# **Chemical Science**

## EDGE ARTICLE



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### Introduction

Utilizing renewable electricity to convert  $CO<sub>2</sub>$ , the primary greenhouse gas, into high-value-added fuels and chemicals presents substantial potential within the industrial landscape. Among the numerous electrocatalytic materials displaying high performance for  $CO<sub>2</sub>$  conversion, stable transition metal porphyrin and phthalocyanine molecular complexes have garnered extensive attention for  $CO<sub>2</sub>$  reduction.<sup>1,2</sup> These complexes boast well-defined active sites and adjustable ligand structures, endowing them with distinct advantages in the realm of electrocatalytic  $CO<sub>2</sub>$  reduction.<sup>3,4</sup> However, the reported turnover numbers (TONs) and turnover frequencies (TOFs) for these molecular catalysts remain relatively low, which is far from the requirements of industrial application. Their catalytic selectivity and stability also encounter significant challenges. Therefore, designing and constructing transition metal-based porphyrin or phthalocyanine catalysts that can operate efficiently and reliably under industrial conditions is crucial for achieving the vision of carbon neutrality.

Functional group modification strategies have been successfully employed to enhance the electrocatalytic

## Graphite conjugated nickel phthalocyanine for efficient  $CO<sub>2</sub>$  electroreduction and  $Zn$ – $CO<sub>2</sub>$ batteries†

Jingwei Han,<sup>a</sup> Qi[ang](http://orcid.org/0000-0003-4487-1820) Xu,<sup>a</sup> Fengkun Ti[an,](http://orcid.org/0000-0003-2531-552X)<sup>a</sup> Hai Sun,<sup>a</sup> Yua[nyu](http://orcid.org/0000-0001-7270-3301)an Qi,<sup>a</sup> Guodong Zhang,  $\mathbb{D}^{\mathsf{b}}$  Jun-Sheng Qin  $\mathbb{D}^{\mathsf{a}}$  and Heng Rao  $\mathbb{D}^{\ast\mathsf{a}}$ 

The linking chemistry between molecular catalysts and substrates is a crucial challenge for enhancing electrocatalytic performance. Herein, we elucidate the influence of various immobilization methods of amino-substituted Ni phthalocyanine catalysts on their electrocatalytic  $CO<sub>2</sub>$  reduction reaction (eCO<sub>2</sub>RR) activity. A graphite-conjugated Ni phthalocyanine,  $Ni(NH_2)_8PC-GC$ , demonstrates remarkable electrocatalytic performance both in H-type and flow cells. In situ infrared spectroscopy and theoretical calculations reveal that the graphite conjugation, through strong electronic coupling, increases the electron density of the active site, reduces the adsorption energy barrier of \*COOH, and enhances the catalytic performance. As the cathode catalyst,  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$  also displays remarkable chargedischarge cycle stability of over 50 hours in a  $Zn-CO<sub>2</sub>$  battery. These findings underscore the significance of immobilization methods and highlight the potential for further advancements in  $eCO_2RR$ . **EDGE ARTICLE**<br> **CONFIDENTIFICATION CONTINUE CONJUGATED MICKEL phthalocyanine for**<br>
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performance of molecular catalysts. It has been observed that electron-withdrawing groups facilitate electron transfer by reducing electron density while simultaneously impeding proton transfer. Conversely, electron-donating groups promote proton transfer but increase the energy barrier of electron transfer.<sup>5</sup> Therefore, it is imperative for functional group modification to strike a delicate balance between electronwithdrawing and electron-donating abilities. As a classical electron-donating group, the amino group  $(-NH<sub>2</sub>)$  offers a unique advantage due to its protonation potential, allowing for tunable electron density.<sup>6,7</sup> For instance, Wang and colleagues reported that electron-donating amino substituents on the phthalocyanine ring significantly enhanced the stability of their catalytic system.<sup>8</sup> Zheng and colleagues developed a donor-acceptor-modified copper porphyrin, wherein amino groups acted as electron donors to induce electron transfer to  $CuN<sub>4</sub>$  sites, resulting in outstanding  $CO<sub>2</sub>$ -to-CH<sub>4</sub> electroreduction performance.<sup>9</sup> However, these molecular catalysts were directly dispersed and physically adsorbed onto the surface of the conductive carbon materials, which is easily deactivated and unsatisfied with the conditions for industrial applications.

Heterogenization of molecular catalysts is another enticing option, offering the potential for elevated current density and enhanced stability through the facilitation of charge transfer. Generally, this is realized through weak physical adsorption or electrostatic interactions that promote  $\pi-\pi$  stacking between catalysts and carbon supports.<sup>10</sup> Unfortunately, the noncovalently attached catalysts may encounter challenges, chiefly pertaining to aggregation at high loadings or leaching

<sup>&</sup>lt;sup>a</sup>State Key Laboratory of Inorganic Synthesis and Preparative Chemistry, College of Chemistry, International Center of Future Science, Jilin University, 2699 Qianjin Street, Changchun 130012, P. R. China. E-mail: rao@jlu.edu.cn

b School of Chemistry and Chemical Engineering, Yangzhou University, Siwangting Road 180, Yangzhou, P. R. China

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issues.11,12 These complications are primarily attributed to the self-stacking effect and the frail connection between the molecular catalyst and its supporting substrate, as reflected by their poor long-term current stability.<sup>13,14</sup> Therefore, covalent linkage employs diverse methods, including amide bonds,<sup>15,16</sup> alkynyl linkages,<sup>17</sup> diazo graftings,<sup>18</sup> click reactions,<sup>19</sup> and substitution reactions,<sup>20</sup> to mitigate the prevalent predicament of self-stacking in molecule catalysts. Nevertheless, it is imperative to recognize that the electronic coupling resultant from such tethers may tend to be feeble, thereby introducing a higher barrier for electron transfer. This, in turn, restricts the rate of electron flow to the active site, curtailing overall catalytic efficiency.<sup>21</sup>–<sup>23</sup> The aromatic linkages with highly delocalized electrons also significantly influence the electronic properties of the active site.<sup>24,25</sup> Surendranath and colleagues demonstrated that graphite-conjugated catalysts (GCCs) greatly enhanced the oxygen reduction activity of molecular catalysts in alkaline media.26,27 Furthermore, GCCs containing Re complexes also exhibited efficient and selective reduction of  $CO<sub>2</sub>$  to  $CO$  with quite low overpotential.<sup>28</sup> The fundamental rationale behind this phenomenon lies in the fact that GCCs anchored via conductive pyrazine bonds exhibit the inner-sphere electron transfer (ISET) reaction, akin to metallic active sites.29,30 This strong electronic coupling significantly promotes the electron transfer between the active sites of the molecular catalyst and the carrier. Edge Article<br>
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Herein, we engineer an amino substituted nickel phthalocyanine as transition-metal active sites for  $CO<sub>2</sub>$  reduction on the edge of graphite planes. This was achieved by condensation of an organic 1, 2-diamine with ortho-quinone moieties native to graphitic carbon surfaces to form a conjugated pyrazine linkage. As a result, the eCO<sub>2</sub>RR performance of  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC  $(Ni(NH_2)_8$ Pc =  $(2,3,9,10,16,17,23,24$ -octaaminophthalocyaninato) nickel( $\pi$ ), GC = graphitic carbon) is dramatically improved compared to catalysts prepared via the non-conjugated covalent method or physical mixing method. Encouragingly, the graphite conjugated catalyst exhibits close to 100% CO selectivity over a wide potential window, in both H-type cell and flow cell setups. In the flow cell,  $Ni(NH_2)_8$ Pc-GC showcases turnover frequency for CO (TOF<sub>CO</sub>) exceeding 250 000 h<sup>-1</sup> and half-cell energy conversion efficiencies of 63% at an overpotential of 775 mV. Furthermore, we assemble a  $Zn-CO<sub>2</sub>$  battery utilizing  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$  as the cathode catalyst, which exhibits a peak power density of 0.73 mW cm−<sup>2</sup> and an faradaic efficiency of CO (FE<sub>CO</sub>) of 98% during the discharge process. This provides a promising approach to electric energy output and chemical conversion.

### Results and discussion

#### Synthesis and characterization of catalysts

The synthesis involved the condensation of ophenylenediamine-containing phthalocyanine synthon,  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc, with *o*-quinone moieties present on the edge planes of graphitic carbon using a modified literature procedure (Scheme 1).<sup>27,31</sup> Specifically, after mixing the dimethyl sulfoxide (DMSO) solution of  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc with the ethanol (EtOH)



Scheme 1 Synthesis of (a)  $Ni(NH_2)_8$ Pc-GC and (b)  $Ni(NH_2)_4$ Pc-GC.

solution of graphite carbon in the absence of  $O_2$ , the resulting mixture was reacted at 130 °C for 24 hours under argon protection. The crude product was then washed with DMSO and EtOH to eliminate any unsupported  $Ni(NH_2)_8$ Pc. Subsequently, the powder was treated with 0.1 M hydrochloric acid for 4 hours to hydrolyze the imine bonds formed on the surface. For comparison, we also prepared a nickel phthalocyanine modified graphitic carbon through non-conjugated amide bonding  $(Ni(NH_2)_4$ Pc-GC,  $Ni(NH_2)_4$ Pc = (tetraaminophthalocyaninato)  $nickel(n)$ ). Initially, graphite carbon was treated with a mixed solution of thionyl chloride  $(SOCl<sub>2</sub>)$  and toluene, facilitating the conversion of carboxyl groups (–COOH) on its surface into acid chloride groups (–COOCl). Subsequently, the treated graphitic carbon and  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc were dispersed in a DMSO solution and reacted under an argon atmosphere at 150 °C for 24 hours. Following the reaction, the product was washed with DMSO and EtOH, yielding  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC. The full synthesis details are described in the ESI.†

The prepared graphitic carbon catalysts were washed three times with DMSO to remove physisorbed active sites until the UV-visible spectra showed no absorption peaks corresponding to the Q-band of the nickel phthalocyanines (Fig. S1†). X-ray photoelectron spectroscopy (XPS) reveals the successful incorporation of phthalocyanine units onto the surface of the graphitic carbon substrate (Fig. 1a and b). In the case of  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC, the N 1s spectrum is deconvoluted into three distinct types of N species with a ratio of 4 : 10 : 1, corresponding to N–C (400.2 eV), N=C (398.8 eV), and pyrazine N (399.8 eV). Regarding  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc-GC, the peak centered at 397.5 eV is attributed to -NH-C. Notably, the Ni  $2p_{3/2}$  spectrum of  $Ni(NH<sub>2)</sub><sub>8</sub>PC-GC$  exhibits a shift towards lower binding energy compared to that of  $Ni(NH_2)_4$ Pc-GC. This shift is attributed to the transfer of electrons from the graphitic carbon to the nickel phthalocyanine. These results substantiate the notion of enhanced electron transfer kinetics between conjugated active sites and carbon substrate, which also aligns with the subsequent DFT calculation results. Considered together, the XPS data support the formation of isolated nickel phthalocyanine active sites conjugated to the surface via pyrazine or nonconjugated to the surface via amide linkages. Furthermore, Raman spectra were employed to gain deeper insights into the structure and molecular interaction of the catalyst. As depicted



Fig. 1 (a) N 1s XPS spectra of  $Ni(NH_2)_8$ Pc-GC and  $Ni(NH_2)_4$ Pc-GC. (b) Ni 2p<sub>3/2</sub> XPS spectra of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC and Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC. (c) Raman spectra of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC, Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC and GC. (d) CV curves in Ar-saturated 0.5 M KHCO<sub>3</sub> solution with a scan rate of 10 mV s<sup>-1</sup> of Ni(NH2)8Pc-GC and Ni(NH2)4Pc-GC.

in Fig. 1c, both  $Ni(NH_2)_8$ Pc-GC and  $Ni(NH_2)_4$ Pc-GC display similar Raman signals to graphitic carbon, indicating that the covalent immobilization process preserves the structural integrity of the graphitic carbon. The peak centered at  $\sim$ 1330 cm<sup>-1</sup> is attributed to the D band, the intensity of which is used to quantify defect concentration. The peak around 1590  $\text{cm}^{-1}$  corresponds to the G band, reflecting the degree of graphitization.<sup>32,33</sup> The ratio of the D to G peak intensity  $(I_D/I_G)$ of the molecule-modified carbon material exhibits a slight increase compared to the pristine graphitic carbon. This increase may arise from the destruction of the surface lattice of the carbon material after undergoing molecular covalent grafting. $32,34$  The Raman spectra and X-ray diffraction (XRD) patterns (Fig. S2†) of catalysts are dominated by graphitic carbon, which obscures the weaker nickel phthalocyanine characteristic signals. This observation is related to either a high dispersion or a low molecular loading of phthalocyanine on the graphitic carbon surface.<sup>35</sup>

The concentration of the conjugated Ni centers was quanti fied by cyclic voltammetry (CV) and inductively coupled plasma optical emission spectroscopy (ICP-OES), with detailed methodologies available in the ESI.† Considering the instability of carbon materials and the activation process of the catalyst during potential cycling (Fig. S3†), we selected the stabilized fourth CV curve for integrating to determine the charge passed in the pyrazine wave. This approach enables an in situ and nondestructive measurement of the surface concentration of nickel active sites that are conjugated to the electrode. As shown in Fig. 1d, the broad redox waves centered at 0.15 V vs. RHE are attributed to the  $2e^-/2H^+$  reduction of the surface pyrazine units to dihydropyrazine.<sup>28</sup> The results of ICP-OES revealed that the content of Ni in  $Ni(NH_2)_8$ Pc-GC and  $Ni(NH_2)_4$ Pc-GC was 0.16 wt% and 0.47 wt%, respectively. Following the method described in the literature,<sup>27</sup> the Ni loading on the  $Ni(NH_2)_8$ Pc-GC electrode surface used in the subsequent study was 15.5

nmol cm<sup>-2</sup>. The Ni content in the electrode prepared with  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC was directly derived from the ICP-OES measurements, which was 39.7 nmol  $cm^{-2}$ . Fig. S4, S5, and S6† show the transmission electron microscopy (TEM) images, scanning electron microscopy (SEM) images, and the corresponding element mappings, illustrating the uniform distribution of phthalocyanine molecules on the carbon surface.

#### Electrochemical reduction of  $CO<sub>2</sub>$  in H-type cell

The electrocatalytic  $CO<sub>2</sub>$  reduction performance of the  $Ni(NH<sub>2</sub>)<sub>8</sub>$  Pc-GC and  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc-GC catalysts was initially evaluated in a three-electrode H-type cell with  $CO_2$ -saturated KHCO<sub>3</sub> solution. The linear sweep voltammetry (LSV) curves reveal that both catalysts exhibited higher current densities in the  $CO<sub>2</sub>$ saturated 0.5 M KHCO<sub>3</sub> electrolyte compared to the Ar-saturated electrolyte, indicating that the as-prepared catalysts are active for  $CO<sub>2</sub>RR$  (Fig. 2a). To more accurately assess the intrinsic activity of the catalysts, we also plotted the LSV curves normalized by the amount of substance in a  $CO<sub>2</sub>$  atmosphere (Fig. S7†). The results indicate that  $Ni(NH_2)_8PC-GC$  exhibits superior  $CO_2RR$  activity compared to  $Ni(NH_2)_4$ Pc-GC. Encouraged by the results of the LSV test, a series of controlled potential experiments were performed within the potential range of −0.48 V to −0.93 V vs. RHE. Gaseous and liquid products were analyzed by gas chromatography (GC) and proton nuclear magnetic resonance spectroscopy (<sup>1</sup>H NMR), respectively. The results indicated that only two gaseous products, CO and  $H<sub>2</sub>$ , were generated at all applied potentials, with no liquid products detected. To investigate the origin of CO, we conducted differential electrochemical mass spectrometry (DEMS) analysis. Our findings reveal that the onset potential for the consumption of  $CO<sub>2</sub>$  and the generation of CO is the same, Chemical Science<br>
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Fig. 2 (a) LSV curves in Ar or  $CO_2$ -saturated 0.5 M KHCO<sub>3</sub> solution with a scan rate of 5 mV s<sup>-1</sup> and 80% iR compensation ( $R_s = 7.4$  Ω). (b) In situ DEMS spectra of  $Ni(NH_2)_8$ Pc-GC in a CO<sub>2</sub>-saturated 0.5 M KHCO<sub>3</sub> electrolyte. The signals of CO<sub>2</sub> ( $m/z = 44$ ), CO ( $m/z = 28$ ) and H<sub>2</sub> (m/z = 2) were captured twice of LSV from  $-0.38$  to  $-0.88$  V vs. RHE. The dashed line represents the potential profile. (c) FEs of CO and  $H<sub>2</sub>$  at different applied potentials. The error bars denote the standard deviations (SD) derived from three independent measurements. (d) Long-term stability tests.

indicating that CO is derived from dissolved  $CO<sub>2</sub>$  in the electrolyte (Fig. 2b and S8†).<sup>36</sup> Additionally, the results indicate that  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC exhibits a lower onset potential for CO<sub>2</sub>-to-CO conversion than  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC (Fig. S9†). The DEMS curves demonstrate a more negative onset potential than the LSV curves, which is attributed to the hysteresis effect of the instrument. Notably, the FE<sub>CO</sub> for Ni $(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC achieves nearly 100% in the potential range of −0.68 to −0.88 V vs. RHE (Fig. 2c). The FE<sub>CO</sub> of Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC is only approximately 92%. Moreover, the current density remains stable across all tested potentials (Fig. S10†). According to Fig. S11,† the CO partial current density  $(j_{\text{CO}})$  of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC and Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC follows a volcano-like trend. The maximum  $j_{\text{CO}}$  for Ni(NH<sub>2</sub>)<sub>8</sub>-Pc-GC is 17.6 mA cm<sup>-2</sup> at -0.88 V vs. RHE, whereas Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC achieves 15.2 mA cm<sup>-2</sup> at the same voltage of  $-0.88$  V vs. RHE. The  $TOF_{CO}$  provides further insights into the intrinsic activity of the catalysts. The maximum  $TOF_{CO}$  of  $Ni(NH_2)_8$ Pc-GC reaches 6.3 s<sup> $-1$ </sup> at  $-0.88$  V vs. RHE, which is 3.2 times that of  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc-GC. For comparison, we also evaluated the eCO<sub>2</sub>RR performance of physically mixed catalysts  $(Ni(NH_2)_8Pc/GC$  and  $Ni(NH<sub>2</sub>)<sub>4</sub>PC/GC$ ). In this method, molecular catalysts are anchored on the surface of carbon supports by  $\pi-\pi$  stacking. The results presented in Fig. S12† indicate that both  $Ni(NH_2)_8$ Pc/GC and  $Ni(NH_2)_4$ Pc/GC exhibit activity for eCO<sub>2</sub>RR. The *i-t* curves of  $Ni(NH_2)_8$ Pc/GC and  $Ni(NH_2)_4$ Pc/GC remain steady across all tested potentials (Fig. S13†). However, compared to  $Ni(NH_2)_8$ Pc-GC and  $Ni(NH_2)_4$ Pc-GC, both FE<sub>CO</sub> and  $j_{\text{CO}}$  values are comparatively lower for Ni(NH<sub>2</sub>)<sub>8</sub>Pc/GC and  $Ni(NH<sub>2</sub>)<sub>4</sub>PC/GC$  (Fig. S14 and S15<sup>†</sup>). The TOF<sub>CO</sub> values of  $Ni(NH_2)_8$ Pc/GC and  $Ni(NH_2)_4$ Pc/GC are found to be 1.2 s<sup>-1</sup> and 0.8 s<sup> $-1$ </sup> at  $-0.88$  V vs. RHE, respectively, as depicted in Fig. S16.† These values are significantly lower than that of  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$ . In a word,  $Ni(NH_2)_8$ Pc-GC demonstrates extremely superior CO2RR activity compared with the recently reported Ni-based catalysts (Table S1†). We then investigated the long-term stability of the catalysts at −0.73 V vs. RHE, as revealed in Fig. 2d and S17.†  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC exhibits optimal stability, maintaining a  $FE_{CO}$  of over 95% in 10 hours. Importantly, the morphology of the catalysts remained intact after the long-term electrolysis (Fig. S18†). This signifies that covalent grafting significantly improves the stability of the catalyst, with the conjugated pyrazine linkage being even more effective than the non-conjugated amide linkage. Edge Article Computer Commonstration-NonCommercial

Electrokinetic studies were performed to explore the effect of different immobilization methods on the reaction mechanism. As illustrated in Fig. S19,† the Tafel slope for  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$ , as fitted with the  $j_{\rm CO}$ , is 152 mV dec<sup>-1</sup>, which is lower than that observed for Ni $(\text{NH}_2)_4$ Pc-GC (179 mV dec $^{-1}$ ). In contrast, the Tafel slopes for the physically mixed  $Ni(NH<sub>2</sub>)<sub>8</sub>PC/GC$  and  $Ni(NH<sub>2</sub>)<sub>4</sub>PC/GC$  are much higher, reaching 194 mV dec<sup>-1</sup> and 216 mV dec $^{\rm -1}$ , respectively (Fig. S20†). The reported Tafel slopes for Ni-based catalysts generally fall within the 80–350 mV dec<sup>-1</sup> range.<sup>33,35,37</sup> The lower Tafel slope for  $Ni(NH_2)_8$ Pc-GC indicates faster reaction kinetics for  $CO_2RR$ , enhancing CO formation.<sup>20,38</sup> Additionally, the Nyquist plots of the catalysts were collected in the range of  $10^{-1}$  to  $10^5$  Hz to compare the charge transfer resistance  $(R_{ct})$  at the catalyst interface. The results,

summarized in Fig. S21 and S22,† reveal that the diameters of the semicircles in the high-frequency region for all catalysts are less than 5  $\Omega$ , signifying excellent conductivity of the catalysts. Notably,  $Ni(NH_2)_8$ Pc-GC exhibits the smallest  $R_{ct}$  and the fastest electron transfer kinetics among all samples. To further understand the intrinsic catalytic activity of the electrocatalysts, the electrochemical active surface area (ECSA) evaluated by comparing the double-layer capacitance (Fig. S23†) was investigated. In the relationship between current density and scan rate (Fig. S24 and S25†), the capacitances for  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$ ,  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc-GC,  $Ni(NH<sub>2</sub>)<sub>8</sub>$  Pc/GC, and  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc/GC were calculated to be 21.85, 11.42, 36.98, and 16.94 mF  $cm^{-2}$ , respectively. These data underscores that the active area is not the decisive factor affecting the catalytic activity, which is consistent with the results obtained from  $N_2$  adsorption– desorption curves (Fig. S26†).

#### Electrochemical reduction of  $CO<sub>2</sub>$  in the flow cell

To further assess the potential of the  $Ni(NH_2)_8$ Pc-GC catalyst for industrial applications, we subsequently investigated its electrocatalytic performance in a flow cell setup. A gas diffusion electrode (GDE) was employed to directly transport  $CO<sub>2</sub>$  to the catalyst surface, effectively overcoming the mass transfer limitation of  $CO<sub>2</sub>$  in aqueous media. The composition of the flow cell and testing details are described in Fig. S27 and the ESI.† According to the LSV curves (Fig. 3a), the current densities are significantly higher compared to those in the H-type cell. Specifically,  $Ni(NH_2)_8$ Pc-GC achieves a current density of 400



Fig. 3 (a) LSV curves with a scan rate of 5 mV s<sup>-1</sup> and 80% iR compensation ( $R_s = 2.0 \Omega$ ) of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC and Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC. (b) Total current densities at different applied potentials. (c) FEs and  $j_{\text{CO}}$  of Ni(NH2)8Pc-GC. (d) Long-term chronopotentiometric stability with a current density of 100 mA cm<sup>-2</sup> of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC. (e) Performances comparison between Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC and Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC at -0.88 V vs. RHE. (f) Comparis of  $j_{CO}$  and TOF for at  $-1.03$  V vs. RHE and recently reported Ni-based catalysts.

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mA cm<sup>-2</sup> at −0.6 V vs. RHE, surpassing that of Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC (23 mA  $cm^{-2}$ ). Furthermore, Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC maintains a stable current density under all applied polarization potentials, reaching an industrial current density of 246 mA  $cm^{-2}$  at  $-1.03$  V vs. RHE (Fig. 3b). The FE<sub>CO</sub> of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC approaches nearly 100% over a fairly wide potential range from −0.43 V to −1.03 V vs. RHE (Fig. 3c). In contrast, the maximum FE<sub>CO</sub> and  $j_{\text{CO}}$  for Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC are 95% at −0.43 V vs. RHE and 16.7 mA cm<sup>-2</sup> at -0.58 V vs. RHE, respectively. (Fig.  $S28$  and  $S29\dagger$ ). Long-term stability in the flow cell is also a crucial criterion for practical applicability. In the case of  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC, the constant-current electrolysis was performed at the current density of 100 mA cm $^{-2}$ . It demonstrated stability for at least 8 hours, with no significant increase in potential and an  $FE<sub>CO</sub>$  close to 100%. However, after 10 hours of electrolysis, the  $FE_{CO}$  drops to 71.8% due to flooding and salting out (Fig. 3d).39,40 Replacing the cathode electrolyte with a fresh one restored the potential but could not be maintained for an extended duration, primarily because of the disruption of the gas-liquid balance on the catalyst's surface after prolonged electrolysis in a strong alkaline solution. However, the current density of  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC remains stable only for a short period. Its FE<sub>CO</sub> declines rapidly after one hour of electrolysis (Fig. S30) and S31†). A more intuitive comparison of performance parameters at −0.88 V vs. RHE highlights the advantages of conjugate coupling in  $CO<sub>2</sub>RR$ , as shown in Fig. 3e. Remarkably, Ni $(\mathrm{NH}_2)_8$ Pc-GC achieves a high TOF $_{\rm CO}$  of 342, 296 h $^{-1}$ , a singlepass  $CO<sub>2</sub>$ -to-CO conversion efficiency of 8.9%, a half-cell energy efficiency of 59% for CO production, and a maximum CO yield of 17.8 L  $\rm{min}^{-1}$   $\rm{m}^{-2}.$  The performance of  $\rm{Ni(NH_2)_8}$ Pc-GC in the flow cell is truly exceptional, outperforming most of the recently reported catalysts (Fig. 3f and Table S2†). Chemical Science<br>
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#### Electrocatalysis mechanism

In situ attenuated total reflectance surface-enhanced infrared spectroscopy (ATR-SEIRAS) was performed to investigate the transformation of intermediates at different potentials in a  $CO<sub>2</sub>$ saturated solution of 0.5 M KHCO<sub>3</sub>. In the in situ ATR-SEIRAS analysis of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC, the peak observed at 1640 cm<sup>-1</sup> corresponds to the H–O–H bend of the adsorbed  $H_2O$  molecules (Fig. 4a, b and S32†).<sup>41</sup> Additionally, two infrared bands at 2321 and 2350  $\mathrm{cm}^{-1}$  represent the adsorbed CO $_2$ . $^{\text{42}}$  The peaks at 1380 and 1545  $cm^{-1}$  are assigned to the symmetric and asymmetric stretching vibrations of \*COOH, respectively.<sup>43</sup> As the reaction potential increases, the peak intensity of \*COOH gradually increases. Furthermore, the signal observed at 2080  $cm^{-1}$  is attributed to the Ni–CO vibration, which gradually broadens as the electrolysis potential becomes more negative.<sup>43</sup> The presence of \*CO and \*COOH as crucial intermediates for CO generation aligns with subsequent theoretical calculations.

We also conducted density functional theory (DFT) calculations to deepen our comprehension of the thermodynamic reaction process. Taking into account the periodicity and edge effect of the carbon unit cell,<sup>26</sup> we developed a model that incorporated amino-modified phthalocyanine coupled with a 14-ring polycyclic aromatic fragment (Fig. S33†). Based on the



Fig. 4 (a) In situ ATR-SEIRAS spectra of  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$  at different potentials in  $CO<sub>2</sub>$ -saturated 0.5 M KHCO<sub>3</sub>. (b) The corresponding 2D colormap surface of (a). (c) Free energy diagrams of  $Ni(MH<sub>2</sub>)<sub>8</sub>PC-GC$ and  $Ni(NH<sub>2</sub>)<sub>4</sub>PC-GC$  for  $CO<sub>2</sub>RR.$  (d) Mulliken charges of Ni atom in Ni(NH2)8Pc-GC and Ni(NH2)4Pc-GC. Different charge densities of (e) Ni(NH2)8Pc-GC (top) and Ni(NH2)4Pc-GC (bottom) when \*COOH adsorbs on Ni site. Color-code atoms represent Ni (lightseagreen), C (gray), N (blue), O (red) and H (white). The isosurface value is 0.05 e Å<sup>-3</sup>. The color blue represents the charge increase and the color yellow represents the charge decrease.

intermediates observed through in situ ATR-SEIRAS, this model allowed us to propose the steps required for the CO generation. Firstly,  $CO<sub>2</sub>$  is adsorbed on the active site, and protons are acquired to form \*COOH. Subsequently, the \*COOH intermediate undergoes a proton-coupled electron transfer (PCET) reaction, resulting in the generation of the \*CO intermediate. Finally, the \*CO intermediate desorbs from the catalyst surface. The adsorption states of two critical intermediates on the catalysts  $Ni(NH_2)_8$ Pc-GC and  $Ni(NH_2)_4$ Pc-GC in the CO<sub>2</sub>-to-CO process are outlined in Fig. S34 and S35.† The free energy of the intermediate states on each model during the  $CO<sub>2</sub>RR$ process was calculated. The free energy diagram (Fig. 4c) confirms that the formation of \*COOH on the Ni site is endothermic, while the generation of \*CO is exothermic. The ratedetermining step (RDS) is the formation of \*COOH for both  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC and  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC, with energy barriers of 2.11 and 2.14 eV, respectively. Moreover, the energy barrier corresponding to  $Ni(NH_2)_8$ Pc-GC (0.86 eV) is significantly lower than that of the CO desorption model for  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC (0.98 eV). Altogether, the conjugated pyrazine linkage activates \*COOH and promotes the desorption of \*CO, giving rise to the favored synthesis of CO.<sup>44</sup>

The Mulliken charges of the Ni atom in  $Ni(NH_2)_8$ Pc-GC and  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC were analyzed (Fig. 4d).  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC exhibits



Fig. 5 (a) Schematic diagram of aqueous rechargeable Zn–CO<sub>2</sub> battery. (b) Charge–discharge polarization curves at a scan rate of 5 mV s<sup>−1</sup>, (c) power density curves, and (d) discharge curves and corresponding FE<sub>CO</sub> values at different current densities of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC and Ni(NH<sub>2</sub>)<sub>4</sub>Pc-.<br>GC. (e) Prolonged discharge curve and FE<sub>CO</sub> at −0.5 mA cm<sup>−2</sup> with Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC cathode. (f) Galvanostatic discharge–charge cycling curves of Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC and Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC at  $\pm$ 0.5 mA cm<sup>−2</sup>. (g) The photograph shows an LED bulb (rated voltage: 1.6 V) powered by two ZCBs connected in series with  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$  cathode.

a higher electron density at the Ni site than  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC, suggesting that the pyrazine linkage facilitates the transfer of electrons from the carbon support to the Ni site.<sup>35</sup> The different charge densities were further examined to understand the impact of the electronic density of the Ni site on  $CO<sub>2</sub>$  activation (Fig. 4e). It was observed that  $Ni(NH<sub>2</sub>)<sub>8</sub>$  Pc-GC converts more electrons (0.046 e<sup>-</sup>) to form \*COOH, in contrast to Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC (0.044 e<sup>-</sup>), lowering the energy barrier for the reaction ( $*$  +  $CO<sub>2</sub> + H<sup>+</sup> + e<sup>-</sup> \rightarrow$  \*COOH).<sup>33</sup> The presence of the pyrazine linkage facilitates enhanced electron transfer, which also improves the adsorption of \*COOH and leads to enhanced activity.

#### Aqueous rechargeable  $Zn-CO<sub>2</sub>$  battery

Considering the benefits of metal– $CO<sub>2</sub>$  batteries in promoting carbon recycling and affordability, significant attention has

been paid to the development of aqueous rechargeable  $Zn-CO<sub>2</sub>$ batteries.45,46 The principle of an aqueous rechargeable battery is displayed in Fig. 5a. In the experimental setup, a hydrophobic carbon paper with a catalyst loading of 0.5 mg  $cm^{-2}$  was used as the cathode, while a polished Zn foil served as the anode. The catholyte comprised  $0.8$  M KHCO<sub>3</sub>, and the anolyte was a mixture of 6 M KOH and 0.2 M  $\text{Zn}(\text{CH}_3\text{COO})_2$ . In the discharge phase,  $CO<sub>2</sub>$  is converted into value-added chemicals while generating electricity. Conversely, during the charging process, the cathode undergoes an oxygen evolution reaction (OER). The efficiency of the OER was tested to confirm the bifunctionality of the catalysts. The LSV curves were recorded in 0.8 M  $KHCO<sub>3</sub>$ over a potential interval of 0.9–2.4 V vs. RHE using a threeelectrode system. As displayed in Fig. S36, $\dagger$  Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC demonstrated a lower onset potential (1.81 V) and a higher current density of 51 mA  $cm^{-2}$  at 2.4 V vs. RHE compared to Ni(NH<sub>2</sub>)<sub>4</sub>Pc-GC (1.92 V, 39 mA cm<sup>-2</sup> at 2.4 V vs. RHE). Furthermore,  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC requires an overpotential of 760 mV to achieve a current density of 10 mA  $\text{cm}^{-2}$ , which is 110 mV lower than that for  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc-GC. According to Fig. S37, $\dagger$  the Tafel slope for OER on Ni $(NH_2)_8$ Pc-GC is calculated to be 265.4 mV dec<sup>-1</sup>, which is much lower than that of  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC, implying that the former achieves faster kinetics in a neutral electrolyte solution.

Based on the efficient  $CO_2$ -to-CO and OER performance, an aqueous rechargeable Zn-CO<sub>2</sub> battery was assembled to explore the potential of  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$  in energy conversion and storage devices. The details of the battery assembly and possible charging and discharging processes were carefully recorded in the ESI.† As presented in Fig. S38 and S39,† the open circuit potential of the assembled battery reaches 1.6 V. When two batteries are connected in series, the open circuit potential reaches as high as 2.6 V. The charge and discharge polarization curves of the assembled  $Zn-CO<sub>2</sub>$  battery were tested (Fig. 5b). It was observed that the charge–discharge gaps increased with higher current densities. Notably, the  $Ni(NH_2)_8$ Pc-GC displays a smaller gap, suggesting superior rechargeability for the battery.<sup>47</sup>–<sup>49</sup> Fig. 5c shows that the maximum power density of Ni $(\mathrm{NH}_2)_8\mathrm{Pc\text{-}GC}$  is 0.73 mW  $\mathrm{cm^{-2}}$  (at a loading of 0.5 mg  $\mathrm{cm^{-2}}$ ), which is comparable to most values reported in the literature (Table S3†). Fig. 5d illustrates the relationship between discharge voltage and the faradaic efficiency of catalysts at different discharge current densities. At a constant discharge current density of 0.5–2.5 mA  $\rm cm^{-2} ,$  both Ni $\rm (NH_2)_8$ Pc-GC and  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC consistently exhibit a FE<sub>CO</sub> above 80%. At the same discharge current density, the discharge voltage of  $Ni(NH_2)_8$ Pc-GC is higher than that of  $Ni(NH_2)_4$ Pc-GC. At a discharge current density of 0.5 mA  $\rm cm^{-2}$ , Ni $\rm (NH_2)_8$ Pc-GC obtains a discharge voltage of 0.37 V and a maximum  $FE_{CO}$  of 98.2%. Therefore, we conducted a long-term discharge test and a galvanostatic charge–discharge cycle test at a current density of 0.5 mA  $cm^{-2}$  to assess the stability of the Zn–CO<sub>2</sub> battery (Fig. 5e, f and S40†). During a 24 hour long-term discharge test of  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$ , the discharge potential and  $FE<sub>CO</sub>$  remain stable, with only a minor attenuation. The potential gap of  $Ni(NH<sub>2</sub>)<sub>8</sub>$ Pc-GC increases from 2 V to 2.17 V after 50 hours of charge–discharge cycling. However, the potential gap of  $Ni(NH<sub>2</sub>)<sub>4</sub>$ Pc-GC increases from 2.52 V to 2.72 V after only 32 hours of cycling, and its discharge potential decays to nearly 0 V. Frustratingly, ZnO precipitation in the anolyte and on the membrane is inevitable, which increases potential gap. Additionally, as a practical application demonstration, two batteries connected in series are shown to power a red LED bulb (1.6 V, 20 mA) (Fig. 5g). Therefore, aqueous rechargeable  $Zn-CO<sub>2</sub>$ batteries are promising in alleviating energy and environmental challenges. Chemical Science<br>
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### **Conclusions**

In conclusion, this study investigated the effect of different incorporation modes between amino-substituted Ni phthalocyanine and graphitic carbon surfaces for the electrocatalytic  $CO<sub>2</sub>RR$ . In both the H-cell and the flow cell,  $Ni(NH<sub>2</sub>)<sub>8</sub>PC-GC$ 

consistently achieves a  $FE_{CO}$  of nearly 100% over a fairly wide potential range, demonstrating superior catalytic performance compared to  $Ni(NH<sub>2</sub>)<sub>4</sub>$  Pc-GC. Especially in the flow cell, Ni(NH<sub>2</sub>)<sub>8</sub>Pc-GC exhibits an exceptional TOF of 342, 296 h<sup>-1</sup> at  $-1.03$  V. Furthermore, in the rechargeable aqueous Zn–CO<sub>2</sub> battery assembled with  $Ni(NH_2)_8$ Pc-GC, a FE<sub>CO</sub> of 94.3% is obtained at a discharge current density of 2.5 mA  $cm^{-2}$ , successfully demonstrating the utilization of  $CO<sub>2</sub>$  fixing in the field of energy storage. Both experimental data and theoretical calculations consistently support the notion that graphite conjugated coupling accelerates the electron transfer kinetics between carbon supports and molecular catalysts, thereby increasing the electron density of Ni center. This study paves the way for the rational design of heterogeneous catalysts with enhanced  $CO<sub>2</sub>RR$  activity, providing a promising avenue for the utilization of  $CO<sub>2</sub>$  in energy-related applications.

### Data availability

All experimental and computational procedures and associated data are provided in the ESI.†

### Author contributions

J. H. and H. R. conceived the project. J. H. prepared materials, performed measurements and analysed the data. Q. X. performed in situ ATR-SEIRAS analysis. F. T. performed in situ DEMS analysis. H. S., Y. Q., G. Z., and J. Q. helped with some of the experiments and characterization. All authors discussed the results and commented on the manuscript. J. H. and H. R. wrote the draft, revised and finalized the manuscript.

### Conflicts of interest

There are no conflicts to declare.

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