbon materials with ultrahigh specific surface area. In general, high surface areas and large pore volumes are highly required for heterogeneous catalysts. Chemical activation followed by carbonization process has been extensively employed to achieve such materials using different chemical activating reagents, typically KOH, H3PO4, or ZnCl2, etc.61,62. The ZnCl2 activation approach can enjoy many advantages of low cost, and minor erosion. During carbonization process, the ZnCl2 salt shows strong interaction or coordination with the carbon π-electron organism. At elevated temperature beyond the boiling point of ZnCl2 (756 °C)62, it can be evaporated away completely, where etching effect toward C−C bonds takes place, leaving numerous carbon atom vacancy and further evolving into abundant nanopores. So it can be concluded that the quantity of ZnCl2 salt used in synthesis has crucial impacts upon the as-synthesized carbon texture. Supplementary Fig. 15, 16 show the microstructure characterizations of the Co-ISA/CNS catalysts synthesized with reduced amount of with ZnCl2 salt (5.5 mmol) and in the absence of ZnCl2 salt, respectively. Although the typical ultrathin 2D morphology can be well maintained, both the samples exhibit much less nanopores. Especially, the Co-ISA/CNS catalysts synthesized without ZnCl2 salt show a relatively smooth surface, indicative of disappearing of in-plane porous structure. The BET surface areas of Co-ISA/CNS catalysts synthesized with 5.5 mmol and without ZnCl2 salt are measured to be 1781 and 1061 m² g⁻¹ (Supplementary Fig. 17).

Considering the capability of a series of metal salt to dissolve silk fibroin in aqueous solution, we extended this strategy to other metal precursors and successfully synthesized a family of analogous M-ISA/CNS (M = Fe, Ni) under identical experimental conditions. As shown in Fig. 3a–d and Supplementary Fig. 18, 19, they all exhibit a similar 2D porous sheet-like morphology with a highly flexible and transparent feature. Figure 3b, e reveal that the corresponding Fe and Ni elements are homogeneously distributed on the whole ultrathin 2D porous N-doped carbon nanosheets, respectively. Aberration-corrected HAADF-STEM images (Fig. 3c–f) and XAFS (Supplementary Fig. 20, 21) further demonstrate the feature of isolated metal single-atomic sites anchored carbon matrix. Therefore, this silk fibroin strategy is very effective and versatile for fabrication of isolated metal single-atomic sites supported on 2D porous carbon matrix, which may open up more opportunities for single-site catalysts in broad applications.

**Catalytic performance for C−H bond activation.** Our catalytic measurements show that the as-synthesized Co-ISA/CNS catalysts could exhibit high activity for the direct catalytic oxidation of benzene to phenol with hydrogen peroxide at room temperature (25 °C). Figure 4a presents a comparison of benzene oxidation catalyzed by Co-ISA/CNS, cobalt phthalocyanine (CoPc), Co nanoparticles/N-doped carbon (Co-NPs), and N-doped carbon (NC). When the reaction is conducted over pure NC (Supplementary Fig. 22), only 0.3% benzene conversion was achieved. Even if loading some cobalt species on NC support in nanoparticles form (Supplementary Fig. 23), only 0.8% benzene conversion is detected. However, when constructing Co-ISA/CNS catalysts, the benzene conversion is promoted to be 68% with 61% phenol yields. To the best of our knowledge, the presented catalytic performance is superior to the most results of heterogeneous catalysts reported in literature.53,54,57,63,64. For comparison, the homogeneous CoPc catalysts exhibit benzene conversion of 1%. The Co-ISA/CNS catalysts show a turnover frequency of 150 h⁻¹ for benzene conversion within the initial 1 h (Supplementary Fig. 24). Moreover, the phenol selectivity over

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**Fig. 3** Microstructure and morphology characterizations of Fe-ISA/CNS and Ni-ISA/CNS. a–d Typical TEM images. Scale bar, 500 and 200 nm. b–e Corresponding EDX elemental mapping. c–f Aberration-corrected HAADF-STEM images. Scale bar, 2 nm.
the Co-ISA/CNS catalysts reaches 97% within the initial 1 h, but decreases progressively with increasing the reaction time and maintains at a value of 90% at the reaction time prolonged to 96 h. The side-product is detected to be hydroquinone, which is over-oxidized from benzene through multiple oxidations during the liquid phase reaction process and similar to that reported in literature63,64. Additionally, as shown in Supplementary Fig. 25, the hydrogen peroxide efficiency over the Co-ISA/CNS catalysts increases from 1.1% to 4.7% in the catalytic reaction process, which is related to the H2O2 decomposition rate and the benzene oxidation rate. It is therefore evident that the Co-ISA/CNS can serve as a new class of fascinating catalyst for efficiently catalytic oxidation of benzene to phenol with hydrogen peroxide at room temperature.

Bal’s work demonstrated that several reaction parameters, such as reaction temperature, composition of reaction medium, amount of oxidant, and type of catalyst, may have important effect on the benzene oxidation reaction in liquid phase over heterogeneous catalysts65. Supplementary Fig. 26 suggests that the benzene oxidation reaction over the Co-ISA/CNS catalyst is highly dependent on reaction temperature. When the reaction carried out at 0 °C, the Co-ISA/CNS catalyst can deliver 14.3% benzene conversion after 24 h, which is higher than that 8.3% of Fe-based catalyst under the identical conditions18, indicative of excellent catalytic performance. With increasing reaction temperature, both benzene conversion and phenol yields increased, but phenol selectivity decreased slightly. Supplementary Fig. 27 shows the effect of the amount of H2O2 in benzene oxidation over the Co-ISA/CNS catalyst. It can be observed that increasing the amount of H2O2 results in higher conversion of benzene. However, when the amount of H2O2 was further increased to 7 ml, the benzene conversion exhibited a little decrease. Supplementary Fig. 28 shows that solvents can also play a very critical role in benzene oxidation over the Co-ISA/CNS catalyst. Among the solvents, the acetonitrile proves to be the best reaction medium mainly due to its excellent capability to well dissolve substrate benzene and offer a highly dispersed system for solid catalyst. When reacted in the highly hydrophobic n-octane, the Co-ISA/CNS catalyst showed the worse activity. In order to insight into the structure-activity relationship of the as-synthesized Co single-site catalyst towards the benzene oxidation reaction, we also measured the catalytic performance of another two samples with different N coordination numbers under the same conditions. As revealed in Supplementary Fig. 29, the Co-ISA/CNS catalyst with CoN4 structure achieved the highest catalytic activity in comparison with Co-ISA/CNS-3 (CoN3) and Co-ISA/CNS-2 (CoN2). The resultant difference in catalytic activity is speculated to be related to their structural stability in practice reaction conditions. In case of the most stable CoN4 structure, the sufficient reactive centers can be effectively build up and maintain the longest lifetime, which could endow the highest catalytic activity. In transition-metal-catalyzed C–H bond activation, the generated metal sites play a key role in achieving desirable catalytic activity. To clarify the effect of metal sites toward catalytic activity, Fe-ISA/CNS and Ni-ISA/CNS catalysts were also used for the benzene oxidation reaction. As shown in

Fig. 4 Catalytic performance and proposed mechanism. a A comparison of benzene oxidation catalyzed by the Co-ISA/CNS catalyst, CoPc, Co-NPs, and NC. b Recycle performance of Co-ISA/CNS catalyst. c, d Proposed mechanism of H2O2 activation and benzene oxidation by DFT calculations over Co-ISA/CNS catalyst and Co-NPs, respectively.
Supplementary Fig. 30, Fe-ISA/CNS and Ni-ISA/CNS catalysts are found to be much less active than Co-ISA/CNS catalyst, indicative of the positive role of Co sites.

In addition to the remarkable catalytic activity, the high recycling stability and reusability of the Co-ISA/CNS catalyst are also an advantage worth addressing. As shown in Fig. 4b, the Co-ISA/CNS catalyst could be easily recycled by centrifugation from the solution and reused at least five times without obvious loss of activity and selectivity. The microstructure and morphology characterizations (Supplementary Fig. 31), as well as EXAFS spectrum (Supplementary Fig. 32) of recycled Co-ISA/CNS catalyst show that their structural integrity and stability are still maintained unchanged. The atomic dispersion of Co species is well preserved after repetitive runs, and no aggregation is observed, unambiguously demonstrating the excellent stability and recycling ability of the Co-ISA/CNS catalyst.

To get more insight into the high catalytic activity of the Co-ISA/CNS catalysts, we performed comparative DFT calculations on the reaction mechanisms of benzene oxidation over Co-ISA/CNS, CoPc, and Co nanoparticles (Co cluster). Figure 4c shows the free energy profile and reaction pathway of benzene oxidation on the Co-ISA/CNS catalyst. Generally, two key steps can be abstracted in the reaction pathway including the initial formation of activated oxygen species by the H$_2$O$_2$ decomposition and the subsequent oxidation of benzene to phenol by the activated oxygen species. The catalytic reaction process originates from the adsorption of oxidant H$_2$O$_2$ molecule on the single Co atomic site. Three adsorption models were proposed, and the corresponding adsorption energy is shown in Supplementary Fig. 33. Interestingly, compared with model $a$ and $b$, here it shows that model $c$ exhibited the lowest total energy (E$_{total}$), indicating the most stable structure. The result suggests that the H$_2$O$_2$ adsorption on the single Co atomic site followed the pathway shown in model $c$, i.e., an oxidant H$_2$O$_2$ molecule was initially adopted on the single Co atomic site and then easily dissociated by forming a Co=O intermediate and releasing one water molecule. After that, another H$_2$O$_2$ molecule was continuously adopted and dissociated on the other side of the Co atomic site, and then an O=Co=O center was established. This process is energetically favorable due to preceding an exothermic step and the moderate energy barrier of 0.86 eV. The generated O=Co=O center could present activated oxygen species to adopt the benzene molecule via the C=O bonding by overcoming an energy barrier of 0.77 eV. The adsorbed benzene was subsequently converted to phenol via an H atom transfer from adjacent C to O with an energy barrier of 0.77 eV. After releasing the phenol from active site, the Co=O site was regenerated in the end of one reaction cycle. The energy barriers to overcome for the three transition states (TS1, TS2, and TS3) are relative low, and the whole reaction pathway was calculated to be exothermic, which are favorable for low temperature reactions.

Previous computational and experimental studies have evidently revealed that the high-valent metal–oxo species generally serve as reactive intermediates in the catalytic oxidation of organic substrates. In particular, for the cobalt-mediated C–H oxidation reactions, the reactive intermediates are frequently implicated to covalent (IV)–oxo species. However, it is difficult to identify the proposed covalent (IV)–oxo intermediates under catalytic conditions, and only extremely rare cases have achieved some direct spectroscopic characterizations. To investigate both coordination environment and oxidation state of cobalt sites in the used Co-ISA/CNS catalysts, the cobalt K-edge XAFS measurements were first performed. As shown in Supplementary Fig. 32, the edge position was found to locate at 7721.64 eV at $\alpha = 0.5$ edge-jump, which is ~0.84 eV blue-shifted from the reported Co(III)–peroxo complexes. Although the observed energy shift is smaller than that valence change from Co(III) to Co(IV), it is still indicative of highly oxidized cobalt sites. Additionally, the used Co-ISA/CNS catalysts delivered higher white line intensity than that of the fresh Co-ISA/CNS catalysts, demonstrating the more positive average oxidation state of cobalt sites in the used Co-ISA/CNS catalysts. The EXAFS curves of the used Co-ISA/CNS catalysts were best modeled with cobalt sites in an about five number N/O coordination environment. These XAFS results strongly indicate that the cobalt active sites in the Co-ISA/CNS catalysts are generated as a square pyramidal Co$^{IV}$–O–O-like species. This conclusion is also well evidenced by the electron paramagnetic resonance (EPR) spectroscopy and XPS data analysis of the Co-ISA/CNS samples before and after the H$_2$O$_2$ treatment. As shown in Supplementary Fig. 34a, EPR spectrum of the Co-ISA/CNS catalyst with H$_2$O$_2$ treatment shows signals at $g = 6.6$ and 2.01, indicating the spin-quartet ($S = 3/2$) ground state. The enhanced EPR signals could be ascribed to the probable formation of Co = O/O = Co = O, leading to Co 3d mixing with O 2p. The interaction between Co atomic sites and activated oxygen species will enhance the unoccupied state of Co species because of the electronegativity of cobalt-bonding O atoms. Supplementary Fig. 34b presents the binding energy variety of Co 2p from the fresh, H$_2$O$_2$–treated and spent Co-ISA/CNS catalysts based on XPS analysis. All spectra exhibit one pair of peaks relative to Co 2p$_{3/2}$ and Co 2p$_{1/2}$. The Co 2p$_{3/2}$ of the fresh Co-ISA/CNS catalysts occurs at 780.8 eV, which is slightly higher than that of Co$^{II}$ but smaller than Co$^{III}$ chemical state. Interestingly, the Co 2p$_{3/2}$ displays a positive energy shift of ~1.1 eV after H$_2$O$_2$ treatment, which is higher than the value of Co$^{III}$ chemical state in literature. Thus, our current findings should support the involvement of the O = Co = O center as the most plausible reactive intermediate in this cobalt-catalyzed oxidation reaction.

For Co nanoparticles, the adopted H$_2$O$_2$ was dissociated into two hydroxyls adsorbing on two adjacent Co atoms (Fig. 4d). After that, one hydroxyl shifted to the other for interaction by releasing one water molecule, this reaction relates a transition state and needs to overcome an energy barrier of 0.82 eV (Supplementary Fig. 35). Undergoing additional elementary reactions, the O = Co = O center on Co nanoparticles was established. However, the subsequent oxidation reaction toward benzene ring is highly endothermic. Additionally, much higher energy barrier over 1.0 eV is required for the last two steps. These results make the benzene oxidation hard to take place at low temperature over Co nanoparticles. Supplementary Fig. 36 shows that benzene oxidation over CoPc undergoes a similar reaction pathway to that on Co-ISA/CNS, but the H$_2$O$_2$ molecule was initially dissociated on Co site forming an OH-Co-OH intermediate. In the H$_2$O$_2$–H$_2$O–CH$_3$CN system, the OH-Co-OH intermediate was easily destroyed for the observation of severely bubbling in the practical oxidation reaction, indicative of a strong tendency to self-decomposition of hydrogen peroxide rather than contributing to hydroxylation reaction, which may break off the following processes and result into the deactivation of Co site. Therefore, the high activity for benzene oxidation to phenol over Co-ISA/CNS catalyst could be ascribed to the intrinsic coordination environment and the atomic dispersion of Co species in Co-ISA/CNS, which provides a novelty in reaction pathway much more superior to that over traditional heterogeneous catalysts.

**Discussion**

In summary, we report a general cocoon silk chemistry strategy to synthesize metal single-site catalysts embedded in 2D porous N-doped carbon nanosheets (M-ISA/CNS, M = Fe, Co, Ni). This synthesis is based on the utilization of amino-group-rich
chemistry and lamella-like sheet structure of silk fibroin. With the assistance of metal salts in solution and pyrolysis process, silk fibroin is transformed into 2D porous N-doped carbon nanoshells for anchoring metal single-atomic sites. The X-ray absorption fine structure analysis and spherical aberration correction electron microscopy demonstrate an atomic dispersion of metal species on carbon matrix. Our experimental measurements show a surprising transformation in activity for direct catalytic oxidation of benzene to phenol at room temperature from inert state to high activity over Co species in the form of metal nanoparticle and isolated single metal atomic sites. DFT calculations discover that the high activity could be attributed to the generated O=C=O center intermediates on single Co sites. Our results offer a reliable approach for configuring metal single-atomic site catalysts and more opportunities for heterogeneous catalytic applications beyond the C-H bond activation in atomic scale.

Methods

Synthesis of Co-ISA/CNS. All silk fibroin used in this work were first extracted from Bombyx mori cocoons through a modified degumming procedure reported elsewhere. 6.9 g of cocoons were boiled for 3 h in a 0.2 M Na2CO3 aqueous solution. The resulting silk fibroin suspension was then rinsed thoroughly with water and purified by centrifugation at 14,000 rpm over two 10-min-long periods to remove the glue-like sericin proteins, and then the degummed silk fibroin was obtained. The degummed silk fibroin was subsequently dissolved in 100 mL aqueous solution of 0.25 M CoCl2 and 0.11 M ZnCl2 under continuously stirring at 80 °C for 5 h, yielding a degummed silk fibroin solution. Subsequently, the solution was evaporated and transferred into a solid mixture. The pyrolysis process was conducted in a tubular furnace under an Ar atmosphere with a heating rate of 3 °C min⁻¹ up to 900 °C for 1 h. After allowing the system naturally cooled down to room temperature, the resulting dark solid powder was transferred into a 1 M HCl solution and refluxed at 120 °C for 12 h to remove soluble salts and generated cobalt nanoparticles. Finally, the Co-ISA/CNS was collected by centrifugation and then dried in oven overnight after washing by distilled water and anhydrous ethanol for several times.

Catalyst evaluation. The catalytic performances were evaluated through a modified process according to the previously published work and the recent publication from our group. Unless otherwise noted, the catalytic benzene oxidation evaluation was carried out as following: the reaction was conducted in a 25 ml sealed glass reactor with 50 mg catalyst, 0.4 ml benzene, 6 ml H2O2 (30%), and 3 ml CH3CN at room temperature (25 °C). After the reaction, the mixture was extracted with ethyl acetate, while the solid catalyst was recovered by centrifugation. The products were analyzed with GC–MS and GC with n-hexadecane as an internal standard. General procedure for recyclability test: After each reaction, the catalyst was separated from the reaction system by centrifugal separator (HERMLE 232 HK) at a 18,400× g spin rate for 10 min. Then the collected catalyst was washed by water and ethanol three times, and then dried at 80 °C for 12 h. The obtained powder was submitted to the next batch of catalytic reaction. The yield of phenol was calculated as (mole of formed phenol)/(mole of initial benzene) × 100%. The selectivity of phenol was calculated as (mole of formed phenol)/(mole of formed phenol + mole of formed benzoquinone) × 100%.

Data availability

The data supporting this study are available from the authors on reasonable request.

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References

1. Yang, X. F. et al. Single-atom catalysts: a new frontier in heterogeneous catalysis. Acc. Chem. Res. 46, 1740–1748 (2013).
2. Liu, T. et al. Photocatalytic route for synthesizing atomically dispersed palladium catalysts. Science 352, 797–800 (2016).
3. Qiao, B. et al. Single-atom catalysis of CO oxidation using Pt3/FeO4. Nat. Chem. 3, 634–641 (2011).
4. Zhang, M. et al. Metal/hydr/oxydes/polymer core–shell strategy to metal single-atom materials. J. Am. Chem. Soc. 139, 10975–10979 (2017).
5. He, F. et al. Zincum–porphin-based metal–organic framework hollow nanotubes for immobilization of noble-metal single atoms. Angew. Chem. Int. Ed. 130, 3551–3556 (2018).
6. Fang, X. et al. Single Pt atoms confined into a metal–organic framework for efficient photocatalysis. Adv. Mater. 30, 1705112 (2018).
7. Yang, M. et al. Catalytical oxygen evolution of Co(OH)$_2$ catalysts stabilized by alkali ions on zeolites and mesoporous oxides. Science 346, 1498–1501 (2014).
8. Chen, Z. et al. Single-site Au catalyst for silane oxidation with water. Adv. Mater. 30, 1704720 (2018).
9. Chen, Y. et al. Discovering partially charged single-atom Pt for enhanced anti-markovnikov alken hydroisylolation. J. Am. Chem. Soc. 140, 7407–7410 (2018).
10. Cheng, N. et al. Platinum single-atom and cluster catalysis of the hydrogen evolution reaction. Nat. Commun. 7, 13638 (2016).
11. Yan, H. et al. Single-atom Pd/graphene catalyst achieved by atomic layer deposition: remarkable performance in selective hydrogenation of 1,3-butadiene. J. Am. Chem. Soc. 137, 10484–10487 (2015).
12. Chen, Z. et al. A heterogeneous single-atom palladium catalyst surpassing homogeneous systems for Suzuki coupling. Nat. Nanotech 13, 702–707 (2018).
13. Ge, J. et al. Atomically dispersed Ru on ultrathin Pd nanoribbons. J. Am. Chem. Soc. 138, 13850–13853 (2016).
14. Yang, J. et al. Efficient and robust hydrogen evolution: phosphorus nitride inside nanotubes as supports for anchoring single ruthenium sites. Angew. Chem. Int. Ed. 57, 9495–9500 (2018).
15. Lin, J. et al. Remarkable performance of Ir/FeO$_4$ single-atom catalyst in water gas shift reaction. J. Am. Chem. Soc. 135, 15314–15317 (2013).
16. Zhang, Y. et al. Atomic iridium incorporated in cobalt hydroxide for efficient oxygen evolution catalysis in neutral electrolyte. Adv. Mater. 30, 1707522 (2018).
17. Lang, R. et al. Hydroformylation of olefins by a rhodium single-atom catalyst with activity comparable to RhCl(PPh3)$_2$. Angew. Chem. Int. Ed. 55, 16054–16058 (2016).
18. Deng, D. et al. A single iron site confined in a graphene matrix for the catalytic oxidation of benzene at room temperature. Sci. Adv. 1, e1500462 (2015).
19. Chen, P. et al. Atomically dispersed iron–nitrogen species as electrocatalysts for bifunctional oxygen evolution and reduction reactions. Angew. Chem. Int. Ed. 56, 610–614 (2017).
20. Zhang, Z., Sun, J., Wang, F. & Dai, L. Efficient oxygen reduction reaction (ORR) catalysts based on single iron atoms dispersed on a hierarchically structured porous carbon framework. Angew. Chem. Int. Ed. 57, 9038–9043 (2018).
21. Gao, C. et al. Heterogeneous single-atom catalyst for visible-light-driven high-turnover CO$_2$ reduction: the role of electron transfer. Adv. Mater. 30, 1704624 (2018).
22. Fei, H. et al. Atomic cobalt on nitrogen-doped graphene for hydrogen generation. Nat. Commun. 6, 8669 (2015).
23. Fei, H. et al. General synthesis and definitive structural identification of Mn$_2$Co$_x$ single-atom catalysts with tunable catalytic activities. Nat. Catal. 1, 63–72 (2018).
24. Cheng, Y. et al. Atomically dispersed transition metals on carbon nanotubes with ultrahigh loading for selective electrochemical carbon dioxide reduction. Adv. Mater. 30, 1706287 (2018).
25. Xue, Y. et al. Anchoring zero valence single atoms of nickel and iron on graphdiyne for hydrogen evolution. Nat. Commun. 9, 1460 (2018).
26. Yang, Y. et al., N-atoms-coordinated Mn cofacitors within a graphene framework as bioinspired oxygen reduction reaction electrocatalysts. Adv. Mater. 30, e1801732 (2018).
27. Chen, W. et al. Rational design of single molybdenum atoms anchored on N-doped carbon for effective hydrogen evolution reaction. Angew. Chem. Int. Ed. 56, 16086–16090 (2017).
28. Chen, W. et al. Single tungsten atoms supported on MOF-derived N-doped carbon for robust electrochemical hydrogen evolution. Adv. Mater. 30, e1800396 (2018).
29. Zhang, B. et al. Stabilizing a platinum, single-atom catalyst on supported phosphonomolybdic acid without compromising hydrogenation activity. Angew. Chem. Int. Ed. 55, 8319–8323 (2016).
30. Bell, A. T. The impact of nanoscience on heterogeneous catalysis. Science 299, 1688–1691 (2003).
31. Nosaka, C., Lugmair, C. G., Bell, A. T. & Tilley, D. T. Synthesis, characterization, and catalytic performance of single-site iron (III) centers on the surface of SBA-15 silica. J. Am. Chem. Soc. 124, 13194–13203 (2002).
32. Kistler, J. D. et al. A single-site platinum CO oxidation catalyst in zeolite KTL: microscopic and spectroscopic determination of the locations of the platinum atoms. Angew. Chem. Int. Ed. 53, 8904–8907 (2014).
33. Li, Z. et al. Sintering-resistant single-site nickel catalyst supported by metal–organic framework. J. Am. Chem. Soc. 138, 1977–1982 (2016).
34. Zhao, C. et al. Ionic exchange of metal–organic frameworks to access single nickel sites for efficient electroreduction of CO$_2$. J. Am. Chem. Soc. 139, 8078–8081 (2017).
35. Guo, C., Jiao, Y., Wacławek, E. R. & Du, A. Single atom (Pd/Pt) supported on graphitic carbon nitride as an efficient photocatalyst for visible-light reduction of carbon dioxide. J. Am. Chem. Soc. 138, 6292–6297 (2016).
36. Lin, L. et al. Low-temperature hydrogen production from water and methanol using Pt-Ni-Mo catalysts. Nature 544, 80–83 (2017).
37. Zhu, C., Fu, S., Shi, Q., Du, D. & Lin, Y. Single atom electrocatalysts. Angew. Chem. Int. Ed. 56, 13944–13960 (2017).
38. Wang, A., Li, J. & Zhang, T. Heterogeneous single-atom catalysts. Nat. Rev. Chem. 2, 65–81 (2018).
39. Liu, J. et al. High performance platinum single atom electrocatalyst for oxygen reduction reaction. J. Alloy. Compd. 636, 275–281 (2015).
40. Qiao, B. et al. Ultrastable single-atom gold catalysts with strong covalent metal-support interaction (CMSI). Nano Res. 8, 2913–2924 (2015).
41. Cui, X. et al. A graphene composite material with single cobalt active sites: a highly efficient counter electrode for dye-sensitized solar cells. Angew. Chem. Int. Ed. 55, 6708–6712 (2016).
42. Zhang, C. et al. Single-atomic ruthenium catalytic sites on nitrogen-doped graphene for oxygen reduction reaction in acidic medium. ACS Nano 11, 6930–6941 (2017).
43. Acharyya, S. S., Ghosh, S., Adak, S., Sasaki, T. & Bal, R. Facile synthesis of Cu3O4 nanoparticle spinel: a recyclable heterocatalyst for the one pot hydrogenation of benzene. Catal. Sci. Technol. 4, 4323–4241 (2014).
44. Acharyya, S. S. et al. Synergistic effect between ultrasmall Cu(n) oxide and Cu3O4 nanoparticle spinels in selective hydrogenation of benzene to phenol with air as oxidant. ACS Catal. 5, 2850–2858 (2015).
45. Acharyya, S. S., Ghosh, S., Siddiqui, N., Konatasha, L. S. & Bal, R. Cetyl alcohol mediated synthesis of Cu3O4 nanoparticle spinels: a green catalyst for selective oxidation of aromatic C-H bonds with hydrogen peroxide. RSC Adv. 5, 4838–4843 (2015).
46. Ramu, R. et al. Mechanistic study for the selective oxidation of benzene and toluene catalyzed by Fe(CIO4)2 in an H2O–H2O2–CH3CN system. Mol. Catal. A 441, 114–121 (2017).
47. Wei, Q., Fan, H., Qin, F., Ma, Q. & Shen, W. Metal-free honeycomb-like porous carbon as catalyst for direct oxidation of benzene to phenol. Carbon N. Y. 133, 6–13 (2018).
48. Bal, R., Tada, M., Sasaki, T. & Iwasawa, Y. Direct phenol synthesis by selective oxidation of benzene with molecular oxygen on an interstitial-N/Re cluster/zeolite catalyst. Angew. Chem. Int. Ed. 45, 448–452 (2006).
49. Tsuji, T. et al. Specific enhancement of catalytic activity by a dicopper core: selective hydroxylation of benzene to benzene with hydrogen peroxide. Angew. Chem. Int. Ed. 56, 7779–7782 (2017).
50. Morimoto, Y., Bunno, S., Fufjida, N., Sugimoto, H. & Itoh, S. Direct hydroxylation of benzene to phenol using hydrogen peroxide catalyzed by nickel complexes supported by pyridylalkylamine ligands. J. Am. Chem. Soc. 139, 857–861 (2017).
51. Schmidt, R. J. Industrial catalytic processes—phenol production. Appl. Catal. A 280, 89–103 (2005).
52. Chen, X., Zhang, J., Fu, X., Antonietti, M. & Wang, X. Fe-g-C3N4-catalyzed oxidation of benzene to phenol using hydrogen peroxide and visible light. J. Am. Chem. Soc. 131, 11658–11659 (2009).
53. Carreño, L. & Silva, A. R. Selective direct hydroxylation of benzene to phenol with hydrogen peroxide by iron and vanadyl based homogeneous and heterogeneous catalysts. Catal. Sci. Technol. 6, 8166–8176 (2016).
54. Borah, P., Datta, A., Nguyen, K. T. & Zhao, Y. VOPO4–H2O encapsulated in graphene oxide as a heterogeneous catalyst for selective hydroxylation of benzene to phenol. Green. Chem. 18, 397–401 (2016).
55. Muniandy, L., Adam, F., Mohamed, A. R., Iqbal, A. & Rahman, N. R. A. Cu+2 coordinated graphitic carbon nitride (Cu-g-C3N4) nanosheets from melamine for the liquid phase hydroxylation of benzene and VOCs. Appl. Surf. Sci. 398, 43–55 (2017).
56. Acharyya, S. S., Ghosh, S. & Bal, R. Fabrication of three-dimensional (3D) raspberry-like copper chromite spinel catalyst in a facile hydrothermal route and its activity in selective hydroxylation of benzene to phenol. ACS Appl. Mater. Interfaces 6, 14451–14459 (2014).
57. Wang, D., Wang, M. & Li, Z. Fe-based metal–organic frameworks for highly selective photocatalytic benzene hydroxylation to phenol. ACS Catal. 5, 6852–6857 (2015).
58. Balducci, L. et al. Direct oxidation of benzene to phenol with hydrogen peroxide over a modified titanium silicate. Angew. Chem. Int. Ed. 42, 4937–4940 (2003).
59. Cho, S. Y. et al. Carbonization of a stable β-sheet-rich silk protein into a beta-sheets-rich silk protein into a beta-sheets-rich free graphene-based photocatalyst. Energy Environ. Sci. 6, 793–798 (2013).
60. Wang, B. et al. Synthesis and reactivity of a mononuclear non-haem cobalt (IV)-oxo complex. Nat. Commun. 8, 14839 (2017).

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Author contributions

Y.Z. conceived the idea, proposed the strategy, designed and performed the experiment, analyzed the results, and wrote the manuscript. W.S. conducted the DFT calculations. J.L. provided AC HAADF-STEM characterization. W.C., L.Z., and J.D. helped with the XAFS characterization and corresponding data analysis. T.C. and Y.H. assisted in HR-TEM, STEM, and EDX elemental mapping characterizations. I.Z. and M.Z. helped with the catalyst evaluation and corresponding data analysis. C.C. and Q.P. helped with data analyses and discussions. D.W. and Y.L. supervised the project, helped design the experiments, evaluated the data, and wrote the manuscript. The results of the manuscript were discussed by all authors.

Additional information

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