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Three dimensional cyclic trinuclear units based metal–covalent organic frameworks for electrochemical CO₂RR†

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A three-dimensional metal–covalent organic framework (3D-MCOF) based on cyclic trinuclear units was synthesized using organic tetrahedral linkers and copper-based cyclic trinuclear complexes. The novel type of 3D-MCOF, named 3D-CTU-MCOF, with the *ctn* topology, is reported herein for the first time. Our study demonstrated enhanced electrocatalytic capacity for CO₂ reduction reaction of 3D-CTU-MCOF compared to independent cyclic trinuclear units.

In reticular chemistry, extended structures are assembled from building blocks linked by strong bonds.¹ Metal–organic frameworks and covalent-organic frameworks have been widely discussed.² It is well known that the dimensions of porous materials influence their porosities and applications.³ For two-dimensional frameworks, the building units are linked to form layers stacked by weak interactions, resulting in one-dimensional channels.⁴ Unlike two-dimensional reticular frameworks, crystalline porous materials with three-dimensional structures generally have interconnecting channels, higher specific surface areas, and porosities.⁵ Therefore, applications for crystalline porous materials with three-dimensional structures have attracted great interest.⁶

Functionality is considered critical to leverage their high specific areas and structural tunability.⁷ metal–covalent-organic frameworks (MCOFs), which combine coordination chemistry and dynamic covalent chemistry, have generated interest in constructing novel porous materials.⁸ In particular,

it is important to incorporate functional building blocks into their frameworks; however, constructing functionalized frameworks often requires complex ligand synthesis. Cyclic trinuclear units (CTUs) have received much attention in developing functional coordination complexes due to their potential in various applications.⁹ MCOFs can act as proper scaffolds to anchor CTUs, which has been shown to endow MCOFs with innovative functions.¹⁰ However, MCOFs containing CTUs are underreported, particularly those with three-dimensional structures.¹¹ While planar copper-based CTUs have been efficiently catalytic in 2D MCOFs, few 2D CTU-based frameworks have been reported and have found applications in catalysis.¹² Three-dimensional MCOFs containing CTU have not yet been reported.

Considering these factors, we introduced a 3D-CTU-MCOF based on *ctn* topology. The 3D-CTU-MCOF exhibited high crystallinity and had abundant copper sites in its framework. It exhibited superior performance to the CTUs(Cu₃L) in electrochemical CO₂ reduction reaction (CO₂RR). To our knowledge, this is the first example of three-dimensional CTU-based MCOF.

The construction strategy of 3D-CTU-MCOFs involves two steps that combine coordination chemistry and dynamic covalent chemistry. In Fig. 1a, copper ions are junction points that link 1*H*-pyrazole-4-carbaldehyde through coordination interactions, producing a planar 3-connected building block named Cu₃L. This structure is not commonly seen in MCOFs. Additionally, we utilize 1,3,5,7-tetraaminoadamantane (TAA, Fig. 1b) as the tetrahedral building unit. The combination of Cu₃L and TAA through condensation results in 3D MCOFs, identified as 3D-CTU-MCOF (Fig. 1c). Notably, the 3D-CTU-MCOF adopts a non-interpenetrated *ctn* net, as depicted in Fig. 1d.

3D-CTU-MCOF was synthesized under solvothermal conditions by transferring Cu₃L and TAA into a 5 : 5 (v/v) mixture of 1,4-dioxane and mesitylene with 6 M aqueous acetic acid as a catalyst. Subsequently, the reaction mixture was heated at 120 °C for 3 days, and yellow microcrystalline powders were collected. The structure of 3D-CTU-MCOF was characterized by a range of measurements. The morphology of the as-synthesized

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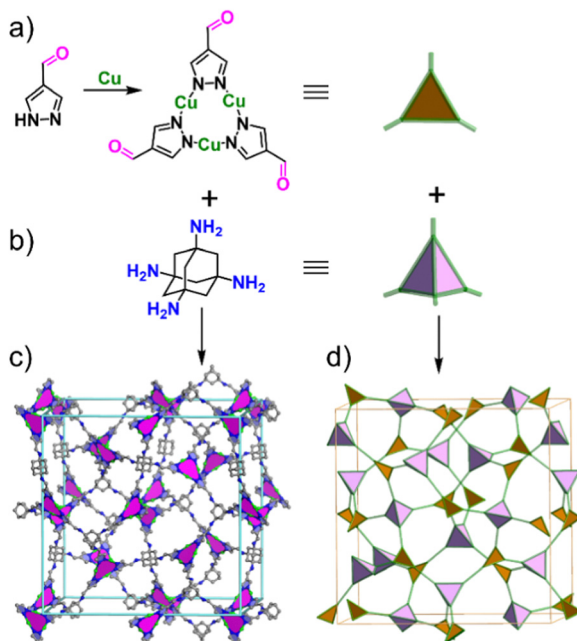


Fig. 1 Illustration of the strategy for building 3D-CTU-MCOF. (a) The mechanism of preparing Cu_3L . (b) The structure of TAA. (c) The crystal structure of 3D-CTU-MCOF (hydrogen atoms are omitted). (d) 3-connected units and 4-connected units to give a **ctn** net of 3D-CTU-MCOF.

3D-CTU-MCOF was studied by scanning electron microscopy (SEM) and transmission electron microscopy (TEM), where rod-shaped crystals of several micrometers were observed (Fig. 2c and d). The integrity of the 3D-CTU-MCOF was investigated by Fourier transform infrared (FT-IR) spectra and solid state ^{13}C cross-polarization magic-angle-spinning (CP/MAS) NMR spectra. The FT-IR spectra (Fig. S1, ESI[†]) revealed the $\text{C}=\text{O}$ stretching modes from Cu_3L at 1670 cm^{-1} , and this peak

became absent after conjugation with TAA. A new peak around 1625 cm^{-1} for 3D-CTU-MCOF indicated the presence of imine bonds.

Additionally, the disappearance of the $\text{C}=\text{O}$ stretching modes indicated the complete transformation of the formyl groups. In the solid-state NMR spectrum, the chemical shift of 153 ppm , attributed to the presence of $\text{C}=\text{N}$, also demonstrated the formation of 3D-CTU-MCOF. The oxidation state of Cu was determined by X-ray photoelectron spectroscopy (XPS), and the binding energy of 932.8 eV implied the presence of $\text{Cu}(\text{i})$ ions in this framework, and the $\text{Cu}2\text{p}$ binding energy of 934.8 eV could be assigned to $\text{Cu}(\text{ii})$ due to partial surface oxidation (Fig. S7, ESI[†]). Additionally, the TGA curve demonstrated that the 3D-CTU-MCOF decomposed at around $177\text{ }^\circ\text{C}$ (Fig. S3, ESI[†]). When the sample was immersed in various organic solvents and water for 24 h , PXRD analyses demonstrated that its crystallinity was still preserved (Fig. S5, ESI[†]).

The crystal structure of 3D-CTU-MCOF was determined by powder X-ray diffraction analysis, and its structural model adopting the **ctn** topology was simulated using Materials Studio software. As a result, it crystallized in the cubic space group, $\text{I}\bar{4}3\text{d}$ ($a = b = c = 37.8218\text{ \AA}$ and $\alpha = \beta = \gamma = 90^\circ$). Moreover, full profile pattern matching (Pawley) refinements were also carried out on the experimental PXRD patterns. The refinement results matched well with the observed patterns ($R_{\text{wp}} = 5.76\%$ and $R_{\text{p}} = 3.59\%$) (Fig. 2a). Peaks at $5.69, 6.82, 7.50, 8.81$ and 11.44° correspond to the $(211), (220), (310), (321)$ and (332) Bragg peaks. We investigated the alternative structure based on the **bor** net. As shown in Fig. S4 (ESI[†]), the experimental pattern did not agree with the simulated one. These results further corroborated that the 3D-CTU-MCOF adopted the expected **ctn** topology. The specific surface area and porosity were characterized by N_2 adsorption and desorption measurements (Fig. 2b). The type I isotherm indicated its microporous character, consistent with the pore size distribution (Fig. S6, ESI[†]). The Brunauer–Emmett–Teller (BET) surface area and pore size were $424\text{ m}^2\text{ g}^{-1}$ and 1.4 nm , respectively, which agreed with the proposed structural model. Notably, the 3D-CTU-MCOF showed a non-interpenetrated **ctn** topology which will facilitate the development of three-dimensional metal covalent organic frameworks.

Encouraged by the interesting results mentioned above, we further explored its performance in electrochemical CO_2RR . It is well known that CO_2 , an abundant source of $\text{C}1$, can be catalytically converted to carbon-based chemicals using copper-containing catalysts. Herein, we propose that 3D-CTU-MCOF with active Cu sites may be suitable for electrochemical CO_2RR . Compared to independent Cu-based CTU, the 3D-CTU-MCOF may have a better activity that can be tuned through the framework formation.

In general, the electrochemical CO_2RR measurements of 3D-CTU-MCOF were performed using a standard three-electrode setup with a carbon paper working electrode, a platinum sheet counter electrode, and Ag/AgCl as the reference electrode. As shown in Fig. 3a, 3D-CTU-MCOF exhibited an enhanced current density in a CO_2 -saturated electrolyte

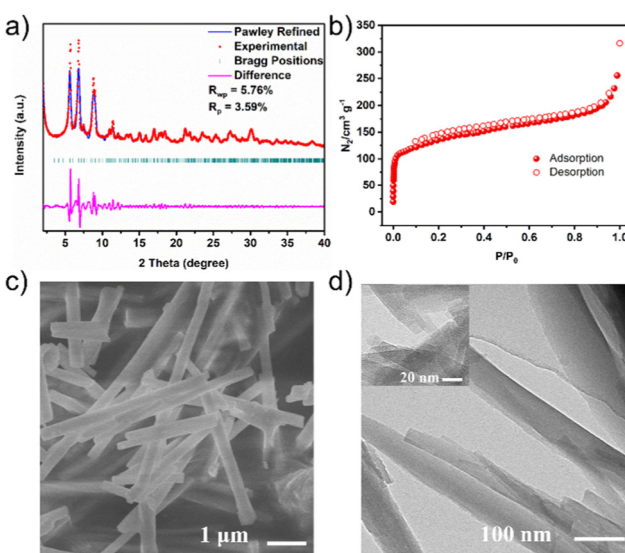


Fig. 2 (a) PXRD patterns of 3D-CTU-MCOF. (b) N_2 adsorption–desorption isotherms of 3D-CTU-MCOF. (c) SEM images of 3D-CTU-MCOF. (d) TEM images of 3D-CTU-MCOF.



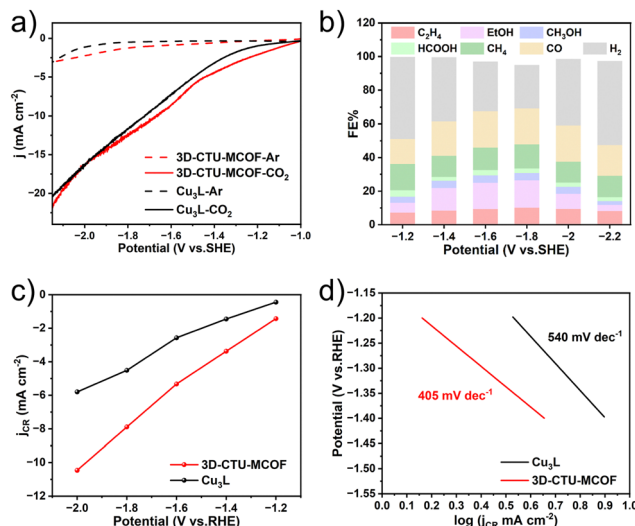


Fig. 3 ECO₂RR activity evaluation of 3D-CTU-MCOF. (a) LSV curves of two catalysts in CO₂/Ar-saturated 0.1 M EmimBF₄-MeCN solutions. (b) ECO₂RR FE% for 3D-CTU-MCOF at different potentials. (c) j_{CR} for two catalysts. (d) Tafel plots for j_{CR} .

compared to that in an argon-saturated electrolyte. Moreover, 3D-CTU-MCOF showed a more significant current density than Cu₃L over a potential range of -1 to -2.15 V vs. SHE in an argon-saturated electrolyte. Besides, in a CO₂-saturated electrolyte (0.1 M EmimBF₄-MeCN solution), the initial potential of 3D-CTU-MCOF showed a positive shift. All results indicated lower reduction overpotential and better performance of 3D-CTU-MCOF compared to Cu₃L.

The product analysis was carried out using gas chromatography and ion chromatography. The total faradaic efficiency (FE) of carbon-based products with 3D-CTU-MCOF as the catalyst remained above 50% throughout the applied potential range (-1.2 V \sim -2.2 V vs. SHE) (Fig. 3b), which was significantly higher than that with Cu₃L as the catalyst (Fig. S9, ESI[†]). The FE of the carbon products reached 69.2% at -1.8 V vs. SHE, with an FE of 26.4% for C₂ products. As shown in Fig. 3c, the partial current densities for both materials were dependent on the applied potential, and the higher density for 3D-CTU-MCOF indicated a higher CO₂ reduction efficiency. The Tafel slope of 405 mV dec⁻¹ for 3D-CTU-MCOF was smaller than that for Cu₃L (540 mV dec⁻¹), implying that 3D-CTU-MCOF was more favourable for CO₂ reduction in the kinetic process than Cu₃L (Fig. 3d). The electrochemically accessible surface area (ECSA) of 1.22 mF cm⁻² for 3D-CTU-MCOF was higher than that of 1.13 mF cm⁻² for Cu₃L, demonstrating the higher catalytic activity of 3D-CTU-MCOF (Fig. S8, ESI[†]). The ICP analysis revealed no copper ions in the filtrate after the electrolysis. In addition, the EIS results also indicated that 3D-CTU-MCOF had a superior kinetic property (Fig. S10, ESI[†]).

In conclusion, we have introduced the copper-based cyclic trinuclear units into three-dimensional metal covalent organic frameworks for the first time. A new CTU-based MCOF adopting **ctn** topology was successfully prepared and showed high crystallinity. Furthermore, 3D-CTU-MCOF displayed better

catalytic activity than Cu₃L in electrochemical CO₂RR. The results suggest combining coordination and dynamic covalent chemistry to anchor active sites in the backbone can improve catalytic activity. Moreover, the successful preparation of 3D-CTU-MCOF paves the way for constructing three-dimensional functionalized crystalline porous materials.

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Conflicts of interest

There are no conflicts to declare.

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