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Author(s): Mahajan, Shreya; Lahtinen, Manu

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Recent progress in metal-organic frameworks (MOFs) for CO₂ capture at different pressures

Shreya Mahajan, Manu Lahtinen

Department of Chemistry, University of Jyväskylä, P.O. Box 35, FI-40014 JY, Finland

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ABSTRACT

Global climate change ensued by the rise in atmospheric CO₂ levels is one of the greatest challenges our planet is facing today. This worldwide distress demands technologies that can contribute to our society toward "negative carbon emissions". Carbon capture and storage (CCS) technologies are in important role for capturing CO2 from existing emission sources, such as industrial and energy production point sources, before new more prominent modifications to the energy infrastructure can be implemented. Recently, alongside point source capture, direct air capture (DAC) processes have emerged as highly sought-after technologies that are able to capture CO2 from the ambient air. Alongside the traditional inorganic adsorbents, a new class of solid porous adsorbents, called as metal-organic frameworks (MOFs) have emerged in recent years also, as a group of potentially very efficient materials to capture CO₂. The promising results of MOF-based adsorbents have already achieved great interest and have contributed to their ever-accelerating research to develop new and even better adsorbents for both point source and DAC recovery technologies. This review highlights the research that has been focused on utilizing MOFs in the carbon capture processes, particularly targeting materials applicable to low CO2 partial pressures but also capturing processes in pure CO2 (1 bar) will be reviewed, because it is a widely used test condition for characterizing sorption properties of MOF adsorbents. Herein, we outline four major approaches, through which the CO2 adsorption capacity and selectivity can be boosted, including targeted modifications of the metal centers, pore size control, proper selection and substitution of linker units, and functionalization of MOFs by amines. The mechanisms of the sorption event are also reviewed from the perspective of both physisorption and chemisorption phenomena. At the end of the review, we briefly examine the variables related to the coordination of technical-economical, process-technical, and physicochemical properties of adsorbents, which both researchers and engineers should consider when developing new adsorbents and recovery processes, with emphasis on material processing, capture capacity, selectivity, regeneration cyclicity, and cost.

1. Introduction

According to the International Energy Agency report "World Energy Outlook 2021" (IEA, 2021, Paris, https://www.iea.org/reports/world-energy-outlook-2021), the worldwide energy demand has witnessed significant 2.3 % growth in the years 2018 and 2019 with an estimate of 1.3 % annual growth until 2040 [1]. The Covid pandemic is estimated to have caused a temporary reduction in energy demand of around 5 % by 2020 but it has already been caught up quickly in following years, as the Covid situation has somewhat eased globally. However, considering the overall picture in the intervening years, comprehensive demands of mankind are mostly being adhered by the consumption of accessible fossil fuels, such as natural gas, petroleum and coal, to produce energy.

The main by-product of this combustion is carbon dioxide (CO₂), lined as a major greenhouse gas [1,2]. Wherefore, the augmented quantity of CO₂ emissions over the years has led to extensive adverse effects on the natural environment, such as global warming, climate change, ocean acidification, rising sea levels, melting of the glaciers, species extinction and ozone depletion, to mention few. Thus, not only threatening humankind and the environment but also the synergy among them [3,4]. Ever since the 19th century, the abrupt and continual increase is witnessed in atmospheric CO₂ concentrations from thereabout 280 ppm (the early 1800 s) to \sim 417 ppm (2022), thus representing an increase of about 50 % compared to the average of 1750–1800. According to the Intergovernmental Panel on Climate Change (IPCC)[5,6] it is predicted that 980 ppm atmospheric CO₂ concentration could be reached by 2100

E-mail address: manu.k.lahtinen@jyu.fi (M. Lahtinen).

^{*} Corresponding author.

[7–9]. As the carbon dioxide content increases, so will the earth's temperature [10]. It is estimated that an increase of 1.4 °C observed in the 1990 s will reach an average temperature increase of 6.1 °C in 2100 if no globally significant actions are taken to mitigate climate change [11].

To address the challenges associated with increased atmospheric CO₂ concentrations it is indispensable that significant and drastic measures are taken promptly, to tackle this inevitably growing problem, by mitigating greenhouse gas concentrations and thereby limiting global warming. To remedy worldwide threats, IPCC has contemplated carbon capture as a simple and important strategy for constraining anthropogenic CO2 emissions [12]. In addition to limiting global warming to 1.5 °C, IPCC prompted expeditious, consequential, and innovative amendments in the energy sector, transport, buildings and their energy management, industrial field and cities to reduce emission levels by 45 % till 2030 [13]. To achieve these objectives, IEA has launched various projects to attain an annual CO2 emission cap of fourteen gigatons to limit the augmentation of global temperature to 2 °C. Also, the IEA regards carbon capture as one of the promising measures to reduce CO₂ emissions, of which these technologies may alone contribute up to eight gigatons of the annual aim of 14 gigatons [14–16].

CO₂ capture is prominently a "negative carbon technology". In general, CO2 capture techniques focus on CO2 emissions from the industrial origins, for instance, cement production plants, power plants, metallurgical industries, and petroleum refineries and can be classified into three main categories: pre-combustion, capture during combustion and post-combustion. Of the above, post-combustion capture is one of the most mature and favorable methods [17]. Typically, depending on the source, the temperature of flue gas is around 40-90 °C and the pressure is close to atmospheric pressure (1 bar), the CO₂ concentration of which can range from 4 % (natural gas combined cycle processes) to 13 - 15 % (coal-fired combustion plant). Main impurities include H₂O (\sim 5 %), and acid gases, such as SO_x and NO_x , in lower amounts [4]. Existing technologies include membrane separation [18,19], amine scrubbing [20-23] and adsorption with solid adsorbents [4,24,25]. Of above. the aqueous amine-based absorption ethanolamine-water solutions) has been the prevailing technique for the post-combustion based CO2 separation despite its well-known drawbacks, like unavoidable loss of amine in course of the ongoing operation, corrosive properties of the solutions and high power consumption over regeneration process due to the heating of large volumes of water with a known high heat capacity [26]. However, adsorption-based alternatives are becoming progressively more competitive.

The adsorption of CO₂ from ambient air by direct air capture (DAC) is a method that has emerged in recent years as an alternative "negative carbon technology". It has already proved to be a promising method to limit global warming. Even though our atmosphere contains already hundreds of gigatons of CO2 it's very dilute concentration (~417 ppm, 0.04 %) is a serious obstacle to the development of successful and efficient CO₂ capturing techniques due to e.g., low diffusion and uptake rates, chemical inertness and targeting selectivity towards CO2. Adsorption under very dilute CO2 causes considerable loss of entropy making a high heat of adsorption as an essential parameter to render adsorption process sufficiently spontaneous [26-29]. Despite the challenge of the adsorption process, alongside with point-source CO2 capturing, the reachability of CO2 from ambient air has encouraged researchers increasingly to develop new applications based on DAC [30, 31]. However, significantly more development is needed on the separation techniques used to capture CO₂ from different low-concentration feeds [32,33].

Research groups of Lackner et al. and Keith et. al, can be considered as significant pioneers of DAC research as they were the first to study carbon capturing properties of ion exchange resins and alkali solutions, respectively [34–37]. Up to date studies have shown how versatile and innovative DAC systems could be, capturing techniques potentially ranging from flexibility of location, being anywhere in the world, to

usability both for point and distributed CO_2 sources, albeit so far only on a small scale. Importantly DAC conditions aid in sustaining the chemical stability of the adsorbent as no exposure to SO_x and NO_x impurities is evident due to their very low concentration in the ambient air, which in contrary is not the case in industrial flue gas emissions [4,38]. Moreover, the atmosphere comprises water vapor, N_2 and O_2 which should be taken also into account as they may induce both positive and adverse effects to the sorption process itself, to target molecule selectivity, as well as to the stability of an adsorbent itself.

Despite the measures already taken the need for efficient ${\rm CO_2}$ capture technology is growing rapidly as global warming keeps accelerating. In recent years aqueous amine solutions are being substituted gradually by solid adsorbents that generally have lower heat capacities in contrast to amine-based solvents, as discussed above [39,40]. Due to the many shortcomings accompanied by amine scrubbing techniques, alternative approaches have been made to develop novel amine-functionalized solid materials both for point source and DAC conditions. Typically for DAC conditions, amine-functionalized solid materials are based on chemisorption which is the prime requirement for DAC adsorbents as it is indicated that with chemisorption stronger CO2 binding and higher selectivity (e.g. CO₂\H₂O) could be obtained due to the covalent interactions between amino groups and CO₂, in contrast to the physisorption that relies on physical less selective weak interactions. The early stage development of amine-appended (tethered/impregnated) CO2 adsorbents originated primarily from old-fashion solid adsorbents that were designed for purification of air indoors, such as in submarines and space crafts [41,42]. In the mid 90 's Leal et al. [43] reported a new CO₂ adsorbing amine-appended silica gel adsorbent. In their work the CO₂ capturing capacities were still rather modest, nonetheless, in the coming years, this new concept drew profound interest [44-50], and thereafter vide variety of amine-functionalized adsorbents have reached an immense growth in their utilizations, in all kinds of areas of carbon capture research. The constantly growing number of new materials have exhibited both improved and unprecedented combinations of properties such as high thermal and chemical stability, remarkable capture efficiency, low sorption heat, good diffusion property, low corrosion affinity, low active component leakage likelihood, and simple packaging and storing requisites [25,39,51-54].

For an exemplary good CO2-binding adsorbent, the following properties can generally be expected: high capacity (>2 mmol CO₂ per g of adsorbent), fast sorption kinetics, selectivity (>100; CO₂/other gases), recyclability (>1000 cycles), thermal stability, ease of handling and storage, as well as low production cost [55]. Of the above, the high affinity that relates to selectivity is an important criterion for adsorbent efficiency, which in part can be amplified by proper structural design of the adsorbent, as well as modification of amine type and content. Entrapment of the CO2 molecules can further be promoted by the fabrication of exclusive geometric channels which encompasses pore size modification and active site allocation. In this context certain traditional inorganic porous materials such as silica gels exhibit amorphous, microporous characteristics but lack periodic uniform porous structures, thus demonstrating low diffusion performance. Also, few Si-based mesoporous materials, like SBA-15 [56,57], MCM-41[58,59], and KIT-6[60,61] possess uniform meso-sized channel structure but the number and exact locations of amine groups remain ambiguous. Explicitly, the incorporation of strong alkaline amines in large enough pores is ideal for the chemical and reversible sorption of ${\rm CO}_2$ molecules. In short, the optimized example system works in our body, in our lungs wherein the complex network of channels with varying dimensions engenders diffusive paths for efficient gas exchange in terms of O2 absorption and CO2 desorption. Similarly, an ideal porous adsorbent should render a lung-akin network of pores. For improving the distribution of amines alongside with optimized pore properties, crystalline porous materials can be utilized, of which metal-organic frameworks (MOFs) are the most studied substances.

MOFs are a group of crystalline materials having periodic one-, two-,

or three-dimensional coordination networks constructed by metal-based nodes (single cations or clusters) bonded (coordinate covalent bond; dative bond) with bridging organic ligands. The nearly endless possibilities to combine different metal and ligand components have currently resulted in over 100,000 MOF structures in the Cambridge Structural Database (CSD) [62]. Many of them comprise exceptional characteristics for instance ultralow densities (lowest so far 0.13 g/cm³) [63], pore volumes with 90 % of free volume [64] and extremely large Brunauer-Emmett-Teller (BET) surface areas up to $10,000 \text{ m}^2/\text{g}[65-67]$. With the exemplified wide range of characteristics alongside with possibilities in their preparation designs, MOFs have already been developed for number of applications, such as gas separation and storage[25, 68], photo catalysis[69], heterogeneous catalysis[70,71], drug delivery [72,73] and medicinal fields[74,75], electrical energy storage and energy conversion[76,77], ion exchange[78], electrochemical applications[79], deterioration of chemical warfare agents[80,81], the extraction of hazardous substances from air and aquatic environments [82,83], stimuli-responsive and energetic materials [84,85], as well as for carbon capture. [11,86] Some recent achievements on carbon capture by MOF adsorbents are highlighted in Fig. 1.

As an example of the utilization of the phenomenal properties of MOFs in gas sorption, the study of Prof. Yaghi et al. [87] in 2005 can be highlighted, wherein MOF-177 was prepared with an excellent high surface area (4500 m²\g) and remarkable CO₂ uptake of 33.5 mmol g⁻ (at 35 bar and 25 °C) corresponding to about two times better CO2 loading capacity at high pressures than compared to much-used zeolites of its time; although at atmospheric pressure (1 bar and 25 °C) only 0.8 mmol g⁻¹ of adsorption capacity was observed. As suggested by Li et al. in their perspective, generally at higher pressures large surface areas seem to lead to better adsorption in contrast to lower pressures, wherein chemical properties of the adsorbent are in more dominant role in defining the CO₂ adsorption performance [88]. As exemplified by the comparative studies of Yaghi [87] and many others[89-91], one of the major benefits of MOF materials is their structural modularity, which allows accurate crystal design of the structures, aiming to consider the structural and geometrical needs on an application-by-application basis [92-94]. The distinct incorporation of appropriate metal nodes and organic linkers eventually lead to a usable structural libraries, which on

other hand has not been materialized for previously known conventional porous materials [88,95–98].

The arena of MOF research is exceptionally diverse, and a significant number of studies have already been published, particularly for CO2 capture and conversion techniques on a general level as well as also focusing specifically on individual methods [11,98,112-114]. In contrast to the previous reviews, this review focuses on recent studies on the development of amine-doped MOFs, especially for chemisorption carbon capture, mainly at CO2 pressures of one bar and below. In addition, the suitability of physisorption mechanisms for the recovery of dilute CO2 concentrations will also be reviewed. The scope of this review is to endow researchers with a detailed layout of the state of the art, areas to probe and ongoing impediments in the field of carbon capture at different CO₂ feeds, thus varying from dilute DAC (417 ppm) to flue gas concentrations (up to 15 % of CO₂), while also viewing the adsorption at 1 bar of CO₂, which is widely used as a condition for comparative testing of MOF adsorbents, mainly in dry conditions but more recently under moist conditions as well. We also go through ways to improve the carbon dioxide capture capacity, such as via targeted modifications of the metal sites, by pore size control, elongation or substitution of organic linker, and by amination of MOFs, also including amine-impregnation of MOFs. For completeness, the consideration of the water effect is also discussed, albeit briefly. The review is centered on the evolution of the MOF-74 family, though parts of the discussion also apply to other benchmarks MOFs. It does not aim to be a full account of all the MOFs reported for CO2 adsorption and separations but rather to highlight specific strategies and trends in the field of point source and DAC conditions. At the end of the review, we briefly examine the variables related to the coordination of technical-economical, process-technical, and physicochemical properties of adsorbents, which both researchers and engineers should consider when developing new adsorbents and recovery processes, with emphasis on material processing, capture capacity, selectivity, regeneration cyclicity, and cost. The review also aims to provide help for the seasoned MOF researchers to become conversant with the fundamental relationship between MOF chemical design, functional groups, and gas uptake capacity.

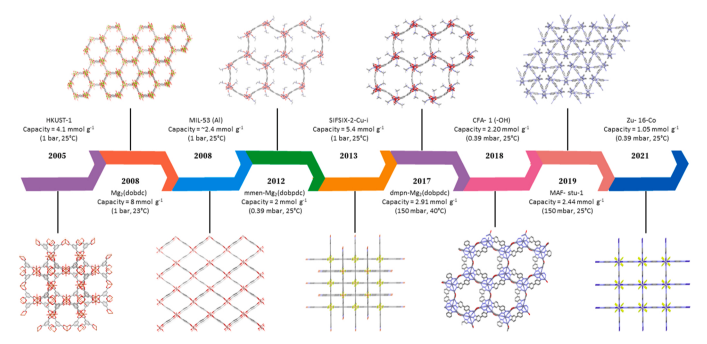


Fig. 1. Illustration of CO₂ capture capacities for some well-known MOFs (a) HKUST-1 [87] (b) Mg₂(dobdc) [99] (c) MIL-53 (Al) [100] (d) mmen-Mg₂(dobpdc) [101] (e) SIFSIX-2-Cu-i [102] (f) dmpn-Mg₂(dobpdc) [103] (g) CFA-1(-OH) [104] (h) MAF-stu-1 [105] (i) Zu-16-Co [106]. CSD entries FIQCEN, VOGTIV, SABVUN, XUXPUD, YEMTIV, ACEWIR, LAWYIV, POVGEP and SUHQUK have been used to create images with Mercury program. [101–111].

2. Approaches to improve CO₂ capture capacity of a MOF

In the field of CO_2 adsorption, with qualifications as mentioned in the Introduction (e.g., high capacity, kinetics, selectivity), the applicable adsorbent needs to bear sufficient resistance against water. Many metals used in the MOF adsorbents are vulnerable to water, which due relatively high affinity of water to the metal ions can cause their hydrolysis, and thereby the chemical deterioration of the network structure. Water has also clearly higher abundance (10–100 times) in the atmosphere compared to CO_2 which in turn has a significant effect on the CO_2/H_2O selectivity. In terms of thermodynamics, the CO_2 binding enthalpy should be considerably exothermic (at least -50 kJ mol $^{-1}$) for an efficient capture process. An adsorbent material should also be chemically and thermally robust to withstand thousands of sorption cycles without loss of activity and chemical integrity [115].

Considering the above criteria, the affinity of a MOF adsorbent for CO2 can further be altered by tailoring the chemical and physical properties of the targeted MOF structure. One of the modification strategies includes the creation of so-called active open metal sites (OMSs) in the network structure. OMSs that are part of the metal-organic framework, can provide potentially higher CO2 intake for the MOF adsorbent, as the inherent rigidity of the framework enables the OMSs to be more accessible throughout the adsorption-desorption events. Also, the pore size of the framework can be controlled via different organic linker unit designs. MOFs sorption properties can also be promoted by functionalizing the organic linkers with nitrogen (N) containing group (s), or via amines that have been appended to the OMSs. By incorporating amine functionalities (e.g., anchored to OMS or as a substitution in the bridging ligand) amidst the pore structure, polar sites (basic property) are generated that in turn interact strongly with CO₂. As a result, capturing properties of the MOF may further be improved due the more selective chemisorption occurring between an amino group and CO2. Studies have also shown that MOFs with pore interiors decorated by abundant nitrogen atoms endow stronger quadrupolar interactions between carbon and the nitrogen atoms, thereby affording higher CO2 binding affinities and selectivity. The above design strategies will be reviewed in this paper under four subsections, namely 2.1. targeted modification of metal centers to form active OMS along with ligand extension approach (reticular chemistry concept) to alter density and accessibility of OMS, 2.3. hybrid ultra-microporous materials, 2.4. amine substituted and N-heterocyclic organic linker ligands, and 2.5. post-synthetic amination and adsorption mechanisms on aminated open metal sites. On subsections 2.2. and 2.6., the impact of water as a constant competitor to CO₂ in the sorption event is also reviewed. Finally, chapter 2.7. represents coating techniques as further measures to increase the tolerance of MOF adsorbents against water. The literature reviewed is generally limited to publications from 2015 to date, and older studies have been included only when they have represented significant advances in carbon capture research in general. For the sake of simplicity, all the metrics in this review are standardized to the following units: adsorption capacity (mmol g-1), temperature (°C) and pressure (mbar or bar).

2.1. Modifications of the metal sites and ligand extension approach

The physisorption on a solid surface is viewed as the interaction of CO_2 preponderantly through intermolecular weak forces like van der Waals or somewhat stronger hydrogen bonding. Fundamentally, in this process, there is no chemical bond formation owing to the absence of chemical interaction. Simple condensation principles are favored which are inclined towards layer-by-layer deposition. The adsorption heat $(-Q_{st})$ of CO_2 is the underlying physical attribute, which prompts the physisorption of CO_2 to a MOF's pore structure. Generally, in a weakly binding physisorbent, the binding energies are considerably less than that of a rational chemical bond, and sometimes intermittently lower than a hydrogen bond. Widely acknowledged boundary value for Q_{st}

chemisorption–physisorption varies between -40 and -50 kJ mol⁻¹[55].

As MOFs are generally highly ordered porous materials, the solvent content in the voids, as well as the ones coordinated to metal centers, can be removed either via a solvent exchange, heat, vacuum, or both to form OMSs without changing the original network structure. MOFs with open metal sites utilize Lewis's acidity, imparted by partial positive charges on metal sites, to interact with CO_2 molecules ($O=C=O\cdots M^{n+}$). Thereby, the aforesaid prerequisite to increase -Qst of CO2 can be fulfilled by forming strong and selective interactions between the host structure and the CO2, via open metal sites. Among the vast family of frameworks, MOF-74 is an extensively studied MOF archetype due to its high density of available open metal sites, and because it can be varied chemically with ease by using a number of different bivalent metals, such as Mg(II), Mn(II), Fe(II), Co(II), Ni(II), Cu(II), and Zn(II) [116]. Structurally MOF-74 also known as M2(dobdc) is consisted of desired metal ion (M) and 2,5-dihydroxy-1,4-benzenedicarboxylic acid (H₄dobdc) which forms a three-dimensional framework having one-dimensional (1D) hexagonal channels parallel to the crystallographic c-axis. In the network structure, solvent molecules are coordinated at every sixth position of the octahedral metal centers, the removal of which generates an open metal site that serves as a Lewis acidic site and eventually can interact with an incoming CO2 molecule [114,116]. Among the series of M-MOF-74 analogues, the Mg-MOF-74 has so far illustrated the highest so far reported CO2 adsorption capacity of \sim 9.9 mmol g⁻¹ under pure CO₂ (25 °C and 1 bar) [117] and adsorption heat of - 47 kJ mol⁻¹ [118] Yazaydin et al. [119] and Caskey et al. [99] made a similar comprehensive screening for CO2 capture performance using different coordination metals in isoreticular series of MOF-74. The study unfolded that with a decrease in the M-O bond length, the CO2 uptake of M-MOF-74 uplifted. It was therefore concluded that the bond lengths seem to define the oxygen affinities of metals in the coordination state (Mg-O (1.969 Å) < Ni-O (2.003 Å) < Co-O (2.031 Å) < Zn-O (2.083 Å)). The obtained adsorption heats for the series were obtained through isotherms that ranged from - 22.1 to - 43.5 kJ mol⁻¹, of which the highest being on Mg-MOF-74 [116]. Based on the experiments carried out, it could be assumed that the highest affinity for CO2 in the case of Mg-MOF-74 was attributable to the enhanced ionic character of the Mg-O bond, which bestows stronger charge-quadrupole interactions between the OMSs and CO₂ [99].

Furthermore, to improve the accessibility and density of OMSs in MOF-74 family, various isoreticular frameworks have been designed by employing the ligand extension approach, wherein a longitudinal dimension of a bridging ligand is elongated but keeping the other properties like the number, type or relative geometrical orientation of functional groups unchanged [101,120,121]. This method utilizes facile incorporation of functionalities by the modulation of the pore size to compensate the moderately small pore size of the native MOF-74 framework that is a limiting factor for the accessibility of OMSs. For instance, McDonald et al. examined pore-expanded MOF-74 variants that were based on the extended analogies of terephthalic acid linkers, such as H₄dotpdc (4,4"-dihydroxy-[1,1':4',1"-terphenyl]-3,3"-dicarboxylic acid), and H₄dobpdc (4,4'-dihydroxy-[1,1'-biphenyl]-3, 3'-dicarboxylic acid) [101]. Also Yoo et al. [121] examined similar pore-expanded structures but using different metal centers such as Ni (II), Mn(II), Zn(II), and Co(II) cations. The adsorption heats calculated from the well-known Clausius-Clapeyron relationship were obtained as -25.0, -33.9, -34.5, and -25.6 kJ mol⁻¹ for Ni₂-, Co₂-, Mn₂-, and Zn₂(dobpdc) MOFs, respectively. The observed trend in adsorption heats implies that there exists a direct relationship between the Lewis acidity of a metal cation and the binding ability of CO₂ molecule. Furthermore, Lim and coworkers [122] illustrated a series of Co-based MOF-74 variants [Co₂(L)(DMF)_x(H₂O)_v]·zH₂O that were made using H₄dobdc, -dobpdc and -dotpdc ligands (L), and microwave-assisted solvothermal synthesis (Fig. 2a).

Three of the frameworks displayed isoreticular structures as

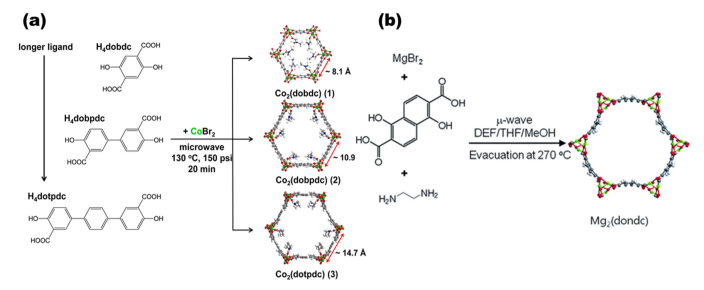


Fig. 2. Synthesis schemes of some (a) Co-, and (b) Mg-based extended MOF-74 structures. Adapted from Ref. [122,123].

validated by powder X-ray diffraction (PXRD) analyses. For the reported systems, Co–Co distances at the edge of pore channels ranged from 8.1 via 10.9–14.7 Å for Co₂(dobdc), -(dobpdc) and -(dotpdc), respectively. These studies emphasized the importance of ligand length and how the interchain interaction is systematically governable by the length of the ligand.

Another well-known strategy for a ligand extension is to utilize fused benzene ring bedded ligands, such as naphthalene and anthracene. Yeon *et al.* reported isostructural member of MOF-74 family, that is Mg₂(dondc) consisted of 1,5-dihydroxy-2,6-naphthalendicarboxylic acid (H₄dondc) [123]. The framework features one-dimensional channels which are nearly 15 Å in size (Fig. 2b), hence settling between Mg₂(dobdc) and Mg₂(dobpdc) with their channel sizes being approximately on the order of 11 and 18.4 Å, respectively. The study concluded that the availability of OMSs in Mg₂(dondc) were the primary binding sites of CO₂. The adsorption capacity of Mg₂(dondc) was somewhat like on Mg₂(dobpdc), thus 4.80 vs. 4.84 mmol g⁻¹, respectively but still smaller than on Mg₂(dobdc) showing 6.1 mmol g⁻¹ capacity at 150 mbar (25 °C). Similarly, Wang and coworkers [124] examined the

effects of the high density of OMSs for carbon capture by designing a porous MOF, $Co_2(tzpa)(OH)(H_2O)_2\cdot DMF$ using 5-(4-(tetrazol-5-yl) phenyl)isophthalic acid (H₃tzpa) consisting of tetrazole and two carboxylic acid groups. For the prepared adsorbent, the experimental gas sorption data illustrated good CO_2/CH_4 selectivity of 31.8 for the gas mixture (1:1 ratio), which was concluded to be the result of high density of open metal sites.

To further investigate properties of the open metal sites, mixed-metal MOFs consisting of two or more different metals have also been prepared. In the case of MOF-74 archetype, the synergistic effect of the bimetallic system demonstrated better gas adsorption and catalytic performances [125–127]. For instance, M/Zn(dobdc) series having evenly distributed metal cations (about 1:1 ratio) in Mg/Zn-, and Ni/Zn-MOF-74 s exhibited adsorption capacities of 6.7 and 6.3 mmol g⁻¹ with adsorption heats of -43 and -41 kJ mol⁻¹, respectively (0 °C and 1 bar) [128]. In Particular, the Ni/Zn-MOF-74 demonstrated a higher heat of adsorption than that of the monometallic Ni-MOF-74 indicating a synergistic effect of mixed metal ions. Similarly, Kim et al. [129] examined bimetallic (M = Mg, Ni, Co, and Mn) MOFs with extended

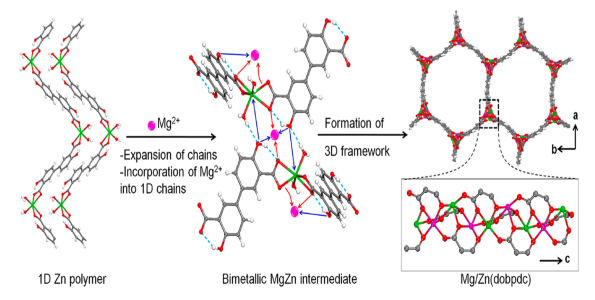


Fig. 3. Suggested mechanism of dimensional transformation from 1D-chain template to a 3D bimetallic Mg/Zn(dobpdc) framework. Adapted from Ref. [129].

ligand (Fig. 3), namely M/Zn(dobpdc). It was found that based on the energy dispersive microscopy (TEM- and SEM-EDS) only the Mg/Zn (dobpdc) showed a 1:1 ratio between the incorporated metals, and higher adsorption capacity than that reported for the monometallic Zn₂(dobpdc), albeit still with a capacity lower than Mg₂(dobpdc). On the other hand, adsorption heat for Mg/Zn(dobpdc) was observed to be in the range of -32.5–44.2 kJ mol⁻¹ (Table 1), hence settling in between Mg₂(dobpdc) and Zn₂(dobpdc) adsorbents.

Table 1 Summary of $\rm CO_2$ adsorption for MOFs with targeted modifications of the metal sites.

MOF	Capacity (mmol g ⁻¹)			-Q _{st}	Temp.	Ref.	
	0.39 (mbar)	150 (mbar)	1 (bar)	(kJ mol ⁻¹)	(°C)		
Mg ₂ (dobdc)	0.088	6.1	~8–9	47	25	[99,	
	\0.14 ^H					116,	
						144,	
						150,	
						153,	
						154]	
Mg ₂ (dobpdc)	0.13	4.85	6.42	44	25	[101]	
Ni ₂ (dobdc)		~4.06 * *	~7.0	39	25	[116,	
En (dobdo)			7.1	22	25	119]	
Fe ₂ (dobdc)			~7.1	33	25	[116,	
Co ₂ (dobdc)		~2.81 * *	~6.90	34	25	151] [99,	
CO2(dobac)		2.01	0.50	54	23	116,	
						110,	
Zn ₂ (dobdc)		~1.30 * *	~5.61	27	25	[99,	
Ziiz(dobde)		1.00	0.01	_,	20	116,	
						119]	
Mn ₂ (dobdc)			~5.9	32	25	[116]	
Cu ₂ (dobdc)			~2.90	22	25	[116]	
Mg ₂ (dondc)		4.80	6.42	34-44	25	[123]	
Mg ₂ (dotpdc)			~5.30	~40	25	[120]	
Mg ₂ (pc-			~6.50	~38	25	[120]	
dobpdc)							
Mn ₂ (dobpdc)		~ 1.95		34	25	[121]	
Co ₂ (dobpdc)		~ 1.88		35	25	[121]	
Ni ₂ (dobpdc)		$\sim \! 2.70$		25	25	[121]	
Zn ₂ (dobpdc)		~1.40		26	25	[121]	
Mg/Zn			5.10	43	25	[128]	
(dobdc)							
Ni/Zn			4.80	41	25	[128]	
(dobdc)			5 00	00.44	0.5	F1.003	
Mg/Zn			~5.90	33–44	25	[129]	
(dobpdc)	o ooH	0.51	0.50	E 4	0.5	F1.4.4	
SIFSIX-3-Cu	0.32 ^H	2.51	2.58	54	25	[144,	
SIFSIX-3-Zn	\1.24 0.13	2.43	2.51	45	25	155] [102,	
31F31A-3-ZII	0.13	2.43	2.31	43	23	144]	
SIFSIX-3-Ni	0.18^{H}	2.47	2.5	47	25	[150]	
SIFSIX-18-Ni-	0.18 *	2.47	2.3	52	25	[148]	
β	0.0			02	20	[1 10]	
SIFSIX-2-Cu-i	0.03^{H}	1.25 * *	5.41	32	25	[102,	
	\0.068					144,	
						155]	
HKUST-1	0.05^{H}	1.59	2.5	34	25	[87,	
			\4.1			150]	
UTSA-16		2.37	4.3	34.6	25	[138]	
MIL-101(Cr)			1.22	21	25	[152]	
ZU-16-Co	1.05		2.87		25	[106]	
TIFSIX-3-Ni	0.67	2.36	2.32	50	25	[106,	
	\1.7 *					146,	
						148]	
NbOFFIVE-1-	1.8 *	2.15	2.2	55.4	25	[145,	
Ni	\1.3					148]	
SIFSIX-14-			4.70	37.7	25	[147]	
Cu-i		${\sim}2.86^{H_{\pm\pm}}$. A F1	20.2	25	F1 407	
dptz-CuTiF6		~2.80 ^^	~4.51	38.2	25	[149]	
MIL-53(Al)			~2.4	30	25	[100]	

^{*} Measured at 1 mbar, ** at 100 mbar, and H humid condition.

2.2. Consideration of water effects

Water is a ubiquitous component of the atmosphere, and its competitive adsorption with CO2, and its influence on the chemical stability of MOFs in carbon capture applications is a serious challenge to be considered. The MOF-74 family and many other MOF designs display lower CO₂ capture capacities under humid conditions due to the competition between water and CO2 at the active binding sites. In addition, water has significant adverse effects on the chemical stability of many MOF materials, which begin to increase especially when attempting to increase the number of adsorbent regeneration cycles from a few to a practical extent of thousands of cycles, i.e., to extend the life and efficiency of the adsorbent. Inevitably, competition between the guest molecules results in the reduction of reactive sites (e.g., OMSs), as water may permanently terminate the sites and eventually can even break the network structure via hydrolysis [130]. Along with MOFs, also conventional inorganic adsorbents are prone to water, as physisorbents such as zeolite 13X and activated carbon have encountered similar adverse effects [9,131]. To improve the water resistance of MOFs, significantly more research is needed in this area, especially for DAC adsorbents, which struggle with the low concentration of CO₂ compared to the concentration of water that is several orders of magnitude higher. For the utilization of OMSs in DAC adsorbents, a better understanding of the adsorption mechanisms is needed for revealing the key factors which may result in better stability and higher selectivity towards CO2 in the presence of competing water [132–134]. In this context, recent studies have demonstrated some progress, e.g., in CO₂ selectivity and dynamics in the presence of water, or other molecules competing for absorption. This positive progress gradually will pave the way towards more water-resistant MOFs, which eventually will supersede the moisture-sensitive MOF materials of the previous generations [135–137].

Xiang et al. have reported UTSA-16 adsorbent (K(H2O)2Co3(cit) (Hcit)) that was prepared using citric acid (H₄cit) which showed reduced affinity for water but still having high density of open metal sites [138]. The experimental data on the CO2 working capacity of the adsorbent showed only minor differences between humid and dry conditions (0.94 and 1.30 mmol g⁻¹, respectively) indicating improved CO₂ separation properties. The observations raised the hope that UTSA-16 could be, at that time, a new efficient adsorbent from both academic and industrial perspectives [139,140]. In addition to the density of OMSs, the pore structure of the adsorbent is also of great importance for the accessibility of the active reaction sites and thus for the total adsorption capacity of the MOF. With a different approach to diminish effects of water, the partial or full substitution of the existing metal centers with metals having a smaller ion radius have been shown to shrink the pore size of MOFs, and causing an increase in -Qst value, which partly defines the adsorption efficiency of the adsorbent [141]. This post-synthetic exchange method offers a promising platform to enhance chemical stability of a MOF and allows to fine-tune its water-resistance while still maintaining the original network structure [142]. For instance, Hu et al. [143] demonstrated a persuasive strategy to functionalize somewhat more water-resistant UiO-66(Zr) MOF by the metalated-ligand exchange (MLE), in which the terephthalic acid (bdc) was exchanged to 1,2,4, 5-benzenetetracarboxylate and its auxiliary sites were occupied by alkali metals (Li⁺, Na⁺ and K⁺). With the described method, the pore size of post-functionalized UiO-66 was able to be altered (for van der Waals interactions) with simultaneous insertion of more polar metal sites (for electrostatic interactions), resulting in to improved CO2\N2 selectivity and CO₂ working capacity. The presented method opens new alternative ways to design water-stable MOFs with potentially higher crystallinity and CO2 separation efficiencies.

2.3. Hybrid ultramicroporous materials (HUMs)

One way to avoid adverse effects of water, are the recently developed

anion-functionalized MOFs consisting of metal ions, organic linkers, and inorganic bridging anions. Even though the potential open metal sites are now being used as coordination sites for anionic species, some of these frameworks have shown exceptional CO₂ uptakes due to the strong interactions between CO2 and the secondary anionic building units of the framework. Shekhah et al. [144] introduced a new strategy producing novel SIFSIX-3-M (MSiF₆(pyrazine)₂.2H₂O) MOFs that contain ultramicropores and have high CO₂ capture capacity and selectivity via physisorption in DAC conditions. The reported MOFs were constructed from MSiF₆ inorganic units (M = Cu, Ni, and Zn) and a pyrazine ligand (Fig. 4). Resulting adsorbents exhibited excellent CO2 sorption energetics and fully reversible physisorption driven adsorption-desorption process under moderate conditions (at 50 °C under vacuum). This was concluded to have occurred, because of the highly electrostatic inorganic anion (SiF_6^2) and the small pore size. Also, in terms of adsorption heat, SIFSIX-3-Cu demonstrated – 54 kJ mol⁻¹ enthalpy in DAC conditions with a steep adsorption isotherm, and 1.24 mmol g⁻¹ capturing capacity (Fig. 4d), thus being significantly better than that of previously reported for Mg-MOF-74. Unfortunately, long-term exposure of SIFSIX-3-M MOFs to water vapor ended up to lower CO2 uptake capacities, limiting the applicability of the adsorbents only to dry adsorption conditions. To address this challenge, Bhatt et al. attempted to develop more water-resistant SIFSIX-typed MOFs for carbon capture by selecting octahedrally coordinating Ni(II) as center metal, which has a shorter M–F bond distance ($d_{M-F}=1.99\ \text{Å}$) than for example in Cu (II)-F. [145] Secondly, the $(SiF_6)^{2-}$ pillaring anion was changed to a more strongly binding (NbOF₅)²⁻ anion, as it practically supersedes the SiF_6^{2-} anion by having bigger ionic radii of the Nb(V) center, resulting in a longer Nb-F bond length compared to Si-F bond (1.899 Å and 1.681 Å, respectively). Secondly, the better nucleophilic characteristics of the (NbOF₅)²⁻ may potentially contribute to better water resistance of the MOF in humid conditions. The prepared NbOFFIVE-1-Ni MOF indeed exhibited similar 1.3 mmol g-1 CO₂ capture capacities both in dry and humid conditions indicating clearly improved tolerance against water vapor. The observed capacity is about 15% higher than that obtained for the analogous SiFSIX-3-Cu. In comparison to the reported physisorption adsorbents at the time, NbOFFIVE-1-Ni MOF turned out to

be the top adsorbent for CO_2 capture both directly from the air and flue gases. Moreover, its straightforward synthesis method has gradually paved the way for economically feasible production of the material in bulk scales (g to kg).

Kumar et al. [146] further investigated SIFSIX analogues using TiF₆² pillars with an aim to enhance CO₂ affinity at lower pressures without worsening their thermal stability and water resistance properties. They synthesized hybrid ultramicroporous material (HUM) TIFSIX-3-Ni which contains TiF₆² (TIFSIX) and pyrazine (3) connected to Ni nodes. The 14-days accelerated stability tests for the adsorbent showed only a minimal decrease in its CO_2 uptake at the end of the runs. The improved thermal stability resulting from the change of an anion type from SiF₆² to TiF_6^{2-} further underlined how the fine-tuning of the pillaring groups remarkably impacted both the physical and sorption characteristics of otherwise isostructural MFSIX-3-M adsorbents. In a similar study, Jiang et al. [147] presented a twofold interpenetrated copper coordinated network with a sequential pocket-like pore structure known as SIFSIX-14-Cu-i containing SiF₆² (SIFSIX) anions and 4,4'-azopyridine (14) ligands (Fig. 5a). Mukherjee et al. [148] represented a new member of HUM, namely SIFSIX-18-Ni-β ([Ni(L)₂(SiF₆)]_n. It was prepared using 3,3',5,5'-tetramethyl-1 H,1'H-4,4'-bipyrazole (L) ligands and containing both high-affinity CO₂ binding sites and enhanced hydrophobicity resulting from pores coated by methyl groups. It exhibited high CO2 uptakes at low pressures (0.5, 1 and 10 mbar) with 0.4, 0.8 and 2.2 mmol g⁻¹ capacities, respectively. It was suggested that the formed hydrophobic pockets in this material resulted in relatively high adsorption heat $(-52 \text{ kJ mol}^{-1})$ and low affinity for H₂O. Water adsorption tests for the material (95% relative humidity) revealed somewhat low uptake of 1.66 mmol g⁻¹ in comparison to the other three HUMs with 7.5, 8.8 and 10.1 mmol g⁻¹ for TIFSIX-3-Ni, SIFSIX-3-Ni and NbOFFIVE-1-Ni respectively. More recently, Liang et al. [149] presented a synthesis method for dptz- $CuTiF_6$ using 3,6-di(4-pyridyl)–1,2,4, 5-tetrazine (dptz) as the ligand (Fig. 5b). The prepared dptz-CuTiF6 MOF exhibited similar $\sim 2.86 \text{ mmol g}^{-1} \text{ CO}_2$ capture capacities (100 mbar, 25 °C) both in dry and humid conditions indicating the CO₂-capture ability of this material is not affected by moisture.

In the year 2021 Zhang and coworkers [106] demonstrated a simple

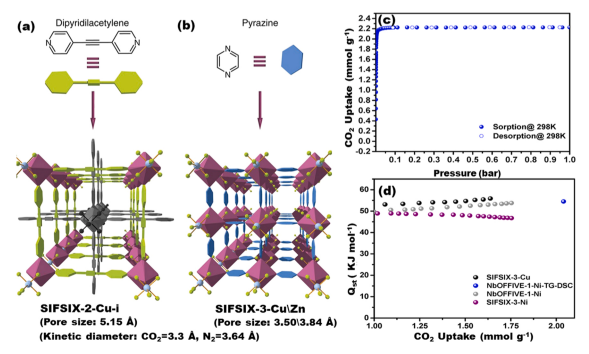


Fig. 4. Crystal structures of (a) SIFSIX-2-Cu-i (b) SIFSIX-3-Cu/Zn, (c) CO_2 sorption isotherm for NbOFFIVE-1-Ni at 25 °C and (d) comparison of CO_2 heat of adsorption for selected HUMs. Adapted from Ref. [144,145].

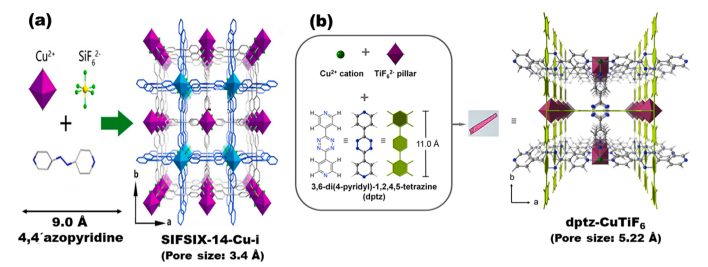


Fig. 5. Crystal structures of (a) SIFSIX-14-Cu-i, and (b) dptz-CuTi F_6 MOFs. Adapted from Ref. [147,149].

synthesis for novel anion-functionalized ZU-16-Co (TIFSIX-3-Co) containing TiF₆² anions and pyrazine ligands (3). ZU-16-Co demonstrated structural similarities with the NbOFFIVE-1-Ni, thus having one-dimensional (1D) pore channels decorated by fluorine atoms. Among the isostructural MFSIX-3 (M = Si, Ti, Ge) family, the reported ZU-16-Co showed high CO2 uptakes both at low and atmospheric pressures (10 and 1000 mbar) with 2.63 and 2.87 mmol g⁻¹ capacities, respectively. Also, a relatively good 1.05 mmol g⁻¹ CO₂ capacity was observed in DAC conditions. The improved CO₂ capture properties were assumed to result from the fine-tuned optimal pore size ($\sim 3.62 \, \text{Å}$) which could arise because of a stronger F···C=O host-guest interaction. Overall, amongst all the HUMs, so far the NbOFFIVE-1-Ni MOF has been found to have the highest CO₂ uptake under DAC conditions, along with reasonably low regeneration energy. It is clear, that more research is needed to understand how changing the pore size and shape affects the water tolerance of pillared MOFs and their selectivity towards CO2 in humid conditions. Selected CO2 capture capacities for the adsorbents presented above, as well as for a few well-known physisorbent MOFs, such as HKUST-1, Fe₂(dobdc), MIL-53(Al) and MIL-101(Cr) found in the references [100,150–152] are shown in Table 1.

2.4. Amine substituted and N-heterocyclic organic linker ligands

Besides the OMSs and HUMs, of which high affinity towards CO2 originates from their Lewis acidic sites, micro-channeled structure, and the electrons of the charged functionalities (e.g., F), the uncoordinated amino groups substituted on the organic linkers can also act as an active polar binding site for CO₂. Amine functionalization can typically be carried out with the help of aromatic or heterocyclic (e.g., N or S containing) organic linker ligands, which are substituted with one or more amino groups. Aliphatic or aromatic free amines can also be added to the framework using post-synthetic methods (PSM), by which the amines can be attached to OMSs, or can be located as free molecules filling the pores of the adsorbent. Along with the amino group, other polar functional groups, such as perfluorinated aliphatic groups (e.g., CF₃), halogens (e.g., Cl, Br) and hydroxyl groups can be used as substituents in the bridging ligands. Typically, the backbone structures of aminated bridging ligands are, for example, aminophenyl, aminobiphenyl, aminonaphthyl, aminotriphenyl and aminoazole groups, of which needs to also contain some bonding groups, such as carboxylic acids or tertiary amines contributing to the formation of the actual framework. By incorporating an amine functionality directly to a linker ligand, has some advantages over the amination of OMS or impregnation of amines. In case of the amine-substituted linker ligand the accessible pore volume for the quest molecules is greater than, if free (poly)amine molecules fill the pores, or are bound to open metal sites of the MOF adsorbent. Also, with aminated linker ligands, an adsorbent can typically be prepared in one or two synthesis steps, whereas with PSM additional preparation steps are needed. For example, aminating OMSs of a pristine MOF includes typically solvent exchange and or further activation steps to remove both the metal coordinated terminal solvents along with solvents in the pores. This is then followed by the actual post-synthetic amination of the freshly created OMSs. Despite of more multi-step synthesis path, PSM is also useful especially when a certain kind of chemical functionality is desired for the adsorbent, but it cannot be a substituent in the bridging ligand in the actual synthesis step of the MOF, for example, due to the chemical reactivity of the group, thermal instability, or unwanted coordination seeking towards the node metal. In the following paragraphs, MOF adsorbents based on various aminated linker ligands and their metrics for CO2 recovery will be reviewed, considering different adsorption conditions, including DAC, flue gases and pure CO₂ that is also used to a significant extent for determining the recovery capacities of adsorbents.

Among countless number of aminated linker ligands, 2-amino-terephthalic acid (NH2-H2bdc) can be highlighted as one of the most widely utilized functionalized organic ligand for preparing aminated MOFs since 2001, when Braun et al. first published MOF-46 made from NH₂-H₂bdc [156]. In their study MOF-2 made of terephthalic acid (bdc) was compared to MOF-46 and it was found that the original framework topology of MOF-2 persisted in the aminated MOF-46. Indeed, since the aminated and the original ligand are often structural analogues, the same synthesis methods can be used for the aminated ligand as for the original ligand. Predominantly, the similarity of the ligand structure allows the topology of the network structure to remain unchanged. In 2002 Eddaoudi et al. [157] reported aminated IRMOF-3(Zn) that belongs to isoreticular series of MOF-5 (bdc with Zn) and, which was tested at the time for methane capture but has later utilized also for carbon capture (\sim 1.4 mmol g⁻¹ at 25 °C and 1 bar CO₂) [158]. Also, some more recent studies for PSM modified IRMOF-3 can be found. For example, Ding et al. reported preparation methods for N-doped porous carbon monoliths (NCMs) via direct carbonization of IRMOF-3.[159] In this process, the partial carbonization modifies the originally microporous structure to contain more meso- and macropores which in turn reveal more amino groups to which CO₂ can readily chemisorb (Fig. 6). Based on the research, it could be concluded that the modified NCM (IRMOF-3/800 in the sorption plot) demonstrated better CO₂ recovery capacity than the pristine IRMOF-3 with 1.64 vs. 0.70 mmol g⁻¹ capacities for NCM and IRMOF-3, respectively at 150 mbar of CO2 (0 °C).

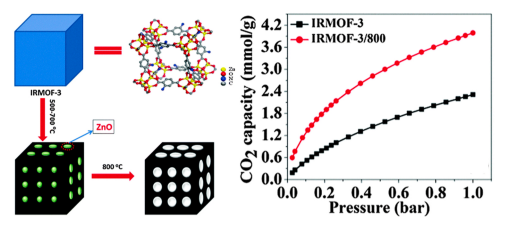


Fig. 6. (Left) Illustration for the transformation of IRMOF-3 to NCMs, and (right) CO₂ sorption isotherms of IRMOF-3 and modified NCMs at 0 °C. Adapted from Ref. [159].

Nearly doubled capacity (\sim 3.99 vs 2.32 mmol g⁻¹) in contrast to IRMOF-3 was observed for NCM at 1 bar CO₂ at 0 $^{\circ}$ C, as can be seen in Fig. 6.

In 2005 Ferey et al. were the first ones to report the nowadays well-known and extensively studied MIL-101(Cr) MOF ([Cr₃F(H₂O)₂O (bdc)₃,nH₂O], ($n\approx25$) which has mesoporous structure, remarkable chemical and thermal stability and, when activated contain OMSs [160]. A number of different aminated MOF analogues have also been prepared from the MIL-101 archetype over the years [161–164], as well as plethora of other MOF topologies which are based on NH₂-bdc ligand, such as NH₂–MIL-101(Al) [165], NH₂–MIL-125(Ti) [Ti₈O₈(OH)₄ (NH₂–bdc)₆] [166], NH₂–MIL-68 (USO-3-In-A) [In(OH)(NH₂–bdc)] [100], NH₂–MIL-53 [Al(OH)(NH₂-BDC)] [167,168], CAU-1 [Al₄(OH)₂(OCH₃)₄(NH₂–bdc)₃] [169], and NH₂–UiO-66 [Zr₆O₄(OH)₄(NH₂–bdc)₆] [170], to mention a few (Fig. 7). Aminated HKUST-1 framework [Cu₃(NH₂–btc)₂] have also been made from 2-amino-1,3,5-benzenetricarboxylic acid (NH₂-btc), as reported by Peikert et al. [171].

Of the above, the NH₂-MIL-53(Al) is another interesting example of NH₂-bdc containing MOF, as it is known for its specific reversible structural elasticity (breathing network) that can be induced by an external stimulus like heat, or the arrival of a guest molecule in adsorption event. Relatively good CO₂ adsorption capacities (e.g., 2.3 mmol g⁻¹ at 1 bar CO₂) with moderate enthalpy of adsorption ($-38.4 \text{ kJ mol}^{-1}$) has been reported for it. These are comparable with the other MOFs shown in Table 2. Furthermore, Dhankhar et al. [174] reported mixed ligand MOF1 ([Zn₂(NH₂BDC)₂(dpNDI)]_n, having N,N´-di (4-pyridyl)- 1,4,5,8-naphthalenediimide (dpNDI) and NH₂bdc ligands. The framework is formed as a 2-fold interpenetrated 3D pillar-layered system with 1D channels decorated by amino groups. The microporous MOF1 demonstrated 1.51 mmol g⁻¹ adsorption capacity and enthalpy of - 46.5 kJ mol $^{-1}$ that were clearly better than that observed

for MOF analogue made with bdc $(-36.0 \text{ kJ mol}^{-1})$. Das et al. [175] designed different mixed ligand MOFs by varying amino, azole and thiophene groups, namely Zn-DAT {[Zn₂(TDC)₂(DATRZ)]·(3 H₂O)· $(DMF)_{n}$, including 2,5-thiophenedicarboxylic acid (H_2TDC) and 3, 5-diamino-1,2,4-triazole (DATRZ) ligands. Whereas Zn-TAZ $\{[Zn_2(TDC)(TRZ)_2]\cdot (DMA)\cdot (MeOH)\}_n$ was made with TDC and 1,2,4-triazole (TRZ) ligands. Interestingly, the amino group containing Zn-DAT showed higher CO₂\N₂ adsorption selectivity and adsorption heat than the simpler triazole containing Zn-TAZ (-39.5 vs. -31.7 kJ mol⁻¹, respectively). In the following study Das et al. [176] reported interesting hydrogen bonded 3D framework, HbMOF1 {[Zn(hfipbba)(MA)]. 3DMF_n made by using 4,4'-(hexafluoroisopropylene)bis-benzoic acid (hfipbba), and amino-rich melamine (MA) ligands. In this case addition of polar (-CF₃) and basic (-NH₂) substitution groups adsorbent proved to be even better than the adsorbents reported earlier by Das et al., as its adsorption heat (-36.7 kJ mol⁻¹) and particularly adsorption capacity (2.21 mmol g⁻¹ at 0 °C) increased significantly.

In addition to the amino group, other N-comprising ligands have been reported to increase the affinity of CO_2 into the pores of the framework <code>[177-180]</code>. For instance, Li et al. <code>[181]</code> prepared $\text{Cu}_4[(\text{C}_{57}\text{H}_{32}\text{N}_{12})(\text{COO})_8]$ consisting of nitrogen-rich triazole-cored octacarboxylic acid ligand (5,5′,5′,5′-((methanetetrayltetrakis(benzene-4,1-diyl))-tetrakis(1 H-1,2,3-triazole-4,1-diyl))-tetraisophthalic acid). The sorption tests revealed remarkable CO_2 uptake of 7.18 mmol g $^{-1}$ at 1 bar (0 °C), and it was concluded that the high capacity of the adsorbent resulted from the synergy between the N atoms and open Cu sites while interacting with approaching CO_2 molecules. Similarly, bio-MOF-11 $[\text{Co}_2(\text{ad})_2(\text{O}_2\text{CCH}_3)_2]$ having adenine (ad) ligands and acetate anions, showed high 6.0 mmol g $^{-1}$ capacity along with excellent CO_2/N_2 selectivity up to 81 at 1 bar (0 °C). The reported MOF consists of mixed acetate/adeninate cobalt paddlewheel units, where

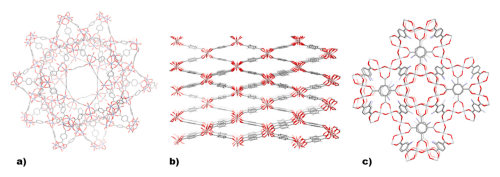


Fig. 7. (a) Partial structure views of MIL-101(Cr), (b) MIL-53(Al) having analogous structure to NH₂-MIL-53(Al), and (c) NH₂-MIL-125. CSD entries OCUNAC, SABWAU01 and MAWBUK have been used to create images with Mercury program Adapted from Ref. [110,111,160,172,173].

Table 2Summary of CO₂ adsorption for MOFs with targeted modifications to linker unit.

MOF	Capacity (mmol g ⁻¹)	-Q _{st} (kJ	Temp.	Ref.		
	150 (mbar)	1 (bar)	mol ⁻¹)	(°C)		
IRMOF-3		~1.4		25	[158]	
IRMOF-3\800	1.64	3.99	29.6	0	[159]	
NH ₂ -MIL-101(Cr)		1.9	~50	25	[161,	
					164]	
NH ₂ -MIL-53(Al)		~2.3	38.4	30	[167,	
					168]	
NH_2 – MIL -125		2.2		25	[11,	
					166]	
CAU-1		3.9	48	25	[169]	
UiO-66 –NH ₂	~1.34	~ 2.65	28	25	[155,	
0 0771.)	$\backslash 1.16^{H} \sim 0.25 *^{H}$	5 06		0.5	170]	
Cu ₃ (NH ₂ btc) ₂		5.26	46.5	25	[171]	
MOF1		1.26 1.39	46.5	25	[174]	
HbMOF 1 IFMC-1		2.7	36.7 30.7	25 25	[176] [211]	
[Zn(btz)]		4.9	31.2	25 25	[211]	
bio-MOF-11		4.1	~45	25	[182]	
NJU-Bai50	0.0018 * \ 0.49	7.1	26.6	25	[195]	
NJU-Bai51	0.13 * \ 0.90		33.3	25	[195]	
IR-MOF-74-III-	0.15 (0.50	2.96	55.5	25	[218]	
CH ₃		2.50		20	[210]	
IR-MOF-74-III-		3.18		25	[218]	
NH_2						
IR-MOF-74-III-		2.09		25	[218]	
CH ₂ NHBoc						
IR-MOF-74-III-		1.91		25	[218]	
CH ₂ NMeBoc						
IR-MOF-74-II-		3.35		25	[218]	
CH_2NH_2						
IR-MOF-74-III-		2.87		25	[218]	
CH ₂ NHMe						
IR-MOF-74-III-		3.00		25	[219]	
(CHNH ₂) ₂						
MUF-17	1.41	2.55	28.3	25	[193]	
Cu-TDPAT		5.9	42.2	25	[197]	
Cu ₃ (TDPAH)		5.2 2.5	33.8	25 25	[213]	
MAF-23	2.05		47.4		[198]	
Zn ₂ Atz ₂ Ox	2.05	3.67	40	20	[196, 214]	
MAF-66		4.4	26.0	25	[216]	
ZJNU-43		4.6	20.0	25	[190]	
ZJNU-44		5.2		25	[190]	
ZJNU-45		4.8		25	[190]	
HP-e	~2.33	~7	35.2	25	[180]	
ZnF(daTZ)	~0.96\0.94 ^H	3.3	33	25	[217]	
ZnF(aTZ)	~0.56\0.50 ^H	3.1	32.3	25	[217]	
ZnF(TZ)	\sim 0.27 \setminus 0.35 H	2.7	23.6	25	[217]	
FJI-H14		6.5	26.6	25	[177]	
CALF-20		4.0 * *	39	20	[221]	
IISERP-MOF4		3.02	29	25	[223]	
HHU-5		~4.78	25.6	25	[178]	
*****	~1.35	~4.0	30	25	[179]	
HHU-2 MAF-stu-1	2.44	3.54	35	25	[105]	

 $^{^{*}}$ Measured at 0.39 mbar, * * Measured at 1.2 bar, and H for humid condition.

nucleobase adenine comprises an aromatic amine, a pyrimidine and diazole moieties, of which the Lewis basic amino and pyrimidine groups can be assumed to be favorable sites for adsorption. Also, the nano-sized channel structure played a significant role in terms of high $\rm CO_2/N_2$ selectivity (30–77) [182,183]. More recently, Chen et al. [184] prepared a moisture stable N-rich cobalt-based MOF, i.e., Co-PL-1 (Co (Imda)(4,4'-bpy) consisting of imidazole-4,5-dicarboxylic acid (Imda), and 4,4'-bipyridyl (4,4'-bpy) ligands. The reported MOF exhibited $\rm CO_2$ uptakes of 1.20 and 2.02 mmolg⁻¹ with high $\rm CO_2/N_2$ adsorption selectivity and good stability under humid conditions both at 150 mbar and 1 bar pressures (25 °C).

In addition to imidazole and triazole groups, an azo group (-N = N -) can have an effect to the selectivity of CO_2 adsorption by increasing the polarity of the pore structure [185–187]. For example, with a mixed-ligand approach, Nagaraja et al. [188] made homochiral

3D-MOFs $\{[M(L-mal)(azpv)_{0.5}]\cdot 2H_2O\}_n$ and $\{[M(L-mal)(bpee)_{0.5}]\cdot 2H_2O\}_n$ 2 H₂O₃, with Co or Ni metal ion nodes (M) and L-malate (L-mal) ligand using two different azo and ethylenic group (-CH=CH-) containing secondary linkers, namely 4,4'-bisazobipyridine (azpy) and 1,2-bis (4-pyridyl)ethylene (bpee). For azpy-MOF the CO2 capacity and the adsorption heat were slightly higher than that of the ethylenic analogue being \sim 4.9 vs. 1.5 mmol g⁻¹ and - 37 vs - 31 kJ mol⁻¹, respectively. It was suggested that the better sorption properties of azby-MOF originated from more polar azo groups inducing considerably stronger adsorbate—adsorbent interactions in comparison to the ethylenic group. Along with the azo group, the basic non-metalated porphyrin cavity can also enhance the selectivity of CO2 and heat of adsorption. For this, an interesting example is the Mn^{II}-porphyrin MOF ([{Mn₂(TCPP)·2 H₂O}· DMF]_n), where 5,10,15,20-tetrakis(4-benzoate)porphyrin (TCPP) ligand was used. The relatively high CO₂ capacity 2.63 mmol g⁻¹ (0 °C, 1 bar) and heat of adsorption of $-32.1 \text{ kJ mol}^{-1}$ values observed for the adsorbent were assumed to occur due to the basic -NH groups of the pyrrole macrocyclic cavity [189].

It is also good to point out that in the formulation and construction of MOFs for CO₂ adsorption, one should not only emphasize the abundance of NH₂ or other N-containing groups but also more critically focus on the impact of the accessibility of active binding sites by approaching CO₂. Song et al. [190] studied array of isoreticular copper-based MOFs, namely ZJNU-43, ZJNU-44, and ZJNU-45 by utilizing three rigid organic ligands, 5,50-(quinoline-5,8-diyl)-diisophthalic, 5,50-(isoquinoline-5, 8-diyl)-diisophthalic, and 5,50-(quinoxaline-5,8-diyl)-diisophthalic acids (Fig. 8). Although all three network structures have almost the same active surface area, they showed pronounced differences in their CO₂ uptakes due to the different number, and dissimilar orientation of N-containing groups inside the framework, which consequently defined accessibility of CO2 toward the available binding sites. In ZJNU-43 the nitrogen atom of the heterocyclic aromatic ring is in the α -position, demonstrating 4.6 mmol g⁻¹ CO₂ uptake. Whereas when located in β-position in ZJNU-44 the CO₂ capture capacity increased significantly to 5.2 mmol g⁻¹ (1 bar, 25 °C). On the ZJNU-45 the number of nitrogen atoms were doubled resulting somewhat lower 4.8 mmol g⁻¹ capacity under the same adsorption conditions. The difference in the determined capacities were also examined by quantum chemical calculations that suggested CO₂ trapping to occur in the narrow windows in case of the structures ZJNU-43 and ZJNU-45. Whereas in ZJNU-44 framework, N-containing sites pointed into the pore centers and by this allowed CO₂ bridging with the organic linkers. The binding energy was found to be highest for ZJNU-44 ($-27.6 \text{ kJ mol}^{-1}$), whereas -18.1- 18.7 kJ mol⁻¹ were calculated for ZJNU-43 and ZJNU-45, respectively. Based on the results, the orientation of the nitrogen-containing sites in the pore structure seem to play an important role in the efficiency of CO2 adsorption. Similar conclusions have been drawn recently by studies in which the subtle changes in the chemical construction and orientations of N-containing sites in MOFs have significantly impacted to the adsorption capacity [191,192].

Recently Qazvini et al. [193] investigated CO_2 capturing properties of MUF-17 ($[Co_5(\mu_3\text{-}OH)_2(aip)_4(H_2O)_2]$, prepared using 5-amino-isophthalic acid (aip) and cobalt ion nodes. In this framework, the primary CO_2 binding sites were positioned at the median of the narrowest channel neck. As can be seen in Fig. 9b, the electrostatic and hydrogen-bonding interactions with short contact distances prevailed between the framework and the CO_2 molecule. It was found that up to 85 % relative humidity, the adsorption efficiency of the MUF-17 remained unchanged. The compatibility of MUF-17 with a polymeric binder, poly-vinylidene fluoride, (PVDF) was also examined. The resulted MUF-17/PVDF composite in form of pellets exhibited similar behavior in gas adsorption experiments as the pristine MOF (Fig. 9c).

With a somewhat different approach, the chelate effect of bridging ligands was investigated by Wang et al. [105]. They reported Zn (imPim), also known as MAF-stu-1 which belongs to the group of metal azolate frameworks (MAF). It consisted of imidazole derivative ligand,

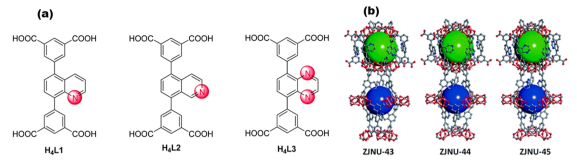


Fig. 8. (a) The organic linker ligands with heterocyclic N-containing core units, and (b) polyhedral nanocages of ZJNU-43, ZJNU-44, and ZJNU-45 MOFs. Adapted from Ref. [190].

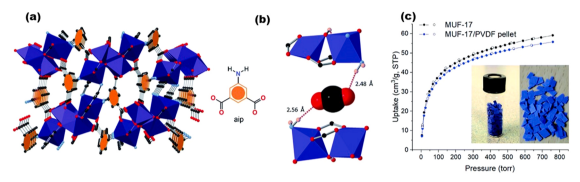


Fig. 9. The structure of (a) MUF-17, (b) Hydrogen bonding interactions with adsorbed CO₂ molecules, and (c) CO₂ adsorption and desorption isotherm of MUF-17/PVDF pellets at 20 °C.

Adapted from Ref. [193].

2-(1 H-imidzol-2-yl) — 3 H-imidazo[4,5-c] pyridine (H₂imPim), which offers binding pockets for CO₂ molecules in a "fit like a glove" manner (Fig. 10). The reported MOF exceeded the earlier benchmarks with a record-breaking thermal (up to 680 °C) and chemical stabilities coupled with a simple synthesis procedure in bulk quantities. Owing to its shape-complemented binding pockets, MAF-stu-1 also demonstrated an excellent capacity (3.54 mmol g $^{-1}$), high CO₂/N₂ (15:85, v/v) selectivity (106), and a low level of energy cost on regeneration (35 kJ mol $^{-1}$). Although MAF-stu-1 is a physisorbent, the obtained metrics are promoting its potential use in CO₂ recovery also in DAC conditions.

One of the available strategies for trapping CO₂ molecules is based on its physical (size and shape) and chemical (polarity) properties. As suggested by Li et al. for instance, by tuning the bridging ligands, molecular

cavities with accurate size and desirable properties can be designed, like a single-molecule-trap (SMT). In their study, the distance of 6.4–8.0 Å was suggested to be optimal for holding CO₂ molecules between two OMSs through electrostatic interaction [194]. This idea was further utilized by Song et al.[195] as they made NJU-Bai50 [Sc₃O(L1)₂(H₂O)₃(NO₃)] $_{\infty}$ and NJU-Bai51[Sc₃O(L2)₂(H₂O)₃(NO₃)] $_{\infty}$ MOFs using 1,10-(4-carboxypyridine-2,6-diyl)bis(1 H-pyrazole-4-carboxylic acid (H₃L1) and 1,10-(4-carboxypyridine-2,6-diyl)bis(3-bromo-1 H-pyrazole-4-carboxylic acid (H₃L2) ligands. To fit CO₂ adequately to pores, the contact distance between two opposite OMSs were tuned from 6.45 Å (NJU-Bai50) to 6.75 Å (NJU-Bai51) by regulating the torsion degree of the ligand. The CO₂ trap of NJU-Bai51 resulted in enhanced host–guest interactions, locking on to a single CO₂ molecule through its two O atoms without chemical bond formation. The increase in CO₂ uptake capacity was observed to be

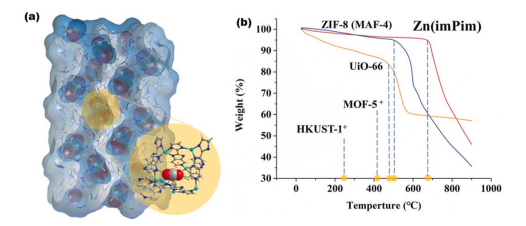


Fig. 10. (a) Partial view of periodic CO₂ binding pockets on Zn(imPim), and (b) Thermogravimetric curves of few selected MOFs compared with Zn(imPim) Adapted from Ref. [105].

0.13 mmol g⁻¹ (72 times higher) and 0.90 mmol g⁻¹ (nearly double) in comparison to NJU-Bai50 (0.0018 and 0.49 mmolg⁻¹) both at 0.4 and 150 mbar (25 °C), respectively. In terms of CO_2/N_2 selectivity (15/85 ratio at 1 bar, 25 °C) NJU-Bai51 showed a significantly better selectivity compared to NJU-Bai50 (545.7 vs. 30.5), most likely due to its custom designed size-matching cavities for CO_2 . In addition to the examples discussed above, other N-containing linkers such as pyrimidines, triazines, and azole derivatives have been utilized in functionalized MOFs, and have also demonstrated intriguing new results [191,196–217].

2.4.1. Consideration of water effects

The affinity, selectivity, and overall CO2 uptake properties of Ncontaining ligands in the frameworks are just as prone to the effects of water as the OMSs are. Fracaroli et al. [218] examined a series of IR-MOF-74-III by incorporating differently functionalized ligands containing -CH₃, -NH₂, -CH₂NMeBoc, and -CH₂NHBoc (Boc = tertbutyloxycarbonyl) functional groups (Fig. 11). On IRMOF-74-III-CH₂NHBoc and -CH2NMeBoc the protective Boc groups were removed by protonation to embody -CH₂NH₂, and -CH₂NHMe in the framework. The conversion was confirmed by ¹H and ¹³C NMR, ESI-MS, and SC-XRD data. Of the tested adsorbents IRMOF-74-III-CH2NH2 revealed the highest CO₂ uptake being 3.2 mmol g⁻¹ (1 bar CO₂), and a combination of carbamic acid (RNHCOOH) and ammonium carbamate (RNHCOO-+H3NR) species were ascertained by solid-state cross-polarization magic angle spinning (CP-MAS) ¹³C NMR spectra to be the main chemisorption products. In addition, the water did not seem to have any effect on the adsorption properties of the adsorbent, because similar retention times were observed with NMR for samples representing both dry and humid conditions. The adsorbent's high selectivity and capacity sustained even at 65 % relative humidity. Thereby it could be assumed that CO2 is likely to bound to the ligand-substituted alkylamine groups rather than to OMSs, as in case of humid condition OMSs should virtually be inaccessible due the water molecules have bound to them.

Furthermore, Flaig and coworkers [219] presented a doubling of the linker-substituted alkylamines in the framework to yield IRMO-F-74-III-($\rm CH_2NH_2$)₂. At lower $\rm CO_2$ pressures, the adsorbent demonstrated about 2.3 times higher $\rm CO_2$ adsorption capacity than IR-MOF-74-III- $\rm CH_2NH_2$ indicating that two alkyl-amines per organic linker further promoted chemisorption. Notably, based on the solid-state NMR ($^{13}\rm C$ and $^{15}\rm N$) data, carbamic acid was observed under dry conditions, whereas under water vapor ammonium carbamate was observed (Fig. 12). The reported study was one of the first ones to demonstrate the actual chemical composition of the adsorption products. It also highlighted the significance of the participation of water in dictating the

adsorption mechanism within the pores of the framework. The different sorption mechanisms of chemisorption are described in more detail in Section 2.5.2.

Along with the MOF-74 family, zeolitic imidazolate frameworks (ZIFs) have also proven to be promising adsorbent candidates for carbon capture due to their intrinsic hydrophobic behavior. Nguyen et al. [220] reported ZIF adsorbents resembling the structural topology of chabazite (CHA), namely ZIF-300 (Zn(2-mIm) $_{0.86}$ -(bbIm) $_{1.14}$), ZIF-301 (Zn $(2-mIm)_{0.94}(cbIm)_{1.06}$, and ZIF-302 $(Zn(2-mIm)_{0.67}(mbIm)_{1.33})$ which were made using 2-methyl-imidazo (2-mIm), 5(6)-bromobenzimidazole (bbIm), 5(6)-chlorobenzimidazole (cbIm), and 5(6)-methylbenzimidazole (mbIm), respectively. For all three adsorbents, remarkably low affinity for water was observed in water adsorption tests with uptake capacities varying from 0.33, 0.32–0.25 mmol g^{-1} , respectively at P/P₀ \approx 0.8 (25 °C). The ZIF-300 proved to be the most effective adsorbent for the dynamic separation of CO₂ from N₂ as the CO₂ adsorption isotherms showed uptakes of 1.3, 1.2, and 0.1 mmol g⁻¹ for ZIF-300, ZIF-301, and ZIF-302, respectively (at 1 bar, 25 °C). The experiments were made under dry and humid conditions, and no loss of activity was observed within three subsequent cycles. Furthermore, Lin et al. reported CALF-20 [Zn₂(1,2,4-triazolate)₂(oxalate)] MOF which demonstrate low water affinity and high uptake of 4.07 mmol g-1 (1.2 bar, 20 °C) but with modest heat of adsorption of -39 kJ mol^{-1} indicating physisorption of CO2. [221] With atomistic grand canonical Monte Carlo (GCMC) simulations, the CO₂ binding sites were found to be in the centers of pores that impede forming of a hydrogen-bonding network between water molecules. The calculated CO_2 binding energy of $-34.5 \text{ kJ mol}^{-1}$ indicated electrostatics and attractive dispersion interactions (85 %) being predominant in interactions between the framework and CO₂. For water adsorption studies, simulations demonstrated that in the absence of CO₂, water uptake (20% RH) was 6 mmol g⁻¹, but with presence of CO2 it was negligible. However, at 40 % RH, water uptake was observed to be 6 mmol g⁻¹ in the presence of CO₂. Thus, below 40 % RH, CALF-20 demonstrates not just preferential CO2 physisorption but also shows suppression of water adsorption by CO2. In addition, adsorbent showed low enthalpic regeneration penalty in flue gas conditions and displayed durability towards steam > 450,000 cycles. [221].

Although with moderately water-resistant MOFs exemplified above and few others found in the literature [222,223], most MOFs will go through structural degradation under humid conditions. The metal centers of the framework, in the long run, end up gradually interacting with a water molecule instead of CO_2 owing to its quadrupole moment [131,224,225]. The long-term chemical stability of the adsorbents against water is an important property, particularly for those designed to function in DAC conditions, wherein water will always be present in

$$(1) \qquad (2) \qquad PdCl_{2}(PPh_{3})_{2}, KOAc \\ dioxane \qquad (2) \qquad PdCl_{2}(dppf) \\ + \qquad (2) \qquad PdCl_{2}(dppf) \\ + \qquad (3) \qquad (4a-4d) \qquad (5a-5d) \qquad R = H, IRMOF-74-III-CH_{2}NHBoc \\ -CH_{2}NMeBoc (5d) \qquad (3) \qquad (4a-4d) \qquad (5a-5d) \qquad R = H, IRMOF-74-III-CH_{2}NHBoc \\ -CH_{3}, IRMOF-74-III-CH_{2}NHBoc$$

Fig. 11. Synthetic route for the functionalized organic linkers incorporated in IRMOF-74-III Adapted from Ref. [218].

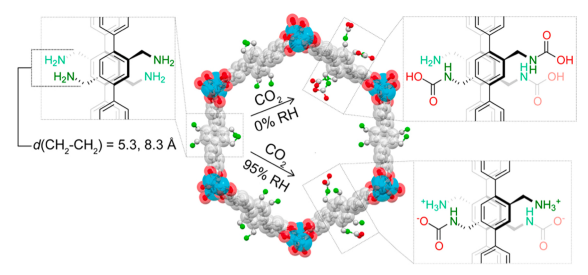


Fig. 12. Modelling of the IR-MOF-74-III-(CH₂NH₂)₂ structure and schematic of the chemisorption species upon reaction between linker-based alkylamine and CO₂ under dry and humid conditions.

Adapted from Ref. [219].

magnitudes several times higher than CO_2 . Selected CO_2 capture capacities for all the adsorbents presented above are shown in Table 2.

2.5. Post-synthetic amination and adsorption mechanisms on aminated open metal sites

Water molecules are in constant competition with CO₂, and the unsaturated open metal sites are particularly attractive to water. To overcome the adverse effects of water, particularly on physisorbent MOFs that are typically based on the adsorption of CO₂ to OMSs, aminecontaining species can be anchored post-synthetically in OMSs. Methods commonly found in the literature are called as amine grafting, tethering, or appending of amines. Two of the well-known MOF-families, namely MOF-74, and MIL-100/101 based MOFs serve again as good examples of recent advances in MOFs that include OMSs (chapters 2.5.1 and 2.5.3). Many of the CO₂ adsorbents of both families are among the best performing adsorbents in dry conditions but many of them, like most other MOF adsorbents, face difficulties in humid conditions due to the gradual deterioration of the chemical structures by hydrolysis [226]. Adsorption mechanisms, which inevitably depend on whether adsorption takes place in dry or humid conditions, are also reviewed in Section 2.5.2.

2.5.1. MOF-74 family

According to Choi et al. [227] OMSs in MOF-74 are capable to form stable bonds with diamine by receiving electrons from the N atom of the first amino group of the diamine, and by this, the second amino group is still available for interaction with CO2. Among the first post-synthetic modification studies, ethylenediamine (en) was tethered into Mg₂(dobdc) to produce en-appended MOF-74. The thermogravimetric analysis (TGA), nitrogen sorption, and DFT calculations of the product revealed that approximately one ethylenediamine molecule was associated per one OMS in each unit cell. After functionalization, the surface area of the modified material was clearly lower than that of the pristine MOF, as it decreased from 1094 to 469 m²/g. However, some 12% increase in CO₂ uptake (from 1.35 to 1.51 mmol g⁻¹) was observed between aminated and pristine MOF-74 under 0.39 mbar CO2. The regeneration tests with four sorption cycles were made for both aminated and pristine MOF-74 and revealed some 21% decrease in the capacity at the end of the cycles for pristine adsorbent, whereas no change in capacity was observed for the aminated one.

Bernini et al. [228] screened several reported and some hypothetical en- Mg_2 (dobdc) MOF variants by Monte Carlo simulations to determine the optimized amine loadings for capturing CO_2 both from flue gas and

ambient air. Ethylenediamine compositions of 3.14, 5.99 and 6.74 wt.-% were incorporated to Mg2(dobdc), and as a result diamine content of about 50% of the available OMSs turned out to be the optimal fill rate for affording the best adsorption capacity. Also, the correlation between thermal stability and amine loadings was investigated by TGA. The superior stability was demonstrated by the lowest amine loading (3.14 wt.-%) followed then by 5.99 and 6.74 wt.-% amine loads. This indicated a clear decrease in thermal stability with an increase in amine loads, as temperature ranges for primary thermal degradation step varied from 278 to 430 via 252-440-228-430 °C, correspondingly. Hydrazine (HZ), the smallest and the lowest boiling diamine (114 °C), was appended in various loads to Mg2(dobdc) by Liao et al. [229] resulting in the highest capture capacity of 3.89 mmol g⁻¹ in DAC (25 °C) with HZ-MOF-74 containing 6.01 mmol g⁻¹ amine load. Interestingly, the thermal stability of the afforded adsorbent was high even though low boiling diamine was incorporated into the framework. The aminated adsorbent was subjected to 5 sorption cycles with desorption at 130 °C (N2), and virtually no capacity loss was observed at the end of the cycles.

As can already be seen from the above examples, amine basicity also plays an important role both in the adsorption and selectivity of CO2. Yeon et al. [123] investigated the effects of amine basicity using an amine-functionalized isoreticular Mg-MOF-74 analogue, namely Mg₂(dondc) made of 1,5-dihydroxy-2,6-naphthalenedicarboxylic acid. Pristine adsorbent showed pore size of about 15 Å, which was then modified using primary, secondary and tertiary amines: ethylenediamine (en), N,N'-di-methylethylenediamine (mmen), and piperazine (ppz), respectively. Due to the relatively small channel sizes of the unmodified adsorbent, the available open metal sites were found to be 75 (en), 55 (ppz), 60 % (mmen) occupied by the amines. Nevertheless, adsorbents exhibited good CO2 capacities ranging from 1.99 (en) via 2.04 (ppz) to 3.20 (mmen) mmol g⁻¹ under post-combustion flue gas conditions. Surprisingly, the adsorption mechanism was attributed to be both physi- and chemisorption, as indicated by the IR spectroscopic analyses. The major weakness of the adsorbents proved to be their poor thermal stability, as leakage of the amines out of the pores was thermogravimetrically evidenced during regeneration cycles. After five cycles, the ethyleneamine containing adsorbent already lost 10% of its initial CO₂ adsorption capacity at 150 °C, and mmen- and ppz-appended ones lost 15% and 14% already at 130 and 120 °C, respectively. For enand ppz-appended adsorbents the uptake losses were associated to 17 and 12 wt.-% amine leakage, whereas mmen-Mg2(dondc) showed only weight loss of 2 %. To examine long term regeneration characteristics,

ppz-Mg₂(dondc) was regenerated 50 times resulting in dramatic 43 % decrease in CO₂ uptake capacity under tested desorption conditions (with Ar at 130 °C). McDonald et al.[101] made similar amination tests but with Mg₂(dobpdc) and the OMSs were post-synthetically aminated by mmen. In DAC the aminated mmen-Mg₂(dobpdc) showed a significantly higher 2.0 mmol g⁻¹ capacity (0.39 mbar, 25 °C) being about 15 times higher than the pristine Mg₂-(dobpdc) showing only modest \sim 0.13 mmol g⁻¹ capacity under the same conditions. The high adsorption heat determined for the adsorbent indicates that CO₂ is chemisorbed to the diamine sites of the adsorbent. Based on the above examples and the literature, it has been suggested that the mmen-Mg₂(dobpdc) adsorbent would be more efficient than, for example, conventional amine-grafted silica-based adsorbents or aqueous amine scrubbers. [230].

2.5.2. Adsorption mechanisms

Regarding the aminated MOF adsorbents already discussed above, the variables related to their adsorption and regeneration process (desorption) can also be considered, of which the actual adsorption mechanism is one of the key factors, and it is dependent on the properties of the available adsorption sites e.g., amines appended on open metal sites, as well as whether sorption conditions are dry or humid and at what temperature. For instance, the mmen-Mg2(dobpdc) has been designated as a 'phase-change' type of CO2 adsorbent, as it shows stepshaped isotherms with a relatively large working capacity at a small temperature increase. However, classical adsorbents (mostly aminefunctionalized inorganic solids) show Langmuir-type isotherms demanding higher desorption temperatures to attain large working capacities. The sudden step-shaped rise in the CO₂ isotherm at a precise pressure signifies cooperative insertion of CO2 amongst the metal centers and the amine clusters. Therefore, it is expected that carbamate species will be created that leads to the generation of ion pairs with neighboring ammonium ions [231]. The foundations for the generation of carbamate species lay on the sorption environment favoring chemisorption in the presence of amino group, wherein water acts as a catalyst and protic solvent for the deprotonation via which the carbamate is generated. When chemisorption governs the CO2 adsorption, the kinetics of CO2 uptake is very fast and highly selective above any other gases [55]. For instance, in the case of tetraethylenepentamine (TEPA) aminated Mg2(dobdc), humid condition promotes CO2 adsorption considerably, which can be seen as an increase in capacity from 6.06 to $8.31 \ \text{mmol} \ \text{g}^{-1}$ on TEPA-Mg₂(dobdc), compared to non-aminated one, due to bicarbonate formation favored by the water-assisted stoichiometry.

Based on theoretical and experimental considerations, it is well-known that amino groups of polyamines, both primary and secondary, prefer the formation of carbamate species with CO_2 , which converts into bicarbonate in the presence of water. In terms of stoichiometry CO_2 capture in dry conditions needs two amino groups per CO_2 to be adsorbed (reactions 1 and 2), whereas in humid conditions only one amino group is needed for adsorption, as can be seen from the reaction formulas 3–5:

$$CO_2 + 2RNH_2 \leftrightarrow NH_4^+ + R_2NCOO^-$$
 (1)

$$CO_2 + 2R_2NH \leftrightarrow R_2NH_2^+ + R_2NCOO^-$$
 (2)

The addition of moisture accelerates the reaction of carbamic acid with water and CO_2 , eventually bicarbonate; is formed (3). There is also a tendency of the amine group to react directly both with CO_2 and H_2O to form a bicarbonate unit (4–5). [232].

$$CO_2 + R_2NCOO^- + 2H_2O \leftrightarrow R_2NH_2^+ + 2HCO_3^-$$
 (3)

$$CO_2 + 2RNH_2 + H_2O \leftrightarrow RNH_3^+ + HCO_3^-$$
 (4)

$$CO_2 + 2R_2NH + H_2O \leftrightarrow R_2NH_2^+ + HCO_3^-$$
 (5)

The study made by Planas et al. [233] verified that an increase in $\rm CO_2$ capture capacity is not merely based on the interaction between a positive carbon and a negative nitrogen atom, in $\rm CO_2$ and amine respectively. They examined $\rm CO_2$ -amine chemisorption mechanism in mmen-Mg₂(dobpdc) using computational chemistry. With density functional theory (DFT) studies, they proposed that N–C bond formation steps are stabilized by cleavage of N–H bonds and the concurrent generation of hydrogen-bonded adducts (Fig. 13). It was also suggested that the hydrogen bonds have a crucial role in preventing unwanted reactions from occurring that may lead to unwanted intermediates and side-product formation.

As can be seen in Fig. 13, the preliminary lowermost-energy state (A_I) comprises two amines allied on a neighboring Mg²⁺ cation. The single H atom of an amine can form a hydrogen bond with N atom of an adjacent amine group. The O atom of confronting CO2 interacts with H atom of the amine and forms a weak early-stage hydrogen bond. After the rotation of the amine group and CO2 molecule, the H-N bond cleaves, consequently followed by the formation of N-C bond between an amine and CO₂, respectively. In this mode, the amine N atom tethers to the C atom and donates its H to the adjacent amine N atom, thereby forming a carbamic unit. Concurrently, the other amine N atom accepts the H atom and forms RNH₂⁺ cation. In accordance with Coulombic force, the positive RNH₃⁺ seeks $N - CO_2^+$ to form new hydrogen bonds (N-H and O-H) with minimal energy states (B_I). Interestingly, it was suggested that with the addition of a second CO2 molecule, hydrogen bond will be formed between the carbamic acid-amine adduct (B_{II} to B_{III}) and CO₂, thus producing a second zwitterionic unit that is stabilized by two-fold hydrogen bonds. Consequent exchange of protons followed by a reorganization of the product leads to the formation of bis (carbamic $\,$ acid), which is stabilized by dual hydrogen bonds in the ultimate structure. Based on the suggested mechanism, the adsorption energy of - 138.25 kJ per 2 moles of CO₂ corresponds to an average adsorption energy of - 69.13 kJ mol⁻¹ which is a reasonably good agreement with the experimental value of – 71 kJ mol⁻¹ for chemisorption of CO₂.[234] The presented mechanism emphasized the importance of the physical length of the amine as the structural constraints of the adsorbent dictate the amine-amine interaction distances.

Encouraged by the presented mechanism Lee et al. further studied CO₂ adsorption in DAC using a system wherein primary amine ethylenediamine was appended into a pore size expanded Mg2(dobpdc) [235]. A significantly higher 2.83 mmol g⁻¹ CO₂ capacity (0.39 mbar, 25 °C) was measured for en-Mg2(dobpdc), and even higher 3.62 and $4.57\ mmol\ g^{-1}$ were obtained at 150 mbar and 1 bar CO_2 pressures in dry conditions, respectively. In regeneration cycles, the capacity remained practically unchanged both in DAC, as well as on CO₂ richer conditions corresponding to the flue gas. En-Mg₂(dobpdc) was also exposed to humid conditions, wherein the capacity also remained unchanged after several cycles. However, the adsorption mechanism proved to be different than the previously suggested carbamate formation, as based on the FTIR and DFT calculations carbamic acid resulted in the adsorption event. In FTIR spectra, the characteristic O-H peak at 3412 cm⁻¹ disappeared with an increase in temperature followed by emerging of N – H stretching at 3337 and 3281 cm⁻¹, therefore indicating adsorption of CO2. The calculated adsorption heat was found to be in the range of $-49-51 \text{ kJ mol}^{-1}$ being consistent with the enthalpy of carbamic acid formation determined by DFT calculations.

In another study, Lee et al. [236] prepared dmen-Mg₂(dobpdc) using N,N-dimethyl-ethylenediamine (dmen) as amine loading. Dmen-Mg₂(dobpdc) showed 4.34 mmol g⁻¹ CO₂ capacity (1 bar, 40 °C) with adsorption heat in the range of -71–75 kJ mol⁻¹ resembling overall features of mmen-Mg₂(dobpdc). The modified adsorbent was subjected to 24 sorption cycles (desorption at 75 °C under Ar) where it showed virtually no capacity loss at the end of the cycles, indicating good thermal stability of dmen-Mg₂(dobpdc) under the tested conditions. Time-dependent in situ FTIR spectroscopic analyses were also made

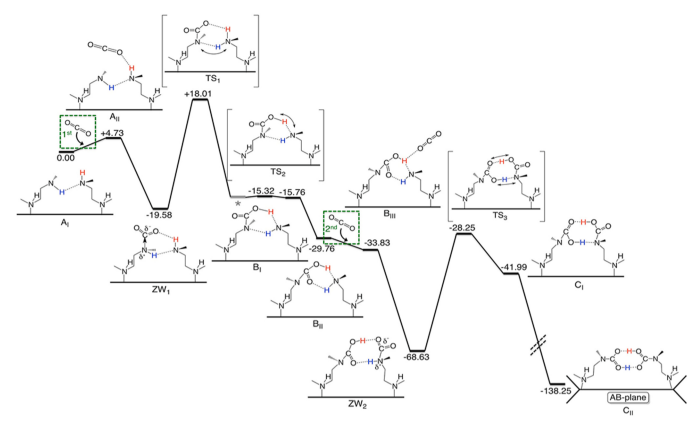


Fig. 13. Possible pathway of stepwise CO₂ adsorption on mmen-Mg₂(dobdc) Adapted from Ref. [233].

under humid conditions using 100 % relative humidity (RH) and 150 mbar CO_2 pressure to simulate flue gas conditions. Notably, under humid CO_2 conditions, the N-H stretching peak at 3397 cm $^{-1}$ appeared in the FTIR spectrum and it was still visible after purging with N_2 . However, the characteristic peaks of CO_2 disappeared with purging. These observations are well aligned with the dmen-Mg₂(dobpdc) examined under similar CO_2 adsorption conditions.

In the studies of McDonald et al. [231] and Lee et al. [236] a somewhat modified adsorption mechanism describing cooperative intake of CO_2 on mmen- and dmen-loaded $Mg_2(dobpdc)$ was suggested. As in earlier models, one end group of a diamine was assumed to be bound with an open metal site, whereas the other terminal amine remained accessible for CO_2 . With exposure to CO_2 , the free diamine group adheres to CO_2 and the resulting carbamic species are formed with similar fashion as suggested by Planas *et al.*, thus via proton transformation from one amine to the adjacent metal coordinated amine N atom (Fig. 14). Based on the DFT calculations, carbamic acid O atom favors binding with Mg over the N atom of amine, originally bound with Mg center. Thus, as illustrated in Fig. 14, highly ordered chains are

generated upon synchronized breakage of the Mg-N bond followed by the consequent forming of Mg-O bond. To validate the suggested model, thorough structural studies were carried out using isostructural series of mmen-M₂(dobpdc) MOFs (M = Co, Ni, Mg, Mn, Fe, and Zn) for which CO2 adsorption isotherms were determined [231]. All the obtained isotherm curves consistently displayed a sharp isothermal step that moved to higher pressure with an increase in temperature, thereby implying a cooperative process wherein CO₂ displace M–N bonds upon its chemisorption. The subsequently formed carbamate and adjoining ammonium ion pairs align in the channels, forming ammonium carbamate chains that are periodically bound to the Mg center via oxygen. This arrangement was suggested to cause sudden increases in the adsorption curves. Adsorption mechanisms were further examined by in situ FTIR and solid-state NMR (13C and 15N) to identify the actual chemical species formed during the sorption process. FTIR spectra displayed a broad peak positioned at 2200 cm⁻¹ and multiple peaks in the range of $1700 - 1630 \, \text{cm}^{-1}$ indicating the presence of ammonium cation and C-O stretching of the amide group and asymmetric deformation of the ammonium cation. These peaks, along with the N-H

Fig. 14. A cooperative insertion mechanism of CO₂ on mmen-Mg₂(dobpdc).

stretching peak at 3397 cm⁻¹ disappeared with the progress of adsorption, implying forming of the carbamate species. NMR experiments suggested interconversion of coordinated and uncoordinated ends of the mmen molecule in mmen-Mg₂(dobpdc), which could be detected as ¹⁵N chemical shifts at 31 and 72 ppm along with CO2 adsorption and can be expected to result from ammonium cation and carbamate species, respectively. On the other hand, with a lack of CO₂, only a single ¹⁵N resonance peak was found for mmen-Mg2(dobpdc). Next, the X-ray absorption near edge structure (NEXAFS) spectroscopy was used for the elucidation of the CO2 adsorption mechanism. A new peak was found at 402.3 eV corresponding to carbamate nitrogen, thus explaining the insertion of carbamate into the metal-nitrogen bond. The reported work is of profound significance as the combination of spectroscopic and diffractometric measurements, including computational studies, provided deep insight into the chemisorption mechanism, and associated chemisorbed species (carbamate or carbamic acid) formed with CO₂ adsorption. Further confirmation of the proposed mechanism was obtained by Drisdell et al. as their detailed study, including computational and spectroscopic measurements on the CO2 adsorption on mmen-Mg₂(dobpdc) also evidenced the formation of either ammonium carbamate or carbamic acid between CO₂, and that the chemisorption with the presence of an amine is clearly favored in contrast to the physisorption adsorption [237].

To investigate adsorption rates and side groups effects on the CO2 desorption mechanism, Jo et al. [238] made a comprehensive study on diamine aminated MOFs compared with number of different types of MOF adsorbents, such as MOF-5, Zn(pyrz)(SiF₆), as well as inorganic adsorbents like zeolites and activated carbon. Branched primary diamines of 1-methylethylenediamine (men) and 1,1-dimethylethylenediamine (den) and linear ethylenediamine (en) were loaded to Mg₂(dobpdc). The modified materials en-, men- and den-Mg₂(dobpdc) exhibited 3.53, 3.6 and 2.15 mmol g⁻¹ adsorption capacities under simulated flue gas conditions, respectively. Based on the correlation diagram determined for the working and capture capacities at 30 mbar CO2, both capacities appeared to be dependent on the structural properties of the incorporated diamines. Several samples were tested and categorized into four groups (Fig. 15). Group I included adsorbents with or without OMSs that showed both working and adsorption capacities to be lower as an example for pristine MOF-74 \sim 1.02 mmol g⁻¹ has been presented (4.5 wt.-%, as illustrated in Fig. 15). Group II comprises primary diamines aminated MOFs demonstrating low working capacity but

high adsorption capacity due to the strong chemisorption between the primary amine and CO2. Group III constitutes hetero-diamine-aminated Mg2(dobpdc) showing high working capacity but low adsorption capacity due to the weaker interactions between the CO2 and the mixed amine groups. Group IV includes den-Mg₂(dobpdc) exhibiting high \sim 2, 49 mmol g⁻¹ adsorption capacity due to the presence of primary amine groups. The dimethyl groups on the ethylene linker in den-Mg₂(dobpdc) were expected to be responsible for the weakening of Mg-carbamate interaction, thus promoting facile desorption and subsequently higher working capacity of ~ 2.77 mmol g⁻¹ was observed. In the reported study, the adsorbent was also subjected to SO_x exposure (500 ppm SO_2/N_2) for a single sorption cycle with adsorption at 40 °C (150 mbar) and desorption at 130 °C under Ar. As the result of test, CO₂ capacity dropped about 6 % under dry, and 13 % in humid conditions. Based on the observed sorption characteristics, den-Mg₂(dobpdc) was suggested to be a suitable adsorbent for a fluidized bed system for continuous CO2 capture.

In the study of Lee et al. [239] diamine aminated Mg₂(dobpdc) adsorbents were made using ethylenediamine (en), primary-secondary diamines, such as N-ethylethylenediamine (een) and N-isopropylethylenediamine (ipen), and various combinations of primary-tertiary and secondary-secondary diamines. MOF adsorbents were then subjected to CO₂ sorption tests under real flue gas conditions. The study demonstrated how the change in diamine chain length or in the number of alkyl substituents define the desorption properties, such as desorption temperature and the temperature difference between the adsorption and desorption processes (ΔT). The amine bulkiness was shown to play an essential role in elevating robustness of the framework when exposed to realistic flue gas conditions which contain some O_2 , SO₂, and water vapor. Interestingly, DFT calculations revealed two-step adsorption behavior for the ipen- $Mg_2(dobpdc)$, which was concluded to be caused by the steric bulkiness of the branched diamines. The presented study clearly emphasized the importance of the proper selection of diamine, as it can have a considerable effect on the durability of the framework and CO₂ uptake efficiency. In this context, Kang et al. [240] presented een-Mn₂(dobpdc) which exhibited at the time the smallest temperature difference between adsorption and desorption processes ($\Delta T = 30$ °C), low regeneration temperature (≤ 100 °C), and high cyclic working capacity of 3.18 mmol g⁻¹ over the tested 600 sorption cycles. The overall efficiency of this adsorbent was further improved by coating the MOF particles with polydimethylsiloxane (PDMS) to enhance water

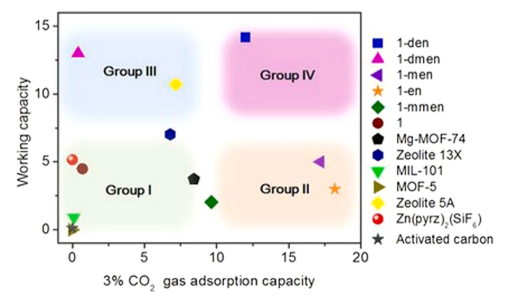


Fig. 15. Correlation diagram for working and adsorption capacities (wt.-%) obtained at 30 mbar Adapted from Ref. [238].

resistance of the adsorbent, and thereby also affecting the adsorption properties of the adsorbent (for coating methods see chapter 2.7).

It has been well-established by now that the cooperative adsorption threshold is governed by two key variables: CO2-inserted sites and the relative stability of an amine bound to the MOF. As a continuation to earlier studies Siegelman et al. [241] investigated the effect of an amine structure on the interaction of the metal-amine bond (M-A), and ammonium carbamate ion-pairing (Fig. 16). Nine diamine aminated Mg₂(dobpdc) adsorbents were prepared, including N-dimethylethyle-(m-2-m), N,N'-diethylenediamine nediamine (e-2-e), N'-diisopropylethylenediamine (i-2-i), N-methylethylenediamine (m-2), N-ethylethylenediamine (e-2), N-isopropylethylenediamine (i-2), N, *N*-dimethylethylenediamine (mm-2),N,N-diethylethylenediamine (ee-2) and N,N-diisopropylethylenediamine (ii-2). Products were examined with combination of single crystal X-ray diffraction, spectroscopic and volumetric gas adsorption methods. In the study, it was found that even small changes in the molecular structure of the diamine can affect the threshold pressure of cooperative CO2 adsorption up to four times at a single temperature. For instance, bulky diamine bound to a metal center would cause steric hindrance, which causes instability of the amine-bound adsorption product (e.g., carbamate) and consequently lowers the step pressure. Also, when substituent groups are coupled on the terminal amine side, the step pressure will increase, because of the weakened ion-pairing interactions in ammonium carbamate chains that are formed by CO2 adsorption. It was also observed that hydrogen bonding and ion-pairing interactions would stabilize CO2-inserted phases, and subsequently lower the step pressure for the continuation of cooperative CO₂ adsorption. The critical principles outlined by the study can be assisted in forthcoming designs of diamine-appended MOFs to achieve increasingly optimized adsorption properties for CO₂ capture.

In the same year, Milner et al. [103] made 3-diaminopropane (pn), 2-methyl-1,3-diaminopropane (mpn) and 2,2-dimethyl-1,3-diaminopropane (dmpn) aminated Mg₂(dobpdc) adsorbents and examined their sorption properties including cyclic stability tests. Of the tested adsorbents in flue gas simulated conditions, dmpn-Mg₂(dobpdc) showed the best step-shaped adsorption at 40 °C (150 mbar) and almost complete desorption at 100 °C (1 bar) with high 2.42 mmol g⁻¹ working capacity. Also, high differential adsorption heat of $-74~\rm kJ~mol^{-1}$ with narrow $\Delta T = 60~\rm ^{\circ}C$ temperature swing window was determined for the dmpn-Mg₂(dobpdc). Based on the extensive sorption and regeneration tests made, the dmpn-Mg₂(dobpdc) demonstrated substantial long-term stability, as only about 3% decrease in amine load was observed without

significant change in work capacity after 1000 regeneration cycles. Due to the fast kinetics observed for the system, the sorption cycles could be run using 5 min adsorption and 1 min desorption steps. The authors concluded that, particularly the dmpn-Mg2(dobpdc) would be a very promising adsorbent for capturing CO₂ from flue gas. In 2019 Choe et al. [242] reported a series of tetraethylenepentamine (TEPA) appended Mg₂(dobpdc) adsorbents containing different amine loadings (with 0.2, 2, 20, and 200 equivalents, samples named T02, T2, T20, and T200, respectively). Surprisingly, on higher TEPA loadings (T20 and T200) a significant increase in the CO2 adsorption capacity from 1.47 to 2.36 mmol g⁻¹ was observed but with an increase in adsorption temperature accordingly from 40° to $80 \,^{\circ}$ C. At first, the obtained results seemed to be slightly inconsistent with the previously investigated aminated MOF adsorbents. Eventually, it was found that observations related to the temperature-dependent behavior of the long-chained TEPA molecules, which at lower temperatures obstructed diffusion of CO2 into the pores but at higher temperatures with increased flexibility of their alkyl chains this obstacle was overcome. Based on the sorption tests TEPA-Mg₂(dobpdc) adsorbs CO₂ via similar cooperative insertion mechanisms as have been observed, for instance on mmen-, and dmpn-Mg₂(dobpdc), and many other diaminated MOF-74 analogues, because the similar type of sharp increases in CO₂ adsorption isotherms

As a natural continuation of linear and branched diamines are cyclic diamines, which Rebecca et al. [243] used to functionalize Mg₂(dobpdc) with 2-(aminomethyl)piperidine (2-ampd) for capturing CO2 in simulated natural gas flue emission stream. The sorption isotherms showed two adsorption steps, and 3.46 mmol g⁻¹ CO₂ uptake was determined for the adsorbent in 40 mbar CO₂ pressure (40 °C). Under humid conditions uptake was further improved, which assumed to be resulted from the hydrogen bonding interactions between water and carbamate chains to form ammonium carbamate species. The sorption tests with 700 cycles under humid conditions at 140 °C for 12 h indicated good stability of the adsorbent as the adsorption capacity remained virtually unchanged. Currently the reported 2-ampd-Mg2(dobpdc) is still one of the best adsorbents among the MOFs developed for the natural gas flue emission technology. Recently, Kim et al. [244] demonstrated the CO₂ adsorption properties of tetraamine functionalized MOFs utilized in emission conditions corresponding to the combustion of natural gas, wherein the emission of CO₂ is only a quarter (~4%, 40 mbar) of the amount released from coal combustion. Again, the ordered, multi-metal coordinated tetramines showed high thermal stability and enabled regeneration of

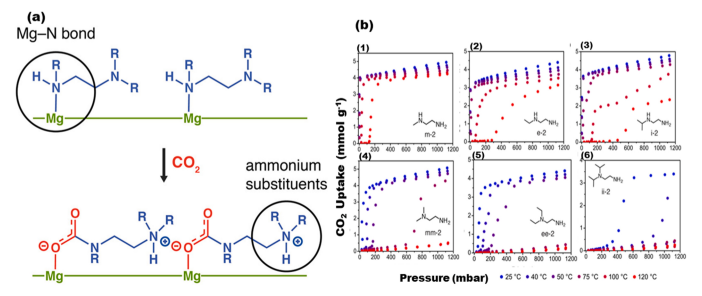


Fig. 16. Illustration of cooperative CO₂ insertion into Mg(II)-diamine sites to form ammonium carbamate chains. Adapted from Ref. [241].

the adsorbent with inexpensive low-temperature steam. A range of tetraamine appended-Mg2(dobpdc) were synthesized by Kim et al. with varying alkyl chain lengths between the adjacent nitrogen atoms in the chain as follows (Fig. 17): triethylenetetramine (2-2-2), N,N'-bis (2aminoethyl) - 1,3-propanediamine (2-3-2), 1,2-bis(3-aminopropylamino)ethane (3-2-3),N,N'-bis(3-aminopropyl) – 1,3-diaminopropane (3-3-3), N,N'-bis(3-aminopropyl)- 1,4-diaminobutane (3-4-3), diethylenetriamine (3-3), and N-(3-aminopropyl)-1,4-diaminobutane (3-4). So far among the various tetramine modified M₂(dobpdc) adsorbents, the 3-4-3-tetraamine appended Mg₂(dobpdc) holds the highest CO₂ adsorption enthalpy of -99 kJ mol⁻¹ with a two-step shaped adsorption curve, breakthrough capacity of 2 mmol g⁻¹ under humid conditions, while also meeting the U.S. Department of Energy (DOE) ambitious target for capturing 90% of CO2 from natural gas flue emissions. Based on spectroscopic analyses, the step-shaped adsorption curve was again caused by the cooperative insertion mechanism originating from the formation of chained ammonium carbamate species. The determined working capacity remains unchanged under humid conditions with over 1000 sorption cycles indicating strong selectivity for CO₂ over water vapor. With the above-described characteristics, the reported 3-4-3-tetraamine appended MOF could potentially be a promising candidate for direct air capture of CO₂ as well.

In 2020, Mao et al. [245] designed alkanolamine and alkoxyalkylamine functionalized Mg2(dobdpc) variants, for which extensive computational and structural studies were combined to evaluate five different possible mechanisms for cooperative CO2 adsorption. Tested alcoholamines included bulky secondary amine groups, such as ethyl (e-2-OH), n-propyl (nPr-2-OH), isopropyl (iPr-2-OH), n-butyl (nBu-2-OH), cyclohexyl (cy-2-OH), isopentyl (iPent-2-OH), and 3-aminopropanol (3-OH) as shown in Fig. 17d. Altogether, with solid-state NMR (13C and 15N) and gas-adsorption studies, they presented the most likely mechanism, wherein subsequently to CO2 adsorption, the hydrogen bonds of the methoxy groups across the ab lattice plane further stabilize the formed ammonium carbamate ion-pairs, and as a result continuous cooperative adsorption propagates along the crystallographic c-axis. Park et al. functionalized Mg₂(dobpdc) with N-methyl-1, 3-diaminopropane (Nmpn), 1,3-diaminopentane

3-diaminopropane (pn), 2,2-dimethyl-1,3-diaminopropane (dmpn) and N,N-dimethyl-1,3-diaminopropane (Ndmpn) and tested their adsorption properties in dilute CO2 concentration (1 mbar) with comparison to many other adsorbent materials (Fig. 18) [246]. The study focused mainly on lowering the regeneration temperature, as for practical real-world applications high desorption temperature is undesirable as it effects adversely to the chemical and structural stability of the adsorbent, and consumes more energy. Interestingly, regeneration of epn-Mg₂(dobpdc) at moderate 70 °C still demonstrated notable 2.88 mmolg⁻¹ adsorption in 1 mbar CO₂ pressure. However, when the similar sorption cycles were made under humid conditions, a significant decrease in the recovery capacity was observed. Authors, attempted to solve the problem by coating the adsorbent particles with a hydrophobic agent polydimethylsiloxane (PDMS) and indeed, with the help of the hydrophobic coating, CO2 uptake decreased only slightly from that of observed in dry conditions, thus being ~2.77 mmol g⁻¹ (for coating methods see chapter 2.7).

As discussed above, for designing new cooperative CO₂ adsorption MOFs for carbon capture it is essential to unravel the interrelationship between the CO₂ insertion mechanism and the chain length of diamine. To this end, Hong et al. [247] made further efforts to clarify CO₂ adsorption mechanisms by functionalizing Mg₂(dobpdc) with primary diamines with varying chain lengths of ethylene- (1-en), propylene-(1-pn), butylene- (1-bn), pentamethylene- (1-pen), hexamethylene-(1-hn), and octamethylene- (on) diamine (Fig. 19). Intriguingly CO₂ isotherms, with the support of FTIR spectroscopy and computational chemistry, evidenced that the CO₂ insertion mechanism is dominant up to the alkyl chain length of four carbons, while the mechanism was not found to occur with longer alkyl chains. With the van der Waals corrected density functional theory (DFT) it was further evidenced that the CO2 insertion process was proved to be dependent on the relative location of diamines across the hexagonal pore structure as based on the calculated models the longer chained 1-pen, 1-hn, and 1-on diamines seem to follow non-cooperative adsorption mechanism. This was particularly clear with the 1-hn and 1-on diamines, as they coordinate across the pore to two distinct Mg²⁺ sites, and thus cannot participate in chemisorption. Surprisingly, bn-Mg₂(dobpdc) displayed three distinct

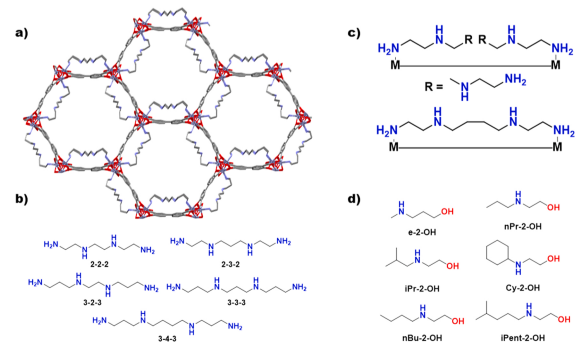


Fig. 17. (a) Illustration of pore structure of amine appended $Mg_2(dobpdc)$ (CSD entry: FUYGUE), (b) different tetraamines appended to $Mg_2(dobpdc)$, (c) structural modeling of 2–3–2 appended and 3–4–3-appended metal sites, and (d) selection of alkanolamines used to functionalize the open metal sites of $Mg_2(dobpdc)$ [110, 244,245].

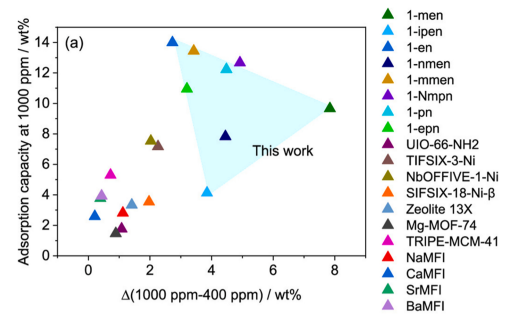


Fig. 18. Comparison diagram of the adsorption capacities at 1 mbar (1000 ppm) CO_2 for selected MOF adsorbents. Adapted from Ref. [246].

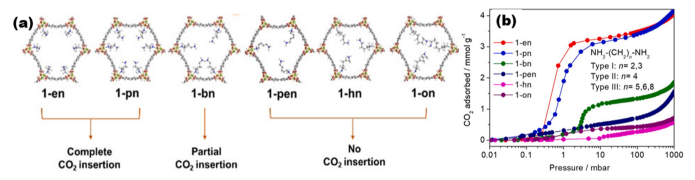


Fig. 19. *n*-alkyl diamine functionalized Mg₂(dobpdc) series (a) CO₂ adsorption mechanism, and (b) CO₂ sorption isotherm measured at 40 °C. Adapted from Ref. [247].

chemical environments for the carbamate species, which are attributed to the varied relative conformations of carbon chains upon CO_2 addition. With this study, new essential chemical insights were acquired for the interrelationship between the CO_2 insertion mechanism and diamine chain length that can further be utilized by the researchers for developing new more efficient amine appended MOF adsorbents for carbon capture.

2.5.3. Post-synthetically aminated MOFs in MIL-family

Although aminated $M_2(dobpdc)$, as well as other MOF-74 analogs have been the subject of intensive research, many other MOF adsorbents with large pore structures have been reported and can also be post-synthetically modified. For instance, MIL-100(Cr) is known for its high thermal stability, large pore size, and has been incorporated with different amines, such as ethylenediamine (en) and N,N'-dimethylethylenediamine (mmen). En-MIL-100(Cr) has demonstrated adsorption capacity of 2.4 mmol g⁻¹ (30 °C, 1 bar CO₂) and adsorption heat of about - 80 kJ mol⁻¹ indicating favoring of a chemisorption mechanism in contrast to the pristine MIL-100(Cr) having 1.6 mmol g⁻¹ and - 63 kJ mol⁻¹ values for capacity and adsorption heat, suggesting physisorption mechanism even though it shows relatively high adsorption heat, no chemisorption capable functional groups exist in the original framework. In the literature, CO₂ capturing efficiency for

ethylenediamine aminated frameworks have been shown to be somewhat higher than on mmen-MIL-100(Cr) having \sim 1.7 mmol g⁻¹ CO₂ uptakes reported under the same adsorption conditions [248].

Due to very large pore structure of MIL-101, besides discrete amines used with the MOF-74 family, also polymeric amines can be used. Polyethyleneimines (PEIs; both low and high molecular weight) are the most widely used of them, due to e.g., for their reasonably high thermal stability. Lin et al. [249] showed in their study that porosity and physical size of the MIL-101(Cr) particles as well as molecular weight and the structural properties of the loaded polyamines, all are important variables to be optimized to afford better CO2 capturing properties for such hybrid materials. In terms of particle size, it was observed that small particle-sized samples enhanced the impregnation of PEI into the inner pores of the adsorbent but at the same time brought down the surface area/pore volumes, and consequently the adsorption capacity decreased. Also, the low molecular-weight PEI diffused more easily into the inner pores compared to the high-weight PEI, which was not fully loaded in, as it more preferably settled on the surface of MOF particles. The highest adsorption capacity of 3.6 mmol g⁻¹ was obtained for PEI-300-MIL-101(Cr) with ultrahigh CO₂/N₂ selectivity at 150 mbar CO₂ pressure (25 °C). In terms of thermal properties, similar thermal stabilities were observed regardless of considerably different MWs as MIL-101(Cr) was impregnated both with linear (MW = 300 Da) and

branched PEI (MW = 1800 and 10,000 Da). Of the above, the linear PEI displayed better regenerability, albeit after 50 sorption cycles the CO2 uptakes were already decreased about 20%. Similarly to PEI- impregnated MIL-101(Cr), Jiang et al. [250] investigated incorporation of PEI into the pores of light-responsive azobenzene-decorated MOF (azoMOF). They used a wet-impregnation technique with various PEI loadings (e.g., 0.39, 0.66, 0.98 and 1.28 mmol g^{-1}), of which the best adsorption capacity was afforded by 0.98 mmol g^{-1} loading of PEI (P_3 /azoMOF). The specific surface determined for the modified azoMOF proved to be significantly lower (324 m 2 g $^{-1}$) than on pristine azoMOF (1275 m 2 g $^{-1}$) but nonetheless, P₃/azoMOF demonstrated ~2.9 mmol g⁻¹ (25 °C, 1 bar CO₂) adsorption capacity. The higher adsorption heat (-64 kJ mol⁻¹) of polyaminated adsorbent referred to chemisorption mechanism, whereas the pristine azoMOF with lower adsorption heat and the capturing capacity (\sim 1.5 mmol g⁻¹) indicated favoring the physisorption. With UV-vis spectroscopy, the effects of light-response on adsorption properties were examined with P₃/azoMOF. In the case of cis-isomer the capacity and adsorption heat were about 33% and 26% lower (2.2 mmol g⁻¹ and -47.5 kJ mol⁻¹, respectively) than that of found for trans-configuration. This suggests cis-P3/azoMOF structure modification to have considerably lower adsorbate-adsorbent interactions than in trans-isomer. For trans-isomer, DFT calculations revealed surface potentials around amine to be - 0.048 eV indicating negligible interaction between trans-azobenzene and amine. On the contrary, cis-isomer showed a significant interaction between cis-azobenzene and amine (-0.025 eV) and with higher interaction, the adsorption sites were more shielded resulting in lower CO2 capacity of cis-isomer. The strategy outlined by the study can be utilized in forthcoming designs of new light-responsive MOFs to achieve increasingly controllable CO2 adsorption separation.

Besides polymeric PEI, various smaller polyamines, such as tris(2aminoethyl)amine (TAEA), triethylenediamine (TEDA), p-phenylenediamine (PPD), tetraethylenepentamine (TEPA) have been appended to the MIL-101 framework. Although in many studies the above amines have been shown to be inferior in thermal stability to polymeric amines like PEI. Darunte et al. [26] examined adsorption properties of tris-(2-aminoethyl) (tren) and branched low molecular weight polyethyleneimine (PEI- 800) loaded to MIL-101(Cr) under DAC conditions. At first modest 0.35 mmol g⁻¹ capacity was observed at a 0.4 mbar CO₂ pressure for tren-MIL-101(Cr). With an increase in the amine loading by wet impregnation, the CO₂ capacity was increased to 2.8 mmol g⁻¹. Unfortunately, due to the high volatility of low boiling tren-amine, adsorbent demonstrated some 13 % uptake loss already after three cycles, whereas only 4 % amine loss occurred on PEI-MIL-101(Cr) in the same conditions. To further examine the optimal amine content on MIL-101, a range of different PEI loadings from 0.18 to 1.8 mmol g⁻¹ were tested. The results unfolded that capture efficiency was found to be directly dependent on amine content until the step-like increase was observed with the amine loads higher than 0.8 mmol g⁻¹. Kyu et al. [251] incorporated polymethacrylamide (PM) into MIL-101 (PM@MOF) via double solvent method and free NH2 groups were further post-synthetically introduced to the material to obtain R-PM@MIL-101. Based on the higher adsorption heat (-50 kJ mol⁻¹) the modified material exhibited chemisorptive CO2 uptake on its amino sites both at low and atmospheric pressures (150 and 1 bar CO2) with 3.6 and 1.4 mmol g⁻¹ capacities, respectively. Further studies on polymeric amine loads were carried out by Sokhrab et al. [252] as they examined intra-channel polymerization of aniline on MIL-101(Cr) to obtain PANI@MIL-101(Cr) hybrid material. With XPS, amine (-NH-), protonated amine (-NH₂⁺-), imine (=N-) and protonated imine (=NH⁺-) signals at 399.2, 400.2, 397.9 and 400.6 eV, respectively were observed. The calculated amine\imine ratio of 2.35 indicated the presence of partially protonated PANI which originates from the intermediate oxidation states between the proto-emeraldine and emeraldine species. The modified material demonstrated CO₂ adsorption capacities of 2.26 and 1.62 mmol g⁻¹ at 0 and 20 °C (1.2 bar CO₂), respectively. Moreover,

high CO_2/N_2 selectivity was observed at 0 °C. Shin et al. [253] incorporated polyvinylamine (PVAm) to MIL-101 (PVAm@MIL-101) via the "ship in a bottle" method. Among the various PVAm loadings tested, the PVAm (0.7)@MIL-101 with the 70% volume filled of full volume by the N-vinylformamide, showed high CO_2 uptake of 1.5 and 3.3 mmol g^{-1} at 150 mbar and 1 bar CO_2 (25 °C), respectively. With N-vinylformamide load, the CO_2/N_2 selectivity of the adsorbent increased approximately 11-fold compared to the unmodified adsorbent.

Babaei et al. [254] conducted a comparative study with amine-impregnated MIL-100 and MIL-101 using p-phenylenediamine (PPD). For the aminated MOFs, the adsorption capacity of MIL-101 was found to be somewhat higher than on MIL-100 with 1.7 vs. 1.3 mmol g⁻¹ at 1 bar CO₂, respectively, as expected due to the higher surface area and pore volume of MIL-101. Zhong et al. [255] surveyed a double-solvent incorporation technique in which the amines are promptly "squeezed" into the MOF pores without collapsing the framework. With the presented technique, MIL-101(Cr) was functionalized using tris (2-aminoethyl)-amine (TAEA), ethylenediamine (en) and triethylenediamine (TEDA). Interestingly, the CO2 uptake of TAEA-MIL-101(Cr) was observed to be 1.5 times higher than for pristine MIL-101(Cr) with uptakes of 5.05 and 3.26 mmol g^{-1} , respectively (at 1 bar CO₂, 0 °C). For these aminated adsorbents, the adsorption heats varied from exceptionally high $-118.7 \text{ kJ mol}^{-1}$ to $-49.2 \text{ kJ mol}^{-1}$ of which the lower values indicate physisorption instead of chemisorption as with unmodified MIL-101(Cr) showing -22.4 kJ mol⁻¹ adsorption heat.

Li et al. took a different approach for amine appending as they aminated MIL-101(Cr) with tris(2-aminoethyl)amine (TAEA) in cyclohexane [256]. The idea behind this concept is that the low polarity solvent will have weaker interaction with the alkylamine than a high polarity solvent. Consequently, the better chemical potential of the dispersed alkylamine molecules will transfer more alkylamine molecules from the solution into the framework (Fig. 20). The prepared TAEA-MIL-101(Cr) exhibited high 4.21 mmol g⁻¹ CO₂ uptake at 150 mbar and 25 $^{\circ}\text{C}.$ Regeneration tests with five cycles, under 15:85 CO₂/He mixture during adsorption (40 °C), and pure He on desorption (90 °C) revealed no appreciable decrease in CO₂ capacity indicating good stability of the adsorbent. The yet different approach was introduced by Li et al., wherein MIL-101(Cr)-SO₃H was loaded with tris (2-aminoethyl)amine (TAEA) via Brønsted acid-base reaction between sulphonic acid and amine groups. CO2 uptakes for the prepared adsorbents varied from 1.12 to 2.28 mmol g^{-1} at 0.4 and 150 mbar (25 °C). In the same study, experimental conditions for adding the amines to the frameworks were also optimized from a thermodynamic point of view. It was concluded that the most practical procedures included a reaction time of about 5 min at room temperature. Authors suggested that with given simple synthetic techniques and economical starting materials, this adsorbent may hold great potential for its large-scale production in the future [257].

Recently, Vo et al. reported amine appended NH₂-MIL-101(Cr) which was further aminated by the incorporation of ethylenediamine (en) into the pores, because of which the recovery capacity of the adsorbent increased by 62% (3.4 vs. 2.10 mmol g⁻¹) compared to that observed for NH₂-MIL-101(Cr)-NH₂ having only amino groups substituted on its linker ligands [258]. Furthermore, Li et al. [259] fine-tuned the hydrophobicity of MIL-101(Cr) by varying the alkyl chain lengths of the primary amines (*n*-propylamine, *n*-hexylamine, and 1-dodecylamine), as shown in Fig. 21. With the longer alkyl chained amines, the initially hydrophilic MIL-101(Cr) was converted into a more hydrophobic one. Of the ones tested, dodecylamine-MIL-101 exhibited the highest hydrophobicity by showing an exponential increase in the contact angle measurements.

Likewise, other MOFs and their aminated variants, such as CuBTTri, HKUST-1, MIL-101(e.g., Fe, Mg), UiO-66, UiO-66-NH $_2$, and MOF-177 [260–267] have been examined for carbon capture but are not reviewed in detail in this review. Selected CO $_2$ capture capacities for all the adsorbents presented above, are shown in Table 3.

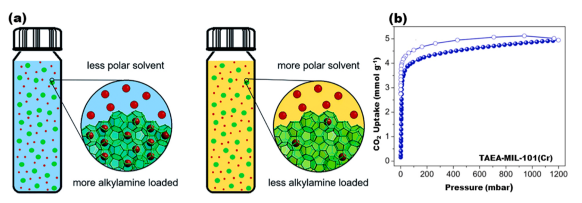


Fig. 20. (a) Illustration of the different alkylamine loading amounts in MOF, and (b) CO₂ sorption isotherm for TAEA-MIL-101(Cr) measured at 25 °C. Adapted from Ref. [256].

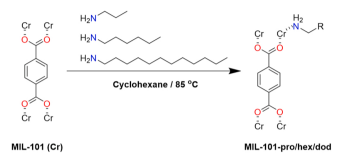


Fig. 21. Illustration of the different alkylamine loading amounts in MIL-101(Cr).

There are also other strategies to design MOFs for carbon capture than using the above exemplified methods. For example, in the recent work of Bien et al. [104] nucleophilic metal-hydroxide (M-OH) functional groups were appended to the MOF. The nucleophilic Zn-OH sites in CFA-1 displayed remarkable performance under DAC conditions with an adsorption capacity of 2.20 mmol g-1 and adsorption heat of - 71 kJ mol⁻¹ indicating CO₂ chemisorption mechanism, wherein Zn-OH converts to Zn-O2COH with aid of inter-cluster hydrogen-bonding interactions that were confirmed by DRIFTS experiments and DFT calculations. The adsorbent was subjected to column breakthrough experiments for both dry and 100% relative humidity at 0.4 mbar CO2. Notably, under dry conditions, CO2 capacity of 2.7–3.2 mmol g⁻¹ was observed in three consecutive cycles but in humid conditions adsorption did not seem to occur at all, limiting the use of sorbent considerably. Furthermore, Bien et al. [268] presented a series of heterobimetallic Co, Ni, or Cu analogues of CFA-1. The Cu-exchanged MOF analogue experienced degradation with PSM while the Ni-OH analogue exceeded the parent Zn-MOF by displaying greater capacity of 2.7 mmol g-1 (0.4 mbar) and high adsorption heat of 84 kJ mol⁻¹. Again, based on DRIFTS experiments, and DFT calculations the Ni-O2COH species did not seem to participate in the inter-cluster hydrogen bonding, unlike in the parent Zn-OH system. Both Zn-OH and Ni-OH variants were subjected to column breakthrough experiments using again both dry and humid conditions (0.25 mbar CO2, 8-13 % relative humidity) resulting in unchanged capacity for Zn-OH in dry conditions, whereas the capacity of Ni-OH variant decreased slightly. However, under humid conditions and after consecutive cycling experiments Ni-OH system appeared to be the more stable one. Overall, it was suggested that MOFs comprising nucleophilic M-OH groups could hold great potential for CO2 capture at low pressures, while some challenges associated with somewhat poor water compatibility and stability of the M-OH-based adsorbents still need to be overcome.

2.6. Confinement of water within MOFs pores

As discussed earlier, water molecules inside the pore structure of MOF can have adverse effects on the desired adsorption process but its presence can also have beneficial effects on CO2 adsorption. In this chapter, we briefly highlight some studies that we believe are benchmarks for describing methods wherein by pre-confining solvents in the cavities of the MOFs will enhance the adsorption of CO2. In 2012, Soubeyrand-Lenoir et al. [269] presented a detailed study for improving carbon capturing properties of UiO-66(Zr), MIL100 (Fe), and HKUST-1, by pre-adsorbing water into the pore structure. Under varied relative humidity (3, 10, 20, 40 %), water vapor had no relevant positive effects on CO₂ uptake in case of UiO-66. While MIL-100(Fe) demonstrated more promising results in comparison to dry conditions, as significant 3- and 5-fold CO2 uptake increments were observed at 20 % and 40 % RH conditions, respectively. This was attributed to the fact that pre-adsorbed water in the mesoporous MIL-100(Fe) forms microporous pockets that will be filled with CO₂ molecules at lower pressures. On HKUST-1, low water content showed an increase in CO₂ uptake but conversely, at higher humidity, the adsorption capacity decreased most likely due to the gradual deterioration of the framework via hydrolysis and saturation of OMSsS by water.

The promoting effect of water on CO2 uptake is evidenced also in more water-stable MOFs that were decorated with bridging hydroxo groups (µ2-OH), like in case of NOTT-400 ([Sc2(OH)2(BPTC)] made of biphenyl-3,3',5,5'-tetracarboxylic acid (H₄BPTC) [270], NOTT-401 ([Sc (OH)(TDA)] made of 2,5-thiophenedicarboxylic acid (H₂TDC) [271, 272], InOF-1 [In₂(OH)₂(BPTC)] [273], MIL-53(Al) ([Al(OH)(BDC)] [274,275], Mg-CUK-1 [Mg₃(OH)₂(2,4-PDC)₂] made of 2,4-pyridinecarboxylic acid (2,4-H₂PCD) [276], CAU-10 [AlO₄(OH)₂BDC] [277], and PCN-250 made of 3,3',5,5'-azobenzenetetracarboxylic acid (H₄ABTC) [278], to mention few. Overall, MOF dependently CO₂ capacity could initially rise, depending on the MOF, substantially in the presence of water but only up to a certain level before it is eventually reduced back to zero. For instance, in the case of the NOT-400 [270], sorption at 20 % RH resulted to a 2.5-fold increase in CO2 uptake but at higher humidity (35 % and 60 %) uptake was no longer observed at all, which could indicate blocking of the pores by water molecules. In a similar fashion with NOTT-401 [272], higher CO2 uptake was achieved reaching 3.2-fold at 5 %, 1.6-fold at 10 % and reducing virtually to zero adsorption at higher than 30 % humidity. Likewise, for indium-containing InOF-1 MOF [273], its capacity doubled at 20 % but above 40 % humidity capture capacity decreased, suggesting again pore blocking by water molecules. Similar trend was observed with CAU-10 [277], Mg-CUK-1 [276], and MIL-53(Al)-TDC containing [Al $_2$ (μ_2 -OH)] units [274,275,279]. It was concluded that $\mu_2\text{-OH}$ interacts with water molecules via strong hydrogen bonds thereby inducing so-called confinement effect [280,281] that causes reduction of the micropores,

Table 3 Summary of CO_2 adsorption for post-synthetically modified MOFs.

MOF	Capacity (mmol g ⁻¹)			-Q _{st} (kJ mol ⁻¹)	Temp. (°C)	Ref.
	0.39 (mbar)	150 (mbar)	1 (bar)			
en-Mg ₂ (dobdc)	1.51				25	[22
Mg ₂ (dobdc)(N ₂ H ₄) _{1.8}	3.89	5.18	5.51	90	25	[22
tepa-Mg ₂ (dobpc)		6.06\8.31 ^H			25	[23
en-Mg ₂ (dondc)		1.99	2.63	7–50	25	[12
ppz-Mg ₂ (dondc)		2.04	3.15	20–47	25	[12
mmen-Mg ₂ (dondc)		3.20	4.13	13–83	25	[12
mmen-Mg ₂ (dobpdc)	2.0	3.13	3.86	71	25	[10
en-Mg ₂ (dobpdc)	2.83	3.62	4.57	49–51	25	[23
dmen-Mg ₂ (dobpdc)	2.03	3.77 ^a	4.34 ^a	71–75	40	[23
men-Mg ₂ (dobpdc)	~0.41\2.19**	3.6 ^a	4.5	65–77	25	[238,24
	-0.41\2.19	2.15 ^a	3.15	65–77	25	[230,24
den-Mg ₂ (dobpdc)	0.06\ 0.02**	3.47 ^a				
ipen-Mg ₂ (dobpdc)	~0.06\0.93**		4.05 ^a	76–85	40	[239,24
een-Mg ₂ (dobpdc)	0 -0 4**	4.04 ^a	5.05 ^a	70–74	40	[23
nmen-Mg ₂ (dobpdc)	~0.76\1.77**	2.92 ^a	3.99 ^a	60–72	40	[239,24
pn-Mg ₂ (dobpdc)	~1.76\2.77**	4.01 ^a	4.49 ^a		40	[103,24
mpn-Mg ₂ (dobpdc)		3.82^{a}	4.14 ^a		40	[10
dmpn-Mg ₂ (dobpdc)		2.91 ^a	3.73^{a}		40	[10
Γ02-Mg ₂ (dobpdc)		2.17 ^a		31–39	40	[24
Γ2-Mg ₂ (dobpdc)		2.53 ^a		77–109	40	[24
T20-Mg ₂ (dobpdc)		1.43 ^a		77-109	40	[24
T200-Mg ₂ (dobpdc)		1.46 ^a		77–109	40	[24
2-ampd-Mg ₂ (dobpdc)	3.47 ^{a,*}			73	40	[24
Mg ₂ (dobpdc)(3–4–3)	~3.40 ^{d,*}			99	90	[24
e-2-OH-Mg ₂ (dobpdc)	5.40	$\sim 2.10^{b}$		81	50	[24
		~1.90 ^b		81	50	
nPr-2-OH-Mg ₂ (dobpdc)						[24
Pr-2-OH-Mg ₂ (dobpdc)		~1.75 ^b			50	[24
cy-2-OH-Mg ₂ (dobpdc)		~1.50 ^b			50	[24
nBu-2-OH-Mg ₂ (dobpdc)		~1.70 ^c			75	[24
Pent-2-OH-Mg ₂ (dobpdc)		$\sim 1.40^{\rm c}$			75	[24
epn-Mg ₂ (dobpdc)	~1.76\2.88**				25	[24
Table 3. Continued.						
Nmpn-Mg ₂ (dobpdc)	~1.76\2.50**				25	[24
1-pn- Mg ₂ (dobpdc)			4.19 ^a		40	[24
1-bn- Mg ₂ (dobpdc)			1.87 ^a	46-62	40	[24
1-pen- Mg ₂ (dobpdc)			1.57 ^a		40	[24
1-hn- Mg ₂ (dobpdc)			0.57 ^a		40	[24
1-on- Mg ₂ (dobpdc)			0.72^{a}		40	[24
en-MIL-100(Cr)			2.4	80	25	[24
mmen-MIL-100(Cr)			1.7	80	25	[24
	0.35		1./	80		
MIL-101(Cr)-TREN	0.35	4.4			25	[2
MIL-101(Cr)-PEI-300		4.1			25	[24
P ₃ /azoMOF			~2.9	64	25	[25
R-PM24 @MOF		1.4	3.6	50	25	[25
PANI@MIL-101(Cr)			1.62 * **	26	20	[2
PVAm (0.7)@MIL-101		1.5	3.3	49	25	[2:
Cr-MIL-101-SO ₃ H-TAEA	1.12	2.28 ^a		87	25	[2
CuBTTri-mmen		2.38		96	25	[20
tepa-NH ₂ -MIL-101			3.1		25	[26
MIL-100-PPD			1.3		25	[25
MIL-101-PPD			1.7		25	[25
ed-MIL-101(Cr)			1.93		25	[2:
ΓAEA-MIL-101(Cr)			2.19		25	[2:
					25	
FEDA-MIL-101(Cr)			1.65	74.0		[2
DETA-MIL-101(Fe)			1.82	74.2	25	[2
PEI-MIL101 (Cr, Mg)			3.04		25	[2
ed-MIL-101(Cr)-NH ₂			3.4		25	[2:
PEI-UiO-66			3.3		25	[20
PEI-UiO-66-NH ₂		2.3	3.1	68	25	[20
ed-HKUST			2.4		25	[2
DETA-MOF-177			2.83		25	[20
ГЕРА-МОГ-177			3.82		25	[26
PEI-MOF-177			2.84		25	[26
CFA-1(Zn-OH)	2.2		2.07	71	25	[10
	2.7					
CFA-1(Ni-OH)	4./			84	25	[26

 $^{^{\}rm a}$ at 40, $^{\rm b}$ 50, $^{\rm c}$ 75, $^{\rm d}$ and at 90 °C, $^{\rm H}$ humid condition, * at 40 mbar, ** at 1 mbar, *** at 1.2 bar.

which further allow the better packing of CO_2 molecules [282,283]. Thus, in the microporous MOF that contains hydroxo (μ_2 -OH) functional groups, it is advantageous to pre-adsorb small amounts of water, as their presence in the pores could increase the adsorption of CO_2 . Nonetheless, more work in experimental and computational research is required to

accurately elucidate the adsorption mechanism under humid conditions in these adsorbents. In general, as discussed above, water presence can also adversely effects on adsorption, because of pore blocking properties, or competitive adsorption to OMSs and amine sites.

2.7. Improved hydrophobicity of MOF adsorbents by coating techniques

As reviewed above hydrophobicity of a MOF can be improved by modifying its internal structure but physical surface structure of the MOF particles can be modified as well. Typical methods used for surface modification are, such as surface coating, post-synthetic thermal annealing, and post-synthetic chemical modification of the surface, as already briefly mentioned in previous chapters. In 2014, Zhang et al. [284] presented a comprehensive polydimethylsiloxane (PDMS) coating technique for amplifying the water stability of the designed MOFs (Fig. 22). Three structurally different MOFs, namely MOF-5, HKUST-1, and ZnBT were coated by PDMS. Surprisingly, hydrophobic PDMS coating induced positive outcomes in humid conditions as adsorption efficiency remained relatively unchanged compared to pristine MOFs. Also, the surface area, crystalline nature and pore texture of the MOF remained reasonably unchanged. Qian et al. [285] deposited a hydrophobic organosilicone layer on the exterior surface of HKUST-1, MIL-125, and ZIF-67 via a facile solution-immersion process without using any heat treatment. The hydrophobic organosilicone layer provided durable protection against exposure to water, and the MOF maintained its porosity for 5 days. Yang et al. [286] successfully established a two-step post-synthetic polymerization strategy to improve the stability of structurally different MOFs, including, HKUST-1, ZIF-67(Co), ZIF-8(Zn), UiO-66(Zr), Cu-TDPAT, Mg-MOF-74 and MIL-100(Fe). This novel technique based on using free base like dopamine as a binder. However, dopamine hydrochloride was used as a precursor to eliminate need of an aqueous alkaline environment for the coating process. The actual process is two-step (Fig. 23), wherein first polydopamine (PDA) was coated on the surface of the MOFs, under a mild oxygen atmosphere, yielding MOF@PDA. In the second step, the new external surface was covalently modified with hydrophobic 1 H, 1 H,2 H,2 H-perfuorodecanethiol (HSF), which eventually improved the water stability of the adsorbent significantly even under harsh acidic and alkaline conditions. In view of the foregoing, this facile strategy could be universally applied to incorporate hydrophobicity into MOFs and to taper their water permeability.

An alternative method for hydrophobic coating is the surface pyrolysis described by Park et al. They synthesized an exceedingly moisture-resistant, black-colored MOF-5 via heat treatment [287]. Although the surface area of the thermally modified MOF-5 was only half of the pristine adsorbent (1740 vs. $3450~\text{m}^2~\text{g}^{-1}$, respectively), the crystallinity, and the remaining porosity were found to be unchanged for two weeks upon exposure to moist ambient air. Hence showing clearly better hydrostability compared to MOF-5. Moreover, Gadipelli et al. [288] tested the effects of post-synthetic thermal annealing on MOF-5 at temperatures lower than its typical thermal decomposition temperature

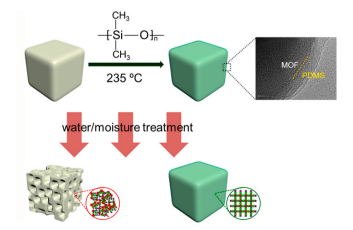


Fig. 22. Illustration of PDMS-Coating on the MOFs Surface. Adapted from Ref. [284].

at around 500 °C. The MOF-5 samples annealed at 380 (also 6 h) and 400 °C for three hours showed higher CO_2 uptakes of 1.5, 2, and 1.1 mmol g^{-1} in comparison to the pristine MOF-5 having 0.8 mmol g^{-1} capacity. Furthermore, in terms of air/water stability, the sample annealed at 400 °C for 3 h maintained its porosity when exposed under ambient air for a maximum 15 days, in contrast to an unmodified adsorbent that showed complete collapse of pore structure under the same conditions.

Sun et al. [289] presented new perspective to preserve structural properties of porous materials under humid CO_2 , and harsh acid/base conditions by chemically coating the exterior of MOF crystals (Fig. 24). The vinyl-functionalized crystalline zeolite imidazole framework ZIF-8-V was employed for the molecular coating. Its external surface was hydrophobically treated with per-fluoroalkyl groups using a thiol-ene Click reaction. The new surface modified ZIF-8 remained structurally unchanged, when exposed to 100% humidity for over 720 h (at 45 $^{\circ}$ C), while the pristine ZIF-8 eventually degraded completely under the same conditions already considerably earlier (240 h).

The study of Deria et al. [290] can be highlighted (Fig. 25) as a functionalization method for improving water-resistance of MOFs by modifying the metal nodes inside the pore structure by introducing functional groups as charge compensating units. In the study, Zr-based MOFs were post-synthetically modified using method called solvent-assisted ligand incorporation (SALI) wherein perfluoroalkane carboxylates of different chain lengths (C1 - C9) were incorporated to NU-1000 [$(Zr_6(\mu_3-O)_4(\mu_3-OH)_4(-OH)_4]_2(TBAPy)_6$] consisting of 1,3,6, 8-tetrakis(p-benzoic acid)pyrene)] (H₄TBAPy). Particularly with perfluorodecanoic acid, NU-1000 preserved both its crystallinity and porosity under 20 cycles of water vapor adsorption and desorption [291], albeit in most cases, porosity was significantly reduced, and the hydrophobically treated Zr₆-oxo centers did not necessarily cooperated directly with the CO2 uptake as desired. Although fluorinated carboxylates somewhat improved the tolerance of Zr₆-oxo centers towards water molecules. Recently, Amato et al. [292] reported MOFs that outperform their Zr-based analogues (MIL-140(Zr) and UiO-66(Zr)). Since the fluorinated MOFs have shown remarkable results with respect to physisorption based capture capacity in DAC, the Amato et al. introduced tetrafluoroterephthalic acid (H2TFBDC) as a linker and prepared Ce-MOFs with MIL-140 and UiO-66 topologies. The adsorption heats were found to be about -39 kJ mol^{-1} indicating physisorption and the evaluated CO2/N2 selectivities were determined to be particularly high among the reported MOFs (>1900).

3. Engineering aspects in CO2 capture by MOFs

Successful implementation of adsorption-based gas separation into a real-life application relies on the recognition of the most suitable adsorbent to match for the practical process conditions. This requires mutual contributions both from the MOF chemistry and engineering communities to constitute a fundamental understanding of their work. It is mandatory that the engineer community is familiar with the basic knowledge of the different adsorbents, and their functionalities while designing optimal process conditions but also keeping in mind the limitations of the adsorbents. Similarly, the chemist or material scientist should be aware of the process requirements and, with those in mind, try to design an adsorbent in a way that its best possible efficiency can be achieved within the technical boundaries set by the process. In this context, to bridge the gap between adsorbents and their engineering aspects, we herein outline the major approaches in the literature for real gas separation characteristics, namely, adsorption heat, gas selectivity, multicomponent gas adsorption, and MOFs post-processing cycles, regeneration, and cost.

3.1. Isosteric heat of adsorption

Isosteric adsorption heat relates to the strength of adsorption of a

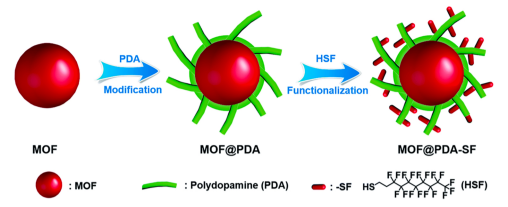


Fig. 23. The modification process used for preparing hydrophobic MOF@PDA-SF composites. Adapted from Ref. [286].

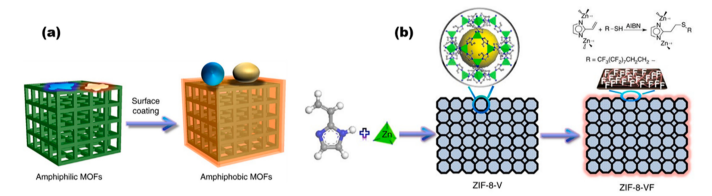


Fig. 24. (a) Amphiphobic surface engineering for MOFs, and (b) schematic illustration to impart amphiphobicity on ZIF-8-V. Adapted from Ref. [289].

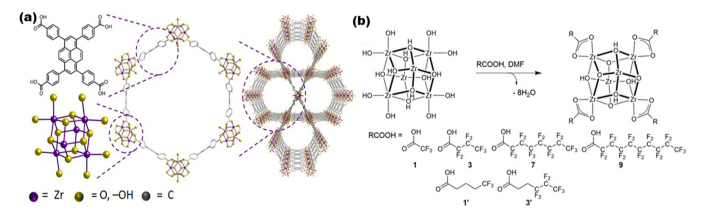


Fig. 25. Crystal structure of (a) NU-1000, and (b) illustration of solvent-assisted ligand incorporation (SALI). Adapted from Ref. [290].

given adsorbate on a given adsorbent. Thus, the energetic interaction strength between the adsorbents and adsorbates is significant in assessing the adsorptive selectivity and the design of regeneration processes [293]. Broadly, the isosteric heat (Q_{st}) can be determined by the well-known Clausius–Clapeyron equation when the following conditions are fulfilled: a) the gas adsorbed obeys the characteristics of an ideal gas, and b) the volume of the gas adsorbed is insignificant [294]. Based on the single-component isotherm curves at divergent temperatures, Q_{st} can be calculated according to the following Clapeyron Eq. (6): [295].

$$ln p = -\frac{Q}{RT} + C$$
(6)

Where, P is pressure correspondent to the same CO_2 uptake under different temperatures, R is the universal gas constant, T is temperature, and C is a constant. Thus, the Q_{st} at each adsorption point is determined based upon the slope of the plot $\ln P$ as a function of (1/T). Also, the other common way to determine the Q_{st} is to fit the temperature dependent isotherms to virial-type Eqs. (7) and (8):

$$ln P = ln N + \frac{1}{T} \sum_{i=0}^{m} a_i N^i + \sum_{i=0}^{n} b_i N^i$$
(7)

$$Q_{st} = -R \sum_{i=0}^{m} a_i N^i$$
 (8)

where P is the pressure (bar), N is the adsorbed quantity (mmol g $^{-1}$), T is the temperature (K), R is the gas constant (8.314 J K $^{-1}$ mol $^{-1}$), a_i and b_i represent the virial coefficients, and m and n exemplify the number of coefficients needed for the appropriate fitting of the isotherms. From the above, researchers have utilized Q_{st} as an important parameter to discover the beneficial changes in the sorbate–adsorbent interaction, and its effect on gas selectivity in functionalized frameworks [296–298]. For instance, Park et al. [299] demonstrated improved adsorption heat, selectivity, and CO_2 capacity of SNU-100 MOF by impregnating different metal ions, namely Li $^+$, Mg $^{2+}$, Ca^{2+} , Co^{2+} , and Ni $^{2+}$ into the pores. The formed electrostatic interactions between CO_2 and the impregnated metal ions, impacted to the adsorption heats which increased from - 29.3 of pristine SNU-100 to between - 37.4 and - 34.5 kJ mol $^{-1}$ depending on the metal ion used for impregnation. Similarly, the adsorption selectivity elevated from 25 to the range of 31–40.

3.2. Ideal adsorbed solution theory (IAST)

The successful implementation of CO₂ capture processes equally demands the study of both a) single component and b) mixture sorption equilibrium data [298,300,301]. Acceptedly, single-component isotherms can be simply obtained by volumetric adsorption experiments. Nevertheless, mixture adsorption equilibrium studies represent a complicated process owing to the participation of different variables in the process [112,302]. In this regard, to resolve these complexities, the IAST method is utilized to describe multicomponent adsorption isotherms under the following three major considerations: a) adsorbate molecules in the mixture have equal accessibility to the whole surface area of the adsorbent, b) the adsorbent surface is homogeneous, and c) the adsorbed phase is an ideal solution in which intermolecular interactions are comparable in strength [303,304]. Such a methodology that explains the multicomponent adsorption isotherms from single-component adsorption isotherms starts from Raoults' law type of relationship between the adsorbed phase and fluid. Utilizing the mathematical fitting outlined by Myers & Prausnitz, the IAST permits the selectivity of an ideal multicomponent mixture to be estimated. [303] In regard to a binary mixture, the selectivity can be obtained by the following Eq. (9):

$$S_{ads} = \frac{q_1 \backslash q_2}{p_1 \backslash p_2} \tag{9}$$

For instance, based on the IAST calculations, McDonald et al. [260] demonstrated significant improvement on selectivity (327 at 25 °C) for mmen-CuBTTri, with a mixture of CO_2/N_2 at 150 and 750 mbar CO_2 , respectively. Similarly, using IAST Hu et al. [305] represented CO_2/N_2 selectivity factor of 346 for MIL-101-DETA at 23 °C. IAST has been broadly employed regarding MOFs to investigate the selectivity for separation of mixtures of industrial concerns, like the gas mixtures of N_2/CO_2 , CH_4/CO_2 and CH_4/N_2 [306–308]. Recently, Billemont et al. [309] utilized IAST to study adsorption of CO_2 from gas mixture like biogas and natural gas using mesoporous MIL-100(Fe) as adsorbent. Overall, IAST emerged with an appropriate model for these promising adsorption processes and based on that MIL-100(Fe) seemed to be potential adsorbent for CO_2/N_2 separation.

3.3. Single-component adsorption experiment

A single component adsorption experiment is often the first screening tool to determine the CO_2 adsorption capacity. Gas sorption

analyzers are categorized in two ways: volumetric (also known as manometric) and gravimetric. Gravimetric CO2 adsorption capacity describes the weight fraction requisite of MOF for an application, whereas the volumetric uptake reveals information about the bed volume of adsorbent, dimensions of required equipment and capturing unit. Data is often reported based on gravimetric analysis, which is the amount of gas adsorbed (in mass or volume, STP or moles) per unit weight of adsorbent. By multiplying this data with the adsorbent density, volumetric capacity can be obtained (cc cc^{-1} or v/v). Particularly for MOFs, crystal density is often employed for this type of conversion. Notwithstanding, it is the highest theoretical value, as in comparison to single crystal, MOF bulk powders are typically loosely packed [310]. For instance, opt-UiO-66(Zr)-(OH)2 exhibits higher crystal density of 1.36 g cm⁻³ than Mg-MOF-74 (0.91 g cm⁻³). Thus, higher volumetric capacity is determined for opt-UiO-66(Zr)-(OH)₂ than Mg-MOF-74 (172 and 165 cm³ (STP) cm⁻³, respectively) at 1 bar and 25 °C. However, in terms of gravimetric capacity under the same conditions Mg-MOF-74 demonstrates higher CO2 capacity than opt-UiO-66(Zr)-(OH)2 (8 and 5.63 mmol g⁻¹, respectively). Few examples of volumetric capacities of known MOFs includes SIFSIX-2-Cu-I (1.24 g cm $^{-3}$, 151 cm 3 (STP) cm $^{-3}$), UTSA-16 (1.66 g cm $^{-3}$, 160 cm 3 (STP) cm $^{-3}$), mmen-Mg₂(dobpdc) (1.16 g cm⁻³, 107 cm³ (STP) cm⁻³), and Mg₂(dobdc)-(N₂H₄) $(1.18 \text{ g cm}^{-3}, 145 \text{ cm}^3 \text{ (STP) cm}^{-3})$ [112]. Thus, the actual volumetric capacity is overestimated and can differ if the interstitial void of the particles and crystal defects are included. Thus, to prevent any incorrect interpretation of the experimental results, it is necessary to mention all the important details, including pressure steps, system response time, sample weight, and representative data demonstrating the suitability of the theoretical model to the experimental response curves [311]. Since the single component adsorption experiment does not consider competitive adsorption among various components, it is therefore, unable to measure the behavior of binary\ternary adsorption. Thus, under real working conditions, dynamic column breakthrough data is suggested, as discussed in 3.5.

3.4. Working capacity

Working capacity is one of the most critical parameters to determine the overall performance of adsorbent, as it takes material regeneration into consideration. Recovery is determined as the fraction between the quantity of product formed over the quantity of product in the feed stream. It is essential to pay attention to fact that the working capacity and the adsorption capacity are different quantity parameters. Working capacity (WC) relates to the difference between the quantities adsorbed at the adsorption and desorption conditions, thus CO_2 uptake capacity (q_{ads}) under adsorption minus CO_2 uptake capacity (q_{des}) under desorption conditions, Eq. (10)).

$$WC_q = q_{ads} - q_{des}$$
 (10)

Working capacity can be determined based on the type of the process, for instance being pressure swing (PSA), vacuum swing (VSA) or temperature swing adsorption (TSA). Increasing the temperature of the column above that of the feed temperature is known as temperature swing adsorption (TSA) and reducing the pressure to a value below that of the feed is known as pressure swing adsorption (PSA). If the desorption is under vacuum, the process is also called vacuum swing adsorption (VSA). In connection with PSA/VSA, the nature of the process is often regarded under isothermal conditions. A single sorption isotherm at a specific temperature T₁, is considered to suffice for calculating the working capacity by merely subtracting the equilibrium uptake at the partial pressure of gas under adsorption conditions (high P), and the uptake at the partial pressure under desorption conditions (low P). However, in the case of TSA, isotherms collected at different temperatures are required, where the working capacity is calculated by subtracting the equilibrium uptake at the partial pressure of gas under adsorption conditions (low T) from the uptake at the partial pressure under desorption conditions (high T) [231,310].

3.5. Dynamic breakthrough experiments

The comprehensive insight of the mixed gas equilibrium adsorption data is crucial for selecting the best adsorbent material, process design, and for substantiating theoretical models of mixture adsorption [112]. This engineering method can be utilized to analyze MOFs in the context of mixture-component adsorption performance and display a screen between idealistic and practicable adsorptions in the industrial applications. Thus, dynamic breakthrough experiments are conducted under conditions like those of a large-scale separation. Since the commercial equipment for breakthrough experiments is not generally available, many breakthrough experiments are conducted using in-house setups. With in-house systems, the measurement accuracy and operational control might vary, and this can lead to erroneous results and interpretations. Therefore, the establishment and running of an experimental breakthrough setup, though straightforward in theory, needs careful consideration and execution practice.

Hu et al. [312] presented a straightforward and comprehensive breakthrough setup for the gas separation. As illustrated in (Fig. 26), the fundamental part of the setup is the stainless-steel adsorption column, wherein an adsorbent is typically loaded in either as powdered or compacted form. The adsorbent bed is introduced to a gaseous stream (generally a binary or ternary mixture), and the mixture composition is continually regulated with an in-line analytical instrument (typically a mass spectrometer). The pressure drop is recorded before and after the adsorbent bed. Additionally, the exit velocity of the breakthrough column is recorded for mass balance accuracy, using argon at a constant flow rate as the internal reference. Thus, the authors presented the comprehensive procedures for conducting the breakthrough experiments so that the change in the gas velocity at the exit, and the pressure drop across the breakthrough column have been considered, which are often overlooked in the literature.

3.6. Gas diffusion

The gas diffusion parameter holds paramount importance in evaluating sorption kinetics in the MOF-based CO_2 capture. The mass transfer coefficient relates the mass transfer rate in separation processes and reveals to the particular preconditions in the design and construction of separation process equipment [313]. Diffusion studies can be categorized into two major classes, those being microscopic and macroscopic [112,314]. Microscopic methods include pulse-field gradient (PFG)-NMR [315], infra-red microscopy (IRM), and interference

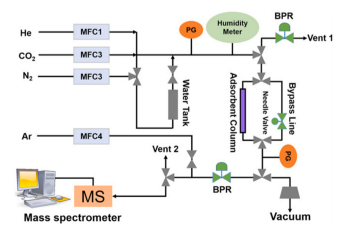


Fig. 26. Breakthrough setup for CO_2 adsorption study. Adapted from Ref. [312].

microscopy (IFM) [316]. The above methods render comprehensive details about the diffusional characteristics at the atomic level, indicating the existence of inhomogeneities, in particular surface barriers that have an impact on the behavior of adsorbates. Unfavorably, methods such as those demand samples with larger particle sizes, likely single crystals, that are commonly not available, as powdered adsorbents dominate in real-life applications. Macroscopic methods include frequency response, chromatographic methods, and zero length column methods (ZLC) [317]. Gas diffusion parameters obtained from macroscopic methods can be evaluated on the basis of the overall gas adsorption behavior of realistic powders like samples and thereby give information that can be useful for the process development [318].

3.7. MOF post-processing cycles and regenerability

As adsorbent materials exhibit a finite uptake capacity of CO₂, thereby continual interaction with ambient air or flue gas conditions will subsequently lead to a thermodynamic equilibrium between the two phases (solid and gas), wherein the adsorption and desorption rates are equivalent [313]. In view of this, it is important that adsorbents are regenerated, and semi-continuous process is developed where the adsorption and regeneration processes are grouped together.

It should also be noted that the post-processing of MOF materials is a crucial step to enable their use in an application. In lab-scale experiments, adsorbents are often supported by fine metal meshes in a fixed bed, prepared on quartz wool, or attached to a substrate. However, utilizing the powdery MOFs in an industrial application can lead to several issues, such as pressure drop, reduced or blocked flow, and abrasion due to powder migration in the system [319]. Thus, they require sufficient post-processing before being secured on a fixed bed. In this manner, the cross-column pressure drop can also be attenuated to increase the column breakthrough dynamics [312].

Darunte et al. [320] explored the possibility of utilizing an amine functionalized mmen- M_2 (dobpdc) (M = Mg and Mn) as secured on a structured monolith contactor under simulated flue gas (Fig. 27). Similarly, Lawson et al. [321] presented polyethylenimine and tetraethylenepentamine impregnated MOF-monoliths that were made with a 3D printing technique using pre- and post-functionalization approaches (Fig. 28a). Recently, Lawson et al. [322] demonstrated HKUST-1 monolith formulation based on the gel-print-grow strategy (Fig. 28b).

As mentioned above, regeneration can be accomplished by different processes or sometimes a combination of one or more methods, such as increasing the temperature, lowering the partial pressure of the adsorbates, reduction in the concentration, or inert gas purging. The objective is to place the adsorbent in a state where its uptake capacity for the adsorbent is lower than in the adsorption step, thus activating the desorption mechanism. In some cases, it is favorable to associate both temperature and pressure/vacuum swing to accomplish more desirable regeneration and, consequently, improved working capacity. After the regeneration of the bed, a new cycle can be initiated. [112].

3.8. Cost of MOFs

Currently, the production cost of MOFs is the key limiting factor for the commercialization of MOF-based adsorbents. The market price of any MOF from commercial suppliers is still exorbitant compared to that of the "conventional" adsorbents like activated carbon, silica, and zeolites. This is largely attributable to the lack of scale-up production methods developed for MOFs. Material costs are dependent mainly on four components that are required in MOF synthesis; solvents (used for reaction media and washing), metal source(s), organic ligand(s), and acid/base catalyst. As an example, material costs for some typical MOFs are shown in Fig. 29. For instance, in the case of MOF-5, producing 1 kg of activated MOF-5 demands 81.30 L of DMF, 1.03 kg of terephthalic acid, and 3.45 kg of zinc acetate dihydrate (with a reaction yield of 63%). These components cost about 414, 34, and 79 USD, respectively. The

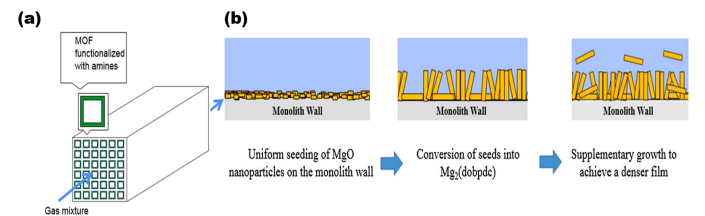


Fig. 27. (a) Monolith -supported MOF system (b) approach to create MOF films Adapted from Ref. [320].

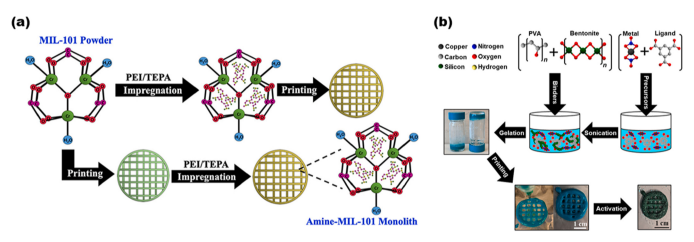
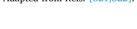


Fig. 28. (a) Manufacturing processes for pre- and post-impregnated MIL-101 monoliths, and (b) illustration of HKUST-1 monolith formulation by gel-print-grow technique.

Adapted from Refs. [321,322].



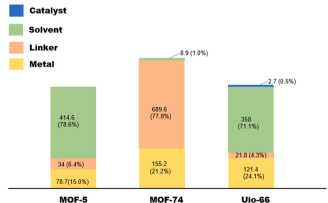


Fig. 29. Material costs of MOF-5, Ni-MOF-74, and UiO-66, given in US \$ [319].

overall product cost is \sim 527 USD kg⁻¹, of which about 79 % arises from the solvent [323]. For activated Ni-MOF-74, the total material cost is reported to be \sim 887 USD kg⁻¹, of which approximately 78 % originates from the ligand [324]. In the case of the UiO-66 total material cost is \sim 504 USD kg⁻¹ wherein solvent accounts for 72 % of the cost [325]. In this regard, substantial efforts need to be made to optimize synthesis methods, including solvent recycling, using water as a (co-)solvent, or using solvent-free mechanochemical synthesis, as well as primarily

utilizing economically inexpensive starting materials. Having said that, MOFs commercialization still requires further intensive research efforts to substantially decrease manufacturing costs and, by that enable the MOF technologies to evolve and prosper on gas sorption-based technologies.

4. Conclusions

Undoubtedly, chemists, materials scientists, and engineers have shown a deep interest in the utilization of MOF for carbon capture to minimize the impact of atmospheric CO_2 on the global temperature rise. The growing number of publications highlights the significant progress that has been already made on CO_2 related applications as reflected by the vast diversity of MOFs known to us, and the good properties they reveal for them to operate already somewhat close to real-life conditions. However, when it comes to their applicability to technical applications, it is important to point out that there are still certain challenges that need to be addressed and require further extensive research efforts.

Firstly, a very limited number of MOFs have been investigated under lower CO_2 concentrations, and mostly single-component CO_2 sorption isotherms are reported instead of real mixtures. For MOFs to be utilized in practical applications, it is important to collect a considerable range of dynamic breakthrough experiment data to establish the constructive relationship between material structure, surface chemical property, and CO_2 capture efficiency. Secondly, the next challenge circles around the MOFs synthesis on a large scale (first at Kg quantities) with reasonable

cost, sufficient phase purity and correct morphology (pellets, monoliths), as well as chemically robust nature, so that they can withstand tens of thousands of loading/unloading cycles. More research efforts are needed to further enhance CO_2 capture capacity and selectivity, particularly under humid conditions.

From a structural point of view, modification of the metal centers on the frameworks, remains a promising strategy and central research direction towards CO_2 capture. However, considering Mg-MOF-74, the best performing MOF under relatively higher pressures and concentrations, is not compatible with DAC owing to its unsatisfactory CO_2 selectivity at lower concentrations. Therefore, physisorption by MOFs is only applicable at relatively higher pressures and concentrations and not in low CO_2 partial pressures. Nevertheless, this is not the case with hybrid ultramicroporous (HUM) mesh-like adsorbents that exhibit physisorption mechanisms.

To support this, Kumar et al.[150] presented a screening of four different classes of porous adsorbents namely SIFSIX-3-Ni, HKUST-1, Mg-MOF-74/Mg-dobdc, zeolite-13X, and TEPA-SBA-15. Predictably, the only chemisorbing TEPA-SBA-15 exhibited superior DAC performance and CO₂/H₂O selectivity in comparison with the other physisorbents. However, even though SIFSIX-3-Ni demonstrated also good efficiency in DAC conditions it still showed poor CO₂/H₂O selectivity, which was also the case for all the other physisorbents. Amongst similar MOFs, the NbOFFIVE-1-Ni has shown the best DAC performance in comparison to a series of similar isostructural materials. In terms of regeneration, HUMs displayed optimal regeneration energy at a relatively low temperature range 90-100 °C under purging with inert gas or vacuum but at humid conditions significant drop in CO₂ uptake, and lower working capacity were observed due to the competitive water adsorption. Thus, it is evident that this class of MOFs is limited only to dry atmospheric conditions and are not good enough to compete with chemisorbents without further development.

These shortcomings can be resolved by functionalization of MOF pores with amines that react with CO2 via chemisorption and convert it into carbamic acid, carbamate, or bicarbonate species. The stronger interaction between amine and CO₂ offers superior CO₂/H₂O selectivity in comparison to physisorbents. In this context, diamine functionalized Mg2(dobpdc) has so far displayed superior CO2 uptake capacity combined with good stability under humid conditions. Further optimization is still required for shaping this MOF into industrially suitable forms (granules, pellets, beads, films, or monoliths). Massive scale-up of production (Mt per year) but with reasonable costs (<\$10 per kg)[326] also needs technical solutions, as well as stability since adsorbent needs to withstand thousands of sorption cycles under practical flue gas conditions and especially in DAC. To fulfill these requirements, DAC technologies must overcome several technical and economic challenges. Although DAC has undergone considerable technical advances in the recent years, the field is still in its infancy. As demonstrated earlier, for efficiently working adsorbent in DAC, the high enough CO2 capacity with high enough adsorption heat to capture CO2 selectively via chemisorption are the essential parameters to develop but also meeting the low enough energy requirement for the desorption process. Future studies should also focus on parameters such as degradation and cycle time of adsorbents. In practical DAC process, both loading\unloading times should be measured instead inert gas purge or gravimetric CO2 uptake experiment. Thus, adsorbent regeneration strategies and capacity measurements based on laboratory scale often encounters new obstacles when they are translated to real-life DAC. More research in the case of DAC adsorbents is needed to evaluate their true potential for commercial carbon capture [38,326-329]. Overall, utilizing MOFs in carbon capture involves many barriers and challenges which are so far unsolved and require both chemists and process engineers to come forward for constructive collaboration in the field of carbon capture. We anticipate that further extensive research efforts and investigations are needed for MOFs to boost them to future applications, and we certainly believe that this progress is possible, as there are diverse possibilities to

formulate, synthesize, and modify MOF-based adsorbents to fulfil the high expectations of an ultimately effective and energy-efficient adsorbents suitable task-specifically for different areas of carbon capture, like for flue gases emissions and DAC.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data Availability

No data was used for the research described in the article.

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