# Binary Atomically Dispersed Metal-Site Catalysts with Core—Shell Nanostructures for O<sub>2</sub> and CO<sub>2</sub> Reduction Reactions

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Engineering atomically dispersed metal site catalysts with controlled local coordination environments and 3D nanostructures effectively improves the catalytic performance for the oxygen reduction reaction (ORR) and the carbon dioxide reduction reaction (CO<sub>2</sub>RR), which are critical for clean energy conversion and chemical production. Herein, an innovative approach for preparing core-shell nanostructured catalysts with different single-metal sites in the core and the shell, respectively, is developed. In particular, as the shell precursors, covalent organic polymers with a thin layered structure that is polymerized in situ and coated on a metal-doped ZIF-derived carbon core are used, followed by a controlled thermal activation. The selective combination and construction of different metal sites increase active site density in the surface layers, promote structural robustness, facilitate mass/charge transfer, and yield a possible synergy of active sites in the core and the shell. The p-FeNC(shell)@CoNC(core), consisting of a polymerized FeTPPCI-derived carbon layer (p-FeNC) on a Co-doped ZIF-derived carbon (CoNC), exhibits remarkable ORR activity and stability in acidic media along with encouraging durability in H2-air fuel cells. Likewise, a p-FeNC(shell)@NiNC(core) catalyst demonstrates outstanding CO2RR activity and stability. Hence, integrating two appropriate single-metal sites in core and shell precursors, respectively, can modulate morphological and catalytic properties for a possible synergy toward different electrocatalysis processes.

# 1. Introduction

dispersed and Atomically nitrogencoordinated single-metal sites in carbon (M-N-C) have emerged as the most promising platinum-group metal (PGM)-free candidates in the field of electrocatalysis.<sup>[1–9]</sup> The unique electronic structure and unsaturated coordination environments of the active sites in M-N-C catalysts have been proven highly active and selective for the oxygen reduction reaction (ORR) and CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR).<sup>[10–15]</sup> However, due to the simplicity and limitation of single-site centers. it is incompetent for addressing and regulating the trade-off between activity and stability toward the ORR.<sup>[16]</sup> For example, highly active single-Fe-site catalysts often deactivate rapidly, especially at the initial stage.<sup>[17]</sup> Previous reports have shown that active site demetallation, carbon corrosion, and H<sub>2</sub>O<sub>2</sub> attack could be primarily responsible for mitigating catalyst stability.<sup>[4,17]</sup>

A catalyst containing two different single-metal sites could expose increased atomic sites and yield a synergistic effect,

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Dr. H. Zhou X-Ray Science Division Argonne National Laboratory Argonne, IL 60439, USA providing a new approach to designing and engineering M–N–C catalysts with optimized morphologies, local geometric structures, and electronic environments.<sup>[18,19]</sup> For example, despite Fe–N–C catalysts exhibiting the most encouraging ORR activity among studied PGM-free catalysts, a serious concern is the Fenton reactions between Fe<sup>2+</sup> and H<sub>2</sub>O<sub>2</sub>, dramatically accelerating the ORR performance degradation.<sup>[5,20–24]</sup> In contrast, as a less active metal for the Fenton reaction, CoN<sub>4</sub> sites can potentially address the severe stability issues, while delivering relatively low kinetic activity in challenging acidic media during the ORR.<sup>[25–29]</sup>

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As another example, single Ni sites convey a high current density for the CO<sub>2</sub>RR toward CO production. However, they suffer from sluggish kinetics of the first proton-coupled electron transfer (i.e., weak adsorption of \*COOH), leading to a high overpotential.<sup>[30]</sup> On the contrary, Fe-N<sub>4</sub> sites have demonstrated low onset potentials for the CO2RR, whereas only maintained moderate activity in a very narrow potential window due to the strong adsorption of \*CO on Fe-N sites,<sup>[31]</sup> resulting in limited CO current density and poor stability.<sup>[32–34]</sup> Therefore, integrating two types of single metal sites in a catalyst would be a promising and feasible strategy to push the performance and durability of PGM-free catalysts beyond their current limitations. However, due to the competitive coordination with N ligands on the same catalyst, it remains a significant challenge to design and integrate atomically dispersed dual-metal sites to catalyze electrochemical reactions effectively.

In this work, we developed a synergistic strategy to design atomically dispersed metal site catalysts with a core—shell nanostructure by allocating different single-metal sites on the core and shell, respectively. The core—shell nanostructure can be tailored by controlling the inner- or outer-layer components for a synergistic effect. In addition, the unique morphological characteristics may be beneficial for increasing the electrochemically accessible active site density and facilitating mass/charge transport during the electrochemical reactions. As the shell precursors, we used covalent organic polymers (COPs) with a thin layered structure obtained by in situ polymerization, which is coated on a metal-doped ZIF-8-derived carbon (MNC), as the core, via a step-by-step self-assembly method, followed by a controlled thermal activation. Carefully modulating the core size and shell thickness in the catalyst significantly increases the exposed density of active sites. Thus, it maximizes the catalytic performance of atomically dispersed M–N–C catalysts for the ORR and  $\rm CO_2 RR$ .

# 2. Results and Discussion

#### 2.1. Catalyst Synthesis, Morphology, and Structure

**Figure 1** shows an effective method for synthesizing core–shell nanostructured catalysts with different atomically dispersed metal sites in the core and shell. First, as the core component, MNC (M = Co, Ni, or Fe) particles with a well-developed porosity and high specific surface area were synthesized via two-stage calcination according to our previous work.<sup>[35]</sup> Next, they were exploited as templates for "in situ polymerization" to synthesize a porphyrin-based shell on their surface. For example, tetraphenylporphyrin iron chloride (FeTPPCI) was chosen as a functional monomer and polymerized on the MNC core using aluminum chloride as a catalyst at 60 °C. A controlled thermal activation was subsequently conducted to obtain core–shell nanostructured catalysts, denoted p-FeNC@MNC, with atomically dispersed Fe sites in the shell and a different metal site (i.e., M) in the MNC core.

The morphological and structural features of the prepared materials were studied with scanning electron microscopy (SEM) and transmission electron microscopy (TEM). For the p-FeNC@CoNC sample, a unique network architecture with granular nodes was formed. The CoNC nanoparticles were randomly anchored and tightly coated with the p-FeNC layer derived from the polymerized FeTPPCl (Figure 2a,b and Figure S1a, S1b, Supporting Information). FeTPPCl was chosen as a novel shell precursor due to its superior properties, that is, well-defined chemical composition, interconnectivity of building units, and versatility in molecular design.<sup>[36,37]</sup> In addition to FeN<sub>4</sub> active sites in the shell layer, the MN4 sites in the core near the outer surface can participate in electrocatalytic reactions.<sup>[38]</sup> On top of the core, a thin shell was formed with uniformly dispersed Fe active sites, which could fully be exposed and easily accessible to the reactants. As a comparison, the heat-treated polymerized FeTPPCl without the CoNC core exhibited severe agglomeration, with uneven micrometer-scale accumulations (Figure S2, Supporting Information). Therefore, as the support



Figure 1. Schematic illustration of the preparation process for the core-shell nanostructured p-FeNC(shell)@MNC(core) (M = Co, Ni, Fe) catalysts.



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**Figure 2.** a,b) SEM, c) TEM, d) high resolution transmission electron microscopy (HRTEM), e) HAADF–STEM image, f) integrated carbon K-edge EELS map, and g) the corresponding p-FeNC@CoNC catalyst spectrum in addition to amorphous and graphitic carbon reference spectra. TEM and HRTEM images of h,i) the p-FeNC@NiNC catalyst and j,k) p-FeNC@FeNC catalyst.

or template, CoNC nanoparticles greatly confined the in situ polymerization of FeTPPCl and effectively restricted the agglomeration during the polymerization and/or the subsequent carbonization process (Figure S3, Supporting Information). In addition, the FeTPPCl-derived shell layer in the presence of the core part could generate increased surface areas and porosities, thus promoting single-site dispersion and sufficient porosities for mass transport.

Our previous density functional theory (DFT) and experimental X-ray absorption spectroscopy (XAS) studies<sup>[39-41]</sup> indicated that, unlike the traditional CoN<sub>4</sub> sites, the CoNC core contained substantial CoN<sub>2+2</sub> sites.<sup>[35]</sup> The single Co site is thermodynamically favorable for the  $4e^-$  or  $2+2e^-$  ORR pathway, which could be beneficial for stability improvement.<sup>[42,43]</sup> As a result, the FeN<sub>4</sub> moieties with high activity in the shell could interact with CoN4 sites located in the core part, likely yielding isolated diatomic Co-Fe metal-nitrogen sites. Therefore, the tailored core-shell structures could modify catalyst morphologies for increasing exposed active site density. Compared with the original shape of the CoNC core, the coating does not cause a significant change in morphologies concerning the shapes and angular edges (Figure 2c,d and Figure S1c, S1d, Supporting Information). The interface between the core and shell parts is apparent. The shell layer is thin ( $\approx$ 10 nm), indicating that the CoNC core support can effectively disperse the p-FeTPPCl shell.

Aberration-corrected high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images

and electron energy-loss spectroscopy (EELS) were used further to reveal the detailed structure and morphology of the p-FeNC@CoNC catalyst. The carbon particles in the catalyst are polyhedrons with an average size of around 130 nm. Unlike the amorphous-dominated carbon structure in traditional Co–N–C catalysts,<sup>[26]</sup> EELS of the carbon K-edge revealed that the catalysts have a graphitic carbon lattice structure (Figure 2e–g and Figure S3, Supporting Information). The graphitic nature of the carbon could facilitate electron transport and enhance corrosion resistance during the electrocatalytic process.

In addition to the p-FeNC@CoNC catalysts, other core-shell nanostructured catalysts with different core metal sites were synthesized using similar procedures. For the p-FeNC@NiNC sample, a well-coated core-shell structure with a medium-thickness shell ( $\approx$ 20 nm) and rounded edges was produced (Figure 2h,i and Figure S4, Supporting Information). Unlike other catalysts, the p-FeNC@FeNC sample displayed significantly different morphology, with the smoothest surface, the thickest shell, and the most obscure boundary between the core and the shell (Figure 2j,k and Figure S5, Supporting Information). Therefore, the morphologies and nanostructures of the catalyst shell are mainly dependent on the metal site of the carbon core. In particular, the Co-doped carbon core possesses the lowest degrees of packing; the Ni-doped carbon forms a medium-thickness shell structure, and the polymerization degree is moderate; the Fe-doped carbon core is the most easily coated, with the thickest shell and the tightest packing. Different metals in the cores likely affect the polymerization degree of the FeTPPCl shell precursors, thus leading to different core—shell structures and morphologies. In addition, in a core—shell structure, the porosity and graphitization degree of the carbon core, which are affected by the doping metal, significantly influence the thickness of the shell and the interaction between the core and shell.

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N2 adsorption-desorption isotherms were conducted to determine the porous features of the different core-shell structured samples (Figure 3a). The template-free p-FeNC sample exhibits a typical type I isotherm attributed to a dominant microporous property with a surface area of  $411.9 \text{ m}^2 \text{ g}^{-1}$ . A remarkable increase in gas uptake at relatively high pressure  $(P/P_0)$ 0.90) reveals the existence of meso- or macropores, likely due to voids and interparticle porosity.<sup>[44]</sup> Similarly, the CoNC core sample also shows a type I isotherm but with a more microporous structure and a higher specific surface area (834.7  $m^2 g^{-1}$ ). After assembling these materials into a core-shell structure, the p-FeNC@CoNC possesses an H3-type hysteresis loop with no major saturation adsorption platform, indicating an irregular pore structure. Its Brunnauer-Emmett-Teller (BET) surface area (493.5  $m^2 g^{-1}$ ) is also decreased relative to the CoNC core, likely due to the coating of the shell layer. Thus, compared with the individual CoNC core and p-FeNC shell, the p-FeNC@CoNC catalyst generated a hierarchical porous morphology with significant microporous/mesoporous structures (Figure 3b). As for desirable single-metal-site catalysts, sufficient micropores can accommodate active sites, meanwhile, mesopores can promote mass diffusion and transport.  $^{[45-47]}$ 

The carbon structures were further studied by Raman spectroscopy (Figure 3c). The prominent peaks at  $\approx$ 1350 and 1590 cm<sup>-1</sup> are designated as the D and G bands, respectively.<sup>[48]</sup> Generally, the intensity ratios of  $I_{\rm D}/I_{\rm G}$  are established to confirm the graphitization degree of carbon.<sup>[49]</sup> The  $I_D/I_G$  ratio value (0.93) of the p-FeNC@CoNC sample is between the singlecomponent p-FeNC (1.06) and CoNC (0.84). The CoNC core provides a higher degree of graphitization. Highly graphitized carbon enhanced electron transport and electrochemical stability but could accommodate less-active metal sites. In contrast, the p-FeNC shell has a relatively low degree of graphitization, exhibiting disordered and amorphous structures, which essentially provide defects to hold dense single-metal sites despite its compromised stability. Therefore, integrating two different carbon structures in a core-shell nanocomposite could be desirable for an optimal activity and stability trade-off.<sup>[50]</sup> In addition, the peaks in the  $2500-3500 \text{ cm}^{-1}$  region suggest ordered structures in the p-FeNC@CoNC sample with inherent characteristics of a 2D layered structure in the p-FeNC shell.

X-ray photoelectron spectroscopy (XPS) elucidated the composition and chemical state of the elements in the surface layers.<sup>[51]</sup> Figure 3d and S6–S8, Supporting Information, show the N 1*s* and other XPS peaks. Tables S1–S3, Supporting Information, summarize the surface elemental composition of different



Figure 3. a) N2 adsorption-desorption isotherms, b) pore size distribution, c) Raman, and d) XPS N 1s analysis for different samples.



catalysts. Compared with the single-component shell or core, the N content in the core-shell nanostructures is slightly reduced due to an additional thermal treatment process (Table S1, Supporting Information). In each of the catalysts, there are four individual peaks, corresponding to the dominant pyridinic N ( $\approx$ 398.5 eV), pyrrolic N ( $\approx$ 399.4 eV), graphitic N ( $\approx$ 401.2 eV), and oxidized N ( $\approx$ 403.5 eV) (Figure 3d). For the p-FeNC@CoNC catalyst, the orbital peaks of the survey spectrum present Co and Fe elements (Figure S8 and Table S1, Supporting Information) and indicate dominant oxidation states of Co and Fe single sites coordinated with N ligands.<sup>[52,53]</sup> The content of graphitic N (51.4%) was found to be increased compared with other individual components (Table S3, Supporting Information), which agrees with the Raman analysis. The graphitized carbon matrix helps enhance carbon corrosion resistance, which is thereby expected to promote catalyst stability.

To further confirm the coordination environment of two different types of metal sites in the core-shell structured p-FeNC@CoNC catalyst, XAS analysis and fitting were conducted (Figure 4).<sup>[54,55]</sup> The Fe K-edge X-ray absorption near-edge structure (XANES) is left of the FePc reference, which indicates that the oxidization state of Fe atoms is lower for the catalyst than for FePc (Figure 4a). Similarly, the Co K-edge XANES confirms that the Co oxidization state in p-FeNC@CoNC is approximately equal to CoPc (Figure 4d). The modeled extended X-ray absorption fine structure (EXAFS) fitting (Figure 4b,e) reveals no metal-metal bonds. The Fourier-transform EXAFS spectra in the R-space of Fe and Co exhibit a primary peak around 1.5 Å, which is attributed to Fe–N and Co–N bonds, respectively. To further confirm FeN4 and CoN4 modification, the modelbased EXAFS fitting was conducted (Figure 4c,f). Compared with the standard FePc and CoPc fitting results (Figure S9

and Table S4, Supporting Information), both Fe and Co in the sample are coordinated with almost four nitrogen atoms. Therefore, the XAS fitting analysis verifies two different single-Fe and single-Co sites coordinated with four N atoms anchored in the shell and the core, respectively.

#### 2.2. Oxygen Reduction Activity and Stability

The ORR activity of the catalysts was evaluated using steady-state staircase linear scanning voltammetry (SCV) measurement in O2-saturated 0.5 M H2SO4 solution on a rotating disk electrode (RDE) (Figure 5a). The metal-free p-NC@NC core-shell structured catalyst, as a control, exhibits poor activity, similar to N-doped carbon catalysts.<sup>[56]</sup> The ORR activity is effectively improved by introducing FeN<sub>4</sub> species on a NC core, revealing FeN4 that sites at the surface are the dominantly active sites for the ORR (Figure S10, Supporting Information). The polymerization of FeTPPCl monomers could avoid metal agglomeration during carbonization, favorable for forming dense FeN<sub>4</sub> active sites. The introduction of the NC support derived from ZIF-8 nanocrystals could further expose more accessible active sites. After further introducing CoN<sub>4</sub> into the core, the core-shell nanostructured p-FeNC@CoNC catalyst only exhibits marginal enhancement of the ORR activity, compared with the FeNC@NC sample with dominant FeN4 sites in the shell. Importantly, these additional CoN<sub>4</sub> sites in the core could remarkably enhance the catalytic stability, as verified by various accelerated stability tests (ASTs).

In particular, the p-FeNC@CoNC catalyst delivers the best ORR catalytic activity, with an onset potential ( $E_{\text{onset}}$ ) of 0.98 V and a half-wave potential ( $E_{1/2}$ ) of 0.87 V versus reversible hydrogen electrode (RHE) (0.6 mg cm<sup>-2</sup> loading; 25 °C; 900 rpm in



**Figure 4.** a-c) Fe and d-f) Co K-edge XANES spectra, fit of the R-space EXAFS, and the Fourier-transform R-space fitting for the sample p-FeNC@CoNC (denoted as FeCo), respectively.



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**Figure 5.** The ORR performance in 0.5 M  $H_2SO_4$  solution. a) Steady-state ORR polarization plots, and b)  $H_2O_2$  yield and electron transfer number of different catalysts at 25 °C. c) Half-wave potentials of shell@core structured catalysts with different metal elements. Potential cycling during low-potential range (0.6–1.0 V vs RHE) for d) p-FeNC@CoNC, e) p-FeNC@FeNC at 25 °C, and f) p-FeNC@CoNC at 60 °C in O<sub>2</sub>-saturated 0.5 M  $H_2SO_4$  solution. g) Potential cycling during the high-potential range (1.0–1.5 V vs RHE) for the p-FeNC@CoNC catalyst. h) Potentiostatic tests at 0.7 V versus RHE and i) 20 h chronoamperometry tests at constant potentials of 0.85 V versus RHE for the p-FeNC@CoNC catalyst.

O<sub>2</sub>-saturated 0.5 M H<sub>2</sub>SO<sub>4</sub> solution), representing one of the highest RDE activities under similar experimental conditions for PGM-free M–N–C catalysts (Table S5, Supporting Information). Furthermore, the core–shell nanostructured catalyst shows a lower H<sub>2</sub>O<sub>2</sub> yield and higher selectivity toward the fourelectron ORR pathway than the single-component catalysts (Figure 5b). Therefore, the deliberate selection of auxiliary polymerization and core supports is crucial for populating accessible active sites and promoting catalytic activity. We found multiple factors are critical to overall ORR activity during catalyst optimization, including metal content, shell-to-core thickness ratio, core sizes, heating temperatures, heating procedures, and core–shell structural effects. They have been carefully studied, and the relevant results are shown in Figure S11–S12, Supporting Information. We also found that the unique core–shell synthesis method is

universal and effective in synthesizing other dual-metal-site catalysts (Figure 5c and S13–S14, Supporting Information).

Catalyst stability was further studied using extensive AST protocols. After 30 000 potential cycles (0.6–1.0 V vs RHE), the p-FeNC@CoNC catalyst only lost 16 mV of  $E_{1/2}$  (Figure 5d). Unlike the continuous activity loss observed in the p-FeNC@FeNC catalyst with a similar initial activity tested under the identical protocol (Figure 5e), the p-FeNC@CoNC catalyst conveys remarkable stability after the initial degradation and better ORR activity after the AST. Previous DFT calculations have elucidated that, after the adsorption of molecular and atomic oxygen under an acidic environment, the Fe-containing center became less stable.<sup>[16]</sup> However, the structural stability of CoN<sub>4</sub> sites is improved according to the calculations of the ion-exchange energy landscapes on single-Fe-site and



single-Co-site models.<sup>[57]</sup> Furthermore, a single Co atom anchored in a carbon matrix would be hard to be demetallated, thereby mitigating the active sites' loss and enhancing catalyst stability. Also, even under a harsher condition at a high temperature (60 °C), the catalyst can maintain relatively high activity ( $\Delta E_{1/2} = 33 \text{ mV}$ ) after 10 000 cycles, indicating promising stability under similar fuel cell conditions (Figure 5f). The corrosion resistance of carbon in the p-FeNC@CoNC catalyst was further assessed by cycling at a high potential range (1.0–1.5 V vs RHE) in a N<sub>2</sub>-saturated 0.5 M H<sub>2</sub>SO<sub>4</sub> solution. After 5000 cycles, the degradation of the catalyst is 25 mV loss in  $E_{1/2}$  (Figure 5g). The remarkable carbon corrosion resistance in the catalyst could be attributed to the high degree of graphitization in the core part.

Moreover, constant potential tests were conducted at potentials of 0.70 and 0.85 V versus RHE in an O2-saturated 0.5 M H<sub>2</sub>SO<sub>4</sub> solution. As a result, the p-FeNC@CoNC catalyst conveys excellent stability due to an insignificant  $E_{1/2}$  (14 mV) loss at 0.70 V after 50 h (Figure 5h and Figure S15, Supporting Information). At 0.85 V, ORR polarization plots were recorded every 5 h, followed by different potential cycles (0-1.0 V vs)RHE) (Figure 5i). During the stability test, the catalyst retains nearly 80% initial current density after 20 h. It presents a partially reversible activity recovery feature, likely due to removing the oxygen functional groups at the catalyst surface through the wide range of potential cycling. However, the overall degradation rates remain unchanged, likely resulting from inevitable demetallization. Overall, benefiting from the advantages of the two types of single-metal sites in the core-shell structure, the p-FeNC@CoNC catalyst provides excellent promise to balance activity and stability trade-off.

The best-performing p-FeNC@CoNC catalyst was further evaluated as a cathode in membrane electrode assemblies (MEAs) (Figure S16, Supporting Information). Under 1.0 bar H<sub>2</sub>-air condition, the open-circuit voltage (OCV) of the p-FeNC@CoNC cathode (0.95 V) is higher than that of the p-FeNC@FeNC cathode (0.90 V). As a result, the p-FeNC@CoNC-based MEA reached a critical current density of 46.4 mA cm<sup>-2</sup> at 0.8 V, which is better than p-FeNC@FeNC (27.0 mA cm<sup>-2</sup>). However, the maximum power density ( $P_{max}$ ) of the p-FeNC@CoNC cathode is  $0.30 \text{ W cm}^{-2}$ , lower than recent Fe/Co-N-C catalysts.<sup>[58,59]</sup> This is likely due to the dense surface of materials, along with the inevitable agglomeration of particles during the MEA preparation process, leading to uneven dispersion of the ionomers and reducing catalyst utilization, which would not be favorable for mass transfer.<sup>[60]</sup> Given the auspicious intrinsic activity studied in an aqueous solution, further MEA design and fabrication optimization are required to achieve outstanding fuel cell performance. However, it should be noted that the MEA yielded a significant enhancement in power density after voltage cycling from 0.6 to 0.95 V for 1000 cycles under  $H_2/$ air, which indicates that more internal active sites were exposed during voltage cycling, thereby gradually improving the performance. As a comparison, the p-FeNC@FeNC cathode, containing only FeN4 in both shell and core, delivered the inevitable performance degradation in MEAs under the same conditions, further confirming that the alteration of Co sites in the core can result in considerable stability enhancement in challenging acidic media.

#### 2.3. CO<sub>2</sub> Reduction Activity

Besides the challenging ORR, we verified the effectiveness of the unique core—shell nanostructured catalysts toward the  $CO_2RR$  through further regulating different active sites in the shell and core. In this case, Ni sites were successfully introduced into the core rather than Co. As a result, the core—shell structured p-FeNC@NiNC catalyst and relevant control samples were further studied for the  $CO_2RR$  using a three-electrode H-type cell in  $CO_2$ -saturated 0.5 M KHCO<sub>3</sub> solution.

As shown in Figure 6a, the p-FeNC@NiNC catalyst with two metal sites shows the highest total current density (around  $100 \text{ mA cm}^{-2}$  at -1.0 V vs RHE) for the CO<sub>2</sub>RR, significantly exceeding that of catalysts with single-metal sites in traditional Fe-N-C or Ni-N-C catalysts. CO was the primary product for the CO<sub>2</sub>RR catalyzed by the electrocatalysts, as analyzed via online gas chromatography. Faradaic efficiencies (FEs) vary with electrode potentials (Figure 6b). We calculated the partial CO production current densities  $(I_{CO})$  by combining the total current density and the corresponding  $FE_{CO}$  (Figure 6c). The p-FeNC electrocatalyst exhibits an increased FE<sub>CO</sub> from -0.3 to -0.5 V versus RHE, which could be attributed to the activation of CO<sub>2</sub>, and then a dramatic decrease at more negative potentials due to the CO2 transport limitation and kinetic competition between CO<sub>2</sub> reduction and hydrogen evolution.<sup>[61,62]</sup> The previous DFT calculations revealed that FeN4 catalysts have a low onset potential for CO production due to the strong adsorption of the \*COOH intermediate on the FeN<sub>4</sub> site for activating CO<sub>2</sub>. However, the strong \*CO binding energy could also suppress the desorption of CO from the active sites under high overpotentials, leading to a dramatic decrease in both CO production rates and selectivity.<sup>[63–65]</sup> In contrast, due to the unique reaction kinetics, the adsorption of \*COOH on the NiN₄ site is thermodynamically more favorable than H\* at high overpotentials, resulting in high FEs toward the CO<sub>2</sub>RR. Also, the \*CO binding on NiN<sub>4</sub> is relatively weak, which inhibits the further CO conversion, thereby producing a high current density for CO production. However, NiN4 sites suffer from sluggish kinetics of the first proton-coupled electron transfer.<sup>[66]</sup> As a result, the FE<sub>CO</sub> on the NiNC catalyst is not prominent in the low-overpotential region. The FE<sub>CO</sub> increased with a continuously elevated overpotential, and reached a maximum value at a potential of -0.8 V versus RHE. Interestingly, the p-FeNC@NiNC catalyst combines the advantages of the two active sites, that is, simultaneously maintaining excellent FEs in both low- and high-overpotential regions. Specifically, it exhibits high CO selectivity with a maximum FE of  $\approx$ 97% at -0.5 V versus RHE and also dramatically outperforms the catalysts with individual metal sites across the entire potential window from -0.3 to -1.0 V versus RHE, suggesting that Fe/Ni sites synergistically inhibit the hydrogen evolution reaction (HER) at high overpotentials. The p-FeNC@NiNC catalyst demonstrated competitive CO formation over a wide potential range and outperformed most reported CO<sub>2</sub> reduction catalysts (Table S6, Supporting Information). For example, at -0.8 V versus RHE, the J<sub>CO</sub> of p-FeNC@NiNC reaches up to 58.5 mA cm<sup>-2</sup>, representing an improvement factor of 5.6 and 1.7 times compared with the p-FeNC and the NiNC samples, respectively. It should be noted that the



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**Figure 6.**  $CO_2$  reduction performance in a  $CO_2$ -saturated 0.5 M KHCO<sub>3</sub> solution. a) Linear sweep voltammetry (LSV) curves, b)  $FE_{CO}$ , c)  $J_{CO}$ , d) Tafel plots, e) TOF for different catalysts at various potentials, and f) stability test for the p-FeNC@NiNC catalyst at -0.5 V versus RHE.

p-FeNC@CoNC catalyst with FeN<sub>4</sub> and CoN<sub>4</sub> sites exhibits an insufficient activity for the  $CO_2RR$ , further verifying the essential roles of the combination of FeN<sub>4</sub> and NiN<sub>4</sub> species in the shell and core, respectively.

To reveal the reaction kinetics, the corresponding Tafel plots were analyzed (Figure 6d) for the CO<sub>2</sub>RR. The p-FeNC@NiNC catalyst exhibits the lowest Tafel slope of 102 mV dec<sup>-1</sup>, revealing that  $CO_2$  reduction proceeds through a mechanism where the first electron transfer is the rate-determining step.<sup>[67]</sup> Also, it shows a smaller Tafel slope than p-FeNC (131 mV dec<sup>-1</sup>) and NiNC ( $118 \text{ mV dec}^{-1}$ ), evidencing the enhancement effect of dual-Fe/Ni sites. The intrinsic activity is further evaluated by calculating the turnover frequency (TOF) (Figure 6e). The calculated TOFs at -0.9 V versus RHE for the p-FeNC, NiNC, and p-FeNC@NiNC catalysts are 4071, 6980, and 12 058 h<sup>-1</sup>, respectively, representing the superiority of the core-shell structure with dual sites in enhancing the CO<sub>2</sub>RR activity. The TOF value of p-FeNC@NiNC is comparable with the best catalysts for electrochemical CO<sub>2</sub>-to-CO conversion (Table S6, Supporting Information).

For designing effective core—shell electrocatalysts for the CO<sub>2</sub>RR, favorable thermal activation, particle size, feed ratio, and Ni content in the precursor are required. For a more intuitive comparison, Figure S17, Supporting Information, shows the FE<sub>CO</sub> and corresponding  $J_{CO}$  for the p-FeNC@NiNC catalyst, presenting a correlation between the single-factor variable and CO<sub>2</sub>RR performance. Furthermore, the p-FeNC@NiNC catalyst maintains 91% of the initial current density (around 15.3 mA cm<sup>-2</sup>) with 95.5% of the initial FE<sub>CO</sub> after a 40 h stability test at -0.5 V versus RHE (Figure 6f), demonstrating

outstanding electrochemical durability. In contrast, the p-FeNC@FeNC catalyst containing individual FeN<sub>4</sub> sites delivers poor stability with  $\approx$ 37.7% *J*<sub>CO</sub> loss after 12 h of the potentiostatic test (Figure S18, Supporting Information). Thus, further engineering two types of single-metal sites in the core and shell could present great promise to improve the catalyst stability for the CO<sub>2</sub>RR significantly.

## 3. Conclusion

In summary, we developed an effective approach to designing and synthesizing atomically dispersed dual-metal-site electrocatalysts with core-shell structures for the ORR and the CO2RR. The two different MN<sub>4</sub> moieties with sophisticated functionalities were separately distributed in the shell and core regions, avoiding coordination competition due to limited N ligands. The thin shell derived from COPs (e.g., in situ-polymerized FeTPPCl) ensures that highly active FeN4 sites are uniformly distributed in the surface layer with favorable mass and charge transports. As the support or template, ZIF-derived MNC nanoparticles can effectively disperse the in situ-polymerized FeTPPCl, suppress the agglomeration of monomers, and promote the maximized exposure of FeN<sub>4</sub> active sites in the shell. Amorphous carbon in the shell essentially helps provide sufficient defects for hosting dense metal sites on a core with a high degree of graphitization, simultaneously achieving a high density of sites and enhanced stability. In addition, the structure and morphology of the core, which is related to the type of incorporated single-metal sites (e.g., Fe, Co, or Ni), greatly affect the thickness of the shell and the interactions between the core



and shell, therefore governing overall catalytic activity and stability. The unique integration of the shell and core containing various metal sites provides a versatile route to producing highperforming electrocatalysts for different reactions concerning selectivity, activity, and stability.

In particular, a variety of such core—shell structured catalysts were carefully examined for two critical electrocatalysis processes:  $O_2$  and  $CO_2$  reductions. Among others, the p-FeNC(shell)@CoNC(core) catalyst exhibited compelling catalytic properties for the ORR and  $H_2$ —air fuel cell performance. Meanwhile, the p-FeNC(shell)@NiNC(core) catalyst yielded excellent  $CO_2RR$  activity and stability for CO production. Although deeper understanding is needed concerning the possible synergy between single-metal sites in the shell, core, and interfaces, this work opens up enormous possibilities for designing advanced atomically dispersed single-metal-site electrocatalysts for critical energy-conversion electrochemical reactions.

## 4. Experimental Section

Synthesis of CoNC Template: Typically, F127 (1 g) was dissolved in methanol (25 mL) and then mixed thoroughly with methanolic solution (25 mL) of zinc nitrate hexahydrate (2.68 g, 9 mmol) and cobalt (II) nitrate hexahydrate (1.16 mg, 4 mmol) at room temperature. Methanol (50 mL) containing 2-methylimidazole (3.2 g, 39 mmol) was subsequently added to the prior solution under stirring and refluxed at 60 °C for 30 min. After that, pink crystal precipitates were collected by centrifugation in fresh ethanol solution three times and vacuum dried at 60 °C overnight. The CoNC template was obtained by carbonization of the as-prepared sample, the program was at 800 °C for 2 h, followed by 1100 °C for 1 h, and the rate was  $30 \,^{\circ}$ C min<sup>-1</sup>.

Synthesis of p-FeTPPCl@CoNC Precursor: Tetraphenylporphyrin iron chloride (FeTPPCl, 50 mg, Sigma-Aldrich), CoNC (100 mg), and chloroform anhydrous (20 mL) were added to a Schlenk flask and stirred for 3 h at room temperature. Anhydrous aluminum chloride (150 mg) was subsequently added into the above solution under stirring for 2 days at 65 °C. After that, the precipitate was collected by filtering and washed with methanol, dichloromethane, tetrahydrofuran, N,N-dimethylformamide, and acetone, respectively. The powder was further purified by Soxhlet extractions for 24 h with methanol and dichloromethane, respectively. After drying in a vacuum oven at  $60 \,^{\circ}$ C for 12 h, the solid p-FeTPPCl@CoNC was obtained as the precursor for subsequent processes.

Synthesis of p-FeNC@CoNC: The as-obtained p-FeTPPCl@CoNC precursor was treated at 1100 °C for 1 h in an N<sub>2</sub> atmosphere with a quick ramp rate of 30 °C min<sup>-1</sup>.

Characterization: Catalyst morphology was determined using SEM on a Hitachi SU 70 microscope at a working voltage of 5 kV. Aberrationcorrected HAADF and bright-field (BF) STEM images and EELS were acquired on a Nion UltraSTEM in the Center for Nanophase Materials Sciences at Oak Ridge National Laboratory. The instrument was operated with an accelerating voltage of 60 kV and was equipped with a Gatan Enfina EELS spectrometer. XPS was conducted using a Kratos AXIS Ultra DLD XPS system equipped with a hemispherical energy analyzer and a monochromatic Al K $\alpha$  source. The source was operated at 15 keV and 150 W; pass energy was fixed at 40 eV for the high-resolution scans. Determination of catalyst surface area and porosity distribution was achieved using the BET analyses of the N<sub>2</sub> isothermal sorption measurement recorded at 77 K on a Micrometritics TriStar II with samples' degassing condition for 6 h at 150 °C under vacuum. Raman spectra of catalysts coated on glass slides were recorded with a wavelength of 514 nm laser at ambient conditions by the Renishaw Raman system. The Gaussian function was used for the fitting of the spectra. Fe and Co K-edge XAS was measured at beamline 12BM, Advanced Photon Source (APS), Argonne National Laboratory (ANL). Data reduction, data analysis, and EXAFS fitting were conducted with the Athena, Artemis, and  $\ensuremath{\mathsf{IFEFFIT}}$  software packages.

# **Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

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# **Conflict of interest**

The authors declare no conflict of interest.

# Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

# **Keywords**

atomic metal sites,  $\mbox{CO}_2$  reduction, core-shell structures, electrocatalysis, oxygen reduction

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