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## Synergistic Cu-based catalysts with multiple active sites for high-efficiency Li–CO<sub>2</sub> batteries

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The lithium–carbon dioxide (Li–CO<sub>2</sub>) battery is a promising energy storage technology that integrates CO<sub>2</sub> utilization with energy storage and conversion. However, its development is hindered by slow reaction kinetics and insulating Li<sub>2</sub>CO<sub>3</sub> discharge products deposited at the cathode, which cause severe polarization and rapid capacity degradation. Herein, novel Cu-based catalysts with multiple active sites anchored on nitrogen-doped carbon (Cu/NC) are developed to achieve highly efficient Li–CO<sub>2</sub> batteries. Metallic Cu nanoparticles facilitate efficient electron transfer during the CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR), while the induced Cu–N/O dual active sites effectively reduce the energy barrier for decomposing discharge products. Furthermore, the optimized N configurations in the NC matrix enhance the intrinsic activity of the catalytic sites. Consequently, the Li–CO<sub>2</sub> battery incorporating the optimized catalyst demonstrates attractive cycling stability over 850 h at 300 mA g<sup>−1</sup>, with a remarkably low overpotential of 1.30 V, showing great potential for low-cost and highly efficient Li–CO<sub>2</sub> batteries. This work provides a strategic route for designing cost-effective multi-active-site catalysts, offering critical insights into the development of high-performance Li–CO<sub>2</sub> batteries.

## Introduction

With the increasing global demand for energy storage, conventional lithium-ion batteries (*ca.* 350 W h kg<sup>−1</sup>) are insufficient to meet the requirements of high-tech applications and long-range electric vehicles.<sup>1–3</sup> In this regard, lithium–carbon dioxide (Li–CO<sub>2</sub>) batteries have garnered significant attention due to their high theoretical energy density (1876 W h kg<sup>−1</sup>) and dual capability of CO<sub>2</sub> capture and conversion.<sup>4–7</sup> Despite their promise as next-generation energy storage systems, Li–CO<sub>2</sub> batteries face critical commercialization challenges. On the one hand, in the discharge process, the progressive accumulation of insulating Li<sub>2</sub>CO<sub>3</sub> invariably induces cathodic surface passivation, electrical conductivity degradation, and catalytic active site blockage.<sup>8,9</sup> In the recharge process, the thermodynamic stability of Li<sub>2</sub>CO<sub>3</sub> necessitates exceptionally high decomposition voltages (typically beyond 4.3 V), resulting in severe side reactions and substantial electrochemical polarization, leading to rapid capacity fade and shortened cycle life.<sup>10–12</sup>

To address these challenges, the development of high-performance cathodic catalysts capable of simultaneously

accelerating CO<sub>2</sub> reduction reaction kinetics and promoting Li<sub>2</sub>CO<sub>3</sub> decomposition has been regarded as an efficient strategy and has aroused significant research interest in recent years.<sup>13,14</sup> Among all the electrochemical catalysts, transition metal-based catalysts are widely applied in various electrochemical catalytic reactions such as the urea oxidation reaction, CO<sub>2</sub> reduction reaction and hydrogen evolution reaction due to their appealing electronic structure tunability, economic efficiency, and resource abundance.<sup>4,12,15,16</sup> In particular, Cu-based catalysts with a wide range of oxidation states (Cu<sup>0</sup>, Cu<sup>I</sup>, Cu<sup>II</sup>, and Cu<sup>III</sup>) and adjustable electronic structure have attracted enormous interest and exhibited great promise for Li–CO<sub>2</sub> batteries.<sup>17,18</sup> For instance, Xu *et al.*<sup>19</sup> combined Cu polyphthalocyanine with carbon nanotubes to obtain a Cu single-atom catalyst anchored on a conductive porphyrin network, realizing a low polarization of 1.8 V at 2000 mA g<sup>−1</sup> in Li–CO<sub>2</sub> batteries. By using the solvothermal method, Cu-tetra(4-carboxyphenyl) porphyrin (Cu-TCPP) nanosheets were demonstrated to be an efficient catalyst, which contributed to sustaining 123 cycles at 500 mA g<sup>−1</sup>. Despite these promising results, Cu-based catalysts still face significant challenges, including slow reaction kinetics, poor selectivity and insufficient stability, and thus require further development.

Inspired by the above-mentioned studies, herein, we developed a novel ultrafine Cu nanoparticle (NP)-decorated nitrogen-doped carbon (Cu/NC) as a high-efficiency catalyst for Li–CO<sub>2</sub> batteries. Comprehensive characterization revealed that metallic Cu nanoparticles facilitate electron transfer during CO<sub>2</sub>

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reduction, while Cu–N/O moieties effectively reduce  $\text{Li}_2\text{CO}_3$  decomposition barriers. Additionally, N-doped carbon with optimized N configurations contributes to improved catalytic activity. Such attractive structural properties endow the Li–CO<sub>2</sub> battery with an optimized Cu/NC catalyst that delivers a low overpotential of 1.30 V and extended cycling stability over 850 h at 300 mA g<sup>−1</sup>, exhibiting great prospects as a low-cost catalyst for Li–CO<sub>2</sub> batteries.

## Results and discussion

Fig. 1a schematically illustrates the formation process of Cu/NC *via* pyrolysis of a precursor containing melamine, cyanuric acid (CA), and copper(II) nitrate hexahydrate under an Ar atmosphere. Specifically, the precursor was prepared by first mixing an aqueous solution of CA, melamine and copper nitrate to form a copper-containing supramolecular complex owing to the hydrogen bond interaction between them.<sup>20</sup> Subsequently, due to the weakly alkaline nature of the melamine solution, the  $-\text{O}^-$  groups in CA molecules coordinated with Cu<sup>2+</sup> ions to form Cu–O bonds,<sup>21</sup> while nitrogen atoms in melamine rings coordinated with Cu<sup>2+</sup> ions, establishing Cu–N coordination.<sup>20</sup> The XRD analysis of the self-assembled precursor (Fig. S1) revealed two new diffraction peaks at  $2\theta = 11^\circ$  and  $28^\circ$  can be detected, which cannot be assigned to a certain product. We infer this may be caused by the periodic arrangements of intraplanar stacking and interlayer aromatic stacking, which demonstrates the well-ordered hydrogen-bonded network within the supramolecular architecture,<sup>22</sup> confirming successful complex formation. The XRD pattern of the precursor of pure NC, which was prepared without copper nitrate, was nearly identical to that of the precursor of Cu/NC, demonstrating that the Cu/NC precursor maintained the structural advantages of the supramolecular template, thereby laying the foundation for the subsequent pyrolysis to prepare highly dispersed Cu/NC catalysts. The Fourier transform infrared (FTIR) spectroscopy data shown in Fig. S2 indicate that the C=O stretching vibrations ( $\nu_{\text{C=O}}$ ) for CA appear at 1781 and 1735 cm<sup>−1</sup> in the Cu/NC precursor, showing an obvious blue shift compared to those in the reported literature (1739 and 1695 cm<sup>−1</sup>), which may be caused by the formation of hydrogen-bonded supramolecular aggregates through N–H···O and N–H···N hydrogen bonds between melamine and CA.<sup>23,24</sup> Furthermore, the Cu/NC precursor exhibits an FTIR spectrum similar to that of the CN (Cu-free) precursor, implying that Cu<sup>2+</sup> incorporation does not affect the formation of the hydrogen-bonded supramolecular assembly. Subsequent pyrolysis at 600 °C for 2 hours under an Ar atmosphere contributed to the formation of N-doped carbon and the firm anchoring of the active sites of the Cu–N/O coordination structure, with most Cu atoms evolving into nanoparticles encapsulated by N-doped carbon to form the Cu/NC catalyst (designated m-Cu/NC). Compared to the precursor, new peaks at 3700–3000 and 1800–1100 cm<sup>−1</sup> have emerged in the FTIR spectra of Cu/NC (Fig. S2), proving the existence of a heterocyclic ring structure.<sup>25</sup> To probe the influence of the Cu-based active sites on the catalytic activity, catalysts with

different Cu contents (0.01 M and 0.05 M) were prepared and named s-Cu/NC and l-Cu/NC, respectively.

The morphology of the catalyst was investigated by using the SEM technique first. As shown in Fig. 1b, the as-formed m-Cu/NC displays uniform nanoparticles with an average diameter of 27.9 nm. These ultrafine nanoparticles are beneficial for increasing the surface area, thus providing more catalytic activity sites and accelerating the reaction kinetics.<sup>26</sup> In contrast, s-Cu/NC presents larger irregular particle agglomerates, and l-Cu/NC shows a flower-like morphology formed by the accumulation of both extremely large and small particles (Fig. S3 and S4). This difference may arise from variations in concentrations. At lower concentrations, fewer nucleation sites form, allowing sufficient time for particle growth and facilitating the formation of larger particles. Conversely, at higher concentrations, explosive nucleation rapidly generates numerous small crystal nuclei, and small particles subsequently attach to larger ones.<sup>27</sup> The TEM images (Fig. 1c and d) further confirm the formation of small nanoparticles uniformly distributed in the amorphous carbon layer. The HRTEM images displayed in Fig. 1e and f distinctly confirm the presence of amorphous carbon and the tight coating of Cu nanoparticles by the carbon layer, which are beneficial for enhancing the structural stability of active sites during repeat electrochemical processes and thus contribute to the long lifespan of the batteries.<sup>28,29</sup> The lattice fringes of 0.208 nm and 0.183 nm were identified, corresponding to the (111) and (200) crystal planes of metallic Cu, respectively, confirming the formation of crystalline Cu. Nitrogen adsorption–desorption isotherms were measured at 77 K to characterize the surface area and pore size distribution of the m-Cu/NC sample (Fig. 1g). The presence of a Type IV isotherm with an H3-type hysteresis loop in the high-pressure region confirms the existence of a mesoporous structure,<sup>30</sup> which helps accelerate the CO<sub>2</sub> gas diffusion and Li<sup>+</sup> transport in the electrolyte, thereby reducing concentration polarization and lowering the charge–discharge overpotential.<sup>31</sup> The BET surface area of m-Cu/NC is calculated to be 64.6 m<sup>2</sup> g<sup>−1</sup> with a dominant diameter of *ca.* 2.58 nm. The larger specific surface area facilitates the exposure of more catalytically active sites, enabling the efficient adsorption of CO<sub>2</sub> reactive species on the external surface active centers and thus enhancing the CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) performance.<sup>32–34</sup>

Fig. 2a shows the XRD patterns of the prepared samples. The diffraction peaks located at 43.3°, 50.5°, and 74.1° can be well matched with the (111), (200), and (220) crystal planes of cubic phase Cu (JCPDS no. 98-000-0172).<sup>35</sup> No obvious diffraction peaks related to carbon can be detected, which may be overlapped by the sharp diffraction peaks of Cu. Fig. 2b displays the Raman spectra of the three samples, where two peaks appear at 1427 and 1594 cm<sup>−1</sup>, corresponding to the characterization peaks of the D-band and G-band of carbon, respectively.<sup>36</sup> The D-band is associated with structural defects or edges in the sample, while the G-band corresponds to sp<sup>2</sup>-hybridized graphitic carbon.<sup>37</sup> From the  $I_{\text{D}}/I_{\text{G}}$  ratio displayed in Fig. 3c, it can be seen that the m-Cu/NC sample exhibits the highest  $I_{\text{D}}/I_{\text{G}}$  ratio (1.01) compared to its counterparts (0.88 for s-Cu/NC and



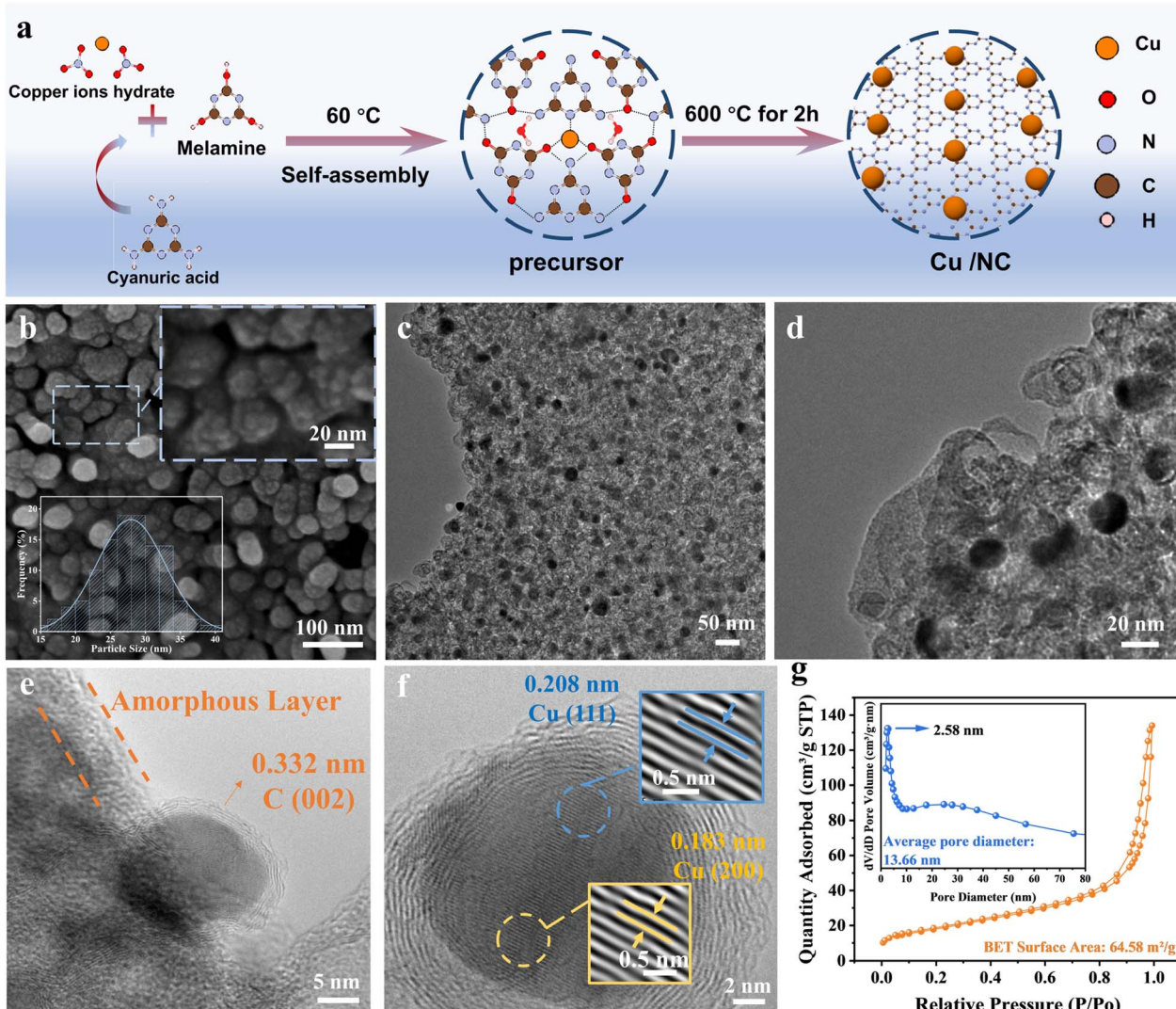


Fig. 1 (a) Schematic illustration of the formation process of Cu/NC. (b) SEM images of m-Cu/NC and the corresponding size distribution of Cu NPs (inset). (c and d) TEM images of m-Cu/NC. (e and f) HRTEM images of m-Cu/NC. (g)  $N_2$  adsorption-desorption isotherms of m-Cu/NC and the related pore size distribution (inset).

0.75 for l-Cu/NC), indicating the formation of more enriched defects, which can serve as active centers that promote the reaction between  $Li^+$  and  $CO_2$ , thereby enhancing the  $CO_2$  performance.<sup>38</sup> Surprisingly, the D-band deviated significantly from the conventional position ( $1350\text{ cm}^{-1}$ ),<sup>39</sup> which may originate from the high-frequency displacement N doping or the interfacial stress between the Cu NPs and carbon matrix, both inducing lattice distortion of the carbon skeleton.<sup>40,41</sup> In addition, one peak appearing at  $1290\text{ cm}^{-1}$  can be detected in these three samples, which can be assigned to the stretching vibration of C–N.<sup>42</sup> The FTIR spectra displayed in Fig. 2d confirmed the presence of C=O ( $1623\text{ cm}^{-1}$ ), C–N group ( $1500\text{--}1000\text{ cm}^{-1}$ ), Cu–N bond ( $530\text{ cm}^{-1}$ ), and Cu–O stretching modes ( $437\text{ cm}^{-1}$ ).<sup>43–45</sup> To further explore the chemical states and coordination environments in Cu/NC, XPS analysis was systematically performed (Fig. 2f and h). The high-resolution C 1s spectrum (Fig. S5) of m-Cu/NC can be divided into three

components: C–C ( $284.8\text{ eV}$ ), C–N ( $286.6\text{ eV}$ ), and C=O ( $288.5\text{ eV}$ ), indicating the formation of N-doped carbon.<sup>46</sup> The peak position of C–N slightly shifts to a higher binding energy compared to that of the standard value (usually *ca.*  $285.6\text{ eV}$ ),<sup>47–49</sup> which may be caused by the electronic coupling effect of Cu NPs.<sup>50</sup> The N 1s spectrum of m-Cu/NC can be deconvoluted into three peaks at  $397.8\text{ eV}$  (16.7 at%),  $398.9\text{ eV}$  (72.7 at%), and  $400.8\text{ eV}$  (10.6 at%), corresponding to Cu–N, pyridinic–N and pyrrolic N, respectively (Fig. 2f).<sup>51</sup> Meanwhile, the N 1s spectra of s-Cu/NCs and l-Cu/NCs indicate that the formed Cu–N, pyridinic–N and pyrrolic N are present in different percentages (2.5 at%, 67.5 at%, and 30.0 at% for the former and 21.3 at%, 60.4 at%, and 18.3 at% for the latter). Compared with the other samples, the m-Cu/NC exhibits the highest pyridinic–N content (Fig. 2e). According to the literature, pyridinic N can serve as active sites for the oxygen reduction reaction (ORR) by modulating the electronic structure of adjacent carbon atoms and



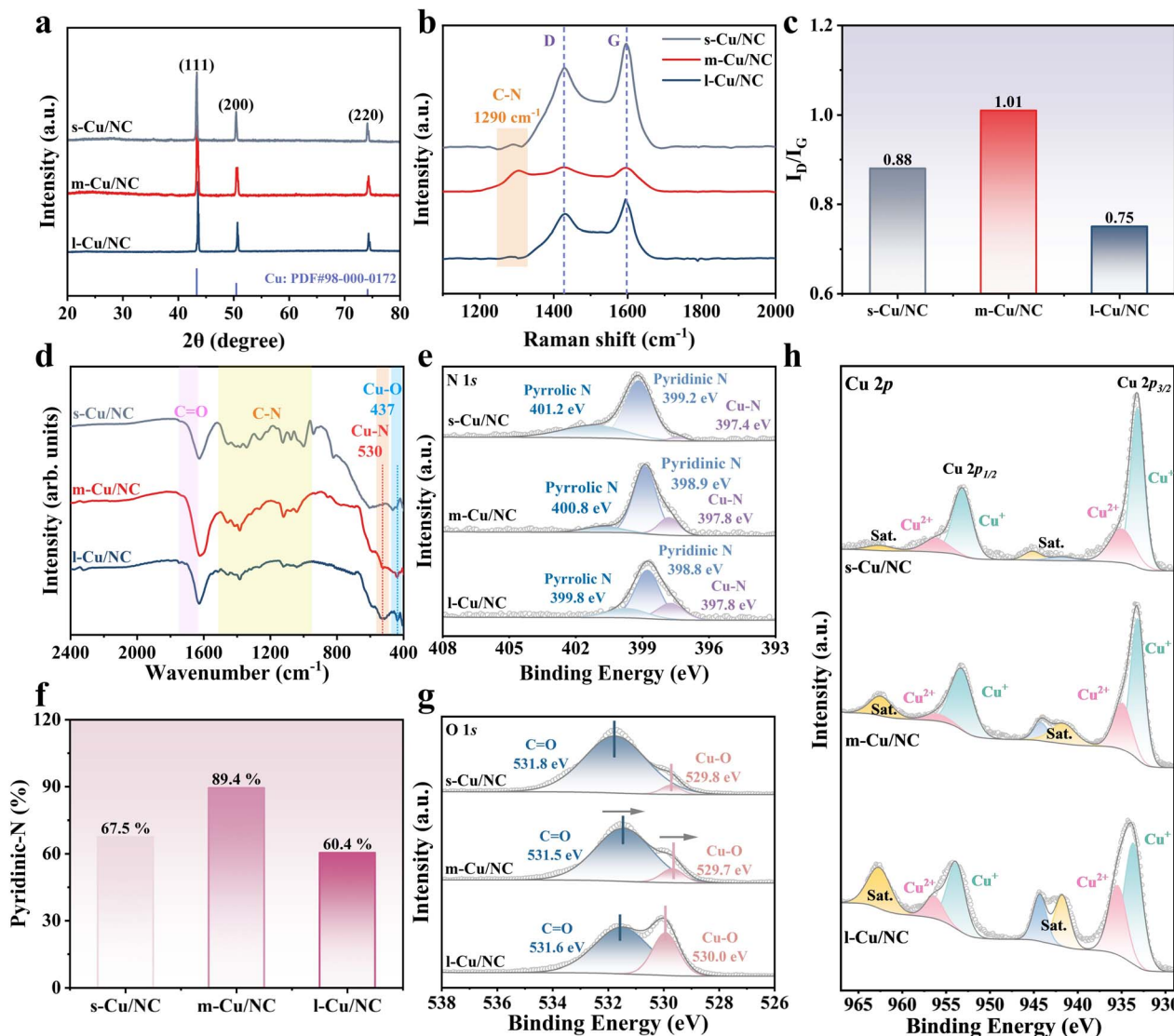


Fig. 2 (a) XRD patterns, (b) Raman spectra, (c)  $I_D/I_G$ , (d) FTIR spectra, (e) N 1s XPS spectra, (f) Comparison chart of pyridinic-N content, (g) O 1s XPS spectra and (h) Cu 2p XPS spectra of the three samples.

enhancing charge transfer efficiency, which can provide additional active sites for  $\text{CO}_2$  conversion in  $\text{Li}-\text{CO}_2$  batteries, leading to enhanced electrochemical performance.<sup>52,53</sup> Thus, m-Cu/NC is believed to show attractive electrochemical performance in  $\text{Li}-\text{CO}_2$  batteries. Furthermore, the O 1s XPS spectrum of m-Cu/NC (Fig. 2g) shows a dominant peak at 531.5 eV related to  $\text{C}=\text{O}$ , and the peak at 529.7 eV corresponds to Cu–O coordination.<sup>33</sup> Meanwhile, Cu 2p XPS spectra (Fig. 2h) displayed characteristic peaks at 933.2/953.4 eV (Cu 2p<sub>3/2</sub> and 2p<sub>1/2</sub> for Cu–N) and 935.0/956.1 eV (corresponding to Cu–O). Satellite peaks at 941.8, 944.3, and 962.7 eV confirmed the  $\text{Cu}^{2+}$  oxidation state.<sup>54,55</sup> Furthermore, XPS quantitative analysis reveals that the Cu atomic content is 17.29% in m-Cu/NC, compared with 2.91% in s-Cu/NC and 21.37% in l-Cu/NC. The moderate Cu content in m-Cu/NC provides an optimal density of surface active sites, effectively promoting  $\text{CO}_2$  reduction for  $\text{Li}_2\text{CO}_3$  formation during discharge. Notably, insufficient Cu content (as s-Cu/NC)

results in limited catalytic activity, while excessive Cu loading (as l-Cu/NC) induces nanoparticle agglomeration, which blocks porous channels and increases the energy barrier for discharge product decomposition, ultimately impairing battery performance.<sup>17</sup> These results conclusively demonstrate the coexistence of dual Cu active centers (Cu–N and Cu–O) in the m-Cu/NC, which can serve as electron transfer channels, optimizing the D-band electron structure of the Cu active center and helping to promote  $\text{CO}_2$  reaction activity and enhance reaction kinetics.<sup>56</sup>

### Electrochemical performance of $\text{Li}-\text{CO}_2$ batteries with the designed catalysts

Considering the attractive structural properties, the electrochemical performance of the  $\text{Li}-\text{CO}_2$  batteries assembled with the prepared catalysts was evaluated. The cyclic voltammetry (CV) curves of the batteries based on Cu/NC catalysts were

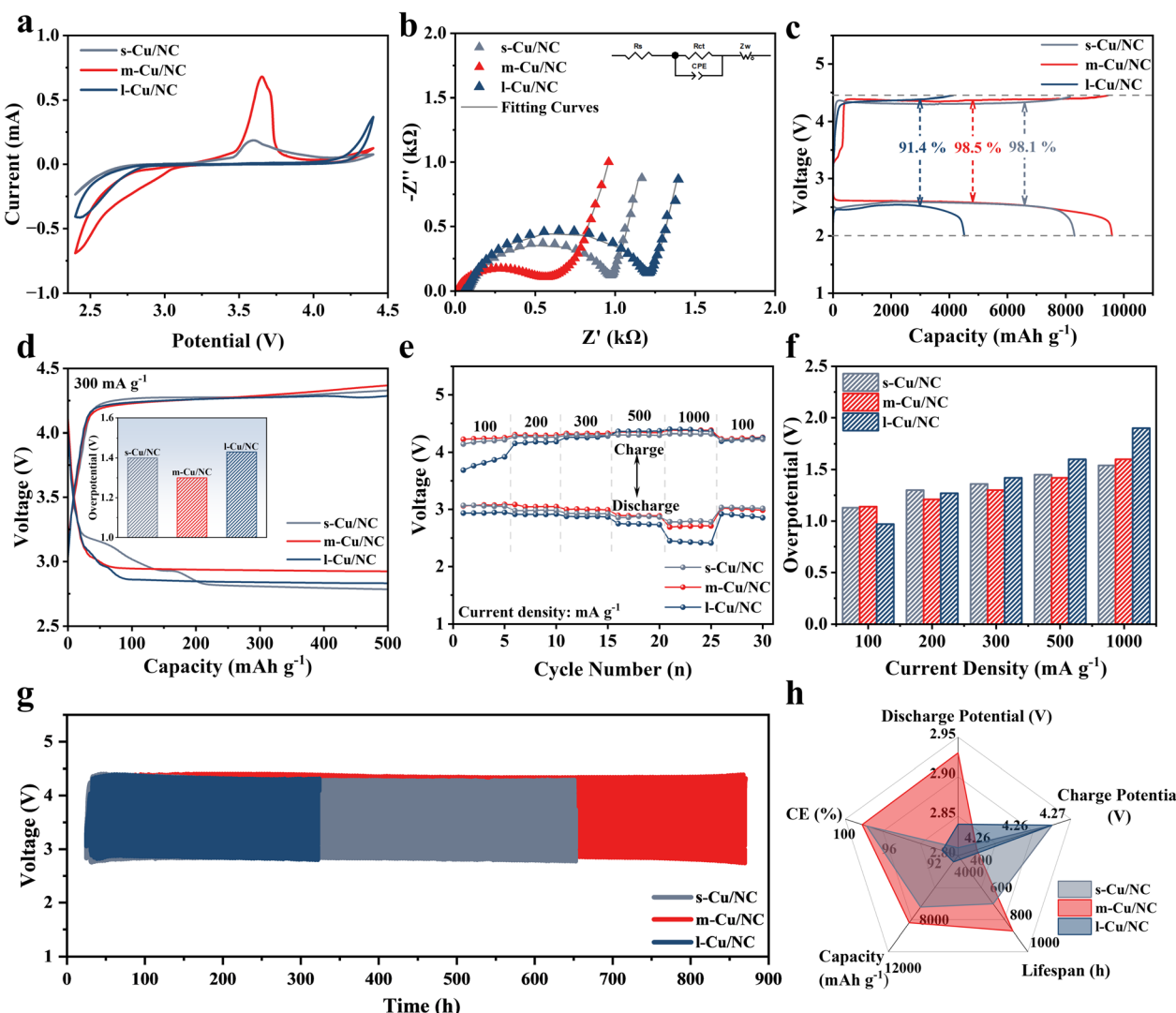


Fig. 3 (a) Cyclic voltammetry curve of the designed catalyst-based Li–CO<sub>2</sub> batteries under a CO<sub>2</sub> atmosphere. (b) Electrochemical impedance spectroscopy of the three samples for Li–CO<sub>2</sub> batteries. (c) Full discharge and charge curves of the three samples at 500 mA g<sup>−1</sup>. (d) Cycle performance of the three samples at a limited capacity of 500 mA h g<sup>−1</sup> at 300 mA g<sup>−1</sup>. (e and f) Rate performances at different current densities. (g) Voltage–time curves of the three samples at 300 mA g<sup>−1</sup> with a limited capacity of 500 mA h g<sup>−1</sup>. (h) Electrochemical performance comparison of the three catalysts.

recorded at a scan rate of 0.1 mV s<sup>−1</sup> over a voltage range of 2.4 V to 4.4 V. From Fig. 3a we can see that the battery with all the catalysts exhibits good activity in the CO<sub>2</sub> atmosphere and shows negligible capacity under the Ar atmosphere (Fig. S6), confirming the necessity of CO<sub>2</sub> as a reactant during redox reaction processes. The m-Cu/NC cathode exhibited more pronounced redox peaks at approximately 2.85 V ( $E_{\text{c1}}$ ) and 3.65 V ( $E_{\text{a1}}$ ),<sup>6</sup> corresponding to the formation and decomposition of the discharge product Li<sub>2</sub>CO<sub>3</sub>, respectively. The observed oxidation peak is lower than the conventional value of 3.8 V (vs. Li/Li<sup>+</sup>), and the reduction peak is higher than the typical 2.8 V (vs. Li/Li<sup>+</sup>), indicating that the m-Cu/NC catalyst effectively reduces the reaction energy barrier for both processes.<sup>10</sup> Compared to the s-Cu/NC and l-Cu/NC curves, m-Cu/NC exhibits the largest peak currents, indicating its superior

activity, which may be ascribed to its uniform spherical structure and good synergistic effect between pyridine N and multi-active sites.<sup>57</sup> Linear sweep voltammetry (LSV) was employed to characterize the reaction kinetics of the investigated systems. As shown in Fig. S7, m-Cu/NC exhibits a significantly higher current density (0.60 mA g<sup>−1</sup>) than s-Cu/NC (0.22 mA g<sup>−1</sup>) and l-Cu/NC (0.24 mA g<sup>−1</sup>), indicating its superior catalytic activity. This enhanced activity will contribute to improving the electrochemical reaction efficiency and accelerating reaction kinetics.<sup>58</sup>

Electrochemical impedance spectroscopy (EIS) was performed to investigate the charge transfer kinetics of each electrode.<sup>59</sup> As shown in Fig. 3b, according to the equivalent circuit fitting results, the  $R_{\text{ct}}$  value of m-Cu/NC was calculated to be 474.8 Ω, which is lower than those of s-Cu/NC (886 Ω) and l-Cu/

NC (1139  $\Omega$ ). The reason for this may be the formed embedded electric field of Cu NPs through the metal-CN interface, resulting in ultrafast charge separation.<sup>17</sup> Additionally, the uniform spherical structure of m-Cu/NC can shorten the transfer path of  $\text{Li}^+$  and thus reduce the  $R_{\text{ct}}$  value.<sup>60</sup>

Fig. 3c and S8 show the full discharge–charge curves and the related coulombic efficiency (CE) of the three catalysts tested at 500  $\text{mA g}^{-1}$ , which reveal that the m-Cu/NC exhibits a superior specific discharge capacity of 9602  $\text{mA h g}^{-1}$  with a higher CE (98.5%) than those of the counterparts (8297  $\text{mA h g}^{-1}$  and 98.1% for s-Cu/NC and 4513  $\text{mA h g}^{-1}$  and 91.4% for l-Cu/NC), and even superior to those previously reported (Fig. S9 and Table S1), suggesting its excellent reversibility. Additionally, the m-Cu/NC catalyst exhibits a higher energy efficiency (EE) value of 59.3% at 500  $\text{mA g}^{-1}$  under full discharge/charge conditions than its counterparts (Fig. S10 and Table S2), demonstrating its favorable energy efficiency.<sup>61–64</sup> When tested at 300  $\text{mA g}^{-1}$ , the

m-Cu/NC-based battery displays a lower polarization voltage of 1.30 V (Fig. 3d), confirming the significant promotion of optimized structure and active sites for  $\text{CO}_2$  transfer kinetics.<sup>65,66</sup>

To evaluate the rate capability, the catalysts were tested at various current densities. As shown in Fig. 3e and f, the m-Cu/NC catalyst exhibits the lowest overpotentials of 1.14, 1.21, 1.30, 1.42, and 1.60 V at 100, 200, 300, 500, and 1000  $\text{mA g}^{-1}$ , respectively, compared to its counterparts, demonstrating its good rate capability across a wide range of current densities. Additionally, the long-term cycling stability of the assembled batteries was investigated at 300  $\text{mA g}^{-1}$ . The overpotential of the m-Cu/NC cathode increased by only 0.60 V after cycling for over 850 h. However, s-Cu/NC and l-Cu/NC show limited cycle life of 652 h and 326 h, respectively. Furthermore, the battery tested at 500  $\text{mA g}^{-1}$  also achieved long-term cycling stability (570 h, Fig. S11). The reason for such attractive performance may be attributed to the synergistic catalytic effect of multiple

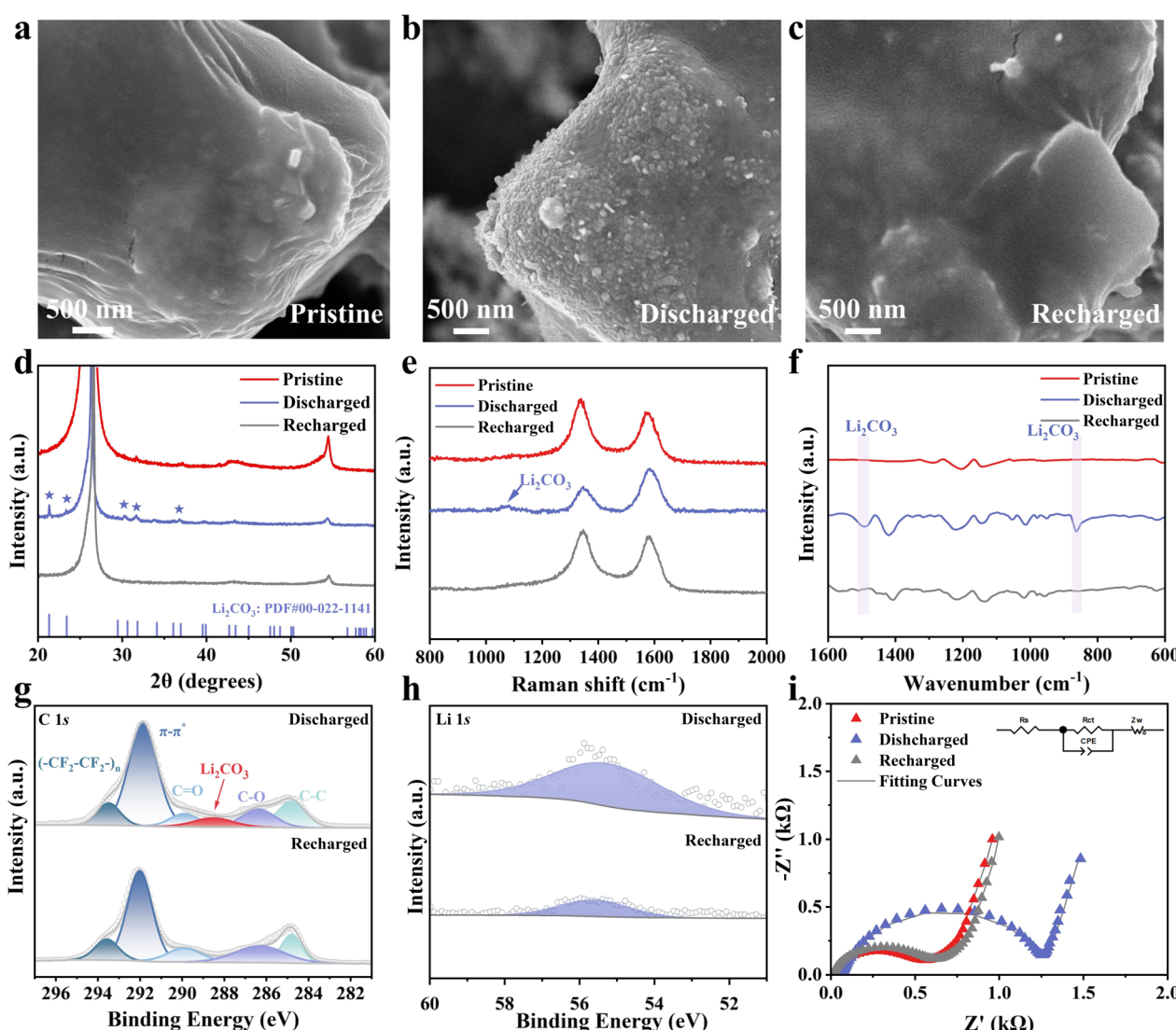


Fig. 4 Characterization of the m-Cu/NC cathode at different states. (a–c) SEM images of m-Cu/NC at the pristine, discharged, and recharged states. (d) XRD patterns, (e) Raman spectra, (f) FTIR, and (g and h) High resolution XPS spectra of C 1s and Li 1s of m-Cu/NC electrodes in the fully discharged state. (i) EIS spectra of m-Cu/NC at different states.



active sites (pyridine N, Cu NPs and Cu–N/O sites) and the NC structure, which can promote the mass transfer of  $\text{CO}_2$  and  $\text{Li}^+$ , avoiding the clogging of discharge products on the electrode surface at high rates.<sup>67</sup> To further explore why the catalyst died after cycling, the battery was disassembled and it was found that the Li anode had completely lost its original metallic luster. When the disassembled m-Cu/NC catalyst was redeployed as the cathode with a fresh Li anode and the electrolyte, the reconstructed Li– $\text{CO}_2$  battery could still achieve 110 cycles (Fig. S12, at 300  $\text{mA g}^{-1}$ ) with an overvoltage of 1.58 V in the final cycle, confirming the stable catalytic performance of the catalyst.<sup>68</sup>

To elucidate the mechanism of the superior electrochemical performance of the m-Cu/NC-based Li– $\text{CO}_2$  battery, the cycled m-Cu/NC electrode at different states was analyzed using various techniques. SEM images (Fig. 4a and b) reveal that the discharged electrode surface is covered with uniformly distributed aggregates composed of small particles, compared to the pristine electrode. These aggregates completely disappear upon recharging (Fig. 4c). We infer that during the discharge process, the large specific surface area enhances  $\text{CO}_2$  adsorption, where Cu NPs and Cu–N/O active sites adsorb and activate  $\text{CO}_2$  molecules, which then combine with  $\text{Li}^+$  to undergo reduction, forming  $\text{Li}_2\text{CO}_3$  and C.<sup>18</sup> XRD analysis further supports this observation. As shown in Fig. 4d, distinct diffraction peaks matching the standard pattern of  $\text{Li}_2\text{CO}_3$  appear on the discharged electrode, confirming  $\text{Li}_2\text{CO}_3$  formation as the discharge product.<sup>10–12</sup> After recharging, these  $\text{Li}_2\text{CO}_3$  peaks vanish, and the XRD profile reverts to resemble that of the pristine electrode, indicating decomposition of the discharge products. Additionally, Raman spectroscopy (Fig. 4e) and FTIR (Fig. 4f) detect characteristic peaks of  $\text{Li}_2\text{CO}_3$  on the discharged electrode, which also disappear completely after recharging. The high-resolution XPS conducted on the electrode at different states (Fig. 4g and h) indicates that the C 1s spectrum shows a distinct peak at 288.5 eV after discharge, attributed to  $\text{Li}_2\text{CO}_3$ , which significantly diminishes after recharging. The other peaks at around 284.8, 286.6, 289.8, 291.9, and 293.5 eV were connected to C–C bonding, C–O bonding, C=O bonding,  $\pi$ – $\pi$  bonding, and  $(-\text{CF}_2-\text{CF}_2-)_n$ , respectively.<sup>69</sup> Additionally, the peak at around 55.6 eV in the spectrum of Li 1s after the process of discharging can be attributed to the formation of Li in  $\text{Li}_2\text{CO}_3$ , which is significantly weakened when the battery is fully recharged, and the peak at 55.6 eV after the process of charging can be attributed to Li in a small amount of remaining lithium salt.<sup>70</sup> In addition, considering that the surface of the catalyst is covered with a wide bandgap insulating  $\text{Li}_2\text{CO}_3$  layer, it will lead to an increase in the charge transfer impedance of the cathode during full discharge. EIS of the cycled m-Cu/NC cathode (Fig. 4i) suggests that the initial impedance value of the battery is approximately 474.8  $\Omega$ . After full discharge, the impedance increases to around 1167  $\Omega$ , which may be caused by the formation of insulating  $\text{Li}_2\text{CO}_3$  and reduced carbon covering the catalyst and lithium surface, hindering electron transport.<sup>71</sup> However, after full recharging, the impedance decreases to 529.1  $\Omega$ , which can be ascribed to the effective decomposition of  $\text{Li}_2\text{CO}_3$ . This value nearly recovers to its initial state but is still

slightly higher, likely due to the presence of trace amounts of undecomposed  $\text{Li}_2\text{CO}_3$  and reduced carbon.<sup>72</sup> These results demonstrate the good reversibility of the battery. Therefore, it can be concluded that the existence of optimized N configuration, small Cu NPs and Cu–N/O promotes the decomposition of discharge products by reducing the decomposition energy barriers. Moreover, the larger specific surface area with enriched mesopores also provides space for the accumulation of discharge products, promoting the effective decomposition of discharge products and thus prolonging the cycle life of Li– $\text{CO}_2$  batteries.<sup>67</sup>

## Conclusions

In summary, we demonstrate a rationally designed Cu–N/O multi-site catalyst integrated with Cu nanoparticles (m-Cu/NC) synthesized *via* facile pyrolysis. This catalyst exhibits attractive bifunctional activity in Li– $\text{CO}_2$  batteries. The synergistic interplay between Cu–O and Cu–N coordination significantly enhances electron transfer kinetics, accelerating both  $\text{CO}_2$  reduction and evolution reactions. The incorporated Cu NPs facilitate efficient electron transfer during the  $\text{CO}_2\text{RR}$ . Simultaneously, the N-doped carbon matrix further enhances catalytic activity for the  $\text{CO}_2\text{RR}$  and  $\text{CO}_2\text{ER}$ , concurrently optimizing electron transport and structural stability. Consequently, the m-Cu/NC-based Li– $\text{CO}_2$  battery achieves stable cycling for over 850 h at 300  $\text{mA g}^{-1}$  with a low voltage gap of 1.30 V, outperforming existing Cu-based catalysts. This work highlights the promising potential of Cu-based materials in metal-air batteries and establishes new design principles for advanced bifunctional catalysts.

## Author contributions

Y. X. and S. X. conceived the idea and wrote the manuscript. S. C. and Y. X. supervised the work and revised the manuscript. S. X. directed the project, prepared the samples, performed the electrochemical performance evaluation, and wrote the paper. G. H., T. Z., and L. Y. analysed part of the data and contributed to the preparation of the manuscript. F. H. revised the manuscript. All the authors discussed and analysed the results, and commented on the manuscript.

## Conflicts of interest

The authors declare no conflict of interest.

## Data availability

The data supporting this article have been included as part of the SI.

Catalyst synthesis, characterizations, and supporting Figures and Tables. See DOI: <https://doi.org/10.1039/d5sc04645a>.

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