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Introduction

Metal-organic frameworks (MOFs) are an emerging class of crystalline porous materials composed of metal nodes and organic ligands.¹ Due to their tunable structures, accessible open metal sites, and extremely large surface areas, MOFs have shown unprecedented performance in a variety of applications.^{2–7} To date, millions of distinct metal-organic frameworks have been synthesized by combining metal ions and organic linkers. However, their coupling is not arbitrary and is usually restricted by Pearson's hard and soft acid and base (HSAB) theory. Hard acids prefer to combine with hard bases and soft acids prefer soft bases.^{8,9} For instance, hard acidic metal ions such as Zr⁴⁺, Fe³⁺, Cr³⁺, and Al³⁺ are used to form stable MOFs with carboxylate-based ligands that act as

Fabrication of a Cu(ı)-carboxylate metal-organic framework by reduction of metal nodes for an azide-alkyne "click" reaction[†]

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Metal–organic frameworks (MOFs) permit significant structural diversity and functional design flexibility due to the various combinations of metal centers and different organic linkers. However, their combinations generally adhere to the classic hard and soft acid and base (HSAB) theory. This makes it impossible to directly synthesize desired MOFs with converse Lewis types of metal ions and ligands. Herein, we present a novel Cu(II) ion cleavage reduction strategy that circumvents the limitations of HSAB theory. We demonstrate this approach by utilizing a metastable CuH₂DOBDC MOF, whose inherent structural instability facilitates structural transformations and phase transitions in response to external stimuli. By leveraging the unique structure of ascorbic acid to reduce Cu(II) ions and cleave the H₂DOBDC^{2–} linkers, we successfully fabricate a new Cu₂H₂DOBDC structure composed of a soft acid Cu(I) and a hard base (H₄DOBDC). The resultant Cu₂H₂DOBDC integrates the characteristics of mesoporosity and hierarchical nanostructures, resulting in excellent mass transfer and abundant accessible Cu(I) active sites, which endows the material with outstanding catalytic activity in the azide–alkyne cycloaddition reaction. This is the first example of fabricating stable single-phase Cu(I)-carboxylic acid MOFs with nearly all Cu(I) ions, and this work offers a new perspective on the creation of a new category of MOFs with soft-acid and hard-base type coordination bonds.

hard bases, such as UiO-66, MIL-101, and MIL-53.¹⁰ Soft acidic metal ions such as Cu^{1+} and Ag^{1+} are used to form stable MOFs with nitrogen- or sulfur-containing soft base ligands, such as MAF-2 and CityU-7.^{11,12} Therefore, it is normally difficult to achieve a class of MOFs based on soft-acid and hard-base type coordination bonds *via* direct reactions of metal ions and organic ligands.

Functionalization of metal-organic frameworks (MOFs) with Cu(1) sites holds significant promise for various applications including adsorption, separation, catalysis, and sensing.13-18 However, according to the above HSAB theory, Cu (I)-MOFs are usually fabricated using ligands with pyridyl N- or sulfhydryl S-centers. Although carboxylate-based MOFs constitute the overwhelming majority, Cu(1)-MOFs fabricated with carboxylate-based ligands are expected to exhibit thermal stability and have broad catalytic applications.¹⁹ Up to now, the direct synthesis of MOFs with Cu(1) metal ions and carboxylate-based ligands has not been reported. Most commonly, Cu(I)-carboxylate MOFs are synthesized using post-treatment approaches. For example, Chen et al. reported that HKUST-1, in the presence of hydroquinone, was reduced to mixedvalence Cu(I)Cu(II)-BTC and could remarkably boost the D_2/H_2 isotope separation.¹⁴ Sun et al. modulated some of the Cu(II) metal nodes in HKUST-1 to Cu(1) by reducing vapor treatment,

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Fig. 1 Schematic illustration of the fabrication of a $\mbox{Cu(i)-carboxylate MOF}.$

thereby significantly enhancing the capture activity of aromatic sulfides.²⁰ To our knowledge, only Qiao *et al.* have recently reported a "molecular scalpel" to chemically cleave conventional CuBDC with a controlled phase transition and synthesize a Cu₂BDC MOF, where the Cu(I)/Cu(I) ratio is 1.²¹ Although the post-treatment approach is an effective strategy for creating new Cu(I)-carboxylate MOFs, achieving a near equal proportion of Cu(I) and Cu(I) sites in the MOF is far from satisfactory. Hence, it is highly desirable to develop an efficient and controllable approach for creating a new MOF with a high proportion of Cu(I) metal ions and carboxylate-based ligands, but this remained a challenge until now.

Herein, we have successfully fabricated a stable singlephase Cu(1)-carboxylate MOF (denoted as Cu₂H₂DOBDC), as shown in Fig. 1. We exploited the unique characteristics of metastable MOF structures, which are prone to undergo structural transformation under external stimulation, to achieve full phase transition. Starting with a reported metastable $Cu(\pi)$ -MOF synthesized from Cu(OAc)₂ and 2,5-dihydroxyterephthalic acid (H_4 DOBDC), we demonstrated that this metastable Cu(II)-MOF could be reduced to construct a novel Cu(I)-carboxylate MOF in the presence of ascorbic acid. A controlled phase transition was achieved by ascorbic acid accurately regulating the chemical state and coordination environment of Cu metal centers. X-ray photoelectron spectrum (XPS) analysis confirmed that nearly all Cu(II) ions were converted to Cu(I) ions. Cu₂H₂DOBDC, with its high chemical stability, mesoporosity, and uniformly abundant Cu(1) exposure sites, exhibits excellent catalytic capability as a recyclable heterogeneous catalyst for the azide-alkyne cycloaddition (AAC) reaction. To the best of our knowledge, this is the first attempt to fabricate a stable single-phase Cu(1)-carboxylate MOF with near-full Cu(1) ions. It is always fascinating to discover new MOF structures, and this work presents a promising step toward the synthesis of desired MOFs with converse Lewis types of metal ions and ligands, which is impossible to realize by conventional methods.

Results and discussion

Structural characterization

The metastable $Cu(\pi)$ -MOF ([$Cu(H_2DOBDC)(DMF)_2$]·DMF)²² (denoted as CuH_2DOBDC) was synthesized by the reaction of

 $Cu(OAc)_2$ and H₄DOBDC at room temperature, according to our group's previous work with a few modifications. Scanning electron microscopy (SEM) images showed that the as-prepared metastable Cu(II)-MOF exhibited a typical rhombohedron morphology with a smooth surface (Fig. 2a). Single-crystal X-ray diffraction indicated that the metastable Cu(II)-MOF crystallized in the triclinic P1 space group and formed a two-dimensional network layer (Fig. S1†). In addition, the powder X-ray diffraction (XRD) pattern of the crystal was consistent with the pattern simulated based on single-crystal XRD data (Fig. S2⁺), verifying the crystal phase purity. Following ascorbic acid treatment, it exhibited a well-defined cube structure with a substantial decrease in size (Fig. 2b). Notably, the surface was no longer smooth and had been replaced by hierarchical nanostructures. The crystallinity of the nanostructure was further confirmed by selected area electron diffraction (SAED) data, which indicated that the cube structure is a single crystal, as shown in Fig. 2c and d.

The XRD pattern showed that the metastable $Cu(\pi)$ -MOF achieved a full phase transition through the reduction of ascorbic acid (Fig. 2e). As the dosage of the reducing agent increased, the new characteristic peaks at 10.65°, 17.02°, 29.02°, and 42.20° gradually increased in intensity, while the metastable $Cu(\pi)$ -MOF characteristic peaks gradually faded (Fig. S3†). By adjusting the addition of ascorbic acid, the controlled synthesis of intermediate samples was achieved. The coexistence of two phases is observed in the XRD patterns.

Thermogravimetric (TGA) analysis was employed to analyse the thermostability, as variations in coordination mode may present different mass loss curves. As shown in Fig. 3a, following treatment with ascorbic acid, the experimentally determined mass percentage of CuO in the final product was 44.63%. The final product exhibited a weight loss of about 6.28% in the temperature range of 30–220 °C, which can be attributed to the desorption of physically adsorbed water and solvent. Furthermore, a weight loss of about 49.09% was observed in the temperature range of 220–330 °C, which belonged to ligand decomposition. TGA analysis showed that the ratio of Cu to the H₄DOBDC ligand significantly changed from 1:1 to 2:1 following ascorbic acid treatment, indicating a potential structural transformation in the material (see detailed analysis in the ESI†). Consequently, the as-synthesized

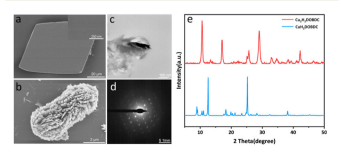


Fig. 2 (a) SEM images of CuH_2DOBDC . (b) SEM images of Cu_2H_2DOBDC . (c) TEM images of Cu_2H_2DOBDC . (d) SAED pattern of the selected area in (c). (e) XRD patterns of CuH_2DOBDC and Cu_2H_2DOBDC .

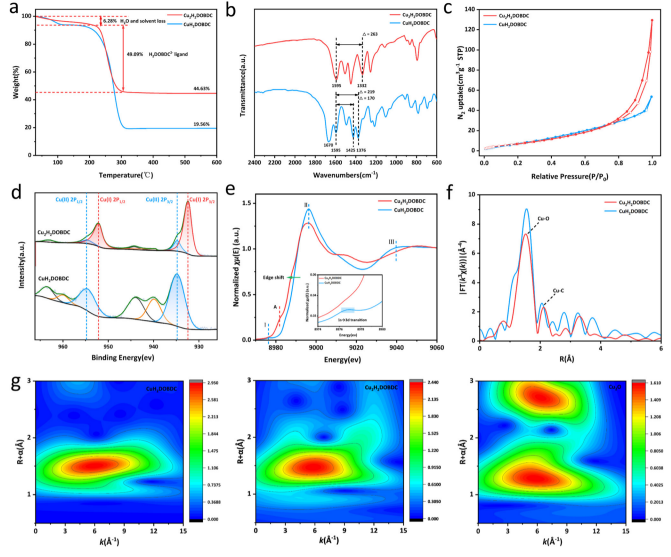


Fig. 3 (a) TGA curves for CuH_2DOBDC and Cu_2H_2DOBDC under an air atmosphere. (b) FTIR spectra of CuH_2DOBDC and Cu_2H_2DOBDC . (c) N_2 adsorption–desorption isotherms of CuH_2DOBDC and Cu_2H_2DOBDC . (d) XPS spectra for Cu 2p. (e) Cu K-edge XANES spectra. (f) Cu K-edge EXAFS spectra. (g) Wavelet transform plots the Cu K-edge EXAFS signals for CuH_2DOBDC , Cu_2H_2DOBDC and the Cu_2O standard.

Cu(I)-carboxylate MOF was denoted as Cu₂H₂DOBDC. Organic elemental analyzer (EA) results further support our conclusions (see detailed analysis in the ESI[†]). In addition, the dropping H₂DOBDC²⁻ ligand was detected by ultravioletvisible (UV-vis) spectroscopy in a reducing solution (Fig. S4[†]). There is a gradual increase in the residual Cu mass percentage observed from CuH₂DOBDC to Cu₂H₂DOBDC, a trend that is further supported by ICP-MS analysis (Table S1[†]). These results further strengthen the conclusion of TGA. The difference thermogravimetry (DTG) analysis suggested that the thermal stability of Cu₂H₂DOBDC is weakened due to the formation of Cu(I) species and the variation of coordination mode (Fig. S5[†]).²⁰ Fourier-transform infrared spectroscopy (FTIR) was employed to investigate the coordination mode between Cu and the carboxylate linker.23 Conclusions about the carboxylate coordination mode are often drawn based on

the splitting value between the asymmetric and symmetric stretches of the carboxylic vibration, where $\Delta = \nu_{as}(COO^{-}) - \nu_{as}(COO^{-})$ $\nu_{\rm s}({\rm COO}^-)$ (Fig. S6[†]).^{21,24} The metastable CuH₂DOBDC MOF contains 2,5-dihydroxyterephthalic acid ligands with two coordination modes: carboxylate groups that connect to a single copper atom through monodentate coordination and those that bridge two copper atoms simultaneously. The FTIR spectra of CuH2DOBDC and Cu2H2DOBDC are shown in Fig. 3b. In CuH₂DOBDC, the asymmetric stretch of the carboxylic vibration peak appeared at 1595 cm⁻¹, while the symmetric stretch peak presented two distinct bands at 1425 and 1376 cm⁻¹.²⁵ Consequently, the observed splitting values of 170 cm⁻¹ and 219 cm⁻¹ in CuH₂DOBDC correspond to bridging and monodentate characteristics, respectively. In addition, the spectra of CuH2DOBDC exhibited a peak at 1670 cm⁻¹ attributed to the carbonyl stretching vibration of

DMF. These results are consistent with the CuH₂DOBDC crystal structure. In contrast, for Cu₂H₂DOBDC, the asymmetric stretch of the carboxylic vibration peak also appeared at 1595 cm⁻¹, but the symmetric stretch peak was only observed at 1332 cm⁻¹. Therefore, a splitting value of 263 cm⁻¹ is consistent with the monodentate mode, indicating that in Cu₂H₂DOBDC, the Cu ions are coordinated to the carboxylate ligands in a monodentate mode. Notably, the absence of a peak corresponding to the carbonyl stretching vibration of DMF in the spectrum suggests that DMF does not participate in coordination within Cu₂H₂DOBDC. From the N₂ adsorption-desorption measurement, Cu₂H₂DOBDC exhibited a type IV isotherm with a distinct hysteresis loop (Fig. 3c), indicating the existence of mesopores,^{26–28} which could be attributed to the ligand loss during the reduction process.²⁰ The pore size distributions shown in Fig. S7[†] reveal that the size of the mesopores is centered at about 17 nm. The SEM images reveal that the surface of Cu₂H₂DOBDC exhibits a hierarchical nanostructure assembled by nanosheets, where the intersections and stacking of the surface hierarchical nanostructures also contribute to the formation of mesopores.²⁹

X-ray photoelectron spectroscopy (XPS) was used for the evaluation of the chemical states and quantitative analysis of Cu species in the samples (Fig. 3d). The Cu 2p spectrum of CuH₂DOBDC exclusively demonstrated the presence of Cu species in the form of Cu(II) ions. However, in Cu_2H_2DOBDC , new characteristic peaks appeared at 932.3 eV and 952.0 eV, which can be attributed to the Cu(1) species. Furthermore, a significant decrease in the intensities of the characteristic peaks associated with Cu(II) was observed, along with the nearabsence of satellite peaks. This significant transformation indicates a reduction process, where nearly all Cu(II) ions in CuH₂DOBDC were reduced to Cu(I) ions following treatment with ascorbic acid. However, two small residual peaks of $Cu(\pi)$ at 934.8 eV and 954.59 eV were observed, which could be attributed to the unavoidable partial oxidation of the sample due to its exposure to air during the reaction or sample processing.^{30,31} The chemical state and coordination environment of the Cu atoms in CuH2DOBDC and Cu2H2DOBDC were further investigated by Cu K-edge X-ray absorption near-edge structure (XANES) spectroscopy and extended X-ray absorption fine structure (EXAFS) spectroscopy. The XANES spectra of the Cu K-edge for both CuH₂DOBDC and Cu₂H₂DOBDC are shown in Fig. 3e. In the spectrum of CuH₂DOBDC, distinct characteristic peaks were observed, including a pre-edge peak I at 8977 eV corresponding to the 1s \rightarrow 3d transition, a white line peak II at 8996 eV, and a resonance peak III at approximately 9042 eV, which indicates that the oxidation state of Cu species is +2.³²⁻³⁴ However, in the XANES spectrum of Cu₂H₂DOBDC, the aforementioned characteristic peaks associated with Cu(II) are absent. Instead, a new peak at 8983 eV emerged, attributed to the 1s \rightarrow 4p transition of Cu(I) (peak A), indicating that the predominant oxidation state of Cu species is Cu(I). Furthermore, the significant shift of the edge position towards the lower energy direction provides further evidence for the predominance of the Cu(I) state.^{33,35} This observation is con-

sistent with the XPS results. In the R-space EXAFS spectra, both samples exhibited a dominant peak at 1.5 Å arising from Cu-O bonding (Fig. 3f). However, for Cu₂H₂DOBDC, the main peak at 1.5 Å possesses a lower intensity, which suggests a different Cu-O coordination environment. Additionally, when compared to CuH₂DOBDC, the Cu-C peak also exhibited a decrease in intensity, which could be attributed to the partial removal of the carboxylate ligand.³³ The EXAFS fitting curves showed that the Cu-O configuration predominantly exists in CuH₂DOBDC and Cu₂H₂DOBDC, but with a change in the coordination number of O atoms around the Cu atoms (Fig. S8 and S9 and Table S2[†]).²⁷ In the wavelet transform (WT) contour plots shown in Fig. 3g, Cu₂H₂DOBDC exhibited an intensity maximum of Cu–O at approximately 5.85 Å⁻¹, which was very close to that of the reference Cu_2O at 5.80 Å⁻¹. Compared with the WT plot of CuH_2DOBDC (6.0 Å⁻¹), this provides further evidence that the formation of Cu(1) species leads to a change in the Cu–O coordination environment.³⁶ Therefore, the above results indicate that the valence state and coordination environment of Cu atoms changed significantly after ascorbic acid reduction, resulting in the formation of a novel Cu(I)-carboxylate MOF.

Based on the aforementioned characteristics, we have formulated a speculation regarding the local structure of Cu_2H_2DOBDC . Current understanding of its structure reveals that the coordination mode of Cu(1) and the ligand is in the monodentate form, with DMF not participating in the coordination. Furthermore, the molar ratio of Cu(1) to the ligand is 2:1. Consequently, we speculate that the local structure comprises a symmetric unit with two Cu(1) ions (Cu1 and Cu2) linked by an H_2DOBDC^{2-} linker (Fig. S10†). Such a possible formation process is shown in Fig. S11.†

Catalytic azide-alkyne "click" reaction

1,2,3-Triazoles are key structural motifs in the preparation of drugs, biochemicals, natural compounds, and functional materials, which are generally obtained by the Cu(1)-catalyzed (CuAAC).37-39 azide-alkyne cycloaddition reaction Cu₂H₂DOBDC, as a heterogeneous catalyst, has abundant welldispersed Cu(I) metal sites and a hierarchical structure, which may endow Cu₂H₂DOBDC with superior catalytic activity. Thus, the catalytic performance of Cu₂H₂DOBDC is evaluated by the azide-alkyne cycloaddition reaction using benzyl azide and phenylacetylene as the initial exploration substrates (Fig. 4a). Initially, the typical reaction between the benzyl azide and phenylacetylene was investigated in the presence of Cu_2H_2DOBDC (10 mg) in acetonitrile at room temperature for 2.5 h. However, a yield of only 26% was attained under these reaction conditions (entry 1 in Table S3[†]). Consequently, the temperature of the catalytic reaction was increased from 25 °C to 60 °C and then to 80 °C. Accordingly, as the temperature increased, the corresponding yields increased to 74% and 99%, respectively (entries 2 and 3 in Table S3[†]). It is noteworthy that only the single catalytic product 1,4-substituted 1-benzyl-4-phenyl-1H-1,2,3-triazole was obtained by utilizing Cu_2H_2DOBDC as the catalyst (Fig. S13 and S14[†]). To further

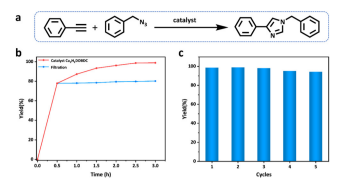


Fig. 4 (a) Schematic representation of the "click" reaction between phenylacetylene and benzyl azide. (b) The time profile of the yield and catalyst filtration test for the "click" reaction between phenylacetylene and benzyl azide. (c) Catalytic recyclability of Cu_2H_2DOBDC for the "click" reaction between phenylacetylene and benzyl azide.

explore the effect of the catalyst amount on the yields, the amount of catalyst was reduced to 5 mg. Nonetheless, a yield of 79% was still achieved (entry 4 in Table S3†). Additionally, experiments were conducted using CuH₂DOBDC along with a blank experiment without any catalyst, resulting in yields of only 39% and 19%, respectively (entries 5 and 6 in Table S3†). Under optimal reaction conditions, Cu₂H₂DOBDC exhibited excellent catalytic performance, with a conversion yield of up to 96% after just 2 h and gradually increasing to 99% after 2.5 h (Fig. 4b). The catalytic performance of Cu₂H₂DOBDC in the CuAAC reaction is superior to many recently reported related materials (Table S4†).

To investigate whether the catalysis was a heterogeneous reaction, the catalysts were separated from the reaction systems by hot filtration after 0.5 h. However, even after continuing the reaction with filtrates, no significant increase in yields was observed (Fig. 4b). The results indicate that Cu₂H₂DOBDC is an excellent heterogeneous catalyst for the AAC reaction. The recyclability of Cu₂H₂DOBDC as a heterogeneous catalyst was investigated. Impressively, Cu₂H₂DOBDC retained satisfactory catalytic activity and could be reused for five consecutive cycles without obvious loss in the catalytic activity, as shown in Fig. 4c. In addition, we further verified the structural stability of the Cu₂H₂DOBDC catalyst after cycling (Fig. S12[†]). The SEM results show that the sample morphology remains and the surface still retains hierarchical nanostructures. XRD analysis confirms that the catalyst maintains its crystallinity and structural integrity after cycling. FTIR spectra show no new peaks or shifts in existing peaks, indicating that there is no change in chemical bonds or functional groups. XPS analysis confirms that the predominant oxidation state of Cu species in Cu₂H₂DOBDC remains as Cu(I) even after undergoing five cycles of reactions. In summary, these results demonstrate the excellent structural and chemical stability of Cu₂H₂DOBDC, ensuring its potential for long-term and reliable catalytic applications.

To further validate the catalytic generality of Cu₂H₂DOBDC, we employed various terminal alkynes with different func-

Table 1 Catalytic reaction of substituted phenylacetylenes and benzyl azide catalyzed by $Cu_2H_2DOBDC^a$

Entry	Alkyne	Azide	Product	$\operatorname{Yield}^{b}(\%)$
1	N ₃	N ₃		99
2	a- () -=	N ₃		94
3	Br-	N ₃		99
4	F	N ₃		99
5		N ₃		95
6	н,со-	N ₃	H ₃ CO-	99
7	0 ₂ N-	N ₃		96
8	№ →=	N ₃		97

^{*a*} Reaction conditions: catalyst (10 mg), azide (1 mmol), alkyne (2 mmol), amyl acetate (1 mmol), acetonitrile (4 ml) and 80 $^{\circ}$ C, 2.5 h. ^{*b*} Isolated yields were calculated by GC, and amyl acetate was employed as the internal standard.

tional groups and azides, including methoxy, nitro, heteroaryl, halogen, or methyl moieties, to investigate the catalytic performance under the optimized conditions. As summarized in Table 1, all the selected substrates could efficiently undergo the Cu₂H₂DOBDC-catalyzed "click" reaction to deliver their corresponding 1,4-disubstituted 1,2,3-triazoles in high yields (Fig. S13–S28†). Cu₂H₂DOBDC exhibits superior catalytic performance and regioselectivity, which may be attributed to the high density of uniformly distributed Cu(1) active sites and the hierarchical MOF structure, which enhances accessibility to the active sites and facilitates fast diffusion of the reactants.

Considering previous reports on the CuAAC reaction, $^{39-42}$ a possible mechanism, as shown in Fig. S29,† has been proposed. Initially, the Cu(1) sites in Cu₂H₂DOBDC interact with the alkyne to form the Cu(1)-acetylide intermediate A. Subsequently, azides react with the Cu(1)-acetylide intermediate A to generate intermediate B. Then, the nucleophilic nature of the terminal N3 in intermediate B makes it attack the alkyne to form metallacycle C, which further shrinks to yield the Cu–triazole intermediate D. Finally, the protonation of intermediate D yields the target product 1,4-disubstituted 1,2,3-triazole, accompanied by the recovery of the catalyst.

Conclusions

In conclusion, we have developed a novel cleavage reduction strategy for fabricating a stable single-phase Cu(i)-carboxylate MOF that circumvents the limitations of HSAB theory. This innovative approach hinges on the distinctive structure of ascorbic acid and the instability characteristics of metastable MOFs. We demonstrated this strategy by chemically cleaving CuH₂DOBDC with ascorbic acid, which precisely cleaves the H₂DOBDC^{2–} linker and triggers a phase transition. We specu-

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lated on the atomic local structure of Cu_2H_2DOBDC and proposed a possible formation process. The resultant Cu_2H_2DOBDC acting as a CuAAC catalyst exhibited superior catalytic performance due to the highly dispersed and accessible open Cu(i) metal sites. This new strategy not only circumvents the limitations of HSAB theory but also opens up a new avenue for the creation of unprecedented categories of MOFs. We anticipate that this research will provide new perspectives to advance the development of MOF materials.

Author contributions

Xiaojiao Hou, Junyi Chen and Yu Fu designed the project, Xiaojiao Hou performed the experiments and wrote the manuscript, Yu Fu supervised this work, Wenxiu He and Xu Zhai collected the characterization data, and Bingbing Chen, Yuanlin Fu, and Liying Zhang contributed to the data analysis. All the authors contributed to the manuscript preparation.

Conflicts of interest

There are no conflicts to declare.

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