Structure—Property—Energetics Relationship of Organosulfide Capture Using Cu(I)/Cu(II)-BTC Edited by Valence Engineering

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ABSTRACT: Valence engineering of copper (Cu) within functional materials is effective to tune the performance and energetics of catalysts and sorbents. Here, by precisely controlling the Cu(II)/Cu(I) ratio employing Na2S2O3 as the reduction agent, we synthesized a family of copper-1,3,5-benzenetricarboxylic acid (Cu-BTC) sorbents for organosulfur compound capture. Integrated X-ray absorption fine structure and density functional theory studies suggest that the atomic-level, short-range order changes caused by Cu reduction lead to simultaneous stability, affinity, and pore topology evolutions in Cu(I)/Cu(II)-BTC, which govern the sorption capacity and binding energetics of dimethyl disulfide capture. This multiscale fundamental study with both experimental and computational results highlights the power of coordination chemistry on materials engineering on atomic level.

INTRODUCTION

Porous materials, such as molecular sieves,1 metal oxides,2 carbon-derived materials,3 and metal–organic frameworks (MOFs),4 play important roles in clean energy production from fossil fuels, pollutant removal, and emission control.5–8 Tuning the metal oxidation states significantly modifies the structure, performance, and energetic stability of MOFs.9 One example is that editing the ratio of Cu(I)/Cu(II) by valence engineering of copper is an effective strategy to enhance the properties of copper-based functional catalytic and separation materials.12–16 Specifically, precise editing in the Cu(I)/Cu(II) ratio of inorganic coordination polymer quantum sheet enables high-efficiency treatment of coking wastewater.17 The Cu(I)/Cu(II) ratio was also confirmed to be a crucial factor in determination of the activity and stability of copper-activated carbon catalysts in gas–solid acetylene dimerization reaction.18 It is also found that the electrochemical performance of Cu-doped α-Fe2O3 samples was a function of Cu(I)/Cu(II) ratio, which significantly modifies the local crystal structure.19 Moreover, Cu valence engineering has been applied to create hierarchical porosity in MOF structures to boost the performance of aromatic sulfide capture.20

Recently, we took advantage of valence engineering by reducing the copper in Cu-1,3,5-benzenetricarboxylic acid (BTC) from Cu(II) to Cu(I) using a series of reduction agents including ethanol, sodium thiosulfate (Na2S2O3), hydroquinone, and glucose. We found that such controllable Cu reduction leads to stronger binding affinity (energy of adsorption), enhanced accessibility, and diffusion for selective organosulfur compound uptake.21–23 However, the atomic level, short-range order evolution of Cu atoms remains unclear, and a comprehensive model based on integrated experimental data and computational prediction is expected to be established.

In this study, integrating X-ray absorption fine structure (XAFS) with density functional theory (DFT), we elucidated the short-range bonding evolutions of the Cu atoms. We intentionally chose dimethyl disulfide (DMDS) as the representative probe molecule for sorbent evaluation considering the multiple challenges in separation of disulfides from light petroleum products, such as low reactivity, poor dissolving activity, and weak affinity. The configurations and energetics of DMDS–Cu-BTC guest–host interactions were computationally evaluated by DFT based on the experimentally refined structural data models of Cu-BTC. Moreover, the impacts of framework topology and affinity of Cu sites on DMDS adsorption were interpreted and discussed.

EXPERIMENTAL METHODS

We systematically synthesized Cu-BTC and Cu(I)/Cu(II)-BTC sorbents using Na2S2O3 as the reduction agent to manipulate the Cu(II)/Cu(I) ratio.21 The synthesis procedure...
is described in the Supporting Information. Here, our samples are labeled \( x \)/Cu-BTC. \( x \) is the S/Cu molar ratio we used in the post-synthesis reduction process. The XAFS data (Cu K-edge, fluorescence mode) were collected at room temperature at the Beamline 14W1 of Shanghai Synchrotron Radiation Facility (SSRF), which was equipped with a Si(311) double-crystal monochromator. Photon energy calibration was performed by setting the first inflection point energy of Cu foil as 8980 eV.\(^{24}\) The storage ring was operated at 3.5 GeV.\(^{24}\)

We carried out DFT simulation using Dmol3 in Materials Studio 6.1, in which high-accuracy Perdew–Burke–Ernzerhof (PBE) functional of generalized gradient approximation was applied.\(^{27,28}\) Additionally, the DFT semicore pseudopods calculation was conducted using the double numerical plus polarization function. The geometric optimization was achieved by converging system energy, force, and displacement without considering symmetry restriction.

We define the energy of DMDS–Cu-BTC interactions, \( E_{\text{int}} \), with eq 1.

\[
E_{\text{int}} = E_{\text{guest-host}} - E_{\text{guest}} - E_{\text{host}} \tag{1}
\]

Here, \( E_{\text{guest-host}} \) is defined as the energy of guest–host interactions. \( E_{\text{guest}} \) and \( E_{\text{host}} \) represent the energy of DMDS and Cu-BTC, respectively. All the three terms share the same unit, eV. Exothermic energy of interactions suggests energetically favorable binding or guest–host interactions.

## RESULTS AND DISCUSSION

We used Na\(_2\)S\(_2\)O\(_3\) as a mild reducing agent for controllable modification of the copper oxidation state in Cu(II)-BTC. Conversion of Cu(II) to Cu(I) results in compositional, structural, and binding affinity modifications leading to largely fluctuant sulfide adsorption.\(^{21}\) Three Cu-BTC samples with different Cu(I)/Cu(II) ratios were synthesized. Other than the parent Cu(II)-BTC with only Cu(II), we prepared two samples by post-synthesis Cu reduction using Na\(_2\)S\(_2\)O\(_3\) as the reducing agent. The samples treated with S/Cu ratios = 0.46 and 2 were denoted as 0.46/Cu-BTC and 2/Cu-BTC, respectively. According to the characterizations results we reported earlier, the Cu(I) to Cu(II) molar ratios of 0.46/Cu-BTC and 2/Cu-BTC are 1.79 and 2.75, respectively (see Figure S1a). The X-ray diffraction (XRD) patterns in Figure S1b suggest that 0.46/Cu-BTC and 2/Cu-BTC retain their characteristic framework topology, while 2/Cu-BTC presents significant structural degradation suggesting an interrupted long-range order caused by severe copper reduction. The Raman results support these structural evolutions (see Figure S1c). The evolving peaks at 275, 448, and 503 cm\(^{-1}\) reflect the Cu–O vibration changes because of reduction from Cu(II) to Cu(I).\(^{29}\) In addition, one additional peak at 31.9\(^{\circ}\) is observed for both 0.46/Cu-BTC and 2/Cu-BTC, the evidence of O-coordinated Cu(I) upon Cu(II) reduction.\(^{29}\)

To further elucidate the atomic-scale transitions of short-range order, XAFS analyses were performed at the Cu K-edge under a continuous gas flow of argon. X-ray absorption near-edge structure (XANES) enables deep insights into the evolutions of coordination and valence of Cu in Cu-BTC upon Na\(_2\)S\(_2\)O\(_3\) reduction (see Figure 1). The XANES of both Cu-BTC and 0.46/Cu-BTC feature steep increase in the white line intensity at about 8980 eV. These sharp peaks at the
absorption edge are associated with the 1s to 3d transition according to data collected on single crystals.\textsuperscript{20} We also employ Cu foil, CuO, and Cu\textsubscript{2}O as reference materials (see Figure 1). Generally, the edge positions of Cu-BTC and 0.46/Cu-BTC match well with those of Cu foil and Cu\textsubscript{2}O, respectively (see Figure 1b,c). This is consistent with the oxidation states of Cu species determined by Raman and XPS.\textsuperscript{21} 0.46/Cu-BTC presents a very similar spectrum as that of Cu\textsubscript{2}O (see Figure 1c) with two peaks at 8995 and 9015 eV corresponding to Cu(I) species, whereas 2/Cu-BTC shows a distinctly different spectrum with disorder in white line and oscillations strongly suggesting a local disorder around Cu atoms caused by excessive dose of reducing agent.\textsuperscript{31} Overall, Na\textsubscript{2}S\textsubscript{2}O\textsubscript{3} reduction leads to decreased Cu oxidation state from divalent Cu(II) to monovalent Cu(I), which causes decreased short-range order around each Cu atom.

EXAFS analysis was applied to determine the coordination modes of Cu species and to reveal the details of local structure evolutions due to reduction. The data obtained are fitted to the shortest backscatter distance around the Cu absorber (first Cu–O shell). We list the coordination numbers (CN), scattering path radius (R), and Debye-Waller factors (\(\sigma^2\) or DWF) of all samples studied in Table 1. Specifically, the first-shell fits of the Cu–O shell EXAFS spectra are plotted in Figure S2. The Fourier transformed spectrum of Cu-BTC in Figure 1d suggests a main backscattering shell at 1.91 ± 0.02 Å, corresponding to the Cu–O shell with an average CN of 3.85 ± 0.4. This confirms that Cu is coordinated to 4 O of the trimesic acid ligands of Cu-BTC. The CNs and interatomic distances are also consistent with the Cu-BTC reference data in Cambridge Crystallographic Data Centre (CCDC).\textsuperscript{32} Once Cu is reduced to Cu(I), the main backscattering of Cu–O decreases to 2.00 ± 0.02 Å for both 0.46/Cu-BTC and 2/Cu-BTC accompanied with significantly decreased average CNs of 3.05 ± 0.3 for 0.46/Cu-BTC and 1.90 ± 0.2 for 2/Cu-BTC (see Figure 1e,f). In addition, the DWFs (\(\sigma^2\)) for the first coordination shells of Cu-BTC, 0.46/Cu-BTC, and 2/Cu-BTC are 0.0005, 0.0019, and 0.0036, respectively. The increased DWFs (\(\sigma^2\)) suggest a decreased short-range order around the Cu atoms as they are reduced, as seen in the XANES and Raman results in Figures 1a and S1c. The atomic-scale transitions in bond length and CN are responsible for the changes in long-range order. Specifically, as CN decreases from 4 for Cu-BTC to 3 for 0.46/Cu-BTC, the number of Cu–O bond in 0.46/Cu-BTC and its length both decrease, resulting in more sites to bind sulfur species. Further decrease in CN to 2 for 2/Cu-BTC increases the number of broken Cu–O bonds, which lead to structural degradation from an open stereoscopic framework (three-dimensional or 3D) to an expanded structure (two-dimensional or 2D) (see Figure 2). Hence, despite increased number of sites for sulfide binding when we create more undercoordinated copper ions, Cu(I) enrichment leads to degradation in long-range order and much less open framework topology, both of which are counterproductive for sulfide removal.

Computationally, we used the distance function of DFT to quantify the distance between Cu and other atoms (see Figure 2). The simulated configurations were correlated with the experimental data. Specifically, for Cu-BTC, a strong absorption band centered at about 1.91 Å in the Fourier transformed spectrum corresponds to O shell. A similar characteristic peak with identical bond length was also observed for the CuO reference (Figure 1d), which strongly suggests that the Cu–O bond within Cu-BTC is similar as that in the CuO reference. Distance function synchronizes with the experimental results by returning a Cu–O bond length of 1.935 Å. The peak of Cu–Cu bond at 2.55 Å in the Cu-BTC spectrum (simulated at 2.47 Å) is in good accordance with that of the Cu foil reference (~2.50 Å).\textsuperscript{33} Moreover, the peaks at 2.82 and 3.11 Å reflect the interatomic distance between Cu and C atoms. These values are consistent with the simulation results, suggesting that the unbonded Cu and C atoms are 2.799 and 3.06 Å apart. Additionally, the peak at 4.20 Å (4.188 Å simulated) is ascribed to the Cu–C interatomic distance in the Cu-BTC structure. Several low intensity peaks representing distances between the Cu atom and O or H atoms due to absorbed water molecules are not discussed. Considering the very minor contributions of these peaks indicating interatomic distances longer than 5 Å, low coordination, and high disorder, we do not focus on them in the EXAFS analyses.

Interestingly, the EXAFS peaks of reduced Cu-BTC samples disappear or shift to new positions (see Figure 1e). Specifically, the Cu-BTC peak at 1.91 Å shifts to 2.00 Å with increased intensity for 0.46/Cu-BTC, which suggests increased Cu–O bond length simulated to be 1.949 Å. The absence of peak at 2.55 Å implies broken Cu–Cu bonds within 0.46/Cu-BTC,

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**Table 1. CN, Path Radius (R), and Mean-Squared Relative Displacements (\(\sigma^2\)) of all Samples Determined by EXAFS**

<table>
<thead>
<tr>
<th>Samples</th>
<th>No. Shells</th>
<th>CN</th>
<th>R (Å)</th>
<th>(\sigma^2) (Å(^2))</th>
<th>R Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu-BTC</td>
<td>1st</td>
<td>3.85</td>
<td>1.91</td>
<td>0.0005</td>
<td>0.0012</td>
</tr>
<tr>
<td>0.46/Cu-BTC</td>
<td>1st</td>
<td>3.05</td>
<td>2.00</td>
<td>0.0019</td>
<td>0.0001</td>
</tr>
<tr>
<td>2/Cu-BTC</td>
<td>1st</td>
<td>1.90</td>
<td>2.00</td>
<td>0.0036</td>
<td>0.0007</td>
</tr>
</tbody>
</table>

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**Figure 2. Structural evolution of Cu-BTC upon Na\textsubscript{2}S\textsubscript{2}O\textsubscript{3} reduction. The length of each bond or interatomic distance are noted (Å).**
isotherm analysis at 77 K. The parent Cu(II)-BTC has type I adsorption isotherm. The Cu(I)/Cu(II) ratios at 293 K are 0.46 for the moderately reduced sample, 2/Cu-BTC, which is consistent with the calculated values of 2.912, 3.398, and 3.840 Å, respectively. Interestingly, a new peak at 4.14 Å emerges, which reflects the distance between the Cu atom and the H bonded to O. Besides, the peak positions of 2/Cu-BTC are nearly identical to those of 0.46/Cu-BTC yet with higher intensity. The optimized structure model of 2/Cu-BTC has a Cu–O bond length of 1.973 Å (see Figure 2) in agreement with the peak observed at 2.00 Å in the Fourier-transformed spectrum. In addition, the simulated interatomic distances between the Cu atom and its adjacent unbonded O, Cu, and H atoms are 2.875, 3.368, and 3.936 Å, mirroring the experimental data at 3.00, 3.40, and 3.87 Å in Figure 1f, respectively. Moreover, the simulated interatomic distances in energy-optimized structure models agree well with the EXAFS data, which further confirm the well-defined structures of 0.46/Cu-BTC and 2/Cu-BTC. In sum, integrating EXAFS and DFT, we demonstrate that employing Na2S2O3 as the reducing agent, we are able to manipulate the oxidation states, CNs, and bonding specificities of Cu species at atomic scale to fine-tune the local environment for expected organosulfur capture.

The impacts in Cu-BTC pore structure as the Cu(I)/Cu(II) ratio varies were evaluated using N2 adsorption–desorption isotherm analysis at 77 K. The parent Cu(II)-BTC has type I isotherm evidencing microporosity (see Figure 3a,b). Na2S2O3 reduction leads to decreased CN for 0.46/Cu-BTC, which enables a more open framework structure. In Figure 3a, a large hysteresis loop is observed on the isotherm of 0.46/Cu-BTC, and the pore size distribution plot suggests its mesopore span from 3 to 4 nm (see Figure 3c). In sharp contrast, because of structural degradation caused by over-reduction, significant porosity reduction is observed on 2/Cu-BTC. The simulated configuration of its collapsed structure is shown in Figure 2. Therefore, Cu(I) enrichment increases the accessibility of binding sites within Cu(I)/Cu(II)-BTC, leading to performance enhancement as we reported earlier, until the collapse of the framework structure.21

According to the DMDS uptakes on samples with different Cu(I)/Cu(II) ratios at 293 K, the moderately reduced 0.46/Cu-BTC has boosted DMDS loading of 146.66 mg-S/g, which is about 17 times that of chemical liquid deposition-modified ZSM-5 and nearly 2 times higher than that of Ag-exchanged ZSM-S.34,35 This value nearly doubles that of the parent Cu(II)-BTC sample (89.23 mg-S/g). In contrast, the over-reduced sample, 2/Cu-BTC, suffers from structural degradation, exhibiting decreased DMDS adsorption capability (17.81 mg-S/g). Very recently, Qi et al. report that vapor reduction is effective to generate mesoscale channels within Cu-BTC crystals.20 Such hierarchical mesoscale porosity was confirmed to facilitate enhanced thiophene adsorption. Here, we observe a similar increase in organosulfur uptake. Furthermore, we found that microporous Cu-BTC samples modified with ethanol and glucose as the reducing agents exhibit significantly increased sorption capacity of sulfide compounds, including DMDS, ethyl sulfide, and 1-propanethiol.21 Such evidence strongly suggests that it is the complex integration of increased binding site affinity from Cu(I) and more open pore structure prior to pore collapse that lead to the sulfide uptake boost on Cu(I)/Cu(II)-BTC. This motivates us to carry out DFT simulation to elucidate the affinity of binding sites toward sulfides, such as DMDS, as the magnitude of Cu(I)/Cu(II) increases.

DFT simulation was performed for DMDS adsorption on Cu-BTC with different Cu(I)/Cu(II) ratios. We constructed a series of Cu-BTC unit cells (see their geometries in Figure S3), each with 24 Cu atoms. Specifically, at Cu(I)/Cu(II) = 0.24, 0.4/20, 4/20, 8/16, 12/12, 16/8, 18/6, and 20/4, their stoichiometries are Cu(II)3(BTC)2, [Cu(I)0.5Cu(II)2.5]H0.5(BTC)2, [Cu(I)-Cu(II)]H2(BTC)2, [Cu(I)1.5Cu(II)1.5]H1.5(BTC)2, [Cu(I)-Cu(II)]H2(BTC)2, [Cu(I)2Cu(II)0.75]H2.25(BTC)2, and [Cu(I)2Cu(II)0.5]H2.5(BTC)2, respectively. Here, Cu(II)3(BTC)2, [Cu(I)2Cu(II)]H2(BTC)2, and [Cu(I)2Cu(II)]H2.5(BTC)2 mimic the parent Cu(II)-BTC, 0.46/Cu-BTC [Cu(I)/Cu(II) ratio = 1.79] and 2/Cu-BTC [Cu(I)/Cu(II) ratio = 2.75] in our experimental studies, respectively. The calculated DMDS–Cu(I)/Cu(II)-BTC interaction energies are plotted against the average ion potential of cations (Z/r, defined as ∑X(Zi/r)/∑Xi), in which X is mole fraction, Z is charge, and r is ionic radius (see Figure 4). As the Cu(I) content increases Z/r decreases. Specifically, Cu(II)3(BTC)2 has the least favorable binding of −20.28 kJ/mol-DMDS, while the DMDS–Cu(I)/Cu(II)-BTC interactions tend to be stronger as Z/r decreases with [Cu(I)2Cu(II)]H2(BTC)2 presenting the strongest binding of −171.87 kJ/mol-DMDS. A subsequent decrease in Z/r results in weakening adsorbate–adsorbent interactions for [Cu(I)2.5Cu(II)0.5]H2.25(BTC)2 (−90.35 kJ/mol-DMDS) and [Cu(I)2Cu(II)0.5]H2.5(BTC)2 (−72.16 kJ/mol-DMDS). Such “volcano plot-like” of guest–host interaction energy as a function of Z/r is consistent with

Figure 3. (a) N2 adsorption–desorption full isotherms at 77 K and (b) microporosity and (c) mesoporosity distribution of all Cu(I)/Cu(II)-BTC samples.
our earlier report on DMDS adsorption into Cu(I)/Cu(II)-BTC, in which we demonstrated that Cu(I) generated preferentially binds to sulfide molecules via strong π-complexation interactions. In this study, we further our understanding with atomic-scale XAFS data and DFT insights, which strongly suggest that increase in Cu(I) concentration leads to enhanced DMDS adsorption capacity on Cu(I)/Cu(II)-BTC. Moreover, DFT simulation suggests increased DMDS–Cu(I)/Cu(II)-BTC interaction energy until the Cu(I)/Cu(II) ratio is too high to maintain the stability of an intact open framework structure. For example, Cu(I)/Cu(II)-BTC with Cu(I)/Cu(II) ratio higher than 2 results in wakened interactions, as observed on [Cu(I)2.25Cu(II)0.75]H2.25(BTC)2 and [Cu(I)2.5Cu(II)0.5]2−(BTC)2−.

### CONCLUSIONS

In conclusion, valence engineering of Cu in Cu-BTC using Na2S2O3 leads to significantly increased DMDS loading. The XRD, Raman, XPS, XAFS results, and DFT simulation all point out that Cu valence engineering results in complex interrelated evolutions in structure–property–energetics relations. In addition, such modification causes increased DMDS adsorption performance. Specifically, 0.46/Cu-BTC with Cu(I)/Cu(II) = 1.79 presents the highest capacity (146.66 mg S g−1). Moreover, XAFS analysis reveals that the increased degree of undersaturation in Cu CN boosts the DMDS adsorption capacity and affinity. Simultaneously, DFT simulation demonstrates that [Cu(I)2Cu(II)]2−(BTC)− has the strongest interactions with DMDS (−171.87 kJ mol−1 DMDS), which is enabled by the synergistic enhanced binding affinity and more accessible mesoporous porosity. This fundamental study demonstrates that transition metal valence engineering is an effective strategy to tune the structures of MOF-based sorbent materials with desired stability and properties.

### ASSOCIATED CONTENT

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.iecr.0c05483.

XPS spectra, XRD patterns, and Raman spectra; shell fits; geometries of a series of Cu-BTC unit cells; and convergence tolerance values (PDF)

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Notes
The authors declare no competing financial interest.

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