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## Pyridyl-containing graphdiyne stabilizes sub-2 nm ultrasmall copper nanoclusters for the electrochemical reduction of CO<sub>2</sub>†

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Developing novel carbonaceous materials with definite chemical structures is conducive to understanding structure–property relationships and expanding their applications in supported metal catalysts. Herein, a brand-new pyridine-substituted graphdiyne (**Py-GDY**) is synthesized through the cross-coupling of 1,3,5-triethynyl-2,4,6-tris(4-pyridyl)benzene, and further applied as a promising carrier in electrocatalysis. Thanks to the precisely introduced pyridyl groups, strong metal–support interaction between the confined Cu species and **Py-GDY** is desirably obtained, resulting in uniformly dispersed Cu sub-nano-clusters (<2 nm) (**Py-GDY-Cu**). Conversely, the Cu size increased dramatically when the pyridyl group of **Py-GDY** was replaced by the phenyl group (**Ph-GDY-Cu**). In a proof-of-concept demonstration of the electrochemical CO<sub>2</sub> reduction reaction, **Py-GDY-Cu** is found to produce CH<sub>4</sub> preferentially to **Ph-GDY-Cu**, owing to the favorable sub-nanocluster size. As a result, an optimum CH<sub>4</sub> faradaic efficiency of 58% is achieved on **Py-GDY-Cu**, which shows a 1.6-fold enhancement compared with that of **Ph-GDY-Cu**. This work broadens the scope of carbonaceous materials for rational metal species immobilization toward efficient catalysis.

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## Introduction

Catalysis not only plays a vital role in the chemical industry but also attracts great attention in the field of fundamental research. Heterogeneous metal catalysts are widely used in diverse heterogeneous catalysis processes. Their supporting materials often determine the performance of metal catalysts by regulating their shapes, morphologies, sizes, and chemical states.<sup>1–3</sup> Carbonaceous materials are a family of supporting materials commonly used in the energy-related research field, and tremendous effort has been devoted to expanding the library of carbonaceous materials.<sup>4,5</sup> Heteroatom-doped carbon materials provide a representative platform for supporting metal catalysts and increasing the dispersion of metal species through strong metal–support interaction (MSI), which is expected to affect the catalytic activity.<sup>6–8</sup> Nevertheless, the current doping

strategies have long been championed by hydrothermal or high-temperature calcination treatment, in which the dopant types and the homogeneity of heteroatoms in carbon materials are difficult to control, putting obstacles in the way of understanding the structure–property relationships.<sup>9,10</sup> Hence, developing heteroatom-doped carbon materials with well-defined chemical structures is highly desirable and remains a challenge in the controllable synthesis of carbon-supported metal catalysts.

Graphdiyne (GDY) has emerged as a new two-dimensional (2D) carbon allotrope prepared *via* the cross-coupling reaction of hexaethynylbenzene.<sup>11</sup> With a conjugated structure composed of benzene rings and diacetylenic linkages ( $-C\equiv C-C\equiv C-$ ), GDY provides defined and uniform anchoring sites for metal atoms.<sup>12–14</sup> A series of single-atom catalysts like Fe, Ni, Cu, Mo, Pd, Pt, Ru, Rh *etc.* has been synthesized on GDY.<sup>12–20</sup> MoO<sub>3</sub> nanoparticles, CuO nanoclusters, and oxidized iridium quantum dots are also successfully supported on GDY.<sup>21–23</sup> The metal atoms are stabilized by the interaction between the d orbitals of metal atoms and the  $\pi/\pi^*$  orbitals of carbon–carbon triple bonds ( $-C\equiv C-$ ). In addition, taking advantage of the bottom-up synthetic strategy, tens of GDY analogues with desired structures have been realized by precisely introducing heteroatoms (B, N, and S) or functional groups ( $-F$ ,  $-Cl$ , and  $-CN$ ) into the corresponding monomers.<sup>24–29</sup> This makes GDY analogues an ideal platform for the precise synthesis of doped carbon materials to support metal catalysts. By the rational

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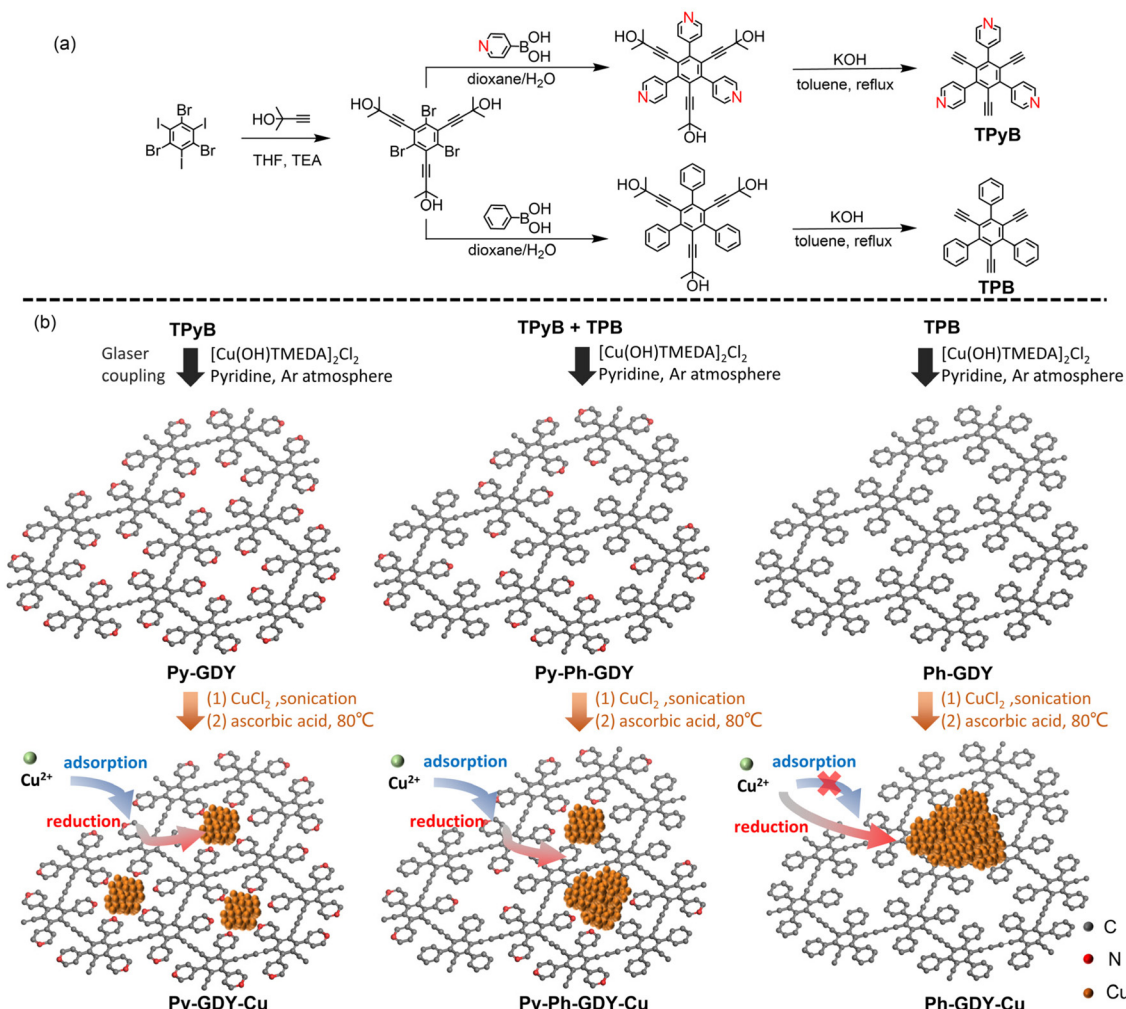


Fig. 1 (a) Synthetic diagram of TPYB and TPB. (b) Schematic illustration for the preparation of Py-GDY, Py-Ph-GDY, Ph-GDY, Py-GDY-Cu, Py-Ph-GDY-Cu, Ph-GDY-Cu.

design of GDY analogues with functional groups, the MSI could be readily manipulated, which may in turn affect the sizes, morphologies, and chemical states of the supported metal catalysts.

Pyridyl groups have been widely used as coordinating ligands in various metal complexes.<sup>30</sup> Adopting the synthetic strategy of GDY, pyridyl groups could be uniformly incorporated into the 2D network of a GDY analogue. The pyridyl groups on the GDY analogue may play a pivotal role in the dispersion of metal atoms *via* strong metal–nitrogen interaction, thus facilitating the construction of nanosized metal catalysts.

Herein, a new pyridine-substituted GDY (Py-GDY; Fig. 1) is successfully synthesized *via* the cross-coupling reaction of 1,3,5-triethynyl-2,4,6-tris(4-pyridyl)benzene (TPYB; Fig. 1). Owing to the merits of the introduced pyridyl group, the strong interaction between the Cu species and Py-GDY leads to the formation of uniformly distributed sub-2 nm ultrasmall Cu clusters (Py-GDY-Cu; Fig. 1). As a comparison, large and uneven Cu nanoparticles were obtained on the other two GDYs, Py-Ph-GDY and Ph-GDY (see their structures in Fig. 1)

with a half content of pyridyl groups and without pyridine groups, respectively. In the electrochemical CO<sub>2</sub> reduction reaction, Py-GDY-Cu showed the highest faradaic efficiency of 58% for CH<sub>4</sub> at -1.0 V vs. a reversible hydrogen electrode (RHE) in 1.0 M KOH, surpassing that of the uneven Cu nanoparticles on Py-Ph-GDY and Ph-GDY. The favorable selectivity for CO<sub>2</sub>-to-CH<sub>4</sub> conversion over Py-GDY-Cu is due to the ultra-small size of the Cu catalyst originating from the presence of coordinating pyridyl groups in the GDY material. The present work not only broadens the scope of the GDY library but also highlights a compelling support material to immobilize metal catalysts toward improved catalysis.

## Results and discussion

### Synthesis and characterization

TPYB was prepared from 1,3,5-tribromo-2,4,6-triiodobenzene in three steps: (i) the Sonogashira reaction; (ii) the Suzuki coupling reaction; and (iii) the base-promoted deprotection reac-

tion. The synthetic procedures are shown in Fig. 1a. All these compounds were thoroughly characterized using  $^1\text{H}$ -NMR,  $^{13}\text{C}$ -NMR spectroscopy, and high-resolution mass spectroscopy (Fig. S1–S10†). Using the Cu-catalyzed Glaser coupling reaction of **TPyB**, **Py-GDY** was produced with a porous framework and evenly distributed pyridyl groups, as shown in Fig. 1b. The transmission electron microscopy (TEM) image of **Py-GDY** (Fig. 2a) reveals a lamellar morphology of **Py-GDY**. Elements of C and N were uniformly distributed, as depicted by the energy dispersive spectroscopy (EDS) mapping images. The X-ray diffraction (XRD) pattern in Fig. S11† displays a broad peak at 21° assigned to the in-plane periodicity, indicating that **Py-GDY** is amorphous but still shows 2D ordering.<sup>31</sup>

The structure of **Py-GDY** was further investigated by the Fourier transform infrared spectroscopy (FTIR), Raman spectroscopy, and X-ray photoelectron spectroscopy (XPS). By comparison of the FTIR spectra of **TPyB** and **Py-GDY** (Fig. 2b), two IR bands at 3290 and 2094  $\text{cm}^{-1}$  are exclusively observed for **TPyB** but not for **Py-GDY**, and these two bands are respectively due to the C–H stretching vibration of  $\text{C}\equiv\text{C}-\text{H}$  and the  $\text{C}\equiv\text{C}$  stretching vibration.<sup>26</sup> Apparently, the terminal acetylenic groups of **TPyB** are converted to, in principle, the diacetylenic linkages *via* the Glaser coupling reaction, leading to the formation of **Py-GDY**. The Raman spectrum (Fig. 2c) of **Py-GDY** displays four bands at 1344, 1546, 1602, and 2185  $\text{cm}^{-1}$ . The band at 2185  $\text{cm}^{-1}$  is attributed to the stretching vibration of conjugated diyne links ( $-\text{C}\equiv\text{C}-\text{C}\equiv\text{C}-$ ),<sup>11</sup> verifying the cross-coupling of the terminal acetylenic groups. For XPS analysis, the C 1s peak of **Py-GDY** (Fig. 2d) is deconvoluted into four peaks of  $\text{C}=\text{C}$  at 284.5 eV,  $\text{C}\equiv\text{C}$  at 285.2 eV,  $\text{C}=\text{N}$  at 285.5 eV,

and  $\text{C}-\text{O}$  at 286.7 eV. The N 1s spectrum of **Py-GDY** (Fig. 2e) depicts only one peak at 398.7 eV, assigned to the N atom of pyridyl groups.<sup>9</sup> All the results above are in good agreement with the proposed structure of **Py-GDY**.

Pyridyl groups are widely used in coordination chemistry for their high tendency to form metal complexes.<sup>30</sup> With its porous structure and abundant pyridyl groups, **Py-GDY** can be a proper support for metal catalysts. To systematically examine the effects of pyridyl groups on stabilizing metal catalysts, we also synthesized another two GDY analogues, **Py-Ph-GDY** and **Ph-GDY**, as shown in Fig. 1b. **Ph-GDY** was prepared by the coupling of 1,3,5-triethynyl-2,4,6-triphenyl-benzene (**TPB**) according to the literature method.<sup>32</sup> Compared with **Py-GDY**, all the pyridyl groups were replaced by phenyl groups in **Ph-GDY**. To further tune the content of pyridyl groups, a mixture of **TPyB** and **TPB** with a molar ratio of 1:1 was used to produce **Py-Ph-GDY**. **Ph-GDY** and **Py-Ph-GDY** were also characterized by FTIR, Raman, and XPS (Fig. S12–S15†). The elemental analysis results show the C/N atomic ratios are 7.7 and 16.6 for **Py-GDY** and **Py-Ph-GDY**, which are close to the values of 9 and 19 calculated from their ideal chemical structures.

After preparing these three GDY analogues, the supported Cu nanocatalysts were further synthesized to investigate the effect of the pyridyl group on the size distribution of the Cu catalysts. In general, **Py-GDY**, **Py-Ph-GDY**, and **Ph-GDY** were respectively dispersed in a  $\text{CuCl}_2$  solution by sonication, and then ascorbic acid was added to reduce  $\text{Cu}^{2+}$  into Cu nanoclusters/nanoparticles.<sup>33</sup> The resulting products are accordingly named **Py-GDY-Cu**, **Py-Ph-GDY-Cu**, and **Ph-GDY-Cu**. According to the EDS spectra (Fig. S16†), Cu is successfully incorporated into **Py-GDY-Cu**, **Py-Ph-GDY-Cu**, and **Ph-GDY-Cu**. But the morphologies of Cu change over the three substrates due to their different MSI, as shown schematically in Fig. 3a. No Cu particles are observed in the TEM image of **Py-GDY-Cu** (Fig. 3b). The aberration-corrected high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) image (Fig. 3c) verifies that Cu nanoclusters with sizes smaller than 2 nm are well distributed throughout the **Py-GDY** support, in line with the EDS spectrum and maps of C, N, and Cu shown in Fig. S16† and Fig. 3d. For **Py-Ph-GDY-Cu**, Cu nanoparticles with sizes of about 10 nm (circled in red in Fig. 3e) are found; besides, Cu nanoclusters with sizes around 1–2 nm are also present on the surface of **Py-Ph-GDY-Cu** as revealed by the aberration-corrected HAADF-STEM image (Fig. 3f). In contrast, the TEM image (Fig. 3h) and the HAADF-STEM image (Fig. 3i) of **Ph-GDY-Cu** disclose the presence of Cu nanoparticles around 100 nm. The distribution of the Cu element is not even as shown in the EDS image of Cu (Fig. 3j). According to the inductively coupled plasma atomic emission spectrometry measurements, the weight percent of Cu was 4.3 wt%, 2.7 wt%, and 2.0 wt% for **Py-GDY-Cu**, **Py-Ph-GDY-Cu**, and **Ph-GDY-Cu** respectively. Both high loading and uniform distribution of Cu nanoclusters are achieved using **Py-GDY** as the support. These phenomena arise from the fact that pyridyl groups on **Py-GDY-Cu** can adsorb  $\text{Cu}^{2+}$  and provide abundant anchoring sites for the nucleation and growth of Cu

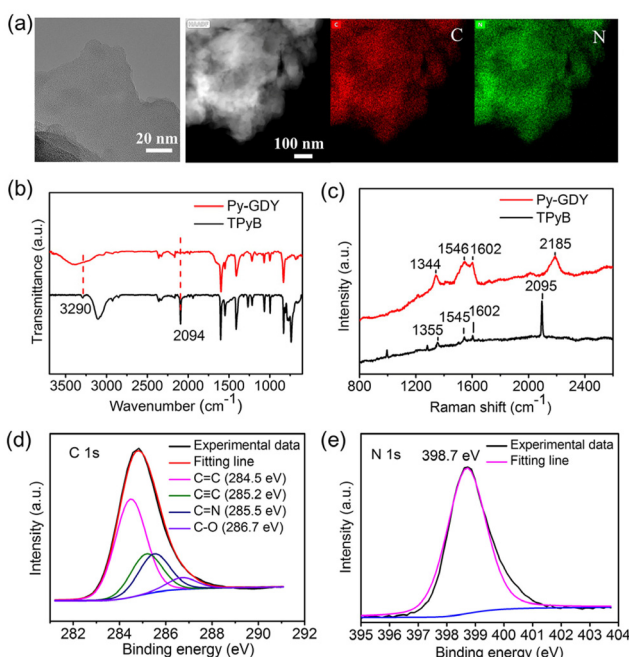
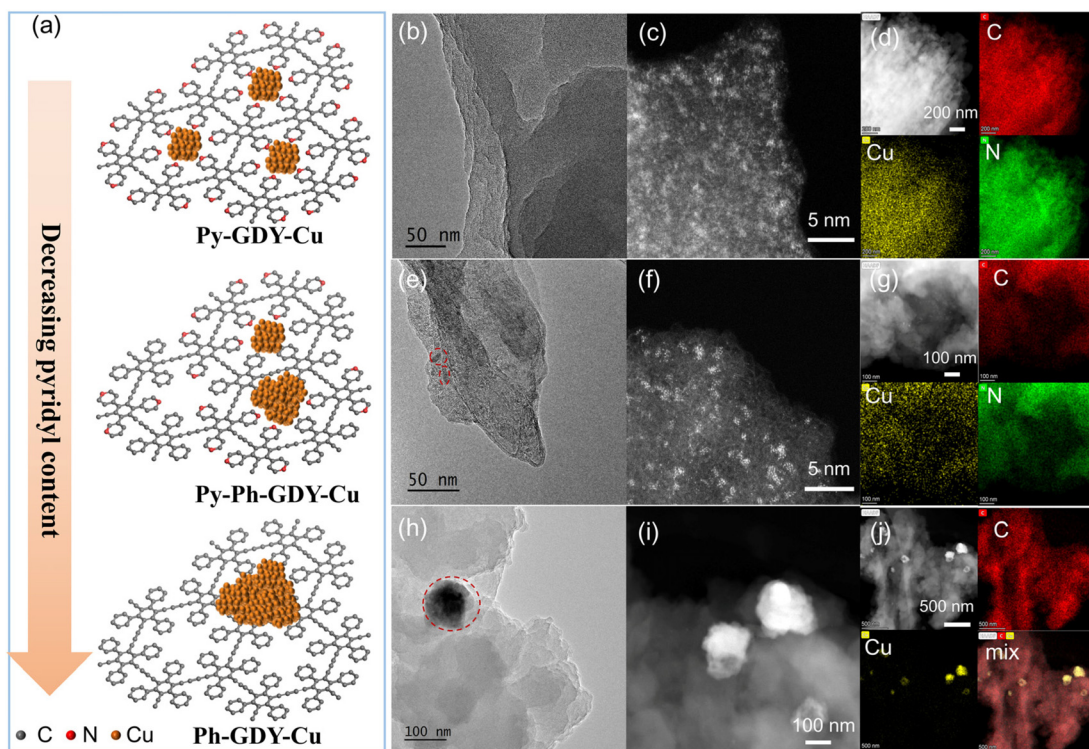


Fig. 2 (a) TEM and EDS mapping images of **Py-GDY**. (b) FTIR spectra of **Py-GDY** and **TPyB**. (c) Raman spectra of **Py-GDY** and **TPyB**. (d) C 1s XPS spectrum, and (e) N 1s XPS spectrum of **Py-GDY**.



**Fig. 3** (a) Schematic illustration of Py-GDY-Cu, Py-Ph-GDY-Cu, and Ph-GDY-Cu. (b) TEM image, (c) aberration-corrected HAADF-STEM image, and (d) EDS mapping images of Py-GDY-Cu. (e) TEM image, (f) aberration-corrected HAADF-STEM image, and (g) EDS mapping images of Py-Ph-GDY-Cu. (h) TEM image, (i) HAADF-STEM image, and (j) EDS mapping images of Ph-GDY-Cu.

nanoclusters. Since phenyl groups are not able to coordinate with Cu atoms, the regular structure composed of both pyridyl and phenyl groups shows a decreased ability to control the size of Cu on the **Py-Ph-GDY-Cu**. Due to the poor interaction between **Ph-GDY** and Cu, Cu easily aggregates into large nanoparticles. The XRD measurement was used to determine the structures of these three samples. In Fig. S17,† no peak of Cu is detected in the XRD patterns of **Py-GDY-Cu** and **Py-Ph-GDY-Cu**, mainly due to the rather small sizes of amorphous Cu nanoclusters.<sup>13,19</sup> **Ph-GDY-Cu** shows three diffraction peaks at  $36.49^\circ$ ,  $43.25^\circ$  and  $50.37^\circ$ , due to the (111) crystal plane of  $\text{Cu}_2\text{O}$  and the (111), (200) crystal planes of metallic Cu. The existence of  $\text{Cu}_2\text{O}$  is due to the oxidation of Cu atoms under an ambient atmosphere.<sup>33</sup>

Further insights into the structural and electronic information on the three samples were obtained from the XPS measurements. In Fig. 4a, the characteristic peaks of C 1s and Cu 2p can be found in all three samples, and only **Py-GDY-Cu** and **Py-Ph-GDY-Cu** possess the signal of N 1s. As shown in Fig. 4b, the Cu 2p<sub>3/2</sub> peaks around 932.9 eV and Cu 2p<sub>1/2</sub> peaks around 952.7 eV in the Cu 2p XPS spectra are associated with  $\text{Cu}^0$  or  $\text{Cu}^+$ .<sup>34</sup> No satellite peaks of  $\text{Cu}^{2+}$  are observed. The Cu LMM Auger spectra of these samples are shown in Fig. 4c. **Py-GDY-Cu** displays the most dominant  $\text{Cu}^+$  peak, while **Ph-GDY-Cu** has the largest fraction of  $\text{Cu}^0$ .<sup>34</sup> This may result from the fact that small Cu nanoclusters are more easily oxidized or that Cu atoms coordinated with pyridyl groups are prone to

oxidation. The N 1s spectra in Fig. 4d demonstrate the interaction between the pyridyl groups and Cu atoms. The peak at 398.7 eV is attributed to the N 1s peak of free pyridyl groups of **Py-GDY**. After decorating Cu nanoclusters or nanoparticles on **Py-GDY** and **Py-Ph-GDY**, a new peak at 399.9 eV appears, indicating that a part of the pyridyl groups coordinates with Cu atoms, leading to a shift in the N 1s peak in the higher energy direction.

#### Electrochemical reduction of $\text{CO}_2$

The electrochemical reduction of  $\text{CO}_2$  has emerged as a promising method for  $\text{CO}_2$  conversion. Cu materials show a unique capability of catalyzing the electrochemical conversion of  $\text{CO}_2$  to various hydrocarbons and alcohols.<sup>35</sup> Improved activity and selectivity of Cu catalysts toward specific products is in high demand. Previous studies show that the “catalytic particle size effect” significantly influences the activity and selectivity of Cu catalysts.<sup>13,36,37</sup> In this work, the Cu nanoparticles ranged from sub-2 to 100 nm by tuning the pyridyl content in the supporting GDY materials. In particular, **Py-GDY-Cu** has pushed the Cu particle size to a sub-2 nm level, where the quantum effect, also known as the “catalytic finite-size effect”, becomes noticeable.<sup>38</sup> With the three supported Cu catalysts in hand, we surveyed their performance in the electrochemical  $\text{CO}_2$  reduction reaction. The experiments were performed using a flow cell reactor, where a  $\text{CO}_2$  stream and the commonly used 1.0 M KOH aqueous electrolyte were fed separately into the

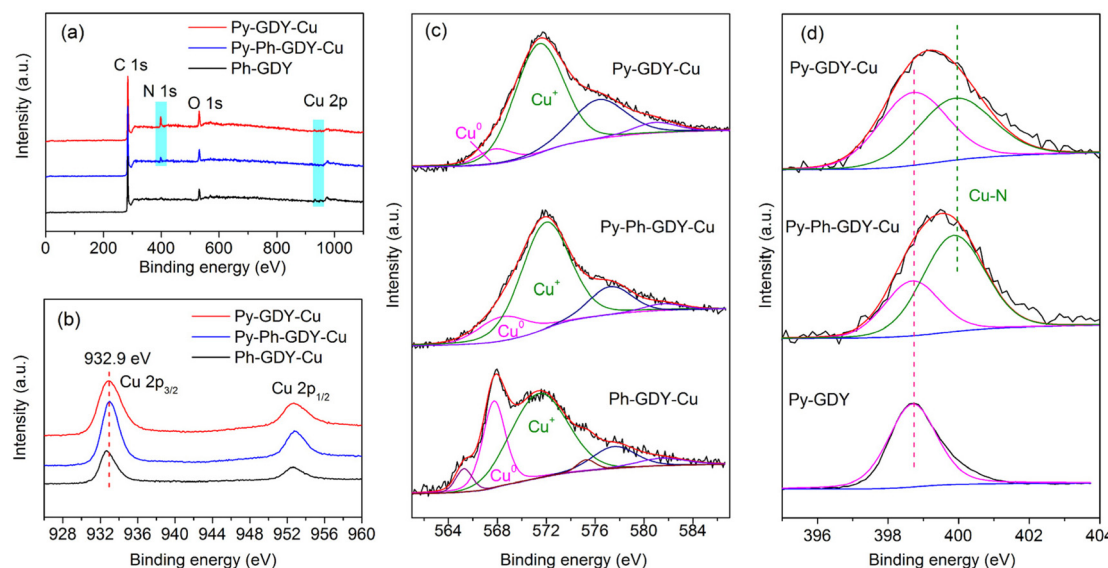


Fig. 4 (a) XPS spectra, (b) Cu 2p spectra, (c) Cu LMM Auger spectra of Py-GDY-Cu, Py-Ph-GDY-Cu, and Ph-GDY-Cu. (d) N 1s spectra of Py-GDY-Cu, Py-Ph-GDY-Cu and Py-GDY.

electrochemical cell through the two sides of the gas-diffusing electrode. Before the electrolysis experiments, electrochemical conditioning was applied for all three catalysts by cyclic voltammetry scanning from 0.5 V to −0.5 V vs. RHE for 60 cycles. After that, steady-state chronoamperometry measurements were conducted to evaluate the products of CO<sub>2</sub> reduction.

As shown in Fig. 5a–c, the electrolysis products consist of CO, CH<sub>4</sub>, HCOOH, C<sub>2</sub>H<sub>4</sub>, CH<sub>3</sub>CH<sub>2</sub>OH, CH<sub>3</sub>COOH, and H<sub>2</sub>. The faradaic efficiency (FE) of CO decreased with applying more negative potentials due to the conversion of CO into other

hydrocarbons.<sup>39</sup> Ph-GDY-Cu showed a higher FE of C<sub>2</sub> products (C<sub>2</sub>H<sub>4</sub>, CH<sub>3</sub>CH<sub>2</sub>OH, CH<sub>3</sub>COOH) than Py-Ph-GDY-Cu and Py-GDY-Cu at all the applied potentials from −0.8 to −1.2 V vs. RHE, as shown in Fig. S18.† The highest FE of 31% for C<sub>2</sub> products was achieved by Ph-GDY-Cu at −1.0 V vs. RHE. The highest FEs of C<sub>2</sub> products for Py-Ph-GDY-Cu and Py-GDY-Cu were 23% and 21% at −0.8 V vs. RHE. As expected, the selectivity of C<sub>2</sub> products increases as the size of the Cu particles increases because the large Cu particles favor the dimerization of \*CO.<sup>13</sup> CH<sub>4</sub> dominates the products under most of the

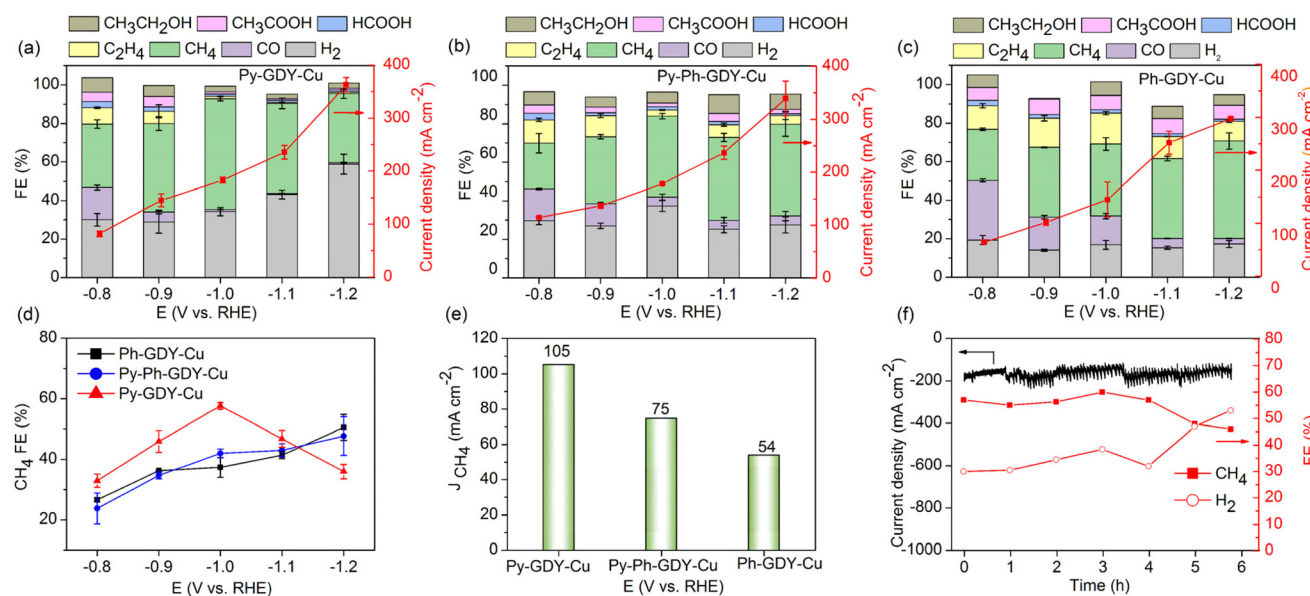


Fig. 5 CO<sub>2</sub> electroreduction performances of (a) Py-GDY-Cu, (b) Py-Ph-GDY-Cu, (c) Ph-GDY-Cu in 1.0 M KOH. (d) FE of CH<sub>4</sub> at −0.8 V to −1.2 V vs. RHE. (e) Partial current density of CH<sub>4</sub> at −1.0 V vs. RHE. (f) Stability test of Py-GDY-Cu and the FE of CH<sub>4</sub>, H<sub>2</sub> measured every 1 h at −1.0 V vs. RHE.

tested potentials. As shown in Fig. 5a, **Py-GDY-Cu** leads to a higher selectivity of  $\text{CH}_4$  than **Py-Ph-GDY-Cu** and **Ph-GDY-Cu** from  $-0.8 \text{ V}$  to  $-1.1 \text{ V}$  *vs.* RHE. The highest FE of  $\text{CH}_4$  reaches 58% for **Py-GDY-Cu** at  $-1.0 \text{ V}$  *vs.* RHE, while **Py-Ph-GDY-Cu** and **Ph-GDY-Cu** show  $\text{CH}_4$  FEs of 42% and 37% at the same potential. On the other hand, **Py-Ph-GDY-Cu** and **Ph-GDY-Cu** show increased selectivity toward  $\text{CH}_4$  with applying more negative potentials; this is due to the increased  $^*H$  coverage and decreased  $^*CO$  coverage at more cathodic potentials, which suppresses the generation of C2 products and facilitates the production of  $\text{CH}_4$ .<sup>40,41</sup> The  $\text{CH}_4$  selectivity of **Py-Ph-GDY-Cu** and **Ph-GDY-Cu** at  $-1.2 \text{ V}$  *vs.* RHE is still slightly lower than that of **Py-GDY-Cu** at  $-1.0 \text{ V}$  *vs.* RHE. In contrast, **Py-GDY-Cu** displays decreased FE of  $\text{CH}_4$  and increased FE of  $\text{H}_2$  at  $-1.1 \text{ V}$  and  $-1.2 \text{ V}$  *vs.* RHE, which is very likely caused by the fact that **Py-GDY-Cu** is more hydrophilic than the other two catalysts, as indicated by their water contact angles in Fig. S19.<sup>†</sup> At  $-1.0 \text{ V}$  *vs.* RHE, the partial current densities of  $\text{CH}_4$  (Fig. 5e) were  $105 \text{ mA cm}^{-2}$ ,  $75 \text{ mA cm}^{-2}$ , and  $54 \text{ mA cm}^{-2}$  for **Py-GDY-Cu**, **Py-Ph-GDY-Cu**, and **Ph-GDY-Cu**, respectively. Previous studies revealed that  $\text{Cu}^+$  species could promote the generation of C2 products<sup>42,43</sup> while small-size Cu catalysts, such as sub-nanometric Cu clusters, favor the production of  $\text{CH}_4$ .<sup>13,36</sup> Which factor dictates the production selectivity of **Py-GDY-Cu**, **Py-Ph-GDY-Cu** and **Ph-GDY-Cu**? In our case, **Py-GDY-Cu** with the largest content of  $\text{Cu}^+$  among these three samples displayed the highest  $\text{CH}_4$  selectivity. Therefore, the product selectivity for our catalysts is not controlled by the Cu oxidation state. In contrast, the size effect could reasonably explain the product selectivity. Small Cu nanoclusters on **Py-GDY-Cu** prefer producing  $\text{CH}_4$  over larger Cu aggregates on **Py-Ph-GDY-Cu** and **Ph-GDY-Cu**, in line with the previous works.<sup>13,36</sup>

The stability of **Py-GDY-Cu** was examined at the potential of  $-1.0 \text{ V}$  *vs.* RHE in the  $1.0 \text{ M KOH}$  electrolyte. In Fig. 5f, no evident decay in the current density is observed after 6 h electrolysis, although some fluctuations exist during the test. The FE of  $\text{CH}_4$  is 57% at the beginning and declines to 46% within 6 h, while the FE of  $\text{H}_2$  increases from 30% to 53%, as shown in Fig. 5f. The change in FE is mainly caused by the flooding of the gas diffusing layer and the salt accumulation in the gas chamber during the electrolysis.<sup>44</sup> The structure of **Py-GDY-Cu** after 6 h electrolysis was characterized by TEM and XRD. As shown in Fig. S20,<sup>†</sup> no XRD peaks of Cu or  $\text{CuO}$  are found, and only the peaks ascribed to  $\text{KHCO}_3$  appear. The aberration-corrected HAADF-STEM image (Fig. S21<sup>†</sup>) shows Cu nanoclusters retained on the surface of **Py-GDY-Cu**. These results indicate good stability of Cu nanoclusters on **Py-GDY** toward the electrochemical reduction of  $\text{CO}_2$ .

## Conclusions

In summary, we have prepared a pyridyl-containing graphdiyne, **Py-GDY**, with evenly distributed pyridyl groups and uniform pores on the conjugated networks. The abundant

pyridyl groups and porous structure of **Py-GDY** provide coordinating sites for anchoring metal atoms, and **Py-GDY-Cu** with sub-2 nm ultrasmall Cu clusters is realized using this novel supporting material. **Py-Ph-GDY-Cu** with decreased content of pyridyl groups possesses both Cu nanoclusters and nanoparticles (about 10 nm). For **Ph-GDY-Cu**, larger and uneven Cu nanoparticles are present. These results indicate the importance of pyridyl groups to restrict the aggregation of Cu and to facilitate the generation of Cu nanoclusters. Regarding the electrochemical reduction of  $\text{CO}_2$ , **Py-GDY-Cu** exhibits the highest FE of 58% for  $\text{CH}_4$  at  $-1.0 \text{ V}$  *vs.* RHE in  $1.0 \text{ M KOH}$ , which is more significant than the highest  $\text{CH}_4$  FE obtained at  $-1.2 \text{ V}$  *vs.* RHE by **Py-Ph-GDY-Cu** and **Ph-GDY-Cu**. The partial current densities of  $\text{CH}_4$  at  $-1.0 \text{ V}$  *vs.* RHE were  $105 \text{ mA cm}^{-2}$ ,  $75 \text{ mA cm}^{-2}$ , and  $54 \text{ mA cm}^{-2}$  for **Py-GDY-Cu**, **Py-Ph-GDY-Cu** and **Ph-GDY-Cu**, respectively. The enhanced selectivity of  $\text{CH}_4$  for **Py-GDY-Cu** is attributed to the small sizes of Cu nanoclusters stabilized by the pyridyl groups. Notably, the sub-2 nm ultrasmall Cu clusters of **Py-GDY-Cu** do not aggregate into large nanoparticles during  $\text{CO}_2$  reduction. Overall, the precise installation of pyridyl groups into carbon-rich materials has been successfully illustrated by taking advantage of the bottom-up wet chemistry synthesis method of graphdiynes. The strong coordination pyridyl groups play a significant role in stabilizing ultrasmall Cu nanoclusters. Our strategy paves the way for synthesizing well-defined nitrogen-containing carbon materials and lays a foundation for the synthesis of ultrasmall metal nanoclusters.

## Author contributions

L. D. conceived the project and designed the experiments. H. D. carried out the experiments and the electrochemical measurements. H. Z., T. S., L. G., C. H., contributed to TEM characterization. S. W., H. L., and H. X. contributed to the synthesis of several compounds and the Raman characterization. H. D., H. Z., and L. D. wrote the manuscript. All authors commented on the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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