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Chemisorption of hydrogen sulfide over copperbased metal-organic frameworks: methanol and UV-assisted regeneration†

Three copper-based metal–organic frameworks (MOFs) with different organic linkers were synthesized for the removal of H_2S gas at room temperature. The synthesized MOFs were characterized by microscopic and spectroscopic techniques to understand their structural, functional, and optical properties. The H_2S adsorption capacity of MOFs calculated by column studies followed the trend: 105.6 mg g^{-1} (CuBDC) > 27.1 mg g^{-1} (CuBTC) > 1.3 mg g^{-1} (CuBDC-N) in dry conditions. The adsorption capacity increased in moist conditions due to an easy dissolution and dissociation of H_2S in a film of water. X-ray photoelectron spectroscopy confirmed the presence of sulfur bound to Cu-sites and sulfate ions. The spent MOFs were regenerated by the successive effect of methanol and low power UV-C radiation. The regenerated CuBTC showed an exceptionally high adsorption capacity of 95.6 mg g^{-1} in the second cycle, which was linked to the reactivation of Cu-sites and improved surface area and porosity. The regeneration process developed in this study is a cost-effective method to recycle chemisorbed MOFs without compromising with their structural and functional integrity.

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1. Introduction

Hydrogen sulfide (H₂S) is a toxic gas with a characteristic rotten egg smell and arises from many anthropogenic sources like crude oil refineries, coal gasification plants, food processing units, livestock farms, and municipal sewage treatment facilities. H₂S emission has been linked to the formation of acid rain, which, in turn, affects human health and the environment. Acute exposure to H₂S at levels of 100-200 ppm can be lethal and have resulted in death.1 Therefore, the removal of H2S from sources has been prioritized to enhance the quality of life and, at the same time, prevent catastrophic events like acid rain.² In this regard, numerous physical/chemical/biological techniques like adsorption, scrubbing, electrochemical decomposition, catalytic oxidation, and bio-filtration have been explored for the removal of H₂S from biogas, feed gases, and flue gases.^{3,4} Currently, adsorption using porous adsorbents is deemed as a lucrative option due to its simplicity, low operational cost, and high performance. Porous adsorbents like activated carbon, metal oxides, zeolites, and composites have been evaluated for H₂S removal and have been shown to have disadvantages like low adsorption capacity and poor reusability.1,4

In the domain of porous adsorbents, a sub-class of organicinorganic hybrids, i.e., metal-organic frameworks (MOFs), is in the limelight for their exceptional surface and pore properties with the easy tuning of physicochemical properties. MOFs are constructed through coordinative interactions between organic linkers and metal ions to form a highly ordered framework with uniform porosity. MOFs are one of the potential candidates for the capture and storage of carbon dioxide,5 adsorptive removal of toxic gases,6 and catalytic conversion of gases into valueadded products.7 In the literature, Cu-based MOFs have been extensively explored for the removal of H2S gas under different experimental conditions to evaluate the adsorption capacity and underlying adsorption mechanism. Though the published works have confirmed Cu-based MOFs as superior adsorbents, among porous materials, the reusability of these MOFs is still a serious concern. Moreover, the literature has been saturated with the studies related to CuBTC (also known as HKUST-1 or MOF-199).8-12 The need to explore other Cu-based MOFs and the lack of proper investigation on the regeneration of spent Cubased MOFs have motivated us to develop methanol and UVassisted regeneration method.

In this study, we have fabricated three copper-based MOFs with 1,4-benzene dicarboxylic acid (H₂BDC), 2-amino-1,4-benzene dicarboxylic acid (H₂BDC-NH₂), and benzene-1,3,5-tricarboxylic acid (H₃BTC) *via* ultrasonication method¹³ to study their applicability in the removal of H₂S gas. The synthesized MOFs were characterized by several microscopic and spectroscopic techniques to understand their structural, functional, and optical properties. The MOFs were studied for

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the adsorptive removal of $\rm H_2S$ in both dry and moist conditions with a focus on understanding the adsorptive mechanism. In this study, attempts were made to regenerate spent MOFs using methanol and UV-C treatment. The effect of regeneration on the structural and functional stability of Cu-MOFs was studied by X-ray diffraction (XRD) and X-ray photoelectron spectroscopy (XPS).

Materials and methods

2.1 Chemicals and reagents

Copper(II) nitrate trihydrate (Cu(NO₃)₂·3H₂O), H₂BDC, H₂BDC-NH₂, and H₃BTC were procured from Sigma Aldrich, Germany. N,N-dimethyl formamide (DMF), ethanol, methanol, and sodium hydroxide (NaOH) were purchased from Samchun Pure Chemicals, Korea. All the chemicals were of analytical grade and used without any further purification. Highly pure H₂S gas (500 ppm balanced with N₂ gas) was procured from Union gas, Korea.

2.2 Synthesis of Cu-MOFs

The fresh copper hydroxide was prepared by adding NaOH solution (82.5 mL, 1.0 mol L $^{-1}$ in deionized water) into an aqueous solution of $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ (10.0 g in 75 mL of deionized water). To it, an H_2BDC solution (6.64 g in 190 mL DMF) was added under ultrasonication (Sonics Vibra-cell 500 W, 20 kHz, 44% amplitude). After 20 min of ultrasonication, the precipitate was collected by centrifugation and washed thoroughly with ethanol. It was followed by drying at 50 °C for 10 days in a hot air oven. The synthesized product was named as CuBDC. Using the same protocol, CuBDC-N was prepared with an $\text{H}_2\text{BDC-NH}_2$ solution (7.24 g in 190 mL DMF). CuBTC was synthesized using an H_3BTC solution (8.50 g in 190 mL DMF). The physical appearance and yield are listed in S. Table 1.† The Cu-MOFs were finely grounded and stored at 70 °C in glass vials.

2.3 Instruments

The surface morphology of Cu-MOFs was analyzed by field emission scanning electron microscopy (FE-SEM) (Hitachi S-4300, Japan). Dried samples were coated with a gold-platinum alloy by ion-sputtering (E-1048 Hitachi ion sputter). The transmission electron microscope (TEM) images were recorded with a field emission TEM (FE-TEM, JEM-2010F, JEOL, Japan). The two dimensional (2D) elemental mapping was done by energydispersive X-ray spectroscopy (EDS) (X-Maxn 80 T, Oxford, UK). Thermal gravimetric analysis (TGA) was done using a Thermogravimetric Analyzer (TG 209 F3, NETZSCH). For Brunauer-Emmett-Teller (BET) analysis, N2 adsorptiondesorption isotherm was measured at -196 °C in a Gemini series Micromeritics 2360 instrument after degassing at 200 °C for 6 h. The diffraction patterns were obtained using Ultima IV (Rigajku, Japan) X-ray diffractometer with Cu Kα and a Ni filter where the scanning speed was set to 3° min⁻¹. Fouriertransform infrared (FTIR) spectra of samples were recorded using KBr pellets over an FTIR spectrometer (Cary670, Agilent).

The recording was done with a single-beam spectrometer with 60 added scans. Ultraviolet-visible light-diffuse reflectance spectroscopy (UV-Vis DRS) spectra of photocatalysts were obtained using a SCINCO S-4100 spectrometer equipped with a photodiode array detector and a diffuse reflectance attachment. The electron spin resonance (ESR) measurement was carried out on a JEOL JES-FE1C X-band spectrometer. For XPS analysis, a K-alpha XPS instrument (Thermo Scientific Inc., UK) with a monochromatic Al Ka X-ray source was used where the pressure was fixed to 4.8×10^{-9} mbar. Spectra have been charge corrected to the main line of the carbon 1s spectrum (aromatic carbon) set to 284.7 eV. Spectra were analyzed using CasaXPS software (version 2.3.14). GL(p) = Gaussian/Lorentzian product formula where the mixing is determined by m = p/100, GL(100) is a pure Lorentzian while GL(0) is pure Gaussian. We have used GL(30).

2.4 Breakthrough experiments

The adsorption experiments were performed by taking 250 mg of a powdered MOF in a Pyrex tube (height: 50 cm, diameter: 1 cm) at 25 °C. The $\rm H_2S$ gas (500 ppm) was passed through it at a flow rate of 0.3 L min⁻¹. The outgoing gas was analyzed by a Gas analyzer (GSR-310, Sensoronic, Korea) with the detection limit of 0.1 ppm. The $\rm H_2S$ concentration was measured every 15 s until the effluent concentration reached 400 ppm. The schematic illustration of the $\rm H_2S$ adsorption system is shown in Fig. 1. For evaluating the effect of moisture, moist air (relative humidity $\sim 60\%$) was blown through the MOF packed in the reactor at a flow rate of 0.2 L min⁻¹ for 30 min before passing $\rm H_2S$ gas.

For the regeneration of spent MOF, exactly 200 mg of the spent MOF was stirred in 10 mL of methanol for 24 h at 300 rpm, followed by separation of MOF by centrifugation and drying at 70 °C for 24 h. The dried MOF was irradiated in a 1.0 L Pyrex tube fitted in an acryl reactor with 4 UV lamps (19 W, UV-C, $I_{\rm max} \sim 254$ nm) for 4 h in the N₂ environment. The adsorption capacity (q, mg g^{-1}) at the breakthrough point (where the effluent concentration reached 400 ppm) was calculated by integration of the area above the breakthrough curve.

$$q = \frac{C_0 Q}{m} \int_0^{t_b} \left(1 - \frac{C}{C_0} \right) \mathrm{d}t \tag{1}$$

where C_0 – initial concentration (500 ppm or 0.697 mg L⁻¹), Q – flowrate, m – the mass of MOF (g), and t_b – breakthrough time.

3. Results and discussions

3.1 Characterization of Cu-MOFs

The morphology of Cu-MOFs was investigated using SEM (Fig. 2a–c) and HR-TEM (Fig. 2d–f). CuBDC was homogeneously distributed as square sheets of size 0.2–0.8 μm and 75–100 nm thickness (Fig. 2a), which was further confirmed by thin sheet-like structures in the HR-TEM image (Fig. 2d). CuBDC-N had a large number of prismatic-shape microcrystallites (1–5 μm) with some nanocrystals (Fig. 2b and e). CuBTC crystals of varying dimensions were observed with no well-defined

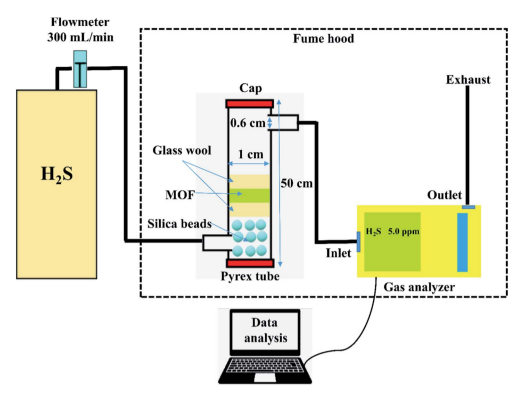


Fig. 1 Schematic illustration of the H₂S adsorption system.

morphology (Fig. 2c and f). The 2D elemental mappings of Cu-MOFs confirmed uniform distribution of Cu in MOFs (S. Fig. 1^{+}).

The surface and pore properties of Cu-MOFs were evaluated by N₂ adsorption–desorption isotherms, as shown in Fig. 3a and b and Table 1. Both CuBDC and CuBTC exhibited type II isotherms with the presence of both micropores and mesopores in the MOFs. The CuBDC-N exhibited low N₂ uptakes at relative pressure less than $p/p_0 \sim 0.2$ with hysteresis loops at $p/p_0 \sim 0.45$ –0.90. These isotherms are described as type IV with characteristics of mesoporous materials. The surface area of CuBDC and CuBTC was significantly higher than CuBDC-N. The formation of comparatively large size particles of CuBDC-N was responsible for its low surface area.

The XRD patterns of Cu-MOFs are shown in Fig. 3c, and assignments are listed in S. Tables 2–4. The diffraction peaks of CuBDC at 10.16° , 12.08° , 13.58° , 17.12° , 17.70° , 20.40° , and 24.82° were assigned to 110, 020, 11-1, -201, 111, 220, and 131 Bragg reflections of CuBDC, respectively. The MOF had monoclinic symmetry (a=11.37 Å, b=14.62 Å, c=7.72 Å, $\beta=108.49^{\circ}$) with C2/m space group. The CuBDC-N was crystallized in the monoclinic symmetry with a slight variation in the lattice parameters due to the presence of amine functionality (S. Table $5\dagger$). The XRD pattern of CuBTC matched well with the HKUST-1, where diffraction peaks at 9.46° , 11.62° , 13.42° , 19.02° , and 25.96° were due to 022, 222, 004, 044, and 355 reflections, respectively. CuBTC crystallized in the cubic symmetry (a=26.36 Å) with $Fm\bar{3}m$ space group. The interval of the cubic symmetry (a=26.36 Å) with a=10.00 where a=10.00 is the sum of the cubic symmetry (a=26.36 Å) with a=10.00 crystallized in the cubic symmetry (a=26.36 Å) with a=10.00 crystallized in the cubic symmetry (a=26.36 Å) with a=10.00 crystallized in the cubic symmetry (a=26.36 Å) with a=10.00 crystallized in the cubic symmetry (a=10.00 cryst

Fig. 3d presents the weight loss of Cu-MOFs when heated from 30 to 900 $^{\circ}$ C at a heating rate of 10 $^{\circ}$ C min⁻¹ in the N₂/O₂

atmosphere. In TGA profiles, the first weight-loss stage (stage I) occurring between 30–110 °C was due to the evaporation of physically adsorbed water. In stage II (110–250 °C), the weight loss was due to the release of chemically bonded water on the metal sites and physically adsorbed solvent (DMF/ethanol). These two losses were significant for CuBDC and CuBTC due to large surface areas and a more open structure of these two MOFs. In stage III (250–370 °C), a 50–60% loss was recorded for all Cu-MOFs, which was attributed to the breakdown of coordination bonds between metal ions and the organic linkers. Around 350 °C, the MOFs were fully decomposed to metal oxides (Cu₂O and CuO).¹⁸

Fig. 4a shows the FTIR spectra of Cu-MOFs. For CuBDC and CuBTC, the band at 3446 cm⁻¹ was due to the O-H stretching vibrations of physically adsorbed water molecules, which was more prominent in CuBTC. Two low-intensity bands at 3052 and 2932 cm⁻¹ were attributed to C-H vibrations of the aromatic skeleton, which were masked by the broad O-H stretching band in CuBTC. Multiple bands in the range of 1200-1000 and 1000-700 cm⁻¹ were assigned to the in-plane and outof-plane C-H bending modes. For CuBDC, acid C=O stretching at 1672 cm^{-1} , C=C at 1511 cm^{-1} , and the symmetric and asymmetric O-C-O stretching at 1396 and 1576 cm⁻¹, respectively, were observed. 19 For CuBTC, the bands at 1710 cm⁻¹ for carboxyl groups was observed along with the bands at 1636 and 1386 cm⁻¹ for asymmetric and symmetric stretching of -C=O, respectively.²⁰ The Cu–O stretching at 468 cm⁻¹ was observed as a low-intensity band.21 For CuBDC-N, distinct bands for N-H (stretching: 3360 and 3476 cm⁻¹; bending: 1579 cm⁻¹) and

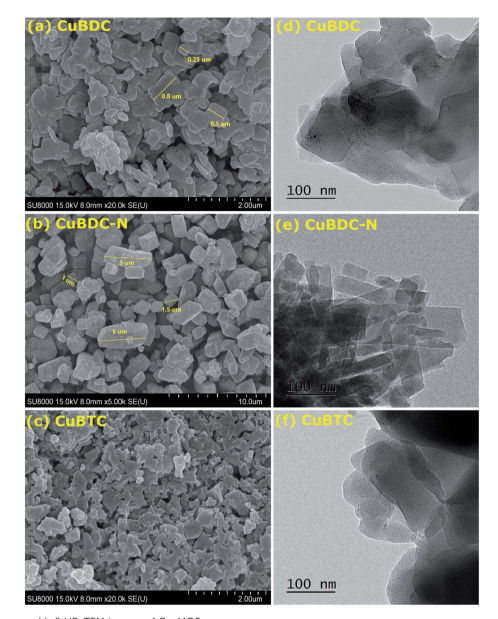


Fig. 2 (a-c) SEM images; (d-f) HR-TEM images of Cu-MOFs.

aromatic C-N vibrations (stretching: 1259 and 1334 cm⁻¹) of BDC-NH₂ linkers were observed.²²

The optical properties of Cu-MOFs were evaluated by UV-Vis DRS analysis (Fig. 4b). All the MOFs absorbed in the entire UV-Vis-NIR region with multiple peaks originating from the ligand-to-metal charge transfer (LMCT) and d–d transitions around the metal centres. The first band in the 270–500 nm could be assigned to the charged transfer from the oxygen in carboxylate to Cu²⁺ ions. The variation in the $\lambda_{\rm max}$ for Cu-MOFs was due to the presence of different organic linkers with varying HOMO energy.²³ The second absorption band at a wavelength greater than 500 nm was due to the d–d transition around the distorted octahedral Cu²⁺ centres. For perfect octahedral geometry, d–d transitions are dipole-forbidden, which leads to weak intensity. The high-intensity d–d band similar to the LMCT band was observed in the present study. A significant distortion in the

octahedral geometry due to the ligands heterogeneity around Cu-sites led to multiple peaks and high intensity of the d-d transition band that should be otherwise forbidden. An insignificant change in the spectra (500–1000 nm) suggested a very similar chemical environment around the metal centres in Cu-MOFs. Cu-MOFs.

$$\alpha h v = A(h v - E_g)^{1/2} \tag{2}$$

where α , hv, A, and E_g represent the absorption coefficient, photon energy (eV), a constant, and bandgap energy, respectively. The bandgap energy of Cu-MOFs, as evaluated from the plot of $(\alpha hv)^2$ versus hv (Fig. 4c), was 2.54, 2.33, and 2.56 eV for CuBDC, CuBDC-N, and CuBTC, respectively.

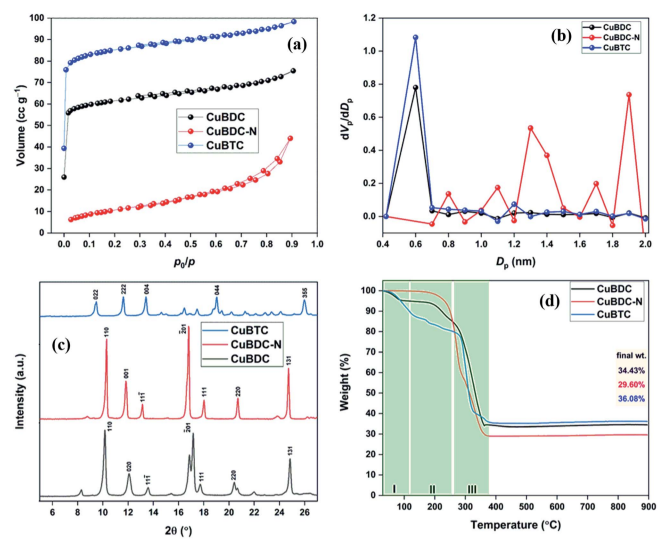


Fig. 3 (a) N₂ adsorption-desorption isotherms at -196 °C; (b) pore size distribution; (c) XRD patterns; (d) TGA patterns of Cu-MOFs.

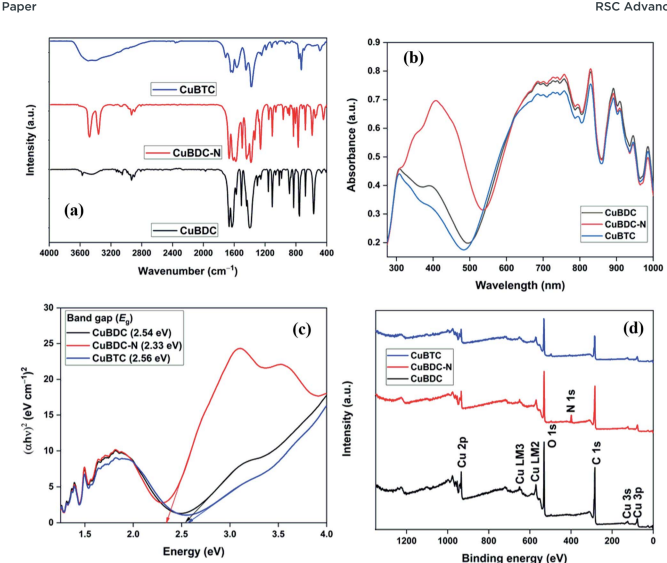
Table 1 The surface and pore properties of Cu-MOFs

MOF	$S_{\rm BET}$ (m ² g ⁻¹)	$V_{\rm t} \left({\rm cm}^3~{\rm g}^{-1}\right)$	D _{p,avg} (nm)	D _{micro} (nm)
CuBDC	217.8	0.117	2.14	0.60
CuBDC-N	38.0	0.068	7.18	1.90
CuBTC	317.0	0.152	1.92	0.60

The full scan XPS survey of Cu-MOFs are shown in Fig. 4d. The peaks for C 1s, O 1s, and Cu 2p at their respective binding energy were observed for all the three Cu-MOFs with one additional N 1s peak recorded for CuBDC-N at ~400 eV. The HRXPS O 1s spectra of Cu-MOFs have two peaks at 531.2 and 531.8 eV, corresponding to Cu-O/C=O and C-O bond, respectively^{19,28} (S. Fig. 2b, d and f†). The HRXPS C 1s spectra of CuBDC had four peaks at 284.7, 286.2, 288.4, and 291.0 eV, which were assigned to C=C, C-O/C-H, O-C=O, and -COOH, respectively (S. Fig. 2a†). For CuBDC-N (S. Fig. 2c†) and CuBTC (S. Fig. 2e†), the similar observations were made with variations

in the binding energy (S. Table 6†). The HRXPS Cu 2p spectra displayed two main peaks at \sim 934 (Cu 2p_{3/2}) and \sim 954 eV (Cu 2p_{1/2}). The Cu 2p_{3/2} peak of CuBDC had two contributions at 932.4 and 934.0 eV for Cu⁺ and Cu²⁺, respectively. The shake-up satellite peaks observed in the range of 935–945 eV further confirmed the presence of copper as Cu²⁺ (Fig. 5a).²⁹ For CuBDC-N (Fig. 5c) and CuBTC (Fig. 5e), the Cu⁺ and Cu²⁺ peaks were observed at similar binding energies (Table 2). From the HRXPS Cu LMM spectra of Cu-MOFs (Fig. 5b, d, f and S. Fig. 3†), the Cu⁺ species were further confirmed with the peak at \sim 570.6 eV, where Cu⁰ species remained absent³⁰ (S. Table 7†).

The ESR spectra of Cu-MOFs recorded at 25 °C are shown in S. Fig. 4.† Two signals, one low-intensity signal at the magnetic field (H) \sim 320 mT (g=2.06–2.08) and the other high-intensity signal at $H\sim$ 470 mT (g=1.42–1.52) were commonly observed in CuBDC and CuBDC-N (S. Fig. 4a and b†). In these MOFs, Cu²⁺ ions were present in two distinctly different coordination environments. The low-intensity signal (g=2.06–2.08) was probably due to the presence of monomeric Cu²⁺ complexes or uncoupled Cu²⁺ pairs in the MOFs. The high-intensity signal at the



(a) FTIR spectra; (b) UV-Vis DRS spectra; (c) Direct bang gap energy; (d) XPS survey of Cu-MOFs

higher magnetic field was assigned to the bridged carboxylatebound two Cu2+ coupled by antiferromagnetic interaction, generating an excited paramagnetic state with total spin S = 1 in the MOF.³¹ Another low-intensity signal at H = 440 mT (g = 1.42) observed for CuBDC might be due to distortions in the MOF structure. The ESR signal of CuBTC at H = 325 mT (g = 2.13) with a peak-to-peak width of about 100 mT matched well with the reported signal for HKUST-1 (S. Fig. 4c†). The signal observed in the spectrum is attributed to Cu^{2+} (S=1/2) paramagnetic centres, presumably due to the monomeric ions complexes ([Cu(H₂O)₆]²⁺) or uncoupled Cu²⁺ pairs in incomplete or stressed paddle-wheels.31

3.2 H₂S breakthrough curves

The H₂S removal performance of Cu-MOFs was evaluated at 25 $^{\circ}\mathrm{C}$ in a fixed-bed reactor. The obtained breakthrough curves using dry and moist Cu-MOFs are shown in Fig. 6. The CuBDC showed the best adsorption performance among all the MOFs with an adsorption capacity of 105.6 mg g⁻¹. The CuBDC-N

showed the lowest adsorption capacity. The trend followed by adsorption capacity was not in line with the surface area. Moreover, the presence of amine-functionalized linkers disfavored the adsorption process as opposed to the reported claim of increased H2S removal over amine-analogues of MOFs.32 Zhang and coworkers have demonstrated that the presence of amine functionalities beyond a certain ratio in MOF-199 disfavored the H2S adsorption process.8 Since the adsorption process of H₂S over Cu-MOFs is governed by the formation of strong Cu-S bond at the expense of weak Cu-O bond,9 the adsorption capacity of MOFs is directly influenced by the Cusite density and nature of Cu-sites. So, the highest copper content of CuBDC could be one of the factors responsible for its exceptional adsorption capacity besides surface area (S. Tables 8 and 9). The adsorption performance of MOFs improved after the addition of moisture for CuBDC-N and CuBTC with a slight loss in the capacity for CuBDC. The presence of water molecules in the MOF structure promoted the adsorption process by the

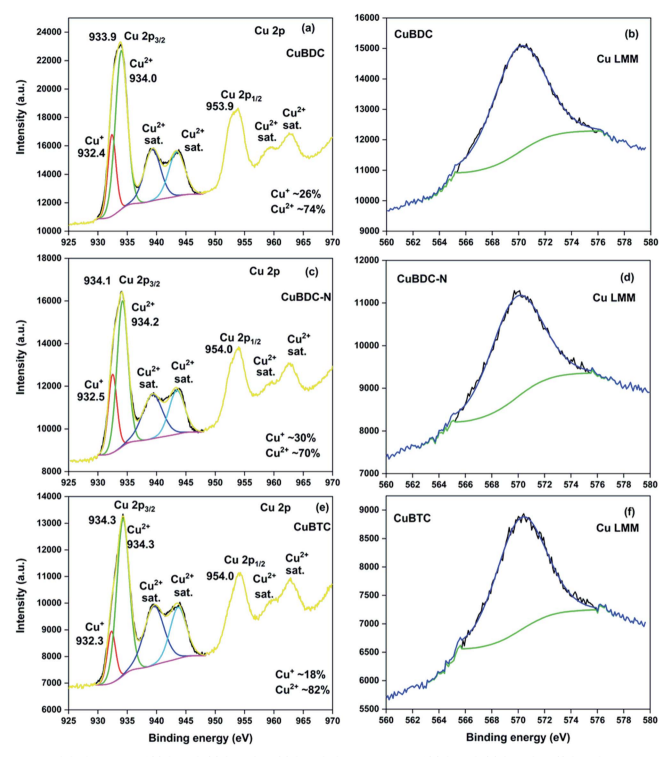


Fig. 5 HRXPS Cu 2p spectra of (a) CuBDC; (c) CuBDC-N; (e) CuBTC; Cu LMM spectra of (b) CuBDC; (d) CuBDC-N; (f) CuBTC.

dissolution of $\rm H_2S$ molecules. The formed $\rm HS^-$ species interacted strongly with the Cu-sites by forming Cu-S bonds. 10

3.3 Regeneration of spent MOFs

The H₂S breakthrough curves of spent and regenerated Cu-MOFs are shown in Fig. 7. The spent MOFs have residual

adsorption capacity arising from the unsaturated zones formed during the column studies. The presence of $\sim\!\!20\text{--}40\%$ adsorption capacity in the second cycle could lead to an overestimation of regeneration efficiency and has been taken into consideration in the present study. Though the regeneration of H_2S -loaded Cu-MOFs is difficult due to the irreversible nature of the chemisorption process, efforts were made to remove

Table 2 The peak-fitting results of Cu $2p_{3/2}$ high-resolution signals of CuBDC, CuBDC-N, CuBTC

Samples	Assignment	BE (eV)	FWHM (eV)	At%
CuBDC	Cu2p _{3/2} Cu ⁺	932.4	1.6	26.2
	Cu2p _{3/2} Cu ²⁺	934.0	2.4	73.8
	Satellite Cu ²⁺	939.3	3.3	_
	Satellite Cu ²⁺	943.4	3.2	
CuBDC-N	Cu2p _{3/2} Cu ⁺	932.5	1.7	30.0
	$Cu2p_{3/2}$ Cu^{2+}	934.2	2.2	70.0
	Satellite Cu ²⁺	939.3	4.0	_
	Satellite Cu ²⁺	943.4	3.2	
CuBTC	Cu2p _{3/2} Cu ⁺	932.3	1.7	18.4
	Cu2p _{3/2} Cu ²⁺	934.3	2.5	81.6
	Satellite Cu ²⁺	939.5	4.0	_
	Satellite Cu ²⁺	943.7	3.3	_

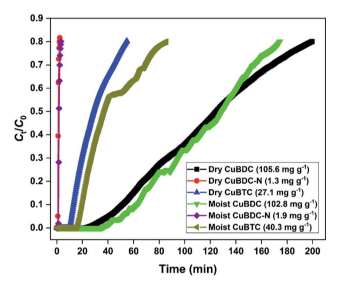


Fig. 6 H₂S breakthrough curve of dry and moist Cu-MOFs.

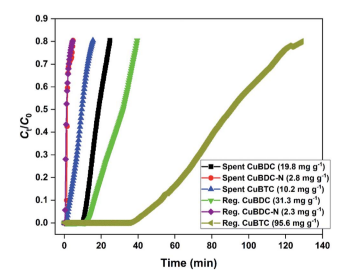


Fig. 7 H_2S breakthrough curve of spent and regenerated Cu-MOFs in the second cycle.

physically adsorbed H2S and generate new binding sites by adopting methanol and UV-assisted regeneration method. It has been reported that in the presence of UV irradiation, MOFs undergo a photolytic-decarboxylation process, which increases their basic nature and favours the adsorption of acidic gases.³³ In the present study, CuBDC-N could not be regenerated by the process. The regenerated CuBDC showed an increased H2S adsorption capacity of 31.3 mg g⁻¹ as compared to the spent CuBDC (19.8 mg g^{-1}). Though the improved performance was recorded for CuBDC, it was only 32% of the fresh CuBDC, suggesting that most of the Cu-sites were bound to sulfide and could not be regenerated. The regenerated CuBTC showed exceptional adsorption capacity where the capacity increased from 10.2 mg g^{-1} (spent CuBTC) to 95.6 mg g^{-1} . The regenerated adsorption capacity of CuBTC was 3.5 times the fresh CuBTC. It could be due to the increased surface area and availability of unoccupied Cu-sites, which were inaccessible in the first cycle.

3.4 Adsorption and regeneration mechanism

The XRD patterns of fresh, spent, and regenerated CuBTC are shown in Fig. 8a. The reactive adsorption of H2S over CuBTC leads to the breaking of Cu-O bonds with the formation of Cu-S bonds, which decreases the crystallinity of the MOF to an extent that most of the diffraction peaks are lost in the XRD pattern.¹⁰ In the present study, the relative intensity ratio of low angle peaks at $2\theta = 9.5^{\circ}$, 11.6° , and 13.4° in spent CuBTC changed significantly as compared to the fresh CuBTC. Moreover, the absence of peaks at $2\theta = 46.2^{\circ}$ (Cu₂S) and 31.8° (CuS) suggested that the Cu-sites remained intact in the MOF.34 Thus, the presence of many of the diffraction peaks in the spent MOF suggested that even after H₂S adsorption, the MOF stability was partially compromised. After the regeneration of spent CuBTC, the relative intensity ratio of low angle peaks was found very similar to that of the fresh MOF. Moreover, all the major peaks were observed in the diffraction pattern of the regenerated MOF, suggesting that chemisorbed Cu-MOFs could be regenerated by the successive effect of methanol and UV-C irradiation.

The XPS full scan survey of spent CuBTC confirmed the presence of sulfur in the MOF as a doublet peak at \sim 160-170 eV (Fig. 8b), which accounted for 3.91% of MOF atomic composition (S. Table 10†). The HRXPS Cu 2p spectrum of spent CuBTC showed a shift in peaks for Cu^{2+} (934.3 \rightarrow 933.9 eV) and Cu^{+} $(932.3 \rightarrow 931.3 \text{ eV})$ sites (Fig. 8c). The redshift in the binding energy of Cu peaks confirmed the interaction of S²⁻ with the Cusites in both +1 and +2 oxidation states.35 The HRXPS S 2p spectrum of spent CuBTC had two peaks at 162.4 and 168.8 eV, which corresponded to Cu-S and SO₄²⁻ ions, respectively³⁶ (Fig. 8e). The analysis confirmed an equal proportion of Cu-S and SO_4^{2-} ions in CuBTC. A large proportion of SO_4^{2-} ions in MOF formed due to the oxidation of HS⁻ or S²⁻ by adsorbed O₂ to elemental S, which was further oxidized to SO₄²⁻ ions.³⁷ The XPS full scan survey of regenerated CuBTC showed a lowintensity peak for S, which accounted for 1.49% of MOF atomic composition (Fig. 8b, S. Table 10†). The decreased S

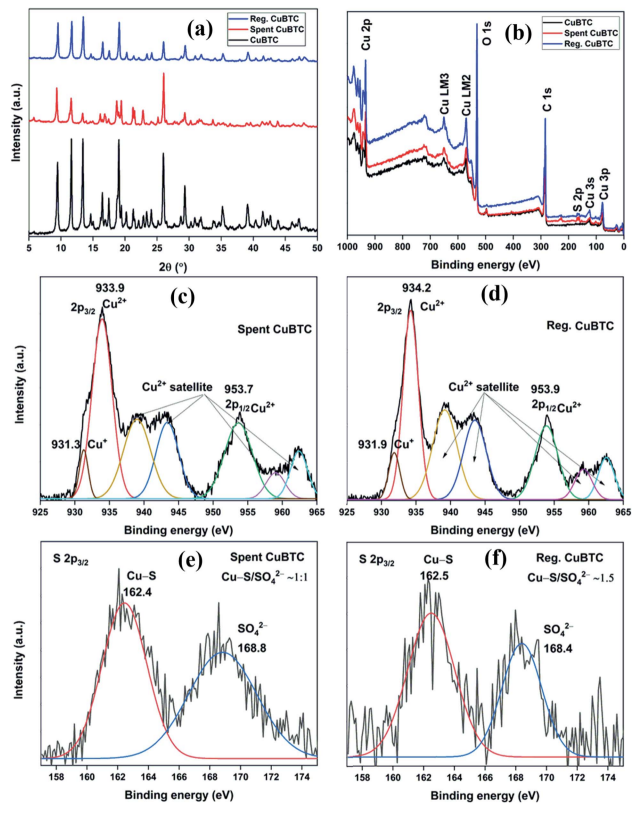


Fig. 8 (a) XRD patterns; (b) XPS full scan survey of fresh, spent, and regenerated CuBTC; HRXPS Cu 2p spectrum of (c) spent CuBTC (moist); (d) regenerated CuBTC; HRXPS S 2p_{3/2} spectrum of (e) spent CuBTC; (f) regenerated CuBTC.

content and repositioning of Cu peaks in the regenerated CuBTC validated the applicability of the regeneration process. The HRXPS Cu 2p spectrum of regenerated CuBTC showed the

repositioning of Cu^{2+} (933.9 \rightarrow 934.2 eV) and Cu^{+} (931.3 \rightarrow 931.9 eV) peaks, which were found close to that of the fresh CuBTC (Fig. 8d). The HRXPS S 2p spectrum of regenerated

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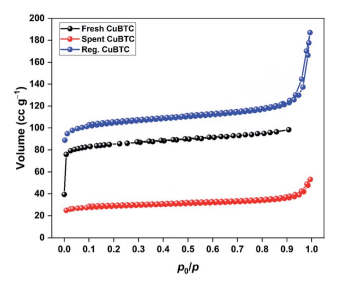


Fig. 9 N₂ adsorption-desorption isotherms of fresh, spent, and regenerated CuBTC.

Table 3 The surface and pore properties of fresh, spent, and regenerated CuBTC

MOF	$S_{\rm BET}$ (m ² g ⁻¹)	$V_{\rm t} \left({\rm cm}^3~{\rm g}^{-1}\right)$	D _{p,avg} (nm)
Fresh CuBTC	317.0	0.152	1.92
Spent CuBTC	110.9	0.078	1.23
Regenerated CuBTC	410.8	0.278	1.15

CuBTC showed the presence of both kinds of sulfur species, but the proportion of Cu-S was higher than that of SO₄²⁻ ions (Fig. 8f). The Cu-S presence in regenerated CuBTC could be assigned to the sulfur bound to the Cu⁺ sites (the Cu⁺ peak was not fully repositioned in the regenerated CuBTC). A decrease in S content and reactivation of Cu-sites achieved through the regeneration process was responsible for an excellent H2S adsorption capacity of regenerated CuBTC.

The N₂ adsorption-desorption isotherms of spent and regenerated CuBTC are shown in Fig. 9. The surface area and pore characteristics of spent and regenerated CuBTC are listed in Table 3. After H2S adsorption, the surface area and pore volume of CuBTC decreased by 65 and 49%, respectively. These structural changes were linked to the cleaving of Cu-carboxylate bonds upon H₂S interactions with the Cu-sites. Such changes are well-documented for stable MOFs like ZIF-8 after H2S exposure.38 The regenerated CuBTC has a surface area of 410.8 m^2 g⁻¹, which was higher than that of the fresh MOF (317.0 m²) g^{-1}). Moreover, the pore volume significantly improved after the regeneration process. Similar observations have been reported for the UV-assisted regeneration of iron-incorporated ZSM-5 zeolite and amorphous silica after the adsorption of volatile organic sulfur compounds.39 These improved surface area and pore properties were largely responsible for the enhanced H₂S uptake of regenerated CuBTC.

Conclusions 4.

Copper-based MOFs with different linkers were synthesized by a rapid ultrasonication method for the removal of H₂S gas at ambient conditions. The MOFs were characterized by different analytical techniques to understand their physicochemical and optical properties. The adsorption capacity of Cu-MOFs, as calculated for the breakthrough study, followed the trend: 105.6 mg g⁻¹ (CuBDC) > 27.1 mg g⁻¹ (CuBTC) > 1.3 mg g⁻¹ (CuBDC-N) in dry condition. Other than the surface area and pore volume, the Cu⁺/Cu²⁺ ratio governed the trend. Stronger interactions of HS⁻ or S²⁻ with Cu²⁺ sites dominated the adsorption process, and CuBDC with the lowest ratio showed the best adsorption capacity. In the moist condition, the adsorption capacity increased due to the easy dissociation of H₂S in water film. The XPS analysis confirmed the formation of Cu-S and ${\rm SO_4}^{2-}$ in the moist condition. The XRD pattern of CuBTC showed a partial loss in the crystallinity due to the formation of sulfuric acid and Cu-S bonds. The regenerated Cu-MOFs were possible using the successive effects of methanol and UV-C irradiation. The regenerated CuBTC showed an exceptionally high adsorption capacity of 95.6 mg g⁻¹, which was 3.5 times the fresh CuBTC. Restoration of low-angle peaks in the XRD pattern confirmed the reversibility of the chemisorption process. Moreover, decreased sulfur content of regenerated CuBTC, repositioning of Cu peaks, and increased surface area confirmed that methanol and UV-assisted regeneration is an affordable method to recycle Cu-MOFs with an insignificant loss in the structural and functional integrity.

Author contributions

N. K. G. was solely responsible for conceptualization, formal analysis, software, writing original draft, and review and editing. N. K. G and S. K. were in charge of data curation, methodology, visualization, and validation. J. B. and K. S. K were responsible for funding acquisition, investigation, project administration, resources, and supervision.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 A. Georgiadis, N. Charisiou and M. Goula, Catalysts, 2020,
- 2 S. L. Malone Rubright, L. L. Pearce and J. Peterson, Nitric Oxide, 2017, 71, 1-13.
- 3 R. Muñoz, L. Meier, I. Diaz and D. Jeison, Reviews in Environmental Science and Biotechnology, 2015, 14, 727-759.

- 4 M. S. Shah, M. Tsapatsis and J. I. Siepmann, *Chem. Rev.*, 2017, 117, 9755–9803.
- 5 R. Maity, H. D. Singh, A. K. Yadav, D. Chakraborty and R. Vaidhyanathan, *Chem.-Asian J.*, 2019, 14, 3736–3741.
- 6 E. Barea, C. Montoro and J. A. R. Navarro, *Chem. Soc. Rev.*, 2014, 43, 5419–5430.
- 7 W.-G. Cui, G.-Y. Zhang, T.-L. Hu and X.-H. Bu, Coord. Chem. Rev., 2019, 387, 79–120.
- 8 H.-Y. Zhang, C. Yang, Q. Geng, H.-L. Fan, B.-J. Wang, M.-M. Wu and Z. Tian, *Appl. Surf. Sci.*, 2019, **497**, 143815.
- 9 M.-H. Lee, K. Vikrant, S. A. Younis, J. E. Szulejko and K.-H. Kim, J. Cleaner Prod., 2020, 250, 119486.
- 10 C. Petit, B. Mendoza and T. J. Bandosz, *ChemPhysChem*, 2010, **11**, 3678–3684.
- 11 N. Bhoria, G. Basina, J. Pokhrel, K. S. Kumar Reddy, S. Anastasiou, V. V. Balasubramanian, Y. F. AlWahedi and G. N. Karanikolos, *J. Hazard. Mater.*, 2020, **394**, 122565.
- 12 C. Petit, B. Levasseur, B. Mendoza and T. J. Bandosz, *Microporous Mesoporous Mater.*, 2012, **154**, 107–112.
- 13 W. Xu, G. Li, W. Li and H. Zhang, RSC Adv., 2016, 6, 37530–37534.
- 14 M. Thommes, K. Kaneko, A. V. Neimark, J. P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol and K. S. W. Sing, *Pure Appl. Chem.*, 2015, 87, 1051–1069.
- 15 C. Bueno-Ferrer, S. Parres-Esclapez, D. Lozano-Castelló and A. Bueno-López, *J. Rare Earths*, 2010, **28**, 647–653.
- 16 M. Shete, P. Kumar, J. E. Bachman, X. Ma, Z. P. Smith, W. Xu, K. A. Mkhoyan, J. R. Long and M. Tsapatsis, *J. Membr. Sci.*, 2018, 549, 312–320.
- 17 A. Ahmed, C. M. Robertson, A. Steiner, T. Whittles, A. Ho, V. Dhanak and H. Zhang, *RSC Adv.*, 2016, **6**, 8902–8905.
- 18 A. Aarti, S. Bhadauria, A. Nanoti, S. Dasgupta, S. Divekar, P. Gupta and R. Chauhan, *RSC Adv.*, 2016, **6**, 93003–93009.
- 19 A. C. Elder, S. Bhattacharyya, S. Nair and T. M. Orlando, *J. Phys. Chem. C*, 2018, **122**, 10413–10422.
- 20 C. Petit and T. J. Bandosz, *Dalton Trans.*, 2012, 41, 4027.
- 21 G. Mahalakshmi and V. Balachandran, Spectrochim. Acta, Part A, 2014, 124, 535-547.
- 22 T. Bunchuay, R. Ketkaew, P. Chotmongkolsap, T. Chutimasakul, J. Kanarat, Y. Tantirungrotechai and J. Tantirungrotechai, *Catal. Sci. Technol.*, 2017, 7, 6069–6079.

- 23 M. A. Syzgantseva, N. F. Stepanov and O. A. Syzgantseva, ACS Appl. Mater. Interfaces, 2020, 12, 17611–17619.
- 24 C. Prestipino, L. Regli, J. G. Vitillo, F. Bonino, A. Damin, C. Lamberti, A. Zecchina, P. L. Solari, K. O. Kongshaug and S. Bordiga, *Chem. Mater.*, 2006, 18, 1337–1346.
- 25 G. Leofanti, M. Padovan, M. Garilli, D. Carmello, A. Zecchina, G. Spoto, S. Bordiga, G. T. Palomino and C. Lamberti, J. Catal., 2000, 189, 91–104.
- 26 H. K. Kim, W. S. Yun, M.-B. Kim, J. Y. Kim, Y.-S. Bae, J. Lee and N. C. Jeong, J. Am. Chem. Soc., 2015, 137, 10009–10015.
- 27 E. Borfecchia, S. Maurelli, D. Gianolio, E. Groppo, M. Chiesa, F. Bonino and C. Lamberti, J. Phys. Chem. C, 2012, 116, 19839–19850.
- 28 N. A. Travlou, K. Singh, E. Rodríguez-Castellón and T. J. Bandosz, *J. Mater. Chem. A*, 2015, 3, 11417–11429.
- 29 M. C. Biesinger, Surf. Interface Anal., 2017, 49, 1325-1334.
- 30 D. P. Oyarzún Jerez, M. López Teijelo, W. Ramos Cervantes, O. E. Linarez Pérez, J. Sánchez, G. del C. Pizarro, G. Acosta, M. Flores and R. Arratia-Perez, J. Electroanal. Chem., 2017, 807, 181–186.
- 31 M. Mazaj, T. Čendak, G. Buscarino, M. Todaro and N. Zabukovec Logar, *J. Mater. Chem. A*, 2017, 5, 22305–22315.
- 32 J. Liu, Y. Wei, P. Li, Y. Zhao and R. Zou, *J. Phys. Chem. C*, 2017, **121**, 13249–13255.
- 33 X. Han, Q. Cheng, X. Meng, Z. Shao, K. Ma, D. Wei, J. Ding and H. Hou, *Chem. Commun.*, 2017, 53, 10314–10317.
- 34 M. Mousavi-Kamazani, Z. Zarghami and M. Salavati-Niasari, *J. Phys. Chem. C*, 2016, **120**, 2096–2108.
- 35 A. Galtayries and J.-P. Bonnelle, *Surf. Interface Anal.*, 1995, 23, 171–179.
- 36 J. Wang, L. Wang, H. Fan, H. Wang, Y. Hu and Z. Wang, Fuel, 2017, 209, 329–338.
- 37 C. Yang, Y. Wang, H. Fan, G. de Falco, S. Yang, J. Shangguan and T. J. Bandosz, *Appl. Catal.*, *B*, 2020, **266**, 118674.
- 38 A. Dutta, N. Tymińska, G. Zhu, J. Collins, R. P. Lively, J. R. Schmidt and S. Vasenkov, *J. Phys. Chem. C*, 2018, **122**, 7278–7287.
- 39 S. Kim, N. K. Gupta, J. Bae and K. S. Kim, *Chemosphere*, 2020, 128943.