



# N-modulated Cu<sup>+</sup> for efficient electrochemical carbon monoxide reduction to acetate

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The electrocatalytic approach of converting carbon dioxide (CO<sub>2</sub>) to valuable chemical commodities and feedstocks provides a promising solution to store intermittent renewable electricity in a high-energy-density way and mitigate CO<sub>2</sub> accumulation in atmosphere [1–3]. Recently, the electrochemical CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) has made remarkable progress in yielding C<sub>1</sub> products (such as carbon monoxide (CO) [4–6] and formate (HCOO<sup>-</sup>) [7–9]) with significantly high current densities and high Faradaic efficiencies (FEs). Meanwhile, the overall performance of CO<sub>2</sub>RR to C<sub>2</sub> products of ethylene [10–12] and ethanol [13–15], has greatly improved on copper (Cu) catalysts. However, acetate (another C<sub>2</sub> product), which is widely used as a raw chemical material for pharmacy, dyestuff, rubber and so on, has an excessively low partial current density (<1 mA cm<sup>-2</sup>) and low FE (<20%) [16,17]. Kanan and coworkers [18] suggested that higher KOH concentrations may facilitate the formation of acetate. However, due to the reaction of CO<sub>2</sub> and KOH forming carbonate, CO<sub>2</sub> is not a proper reaction gas in alkaline reactors.

Given the inertness of CO in alkaline electrolytes, the CO reduction reaction (CORR) instead of CO<sub>2</sub>RR can be conducted in alkaline reactors, which could increase the charge transfer rates, suppress the competition of hydrogen evolution reaction (HER) and improve the acetate FE [19,20]. Moreover, \*CO is known as a vital intermediate for the formation of multi-carbon (C<sub>2+</sub>) products in CO<sub>2</sub>RR [21–23], and thus CORR is potentially more direct for C<sub>2</sub> electroproduction.

To date, copper is the state-of-the-art catalyst for the electro-conversion of CO<sub>2</sub>/CO to C<sub>2</sub> products. Lots of strategies, including morphological modification [24,25],

grain boundary control [26], reaction condition optimization [19], introducing various elements as modulators [14] and modifying the local oxidation state of Cu [10,27], have been taken to enhance the selectivity of C<sub>2</sub> products and the current density. Surface Cu<sup>+</sup> active sites modulated by non-metal elements have been proved theoretically and experimentally to greatly improve the selectivity of C<sub>2</sub> products [10,12,28]. However, the oxygen-modulated Cu<sup>+</sup> species, always derived from cuprous oxide (Cu<sub>2</sub>O) and copper oxide (CuO), is extremely unstable and tends to be reduced to Cu<sup>0</sup> species under CO<sub>2</sub>/CORR conditions, leading to decreased catalytic performance [29].

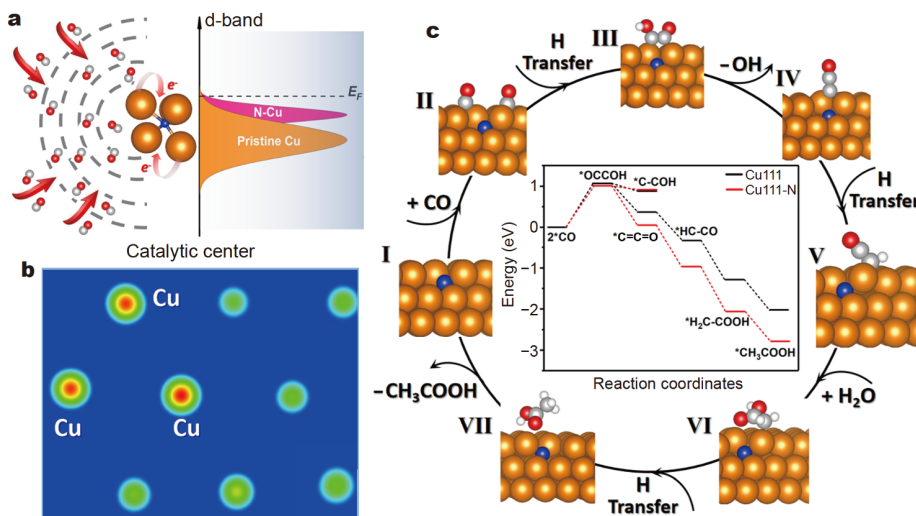
In contrast to the instability of oxygen-modulated Cu<sup>+</sup>, copper nitride has been reported to be stable during CO<sub>2</sub> reduction [30,31]. Negligible degradation of high C<sub>2+</sub> product FEs was achieved on a Cu-on-Cu<sub>3</sub>N catalyst in 30 h, suggesting that nitrogen (N) could be a competent stabilizer to maintain the Cu<sup>+</sup> state [30]. In addition, doping N also provides a more favorable pathway for the reduction of CO<sub>2</sub> to acetate, and a remarkable FE of acetate (FE<sub>acetate</sub>) (nearly 78%) was achieved on a N-doped nanodiamond/Si rod array [32], but the acetate partial current density was lower than 1 mA cm<sup>-2</sup>. Inspired by the above studies, we therefore explore whether incorporating N into Cu catalysts could preserve Cu<sup>+</sup> species and further improve the current density and FE<sub>acetate</sub> in CORR.

We firstly sought to use density function theory (DFT) to testify the existence of Cu<sup>+</sup> species modulated by incorporating N into Cu and then investigated its effect on the selectivity of products in CORR. Bader charge analysis, as shown in Fig. 1b, presents the obvious difference

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**Figure 1** Schematic illustration of acetate electrosynthesis process. (a) Schematic illustration for the synthesis process of acetate using N-Cu catalyst, and the comparison of DOS between pristine Cu and N-Cu systems, where  $E_F$  refers to Fermi level. Color code: Cu (brown), N (blue), C (light gray), O (red). (b) Bader charge analysis of the first layer Cu atoms from the constructed N-Cu model. (c) Free energy profiles of acetate formation on Cu (111) and N-Cu (111) surface, respectively. The optimized structures of key intermediates during CORR on N-Cu (111) are shown in the reaction cycle.

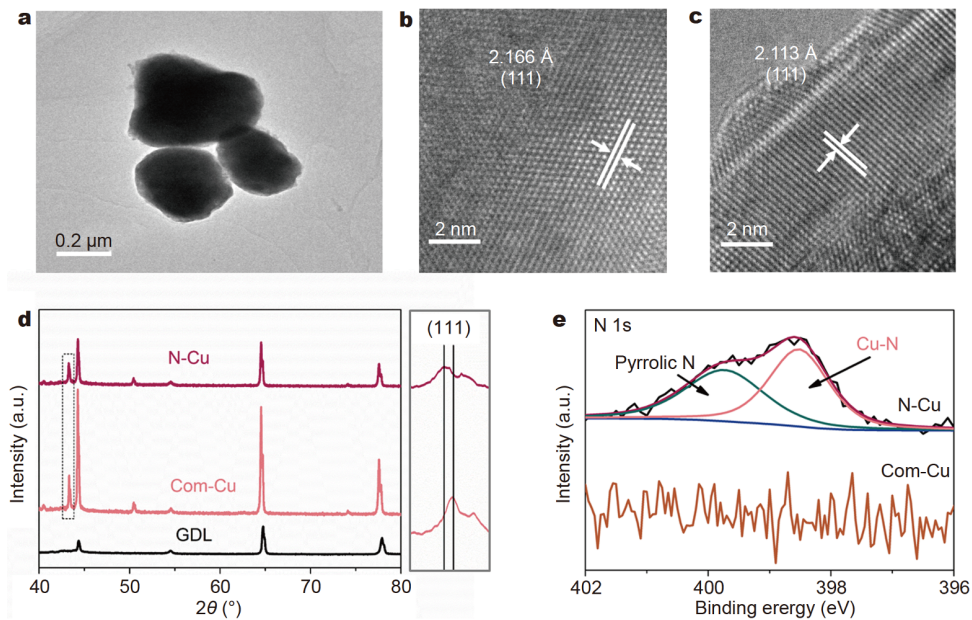
of charge density with respect to the first layer Cu atoms in N-modulated Cu (denoted as N-Cu), indicating that the presence of N withdraws electrons from nearby Cu atoms, further resulting in three positively charged  $\text{Cu}^+$  with formal charge from 0.25 to 0.35 e (Fig. S1). The existence of N also shifts the electronic density of state (DOS) toward Fermi energy (Fig. 1a), referring to a decrease of work function, which is consistent with the charge analysis.

Indeed, from our DFT calculations, we predicted that the presence of N decreases the formation energy of  $^*\text{OC}-\text{COH}$ , as the potential determining step (PDS) of  $\text{C}_2$  products, by  $\sim 0.1$  eV, as shown in Fig. 1c. The selectivity among different  $\text{C}_2$  products, such as ethylene, ethanol and acetate, is determined by the branching of reaction pathways after  $^*\text{OC}-\text{COH}$ . Cheng and coworkers [33] found that the branching of ethylene and acetate is in the early stage after forming  $^*\text{OC}-\text{COH}$ , and distinguished  $^*\text{C}=\text{C}=\text{O}$  as a key intermediate toward acetate formation. The formation of  $^*\text{C}=\text{C}=\text{O}$  is likely from the dehydration of  $^*\text{HOC}-\text{COH}$  via a non-electrochemical reaction. Meanwhile,  $^*\text{HOC}-\text{COH}$  can be electrochemically converted to  $^*\text{C}-\text{COH}$ , which is the precursor of ethylene and ethanol formation [34]. Thus, we considered the reactions from  $^*\text{HOC}-\text{COH}$  to  $^*\text{C}-\text{COH}$  or  $^*\text{C}=\text{C}=\text{O}$  in our DFT simulation to predict the selectivity of acetate on N-Cu.

As demonstrated in Fig. 1c, compared with pure Cu, N dopant significantly lowers the formation energy of

$^*\text{C}=\text{C}=\text{O}$  by 0.36 eV, while the formation of  $^*\text{C}-\text{COH}$  is slightly destabilized by 0.066 eV. Both these trends work in the same direction of promoting the selectivity towards acetate formation. We also calculated the following reactions to complete the reaction pathway of acetate formation. All the remaining reaction steps are exothermic. Thus, according to DFT calculations, we predicted that introducing N to Cu metal to form  $\text{Cu}^+$  could be a feasible strategy to enhance acetate production.

Inspired by the DFT prediction, we sought to incorporate N atoms into Cu lattice *via* a solvothermal synthesis. The as-prepared catalyst was firstly characterized by transmission electron microscopy (TEM) (Fig. 2a). The catalyst demonstrated an irregular morphology, with a particle size of 100–200 nm. The elemental mapping obtained by energy-dispersive X-ray spectroscopy (EDX) (Fig. S2) indicated that N was homogeneously distributed in the Cu. High-resolution TEM (HRTEM) results showed that the inter-planar spacing of N-Cu is 2.166 Å, slightly larger than that (2.113 Å) of commercial Cu (Com-Cu). As the N anion possesses a larger ionic radius than Cu ion, the insertion of N may cause the lattice expansion [35]. These structural distortions were further validated by the powder X-ray diffraction (XRD). As shown in Fig. 2d, the (111), (200) and (220) planes of cubic copper correspond to the diffraction peaks at around 43.3°, 50.4° and 74.1°, respectively. While the diffraction patterns of Com-Cu and N-Cu are very similar, the diffraction peak of N-Cu



**Figure 2** The structural characterizations of N-Cu and Com-Cu. (a) TEM image of N-Cu. (b, c) The HRTEM images of N-Cu (b) and Com-Cu (c). (d) XRD patterns of Com-Cu and N-Cu and the partial magnification of the corresponding diffraction peaks of the (111) facet. (e) XPS N 1s spectra of Com-Cu and N-Cu.

corresponding to (111) facet has slightly shifted to the lower angle region with the incorporation of N. This slight deviation suggests that N doping does not alter the intrinsic structure of metallic Cu but results in a mildly enlarged inter-planar distance. These results point out that N has been successfully introduced into the Cu lattice.

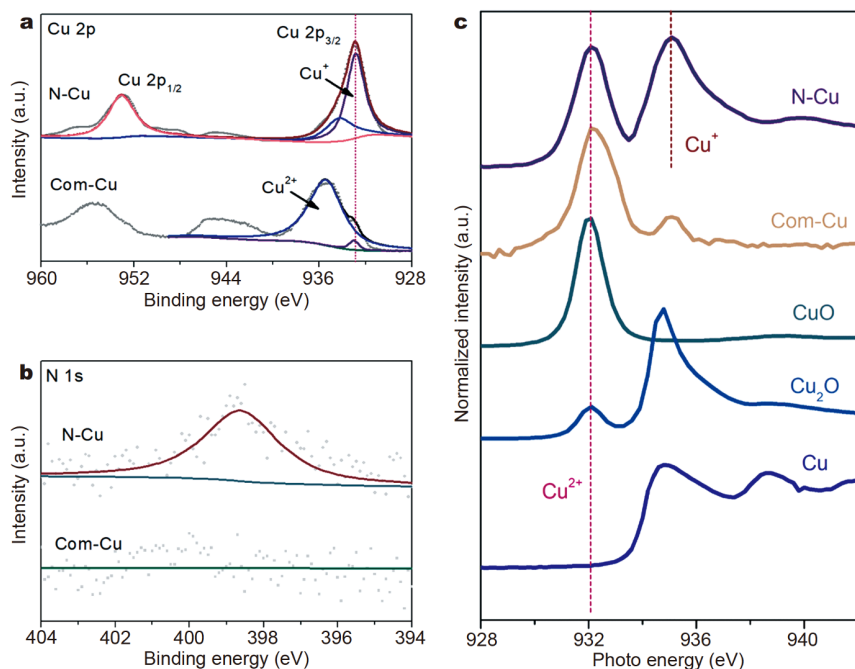
We further conducted X-ray spectroscopy to investigate the chemical states of surface Cu and N atoms. The X-ray photoelectron spectroscopy (XPS) spectrum of N 1s (Fig. 2e) in N-Cu exhibited that the peak located at 398.5 eV is indexed to the Cu–N bond [31]. Meanwhile, soft X-ray absorption spectroscopy (sXAS) was also conducted to examine the existence of N. From the N K-edge sXAS profile of the sample (Fig. S3), the obvious peak in N-Cu verified the presence of N.

To further explore the chemical state of Cu after CORR, we conducted the *ex-situ* XPS and Cu  $L_3$ -edge sXAS spectra of N-Cu and Com-Cu as control samples. In the typical Cu 2p XPS spectra, two peaks at 933 and 952.8 eV are ascribed to the Cu 2P<sub>3/2</sub> and Cu 2P<sub>1/2</sub> of Cu<sup>0</sup>, while the relative two Cu<sup>2+</sup> peaks of Com-Cu are at 934.5 and 955 eV. But the two peaks of N-Cu are obviously broader (Fig. 3a), confirming that its surfaces are enriched with Cu<sup>+</sup> sites [30]. The residual N in N-Cu also verifies the stable structure of N-modulated Cu<sup>+</sup> during the CO reduction (Fig. 3b). As depicted in Fig. 3c, the larger peak

located at 934.8 eV of N-Cu ascertains the presence of surface Cu<sup>+</sup> [36–38]. The above results prove the existence of surface Cu<sup>+</sup> modulated by N dopant, which is consistent with the DFT simulation results, indicating a promise for improving the production of acetate.

To further verify the effect of N-modulated Cu<sup>+</sup> active sites on CORR performance, we conducted chronopotentiometry experiments on N-Cu (Figs S4–S6), and chose Com-Cu (Figs S7–S10) and Cu<sub>2</sub>O nanoparticles as control samples (Figs S11–S14). All tests were carried out in a three-electrode flow cell system in 2.0 mol L<sup>-1</sup> KOH aqueous electrolyte to eliminate the mass transport limitation [39]. A maximal FE<sub>acetate</sub> of 42% was obtained on N-Cu at the potential of -1.27 V vs. reversible hydrogen electrode (RHE), while the Com-Cu exhibited an FE<sub>acetate</sub> of 29% (Fig. 4a, b). Furthermore, the highest FE<sub>acetate</sub> of Cu<sub>2</sub>O (Fig. S13) was 12%, suggesting that Cu<sub>2</sub>O had significantly poorer selectivity of CORR to acetate than N-Cu. However, the FEs of other C<sub>2+</sub> products of Com-Cu and Cu<sub>2</sub>O at lower potentials were all higher than the equivalents of N-Cu. As reflected, this N-modulated Cu<sup>+</sup> strategy dramatically improves the selectivity of acetate by suppressing the formation of ethylene and ethanol, in comparison with Com-Cu and Cu<sub>2</sub>O.

To further investigate the CORR performance, we conducted the cyclic voltammograms on the catalysts in 2 mol L<sup>-1</sup> KOH aqueous electrolyte at the scan rate of



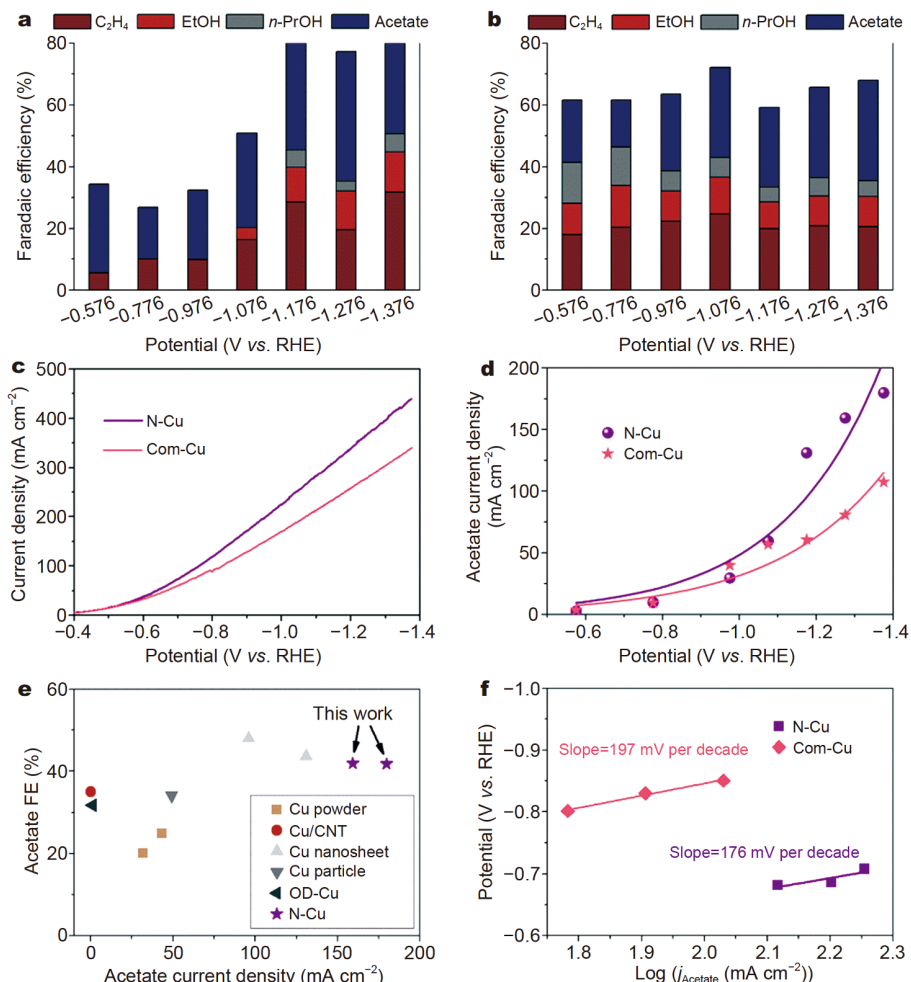
**Figure 3** Characterizations of the surface chemical state of Cu in catalysts after reaction. (a, b) XPS Cu 2p and N 1s spectra of Com-Cu and N-Cu after 30-min reduction of CO, respectively. (c) Soft XAS of Cu  $L_3$ -edge for Com-Cu and N-Cu after CORR for 30 min.

50 mV s<sup>-1</sup>. The linear sweep voltammetry curves in Fig. 4c reveal that the cathodic currents of the two samples quickly increase as the applied potential is more negative than -0.5 V vs. RHE. In addition, compared with Com-Cu, a larger increase of current density for N-Cu was acquired, indicating that N-Cu exhibited higher CO reduction activity. As illustrated in Fig. 4d, the acetate partial current densities of N-Cu and Com-Cu both increase considerably, demonstrating remarkable transfer of CO to the triple-phase reaction interface. Moreover, the highest acetate partial current density of 180 mA cm<sup>-2</sup> was achieved at -1.37 V on N-Cu, which is superior to the performance on Com-Cu, Cu<sub>2</sub>O (Fig. S12) and the state-of-the-art catalysts [19,20,39–41] (Fig. 4e, Table S1), indicating that surface N-modulated Cu<sup>+</sup> active sites remarkably boosted the catalytic activity of Cu. Furthermore, the acetate turnover frequency (TOF) [42] exhibited the same tendency for the N-Cu. The TOF of N-Cu is ~0.048 s<sup>-1</sup> per total Cu atoms at -1.37 V, approximately twice higher than that for Com-Cu. In sum, the highest FE<sub>acetate</sub>, highest acetate partial current densities and the TOF indicate that the Cu<sup>+</sup> species on N-Cu present a higher reduction rate and higher catalytic activity towards acetate.

To further understand the experimental mechanistic pathways for the CORR in the N-Cu catalyst, the Tafel

slope derived from three partial current densities was investigated. Fig. 4f shows the slope values of N-Cu and Com-Cu, which are 176 and 197 mV per decade, respectively. For both samples, their slope values are close to 118 mV per decade, indicating that the rate-determining step of the CORR is the dimerization of two surface-adsorbed \*CO [41,43]. In comparison with Com-Cu, N-Cu owned a lower Tafel slope, revealing superior CO reduction catalytic kinetics on N-Cu. Additionally, to further examine the stability of N-Cu, we tested its performance at a constant current density of 250 mA cm<sup>-2</sup> in 2 mol L<sup>-1</sup> KOH electrolyte (Fig. S15). The applied potential changed little over the course of 1 h of electrolysis with gradual increase and sudden decrease, which arose from the gradual formation and sudden disappearance of gas bubbles in the liquid catholyte chamber.

In summary, with the help of DFT prediction, we proposed a N modification strategy to modulate the electronic structure of Cu-based electrocatalysts. The N-modulated Cu<sup>+</sup> catalyst was highly active and can switch the intermediate transfer from \*COCOH towards \*CCO—the vital intermediate dominating the acetate formation. XPS, XAS and electrochemical studies well elaborated the link between highly efficient conversion of CO to acetate and the positively charged copper. Our N-doped Cu catalyst achieved an FE<sub>acetate</sub> of 42% with a



**Figure 4** CO reduction performances of N-Cu and Com-Cu in a flow cell system. (a, b) FEs of CORR products on N-Cu (a) and Com-Cu (b) at various applied potentials in 2 mol L<sup>-1</sup> KOH electrolyte. (c, d) The electrochemical reduction of CO polarization curves (c) and acetate partial current densities *versus* applied potentials (d) of Com-Cu and N-Cu. (e) The comparison of FEs and partial current densities for acetate between this work and state-of-the-art CO reduction systems [19,20,39–41]. (f) Tafel slopes derived from partial current densities for Com-Cu and N-Cu.

superior partial current density of 180 mA cm<sup>-2</sup>. These findings may improve the electrocatalyst design in the future and promote the industrial application of CO<sub>2</sub>RR/CORR technologies.

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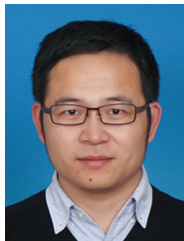
**Author contributions** Peng H, Zhang B and Cheng T supervised the project. Yang H and Cheng T performed the DFT simulations. Ni F and Wen Y carried out the synthesis of catalysts and electrochemical measurements. Bai H and Zhang L performed the TEM experiments. Cui C, Li S and He S performed the sXAS experiments. All authors discussed the results and helped with the manuscript preparation.

**Conflict of interest** The authors declare that they have no conflict of interest.

**Supplementary information** Experimental details and supporting data are available in the online version of the paper.



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## 氮调控正价态铜用于高效电化学还原一氧化碳制备乙酸

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**摘要** 利用间歇性可再生能源电转化二氧化碳和一氧化碳制备多碳工业原料和燃料能够完善碳循环。然而, 二氧化碳转化为乙酸的法拉第效率难以满足大规模应用的要求。研究发现, 调控铜的局域电子结构至正价态, 以及使用一氧化碳作为反应气均能够提高多碳产物的选择性。因此, 我们提出杂原子调控策略, 用于调节催化剂表面活性位点的氧化态。密度泛函理论研究表明, 掺杂氮原子调控正价态铜位点, 在热力学上有利形成中间体\*CCO, 进而生成乙酸。我们进一步利用溶剂热法合成氮掺杂的铜催化剂, 并通过软X射线吸收谱和X射线光电子能谱证实催化剂表面存在大量正价态铜位点。同时, 该催化剂在一氧化碳还原反应中实现了42%的乙酸法拉第效率和已知最高的乙酸偏电流密度180 mA cm<sup>-2</sup>。