Zeolite-like metal–organic frameworks (ZMOFs): design, synthesis, and properties

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This review highlights various design and synthesis approaches toward the construction of ZMOFs, which are metal–organic frameworks (MOFs) with topologies and, in some cases, features akin to traditional inorganic zeolites. The interest in this unique subset of MOFs is correlated with their exceptional characteristics arising from the periodic pore systems and distinctive cage-like cavities, in conjunction with modular intra- and/or extra-framework components, which ultimately allow for tailoring of the pore size, pore shape, and/or properties towards specific applications.

1. Introduction

In recent years, hybrid organic–inorganic materials, especially metal–organic frameworks (MOFs),1 have developed rapidly due, in part, to their endlessly modular and versatile nature, which is evident from the numerous reported metal-ion or metal-cluster types in combination with a continuously expanding library of multi-functional organic ligands. In addition, MOFs, which vary in dimensionality from two- to three-periodic extended frameworks, including open, permanently porous structures, are efficiently generated through typically mild synthetic techniques, resulting in highly crystalline materials, ideal for in-depth characterization of their structures. As such, correlations have been drawn between their structure(s) and properties, indicating their outstanding potential in many applications (e.g., gas storage/separation/sequestration, catalysis, sensing, magnetism, non-linear optics, and more).2–14 In this context we see another key feature contributing to the precipitous advancement of MOFs, the potential for designing methods towards tailored functional materials.

Numerous rational approaches to target particular MOF structures have been devised and systematically developed over the past couple of decades. A major advancement is attributed to the molecular building block (MBB) approach,15–23 an approach that views certain discrete components with known features as individual building blocks for the construction of a final structure; essentially, the effective coordination geometry of single-metal ions and/or inorganic clusters, as well as the shape of the corresponding multifunctional organic ligands, directs the MOF formation, usually based on known, targeted network topologies. This strategy offers a prospective avenue toward not only the design and construction of materials, but also designed functional materials, as desired functions/properties can be incorporated at the design (i.e., building block) stage. For the primary construction of the targeted structures, it is necessary to utilize MBBs that possess rigidity and desired directionality prior to the assembly process. As the inorganic MBBs are typically formed in situ, it is fundamentally important to identify the appropriate reaction conditions under which they are consistently generated. Once this aspect is realized, desired frameworks can be targeted by a combination of the inorganic MBB and judiciously selected organic ligands (which may serve merely as bridging linkers or as additional rigid, directional MBBs, depending on the desired framework).2,3

One ideal type of structure to target is the group of purely inorganic materials known as zeolites, which represent a benchmark in the area of porous solid-state materials, owing this status to their notable commercial significance.24 These materials are comprised of Si and/or Al tetrahedral metal ions (T), linked by oxygen atoms (O, technically oxide ions), at approximately 144° T–O–T angles. The attractiveness associated with these compounds relies, in part, on their porosity, with homogeneously-sized and -shaped openings and voids, and forbidden interpenetration. The confined spaces permit their conventional use par excellence as shape- and size-selective catalysts, ion exchangers (ion removal and water softening), adsorbents (separation and gas storage), etc.24–37 The diversity...
of these compounds is reflected in the extended number of framework types (there are currently 225 zeolites, as recognized by the International Zeolites Association), each differentiated by a specific profile, such as the size of member rings (MR), window/aperture opening, cage dimensions, charge density, framework density (FD, the number of tetrahedral vertices per 1000 Å³), and types of pores. Thus, the access to a multitude of networks makes zeolite-like (also sometimes referred to as zeolitic, zeotype, or zeotypic) materials highly valuable in function. The diverse nature of these materials is often influenced by the synthetic conditions, and by the use of structure directing agents (SDAs). However, limitations in their design and tunability restrict these functional materials to certain pore sizes and, consequently, to smaller molecule applications. In addition, a general trend implies that increasing pore sizes may lead to unidimensional pore systems and, hence, limit the applications.

In this context, and considering the relevance of the functions associated with solid-state porous materials on a societal and industrial level, the pursuit of novel materials, like MOFs, based on, and expanding, zeolitic networks has become a prominent and encouraging avenue. Consequently, this review aims to portray the state-of-the-art in the emerging area of MOFs related to zeolite nets. The focus will be placed on the breadth and efficacy of design routes (Fig. 1), delineating avenues toward the construction of zeolite-like MOFs (ZMOFs):

(i) based on the “edge-expansion” of traditional zeolites;
(ii) assembled from hierarchically superior building units, such as metal–organic cubes, regarded as d4Rs in zeolites;
(iii) derived from enlarged tetrahedral building units; and
(iv) built via organic tetrahedral nodes.

The most prominent examples of each strategy are to be briefly outlined. The concluding outlook will summarize the advancements in this field, with emphasis on the potential of pertinent applications.

2. Design strategies and synthetic challenges

The construction of MOFs from MBBs has facilitated the process of design and has set necessary conditions for the assembly of targeted networks. In particular, carboxylate-based metal clusters have proven effective at generating intended MBBs in situ, which has allowed access to expected, as well as novel, frameworks. Indeed, by gaining adequate control over these design tools, a new generation, an array, of novel materials has been pursued and detailed. Among metal–organic assemblies, primary emphasis has been placed on 3-periodic nets due to their potential for applications. Analysis of the literature occurrences of 3-periodic MOFs supports that the most prevalent framework types are based on 4-connected nodes, such as dia, nbo, eds, and lv (none of which are zeolitic). These reference three-letter codes are generally associated with the structural features/building blocks of particular networks, as implemented by O’Keeffe. In the context of this review, the topological identity of inorganic zeolites will be identified with uppercase three-letter codes (e.g., RHO), as implemented by the IZA, while their metal–organic analogues will be expressed by the corresponding bold lowercase three-letter code (e.g., rho). It should be noted that there are some cases where the three-letter codes are not the same for the IZA and the RCSR (O’Keeffe) analogues (e.g., BCT and crb, respectively).

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The rational construction of 4-connected, specifically tetrahedrally connected, porous materials, related in their topology and function to zeolites, with enlarged cavities and periodic intra- and/or extra-framework organic functionality, is an ongoing synthetic challenge, and it is of exceptional scientific and technological interest. The large and extra-large cavities offer great potential for innovative applications (serving as nano-reactors, becoming a platform for a variety of alternative applications pertaining to large molecules, nanotechnology, optics, sensor technology, and medicine, for example), enhancing...
the correlation between structure and function. Within the last few years, MOFs with zeolitic topologies (ZMOFs)[15] have become a major focus for research groups in the materials chemistry community, who are particularly interested in the attractive properties associated with this unique subset of MOFs. Of particular interest, these materials inherently lack interpenetration (a feature that often affects the pores of would-be open MOFs); hence the accessibility to their porous channel systems is fully exploitable.

Expansion and/or decoration of tetrahedrally connected open networks, specifically zeolite-like nets, using the MBB approach provides the material designer with a prospective method to systematically construct functionalized porous materials with tunable and enlarged cavities by decorating the net and/or expanding the edges with a longer linker, or by substituting the tetrahedral vertices with larger super tetrahedral building units. To date, the synthesis of zeolite-like MOFs has proven to be more than trivial, as the complexity associated with these structures cannot be easily overcome. Moreover, the assembly of simple tetrahedral nodes correlates most often with the formation of the aforementioned dia (cubic diamond) topology, the so-called “default” structure for this type of node, which is not zeolitic.[45]

Therefore, multiple routes have been explored for targeting “smarter” tetrahedral building blocks, associated with the intended angle of connectivity in order to access non-default nets, and furthermore to generate MOFs with zeolite topologies. Amongst MOF analogues to zeolites, the sodalite (SOD, sod) net has the highest occurrence, as the structure accommodates a wide range for the T–O–T angle.[38] Over the years, other MOFs with zeolite-like topologies have been synthesized, such as aco,[46] ana,[47,48] crb (BCT),[47,49] dtt,[47,49,62–64] gis,[47,49,59,65–70] gme,[47] lta,[47] mer,[47,49] mtn,[72–75] sod,[47,49,76–80] and rho,[47,49,81–88] but only recent studies consider an in-depth, systematic approach for the construction of these materials. Of those zeolitic networks targeted, the number of experimentally encountered frameworks can be considered limited. In order to access a larger number of zeolite-like frameworks, including unrevealed (e.g., hypothetical) topologies; it is necessary to consider multiple variables, including SDAs and the nature of the tetrahedral or supertetrahedral building blocks, along with the angularity/additional functionality of the organic component. Theoretically, the number of possible structures to construct with these set conditions is yet vast, as reflected in the high number of zeolite-like networks from hypothetical databases.[89]

3. The edge-expansion approach to zeolite-like metal–organic frameworks (ZMOFs)

3.1. ZMOFs from angular ditopic N-donor ligands: pyrimidine-, imidazole-, and tetrazole-based linkers

The aim to construct functional hybrid solid-state porous materials with topologies akin to inorganic zeolites has been pursued by implementing a top-down, bottom-up approach. That is, by deconstructing the nets into small components, it is revealed that, as mentioned above, the materials are built from cationic, corner-sharing tetrahedra (T), bridged by an O²⁻ anion (with an average T–O–T angle in the range of ~141°). “Edge-expansion” refers to a principle that consists of replacing the oxide ion with an organic functionality that preserves the intended angle of connectivity, and that is capable of rendering a material with similar features, only on an expanded scale.[88]

The original strategy is based on choosing single-metal ions with preferred tetrahedral geometry, in combination with angular ditopic N donor organic ligands. Such candidates have been aromatic nitrogen heterocycle-based linkers such as imidazole, pyrimidine, or tetrazole molecules, and some relevant paradigms are briefly outlined in this section.

One of the earliest examples is reflected in the work of Keller, in 1997,[90] where the potential offered by pyrimidine ligands to afford crystalline materials with structures and properties related to porous inorganic materials is considered. In this instance, the compound reported is based on tetrahedral copper(II) centers coordinated by four pyrimidine molecules, and where BF₄⁻ anions are ensuring the charge balance of the assembly. The 3-periodic net has pcl (paracelsian) topology, consisting of 4-, 6- and 8-MRs, possessing structural features closely related to the ones observed in the feldspar material family, a group of silicate minerals. This early reference is of great importance, as it clearly delineates the use of intended organic ligands as potential mediators for the synthesis of metal–organic analogues of zeolites.

The same year brings a report of an interesting material also derived from a pyrimidine derivative, namely 2-amino-5-bromo-pyrimidine, yielding an early relevant example of a MOF with a true zeolite-like topology. The structure consists of copper(n) metal ions, with slightly distorted tetrahedral geometry, that are bridged by bromide ions, in addition to coordination to the organic moieties (i.e., each copper forms two coordination bonds with two nitrogen atoms of the pyrimidine ligands, along with two other bonds with two bromide anions). The resulting 3-periodic framework possesses crb (BCT) zeolitic topology (Fig. 2) with unidimensional channels consisting of alternating cavities, one in which the amino groups are pointing to its interior (5.223 Å point to point, not considering the van der Waals (vdW) radii of the nearest atoms), while the other has the corresponding bromo-functional groups positioned inside the rather inaccessible cavity (3.695 Å point to point).

Given these guidelines, subsequent paradigms outline derivations of the approach described above. One example of a MOF with sod topology was reported by Tabares et al. in 2001, where 2-hydroxypyrimidine (2-Hpymo) organic ligands are bridging square planar copper(n) metal ion centers to construct the 3-periodic net.[79] Although the framework exhibits neutral characteristics, the authors delineate the selectivity of the material with regard to the salt or the ion pair, AX (where A = NH₄⁺, Li⁺, K⁺, or Rb⁺ and X = ClO₄⁻, BF₄⁻, or PF₆⁻), recognition. The flexibility of the material is also mentioned, as it undergoes a reversible phase transformation, from a rhombohedral to a cubic phase upon immersion in a MeOH–H₂O solution.[79]
Interactions at low temperatures of methylimidazolate and triazolate complexes.\(^{52}\)

Interestingly, one of the compounds reported, based on an imidazole derivative, yields a 3-periodic ZMOF. Reaction between ferrocene and 2-methylimidazole results in tetrahedral iron-met-metal ion nodes, which, in conjunction with the organic linker, afford a material with crb (BCT) zeolitic topology. The unperiodic channels accommodate ferrocene molecules.\(^{52}\)

Furthermore, in 2001, Sironi and co-workers reported a series of polymorphs constructed from copper and imidazole. Among the supramolecular isomers, with 1:2 metal to ligand stoichiometry, a compound with sod topology (Fig. 4) is accounted.\(^{78}\)

The framework exhibits small unidimensional channels, 7.8 Å \(\times\) 5.8 Å, distances measured point to point without considering the vdW radii of the nearest atoms.

Although valuable examples, the two compounds detailed above were not deliberately targeted as conceptual means to yield materials that mimic inorganic zeolite topologies. However, soon after, in 2002, Lee et al. emphasized the importance of the geometric attributes of this linker, and its capability to yield zeolite-like MOFs.\(^{92}\)

Their work evidences the potential offered by imidazolates to facilitate the synthesis of non-default MOFs based on tetrahedral nodes. The authors report on the synthesis of a compound derived from tetrahedral cobalt(II) metal ions, coordinated by four imidazole units, resulting in a 3-periodic net with nog topology. In spite of possessing zeolite-like features, the structure is not very open and its functionality is limited as a result of its structural features, an observation also valid for the previously discussed imidazolate-based MOFs.

Later studies conducted by Yaghi and co-workers,\(^{47,49}\) among others,\(^{48,63,82}\) further reinforce the ability of imidazolato-based linkers to yield MOFs with zeolitic topologies and properties. Synthetically, a challenge is associated with the corresponding frameworks; they are neutral, which affects the reaction environment by limiting the ability to utilize SDAs, thus limiting the variety of the zeolite-like topologies that can be derived solely from imidazole. An alternative route in favor of structure diversity is portrayed by linker functionalization (i.e., imidazole derivatives).

In accordance with this approach, Yaghi et al. report on the synthesis of various ZMOFs, specifically referred to as zeolitic imidazolate frameworks (ZIFs), including ana, crb (BCT), dft, gme, gir, ita, mer, sod and rho.\(^{93}\) The overall topological features resemble the ones encountered in the traditional inorganic zeolites, however on a larger scale, as a result of the aforementioned edge-expansion (i.e., replacing the O ion with the angular imidazolate organic ligand).

When using benzoimidazole, two materials with zeolitic sod and rho topologies are obtained; however, by replacing the carbon atom with a nitrogen atom in the 4-position of benzi-midazole, the ubiquitous diamond structure is favored, which highlights the difficulty of synthetically avoiding this default topology. Conversely, by replacing the carbon atom(s) with a nitrogen atom(s) in 5- or 5- and 7-positions on benzimidazole, a framework with ita (LTA, Linde type A, or zeolite A) topology is constructed, consisting of two types of cages, truncated cuboctahedra (2 cage) and truncated octahedra (β cage) (Fig. 5).\(^{74}\)
In contrast, this result demonstrates the potential effectiveness of organic ligand functionalization, as access to hierarchically complex structures with more than one type of cage remains a challenge. The inorganic LTA material has an internal free diameter of 11.4 Å, while in its metal–organic analogue, it increases to 15.4 Å. The material exhibits accessible porosity as evidenced by Ar, H₂, CO₂, and CH₄ gas sorption studies, possessing an estimated Langmuir surface area of 800 m² g⁻¹. Hydrogen, carbon dioxide, and methane sorption studies were performed, and the potential for gas separation (CO₂–CH₄ mixtures) has also been investigated.

Additionally, this approach allows access to compounds possessing unprecedented zeolite-like features and extra-large cavities.
Two examples of such materials are ZIF-95 and ZIF-100, generically termed \textit{poz} (ZIF-95) and \textit{moz} (ZIF-100).\textsuperscript{94} The first compound encompasses four different types of cages, two having impressive dimensions: \textit{poz} A with accessible pore sizes of 25.1 Å $\times$ 14.3 Å and \textit{poz} B, 30.1 Å $\times$ 20 Å; similarly, \textit{moz} is constructed from cages that have up to 35.6 Å internal exploitable voids. Their estimated Langmuir surface areas are 1240 m$^2$ g$^{-1}$ and 780 m$^2$ g$^{-1}$, respectively, values much larger than the ones encountered for zeolite materials. The evaluation of the gas sorption related properties revealed that both materials selectively retain CO$_2$ in the pores in 50 : 50 CO$_2$/N$_2$, CO$_2$/CH$_4$, or CO$_2$/CO mixtures.

To access additional ZIF structures,\textsuperscript{95} high throughput synthesis, a method inspired by similar techniques for testing pharmacology products, was implemented.\textsuperscript{95} Introducing mixed organic ligands in the synthesis leads to the complementation of the conventional synthetic approaches. This approach has permitted the production of a multitude of materials based on tetrahedral nodes, including the default diamond structure, along with desired zeolite-like compounds, some previously synthesized through standard methods.

Another way to construct new imidazole-based zeolitic MOFs is to take advantage of the easy formation of tetrahedral M$^{2+}$-imidazole chemical bonds to design new imidazole-based ligands that can react as larger building blocks. Sun \textit{et al.} reported a Co-based ZMOF built up from a novel, rigid 3-imidazole-containing ligand, 1,3,5-tri(1H-imidazol-4-yl)benzene. The structure is a binodal, (4,4)-connected (i.e., (Co-ligand)-connected), porous net displaying a zeolitic \textit{bct} topology.\textsuperscript{96}

A similar concept was used to develop a “lightweight” version of ZIFs, referred to as zeolitic boron imidazolate frameworks (BIFs). Zeolite-like nets were targeted from predetermined tetrahedral boron-imidazolate complexes (from imidazole or imidazole derivatives).\textsuperscript{97} These complexes are synthesized prior to the MOF process and then are further linked by monovalent cations (such as Li and Cu) into extended nets. For the creation of four-connected topologies from these complexes, the authors used a strategy similar to the one that led to the discovery of microporous AlPO$_4$ by analogy with porous silica. Just as Al$^{3+}$ and P$^{5+}$ ions can replace two Si$^{4+}$ sites in a porous silicalite, Li$^+$ and B$^{3+}$ can replace two Zn$^{2+}$ sites in a Zn(im)$_2$ ZMOF framework (Fig. 6). The strategy affords materials with zeolitic topologies (\textit{sod}), but also other types of 4-connected nets.\textsuperscript{98} In some cases, the boron-imidazolate precursors are 3-connected, resulting in mixed 3,4-connected nets, or materials solely based on nodes of 3-connectivity. It is worth mentioning that Leoni \textit{et al.} have predicted and studied the stability of 30 topologically different BIFs by DFT calculations, and have concluded that the \textit{fau}, \textit{rho}, and \textit{gme} nets are the most promising candidates for hydrogen storage applications.\textsuperscript{99}

One drawback of BIF materials comes from the short B–N distance bond ($\sim$ 1.5 Å) that implies a closer contact and stronger steric repulsion between imidazolate bridges, making the tunability...
of the framework challenging. Hence, Feng et al. have developed a new series of materials based only on 4-connected lithium nodes (Li–N ~ 2.0 Å) using a mixed-ligand strategy.\textsuperscript{100} It should be noted that the total charge of the resultant 4-connected framework would be negative, if B\textsuperscript{17}/Li\textsuperscript{17} is replaced with Li\textsuperscript{17}/Li\textsuperscript{17} and no change is made in the imidazolate ligands.\textsuperscript{100} Accordingly, half of the negatively charged imidazolate ligands were replaced by neutral ligands, giving rise to new materials, named MVLIF-1 and MVLIF-2 (MVLIF stands for mixed valent ligand lithium imidazolate framework), that display non-zeolitic 4-connected topologies, qzd and dia, respectively.

As mentioned for some of the imidazolate-based ZMOFs, and evidenced by the incredible number of studies reported in this field, one of the most promising applications for MOF materials is gas adsorption. Long et al.,\textsuperscript{101,102} among others,\textsuperscript{103,104} have developed a series of sod-like materials from yet another promising type of N-donor ligand, a tetrazolate (in this case, 1,3,5-benzenetristetrazolate linker that also incorporates an imidazole-like, triazole,\textsuperscript{107} and BTC (and expanded) derivatives.\textsuperscript{108–113}

Recently, important efforts have been dedicated to developing new materials for the capture and storage of greenhouses gases, such as CO\textsubscript{2}.\textsuperscript{114–116} Many strategies to enhance carbon dioxide adsorption have been introduced, such as the use of coordinatively unsaturated metal centers,\textsuperscript{40,117} the optimization of the pore size,\textsuperscript{118,119} or incorporation of alkyamines.\textsuperscript{107,120–122} A third approach to increase CO\textsubscript{2} adsorbent amounts is the presence of amine-functionalized aromatic linkers.\textsuperscript{123,124} Lan et al. have investigated the impact of the utilization of a N-rich aromatic ligand (without NH\textsubscript{2} groups) by fabricating a sod-ZMOF (Fig. 7) based on another tetrazolate linker that also incorporates an imidazole-like, triazole core (4,5-di(1H-tetrazol-5-yl)-2H-1,2,3-triazole); this zeolitic framework demonstrates the achievement of high uptake capacity for CO\textsubscript{2}, even in the absence of primary amines and open metal sites.\textsuperscript{125}

To conclude this subchapter, it becomes apparent that the use of non-linear N-based ditopic or polytopic linkers, in conjunction with appropriate metal ion coordination geometry, successfully qualifies for the synthesis of MOFs with structures and functions closely related to zeolites. The identity of the linker, accompanied by various functional groups, influences the structural diversity and tunability of the materials.

### 3.2. ZMOFs constructed from the single-metal-ion-based MBB approach

Meanwhile, another approach to zeolite-like metal–organic frameworks was developed by our group, implementing a single-metal-ion-based MBB approach, which focused on the introduction of a higher degree of information at the single-metal ion level, which is crafting “smarter” predetermined building blocks.\textsuperscript{15,86} The concept involved the use of organic ligands, like and including imidazolates, that have angular N-donors, but also must include secondary donors, such as O-donors; together the N- and O-donors form heterochelating moieties (e.g., the nitrogen atom is positioned on the aromatic part of the ring, having carboxylates located in the x-position relative to the nitrogen).

The main advantage of this approach as compared to solely carboxylic acid or nitrogen-based ligands is the rigidity and directionality reinforced by the chelating ring that locks the metal in its position and maintains the geometric requirements that facilitate the design of targeted frameworks. Within the net-to-be, the nitrogen atoms direct the topology (i.e., angular), while the carboxylates lock the metal in its position. Hence, the main attributes of this approach are the rigidity and the directionality embedded in these single-metal-ion based MBBs, which preserve the geometric specificities of the organic ligands utilized.
The polytopic nature of such ligands has the leading role as to fully saturate the coordination sphere of the single-metal ion, in such a way that it precludes coordination of unwanted solvent or guest molecules. The hetero-functionality provided by the organic ligands results in the generation of MBBs of the type MN\(_x\)(O\(_2\)C\(_y\))\(_z\) (where \(x\) represents participating angular N donors, typically involved in a ring of chelation, while \(y\) is translated to the additional carboxylate functionalities (often O-chelating) that complete the coordination sphere at the available metal sites; \(M\) is typically a 6-, 7-, or 8-coordinate metal ion).\(^{126-128}\)

As such, the single-metal-ion-based MBB approach was quickly realized as a suitable method for targeting ZMOFs.\(^{15,87,88}\) In addition to those zeolitic MOFs mentioned above, our ZMOFs represent a unique subset that is not only topologically related to the purely inorganic zeolites, but also exhibits similar properties:

(i) forbidden self-interpenetration, which permits the design of readily accessible extra-large cavities;

(ii) chemical stability, where the structural integrity is maintained in water (a feature not commonly encountered in MOFs), and allows for ZMOF applications for heterogeneous catalysis, separations, and sensors, especially in aqueous media;

(iii) anionic ZMOFs possess the ability to control and tune extra-framework cations toward specific applications such as catalysis, gas storage, the removal/sequestration of toxic metal ions, etc.

As with some of the previously mentioned approaches, this method involves edge-expansion of zeolite-like nets toward the design and synthesis of very open ZMOFs. The key factors are related to the ability to generate rigid and directional single-metal-ion-based MBBs that serve as the tetrahedral nodes (T) or tetrahedral building units (TBUs), which are to be positioned and locked at the intended angle, through the aid of the pre-designed heterofunctional organic ligands (concept schematically depicted in Fig. 8).

Therefore, non-default structures for the assembly of TBUs, such as zeolites, can be more easily targeted by judicious selection of the appropriately shaped rigid MBBs and linkers. Consequently, it is evident that introducing information into the MBB is vital, and it is of broad interest to use the MBB approach, based on rigid and directional single-metal-ion TBUs, as a solid platform and basis for developing new design strategies to construct and functionalize novel ZMOFs for specific applications.

Our approach allows for the preferential targeting of anionic ZMOFs, allowing for the utilization of different SDAs, suggesting that this strategy has little limitations in terms of the range of materials that it can generate. As in zeolite systems, the role played by SDAs in MOF systems enhances their potential for diversity, as has been previously demonstrated with the synthesis of supramolecular isomers derived from indium metal ions, 2,5-pyridinedicarboxylic acid (H\(_2\)PDC), and different SDAs: a discrete octahedron,\(^{126}\) a 2D layered Kagomé structure,\(^{126}\) and a 3D diamondoid net.\(^{127}\) The same method has been utilized to target other metal–organic polyhedra (MOPs), like metal–organic cubes.\(^{128-130}\)

In addition, in contrast to most zeolites, and along with green chemistry concepts, the synthesis of our ZMOFs is performed under mild conditions, which also permits the conservation of the structural integrity of the organic components.

Based on the single-metal-ion MBB conditions (possessing both desired angularities and heterofunctionality), imidazolodiacarboxylates and pyrimidinecarboxylates represent potential attractive candidates for targeting the desired ZMOFs.\(^{15,87,88}\) From a metal ion choice perspective, those metals that have primarily six to eight available coordination sites are targeted (although a higher number of sites can be utilized as well), ions that should allow the formation of the intended building blocks of the type MN\(_4\)(O\(_2\)C\(_2\))\(_2\), MN\(_2\)(O\(_2\)C\(_2\))\(_4\), MN\(_4\)(O\(_2\)C\(_2\))\(_6\), or MN\(_2\)(O\(_2\)C\(_2\))\(_8\), to ultimately render MN\(_4\) or MN\(_2\)(O\(_2\)C\(_2\))\(_2\) directing units, all capable of being translated into TBUs.

According to these criteria, 4,5-imidazolodiacarboxylic acid, H\(_4\)ImDC, is well-suited to target MOFs with zeolite-like topologies, since it concurrently possesses two N\(_\text{3D}-\text{O-hetero-chelating moieties with a potential angle of 144}^\circ\) (directed by the M–N coordination). Additionally, if four HImDC ligands saturate each single-metal ion coordination sphere (divalent or trivalent), an anionic ZMOF can be realized. As in the 2,5-H\(_2\)PDC-based supramolecular isomers mentioned previously, the anionic nature allows for the utilization of cationic SDAs, as well as the exploration of applications akin to traditional zeolites (e.g., ion exchange).

A reaction between In(NO\(_3\))\(_3\)\(_2\)H\(_2\)O and H\(_4\)ImDC, in the presence of different SDAs does, in fact, yield different ZMOFs (Fig. 9).\(^{15,88}\) Specifically, imidazole (HIm) leads to a sod-ZMOF, while 1,3,4,6,7,8-hexahydro-2H-pyrimido[1,2-\(a\)]pyrimidine (HP) yields rho-ZMOF-1, where both materials possess volumes up to 8 times larger than their inorganic analogues. In the In-HImDC sod-ZMOF-1, each 6-coordinate In\(^{3+}\) ion is hetero-chelated by
two HImDC\(^{2-}\) ligands and coordinated by the ancillary nitrogen-donor from two other HImDC\(^{2-}\) ligands, resulting in the desired InN\(_{4}(O_{2}C-)\)\(_{4}\) MBBs (InN\(_{4}\) TBUs). In rho-ZMOF-1, each single-indium ion is 8-coordinate, saturated by the hetero-chelation of four HImDC\(^{2-}\) ligands to give the intended InN\(_{4}(O_{2}C-)\)\(_{4}\) MBBs (InN\(_{4}\) TBUs). This anionic rho-ZMOF-1 was the first material ever to contain an organic component and have a zeolite RHO topology. Unlike inorganic RHO zeolite and other RHO analogues, rho-ZMOF-1 requires twice as many positive charges, 48 (24 doubly protonated HPP molecules) vs. 24, to neutralize the anionic framework. Also, the extra-large cavities can accommodate a sphere of 18.2 Å in diameter inside each cage, outlining a benefit from edge-expansion of the aluminosilicate analogue evident by the doubling (3.10 nm vs. 1.51 nm) of the unit cell. In addition, the sod-ZMOF-1 represents the first example of a MOF with an anionic framework based on the sod topology, although some other examples of neutral or cationic sodalite-like MOFs have been synthesized previously, as detailed above.\(^{15,48}\)

Additionally, our group’s findings lead to the discovery of a novel zeolite-like net, usf-ZMOF (Fig. 10),\(^{131}\) with an unprecedented topology at the time of its synthesis; independently, the topological data were reported in one of the hypothetical zeolite databases by the time of publication, and now it is referred to in the RCSR database as med topology. The synthetic protocol involves similar reagents as for sod-ZMOF-1 and rho-ZMOF-1 detailed above, yet in the presence of a different SDA, 1,2-diaminocyclohexane (1,2-H\(_{2}\)DACH). Each indium metal ion is coordinated to four nitrogen atoms and four oxygen atoms of four separate HImDC ligands, respectively, to form an eight-coordinate MBB, InN\(_{4}(O_{2}C-)\)\(_{4}\) (InN\(_{4}\) TBUs). The anionic nature of usf-ZMOF is neutralized by 40 doubly protonated 1,2-DACH molecules, accommodated by a unit cell with a volume (45 245 Å\(^3\)) that is 9.55 times higher than its analogous yet-to-be-constructed zeolite (4735 Å\(^3\)).

Given the anionic nature of our ZMOFs, a diverse range of applications is exploitable. The zeolite-like nature favors facile ion exchange capability of the organic cations in the cavities. To demonstrate, rho-ZMOF-1 was utilized, and it was found that the counter cations can be fully replaced at room temperature after 15 to 24 hours (depending on the inorganic cation used); the fully exchanged compounds retain their morphology and crystallinity. In a recent study, the effect of several extra-framework cations (as-synthesized sample, containing dimethylammonium cations, ...
DMA$^+$, and the ion-exchanged Li$^+$ and Mg$^{2+}$ samples) on the H$_2$
sorption energetics and uptake is reported.$^{132}$

Findings reveal that molecular hydrogen in ion-exchanged
ZMOFs (Fig. 11) clearly demonstrates that the presence of an
electrostatic field in the cavity is largely responsible for the
observed improvement in the isosteric heats of adsorption in
these compounds, by as much as 50%, relative to those in neutral
MOFs. The extra-framework cations are fully coordinated by aqua
ligands, and are not directly accessible to the H$_2$ molecules; thus,
open-metal sites do not contribute to a dramatic increase in the
overall binding energies. However, these results are promising
and may be regarded as the first of several steps towards
improving binding energies to values around 15–20 kJ mol$^{-1}$.

ZMOFs offer great potential for reaching this goal by tuning
the accessible extra-framework cations and/or introducing
open-metal sites, along with a reduction in pore size and
functionalization on the organic links.

At the same time, the large pores of ZMOFs are well-suited to
adsorb not just metal ions, but also larger molecules, like cationic
fluorophores for sensing-related applications, for example. The
double eight-member ring (d8R) cages of In-HImDC rho-ZMOF-1
represent ~9 Å windows that allow access to the extra-large
cavities, $\alpha$-cages, with an internal diameter of 18.2 Å. The cationic
fluorophore, protonated acridine orange (AO), is of the appro-
priate size, and can be diffused into the $\alpha$-cage cavities, where the
electrostatic interactions with the framework preclude further
diffusion of cationic AO out of the cavities/pores, essentially
anchoring the fluorophore (Fig. 12).$^{15}$ The extra-large dimensions
allow for the diffusion of additional neutral guest molecules,
and the anchored cationic AO is utilized to sense a variety
of neutral molecules, such as methyl xanthines or DNA nucleo-
side bases.$^{15,88}$

Adsorption of large molecules for sensor applications in
In-HImDC rho-ZMOF-1 opens up the possibility of evaluating
its extra-large cavities as hosts for large catalytically active
molecules, and its effect on the enhancement of catalytic
activity. In recent studies, the successful encapsulation of free
base porphyrin [H$_2$TMPyP][p-tosyl]$^{4+}$ was probed, followed by
post-synthetic metallation by various transition metal ions to
produce a wide range of encapsulated metalloporphyrins (Fig. 13).$^{133}$

The catalytic activity assessment consists of cyclohexane oxidation,
performed in the presence of Mn-TMPyP. After 24 h, based on
the amount of oxidant present in the initial reaction mixture, a
total yield (from cyclohexane to cyclohexanol/cyclohexanone)
of 91.5% and a corresponding turn over number (TON) of 23.5
(catalyst loading of 3.8%) are noted, a noticeably higher yield
compared to other systems of supported metalloporphyrins
(e.g., zeolites or mesoporous silicates).

More recently, the single-metal-ion-based MBB approach to
design and synthesize a variety of ZMOFs has been successfully
supported by work conducted with other angular hetero-functional bis-chelating bridging ligands, namely pyrimidinecarboxylates. Reaction between In(NO₃)₃·C₅H₂O and 4,6-pyrimidinedicarboxylate (4,6-PmDC) under hydro-solvothermal conditions yields another anionic sod-ZMOF, (Fig. 14 left), from InN₄(O₂C–)₄ MBBs (InN₄ TBUs). Reaction between 2-cyanopyrimidine (where the 2-pyrimidinecarboxylate (2-PmC), was generated in situ) and Cd(NO₃)₂·C₄H₂O, in the presence of piperazine (Pip) and under hydro-solvothermal conditions, produces another rho-ZMOF, (Fig. 14 right), from CdN₄(O₂C–)₄ MBBs (CdN₄ TBUs).

The inherent properties of these ZMOFs allow for the evaluation of a breadth of properties. Full ion exchange in water based solutions at room temperature over short periods of time (less than 24 hours) was conducted on the anionic In-PmDC sod-ZMOF-2 (results confirmed by atomic absorption). The structural integrity is maintained as evidenced by PXRD analysis. Gas sorption (H₂, Ar, N₂) studies were also performed, evidencing accessible porosity with apparent Langmuir surface areas estimated to be 616 m² g⁻¹ for In-PmDC sod-ZMOF-2, and considerably higher for Cd-PmC rho-ZMOF-2, 1168 m² g⁻¹. The deliberate enhancement of the framework-hydrogen interactions is an ongoing challenge. In this context, an increase in the isosteric heat of adsorption was observed in anionic ZMOFs, due to the presence of high local charge density. These unique compounds are therefore suited to outline a functional platform for investigation of the effect of pore size and charge, as well as the effect of various extra-framework cations, upon the hydrogen isosteric heats of adsorption and uptakes.

Another advanced single-metal-ion-based strategy that may allow access to new, open ZMOFs involves the use of chelating carboxylate ligands (L), where, specifically, three-member rings are preferentially produced in situ. It has been theorized that this specific three-member ring building unit, which has the smallest possible ring size, can be utilized to create new topological types with low framework density, and is highly recommended in developing strategies for construction of new porous materials.

Along this route, Bu et al. reported the use of the MBB approach, with indium-based TBUs, to target new zeolitic MOFs built up from In₃L₃ three-member rings by using a mixed-ligand strategy. Indeed, mixed dicarboxylate linkers with 120° and 180° angles were employed with indium ions to generate a series of 3D crystalline porous materials (CPM) displaying zeolitic npo topology. The structure of CPM-2-NH₂ consists of three crystallographically independent In³⁺ ions coordinated by four bidentate carboxylates (InO₈ to give InO₂C–¢₄ TBU) from two bent dicarboxylate ligands (L) and two biphenyl dicarboxylate (bpdc) ligands (L'), giving rise to triangular (In₃L₃) groups acting as SBUs. Each SBU is connected to six neighboring SBUs through six linear ligands resulting in the formation of a 3D zeolitic npo framework (Fig. 15). The MBB approach based on rigid and directional single-metal ion TBUs derived from heterochelating organic ligands, as well as chelating ligands, has been proven to be an effective and versatile strategy for the construction of ZMOFs. The resemblance to traditional zeolitic materials is outlined from a topological viewpoint, as well as in the anionic nature and synthesis (use of SDAs), and properties related to these materials. Furthermore, the superior ability of ZMOFs to fine-tune was proven, as the pore size, charge density, and surface areas were readily altered, leading to the development of a solid platform for relevant applications.
Fig. 13 Schematic representation of the encapsulation and metallation of the [H₂TMPyP]⁺⁺ porphyrin ring enclosed in the \textit{rho}-ZMOF-1 ρ-cage (left); the pink spherical cage schematically represents the average localization of the encapsulated porphyrin in the \textit{rho}-ZMOF-1 ρ-cage. Catalytic activity from cyclohexane to cyclohexanol/cyclohexanone in the presence of the ZMOF-metalloporphyrin (right).

Fig. 14 Pyrimidinecarboxylate-based ZMOFs: sod-ZMOF-2 (left) and rho-ZMOF-2 (right).
4. ZMOFs derived from metal–organic cubes (MOCs)

An alternative route toward intended ZMOFs is probed by means of the introduction of a superior level of built-in information prior to the assembly process, beyond even single-metal-ion-based MBBs, a condition made possible by utilizing MOPs as supramolecular building blocks (SBBs), as previously demonstrated by our group. Among others, a suitable approach to access ZMOFs is based on the assembly of metal–organic cubes (MOCs), which resemble d4R composite building units in inorganic zeolites. By utilizing the previously delineated single-metal-ion-based MBB approach, our group reported the synthesis of a metal–organic cube, MOC-1. In this robust assembly, the ditopic heterofunctional imidazoledicarboxylate ligands represented the edges of the cubes, while the single-metal-ion MBBs occupied its vertices.

The unique structure of this MOC, among others, leads to peripheral functional groups/coordination sites, which offer the potential for external coordination to metal ions and/or hydrogen bonding. In this context, MOCs are equivalent to the d4R composite building units in zeolites, and hence could also be regarded as suitable SBBs to construct structures based on the connection of d4Rs. Specifically, these MOC-based SBBs are suitable for the targeted synthesis of non-default nets, such as some of the 8-connected edge-transitive nets (where the cube is the vertex figure of an 8-connected node, and edge-transitive refers to the fact that structures contain only one type of edge). Moreover, the augmentation of edge-transitive nets, such as buc, flu, scu and reo, outlines a close relationship with some corresponding zeolite-like frameworks, namely, ACO, AST, ASV, and LTA. The MOC SBBs can be linked to form extended frameworks in two different ways (as depicted in Fig. 16): (1) through linear connections, where nets based on zeolites ACO and LTA can be accessed or (2) cross-linked through additional 4-connected (tetrahedral) nodes, to result in materials with AST and ASV-like zeolitic topologies.

Indeed, reaction of In(NO₃)₃·5H₂O with 4,5-dicyanoimidazole (4,5-DCIm) in a DMF solution and in the presence of pip affords pale yellow homogeneous microporous material with dendrimer-like morphology, referred to as aco-ZMOF or MOC-2. The cubes are linearly connected vertex-to-vertex via intermolecular hydrogen bonds, O–H⋯O, 2.786 Å. Each metal–organic cube concomitantly connects to eight neighbouring cubes through 24 hydrogen bonds, as the oxygen atoms pointing outward of each cube form three intermolecular hydrogen bonds with the corresponding oxygen atoms of the neighbouring cube. Consequently, the periodic arrangement of the discrete molecules results in an open framework that resembles the ACO zeolite topology (Fig. 16 left column).

The framework possesses accessible channels that can accommodate a sphere with a maximum diameter of 11.782 Å, considering the vdW radii of the nearest atoms, and an estimated Langmuir surface area of 1420 m² g⁻¹. The material is exceptionally robust, in the context that the MOCs are solely sustained by hydrogen bonds, and stores up to 2.15% weight H₂ at 77 K and atmospheric pressures.

In addition to the previous example, the solvothermal reaction of Cd(NO₃)₂·4H₂O and H₃ImDC, in the presence of Na⁺ ions, gives rise to a material in which each cube is vertex-to-vertex linearly linked to eight other MOCs. In this assembly, half of the total number of vertices is connected through four sodium atoms, while the other four vertices are bridged by hydrogen bonded water molecules (Fig. 16 middle column). The framework of MOC-4 possesses an overall a structures, where the a cage is based on 12 MOCs, while 6 other cubes gain rise to the formation of the b cage. The largest sphere that can fit into the cavities without interacting with the vdW surface of the framework has an approximate diameter of ~32 Å for the a cage and ~8.5 Å for the b cage.

Examples of tetrahedrally connected MOCs were also achieved experimentally. The reaction between In(NO₃)₃·5H₂O and 4,5-DCIm in an EtOH solution yields colourless polyhedral crystals. The discrete cubes are connected by ammonium cations, which are linking four MOCs in a tetrahedral arrangement through N–H···O hydrogen bonds to generate a structure...
Fig. 16 Prototype scheme for MOC-to-ZMOF linkage: connectivity through (left and middle columns) linear linkers, resulting in aco-ZMOF and lta-ZMOF, as well as through tetrahedral nodes (right column), leading to the assembly of ast-ZMOF.

with the zeolite AST topology, ast-ZMOF-1, where the Langmuir surface area is estimated to be 456 m² g⁻¹.

Another example based on ast topology, ast-ZMOF-2, is afforded by the reaction of H₄ImDC and Zn(NO₃)₂·6H₂O in a DMF–H₂O...
solution, in the presence of excess zinc and guanidinium cations, (Fig. 16, right column). Under these conditions, the MOCs are concomitantly connected through their edges and vertices to zinc and guanidinium ions, respectively, and further extend to twelve adjacent MOCs to yield yet another material with a zeolitic topology and features.

More recently, a systematic investigation using 2,2’-(1H-imidazole-4,5-diyl)di-1,4,5,6-tetrahydropyrimidine and different M²⁺ metal ions was conducted to elucidate the effect of different parameters, such as the nature of metal ions, counterions, solvent systems, solution pH, and temperature, on the nature of isolated MOC products. It was demonstrated that the formation of a given targeted MBB requires the careful choice of reaction conditions, i.e. reactant concentration, temperature, solvent mixture, metal ion, counterion, and pH. A properly chosen solvent system should promote desired ligand-to-metal coordination, as well as facilitate nucleation and crystallization. Counter-ions with limited interference to ligand coordination are vital for the metal–ligand nucleation and growth to yield crystals manageable for characterization. Counter-ions should promote desired ligand-to-metal coordination, as well as facilitate nucleation and crystallization. Counter-ions with limited interference to ligand coordination are vital for the metal–ligand nucleation and growth to yield crystals manageable for characterization.

In summary, this approach has proven to be pertinent to the construction of zeolite-like materials. The strategy conveys a superior route towards novel ZMOFs, outlining a hierarchical pathway initiated from single-metal ions with anticipated coordination geometries, and predesigned to act as rigid and directional vertices, into MOCs that can be utilized as d4Rs, to ultimately result into intended zeolitic frameworks, ZMOFs.

5. ZMOFs built from supertetrahedral (ST) building blocks

Metal–carboxylate clusters generated in situ represent MBBs commonly employed to target MOFs, since they possess metal–oxygen coordination bonds that result in the generation of rigid nodes with fixed geometry. Accordingly, they aid the formation of robust (and possibly, permanently porous) 3-periodic frameworks, as well as hold potential for open (or coordinatively unsaturated) metal sites that are of interest for various applications. Various organic linkers can be employed to connect such clusters that ultimately can result in solid-state materials with pore sizes that may go beyond the microporous regime, which is characteristic of the majority of MOFs.

Metal–carboxylate clusters can also be utilized as a possible pathway to tetrahedral-based zeolitic MOFs; ZMOFs can be attained through access to tetrahedral-like building blocks based on metal–carboxylate clusters, which, in combination with linker connectivity, comprise the necessary requirements to be translated into supertetrahedral (ST) building units. The ST building block can be viewed as the enlarged version of a simple tetrahedral vertex, having, as a final outcome, a material with analogous topology, only on an expanded scale, albeit a different method from the aforementioned edge-expansion.

In this context, investigations carried out by Férey and co-workers resulted in two porous solid-state materials, MIL-100 and MIL-101, that outline the decorated and augmented mtt zeolite topology. The group’s approach is focused on rendering rigid carboxylate-based metal clusters that maintain, unaltered, their pre-designed function throughout the assembly process. Specifically, they utilized building blocks derived from trimeric units constructed from three metal octahedra meeting at a vertex (V), µ₃-O (of the type M₆O₃(C₆H₄), to generate an overall ST building unit, where, in MIL-100, the carboxylates come from trimesic acid, or 1,3,5-benzenetricarboxylate (1,3,5-BTC), ligand (L),...
which lies on the four faces of the ST \([(M_3O)_4L_4\) or V$_4$L$_4\)], while the trimers occupy its vertices. By linking the corners of the STs, the material possesses decorated and augmented mtn topology (by definition, augmentation being referred to as the replacement of each vertex of an N-connected net by N-vertices, i.e., in this case, a tetrahedron by a ST). The structure exhibits two different types of cages: the smaller cage (\(~25\text{ Å in diameter}\)) consists of 20 ST, while the larger cages are built from 28 ST (\(~29\text{ Å in diameter}\)), highlighting an apparent Langmuir surface area of 3100 m$^2$ g$^{-1}$.

Soon after this discovery, Férey et al. reported the synthesis of MIL-101, a material also based on M$^{III}$ trimer inorganic clusters (the same as in MIL-100), but in the presence of a different linker, namely 1,4-benzenedicarboxylic acid (1,4-BDC). Similar to MIL-100, a ST is generated by the arrangement of the trimers on its vertices; however, in this instance, the linker represents the edge of the ST \([(M_3O)_6V_4L_6\)], bridging the vertices in the intended tetrahedral arrangement (Fig. 18). The two types of cages are once again comprised of 20 ST (\(~29\text{ Å in diameter}\)) for the small cage and 28 ST for the large cage (\(~34\text{ Å in diameter}\)), with an exploitable outstanding estimated Langmuir surface area of 5900 m$^2$ g$^{-1}$. Very recently, a series of isoreticular analogues of MIL-100 and MIL-101 showing an exceptional pore size, of up to 68 Å, have been synthesized.$^{145,146}$

Gas (hydrogen, CO$_2$, and CH$_4$) sorption capabilities were evaluated on both MIL-100 and MIL-101.$^{117,147}$ In the context of reducing the effects of greenhouse gases, it was shown that both compounds exhibit remarkable results in storing large amounts of CO$_2$ and CH$_4$ at relatively high pressures (> 5 MPa).
and ambient temperatures. Specifically, at 6 MPa and 300 K, MIL-100 adsorbs 9.5 mmol g\(^{-1}\) CH\(_4\) and 18 mmol g\(^{-1}\) of CO\(_2\) at 5 MPa and 300 K. As expected, MIL-101 performs superior to MIL-100 adsorbing 13.6 mmol g\(^{-1}\) methane at 6 MPa and 300 K and 40 mmol g\(^{-1}\) of CO\(_2\) at 5 MPa and 300 K. Further evaluations illustrate the potential for drug delivery applications. Studies confirmed the uptake and controlled release of ibuprofen (IBU), among others, from the extra-large pores of MIL-100 (apertures of 4.8 Å and 8.6 Å) and MIL-101 (apertures of 12 Å and 16 Å), with 0.347 g IBU per gram of dehydrated MIL-100 and 1.376 g of IBU per gram of dehydrated MIL-101.

Another example pertinent to this approach was reported by Kim et al., where the framework is built from truncated ST formed by four terbium metal ions that constitute its vertices. The arrangement of these units, aided by the linkage provided by the triazine-1,3,5-tribenzoic acid (H\(_3\)TATB), contributes to the generation of a mesoporous material also with mtn zeolite topology. The two mesoporous cages are denoted S (20 truncated STs formed by twelve 5-MR, 39.1 Å) and L (28 truncated STs with twelve 5-MR and four 6-MR windows, 47.1 Å), and generate a Langmuir surface area of 3855 m\(^2\) g\(^{-1}\), after sample activation at 160 °C. High-pressure CO\(_2\) gas sorption studies confirmed considerable amounts of gas stored under these conditions (18 mmol g\(^{-1}\) at ca. 45 bar and ambient temperature).

A variant strategy towards constructing hybrid materials with zeolitic topologies derived from ST-based building blocks was undertaken by a computational driven approach. The concept involves organic linkers (such as 1,4-benzenedicarboxylate and imidazole), where the tetrahedral single-metal ion center is replaced by Zn capped e-type Keggin polyoxometalates (POMs), targeting the assembly of zeolitic polyoxometalate-based metal-organic frameworks (Z-POMOFs). Theoretical studies validate this strategy by predicting a series of structures with zeolitic topologies based on such “exotic” building blocks. Experimentally, only two examples were accessed using this method to date; however, none possesses zeolitic features. In one instance, the default diamondoid net is reported, while a subsequent example exhibits a layered structure. Once again, this approach outlines the difficulty associated with the synthesis of highly porous zeolite-like nets. At the same time, it stresses the importance of concrete design concepts mediated by modelling approaches.

6. ZMOFs constructed via organic tetrahedral nodes

The final method to construct ZMOFs that we will describe is based on a so-called “inverted” approach, in which the organic molecules act as tetrahedral nodes. This concept involves metal ions with various coordination numbers required to act as the angular ditopic linkers, while auxiliary, weakly-coordinating ligands satisfy the remaining coordination sites of the metals. Ultimately, the organic TBUs have to be positioned at suitable angles (average of ~144°) in order to facilitate structures with zeolite-like features.

The four nitrogen atoms in hexamethylenetetramine (HMTA), situated in a tetrahedral arrangement, qualify this molecule as a potential organic tetrahedral node. From the work of Qiu et al., comes an example of a zeotype material with mtn topology (Fig. 19), constructed from the bridging of HMTA ligands via the 2-connected trigonal bipyramidal Cd(II) cations. The resulting cationic 3-periodic net (where the charge balance is ensured by chloride anions) is constructed from two types of cages: the large cage consists of four 6-MR, 12.3 Å × 13.1 Å, and twelve 5-MR, 10.4 Å × 10.4 Å in diameter, while the smaller cage consists of twelve 5-MR. The overall volume of the unit cell is 117 225 Å\(^3\), as compared to 7920 Å\(^3\) of the corresponding inorganic analogue. Anionic exchange of Cl\(^-\) with SCN\(^-\) was carried out, probing favourable retention of the framework profile upon such treatment.

More recently, the design and synthesis of novel tetracarboxylic acid ligands based on isophthalic moieties have allowed Xu et al. to construct a new ZMOF. The so-called 4+4 strategy has been employed using 5-[bis(4-carboxyphenyl)aminol]isophthalic acid, where the central N atom plays a crucial role in the conformation, as well as the indium cation that is well-known to coordinate to four carboxylate groups in a bidentate fashion, In(O\(_2\)CR)\(_4\) (as highlighted previously in this review). The combination of these two kinds of tetrahedral nodes, In(O\(_2\)CR)\(_4\) and the ligand, respectively, leads to the formation of an overall 3D anionic framework.

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Fig. 19 mtn-like framework based on organic (HMTA) tetrahedral nodes; structural and tiling representation of mtn topology with 2B\(^{12}5\)\(^+\) + [5\(^12\)-6\(^1\)] tiles.
adopting a sod topology (Fig. 20). The structure delimits sodalite cages with dimensions of $26 \times 26 \times 8.2$ Å in which are localized DMA$^+$ cations that ensure the overall charge balance of the structure.

7. Summary and outlook

In the continuous endeavour to produce functional materials for targeted applications, MOFs are becoming strong candidates for meeting current societal and technological needs. In this review, the focus was placed on identifying possible strategies and relevant examples of MOFs that possess periodic intra-framework organic functionality and zeolite-like topologies and properties. The portrayed examples demonstrate that complex structures, based on non-default nets, such as zeolite-like MOFs (ZMOFs), not only can be discovered serendipitously or through high throughput methods, but also, more importantly, can be designed and assembled by the rational choice of rigid and directional building blocks containing the required hierarchical information. The presented strategies offer great potential to access complex structures that are not readily constructed from the conventional assembly of simple building blocks, aiding the advancement in the design and synthesis of functional materials.

Solid-state materials with large and extra-large cavities, such as ZMOFs, allow for a multitude of diverse studies that complement areas where traditional zeolite materials encounter limited tunability. Ipso facto, innovative functions are arising from the specificity associated with their porous nature. Most concerted efforts highlight gas sorption/separation studies (e.g., hydrogen storage as a clean energy source for mobile applications, and selective storage and separation of greenhouse gases (CO$_2$ and CH$_4$)). The uptake and controlled release of drugs gear their functions towards other sectors, as well. Recent results demonstrate the ability of anionic ZMOFs to serve as (host–guest)–guest sensors, where the ZMOF portrays a periodic porous platform for fluorescent cations that act as the sensors, as well as supporting catalytic activity mediated by the encapsulation and metallation of metalloporphyrins.

Nonetheless, as the ability to engineer functional porous solid-state materials has been greatly developed through rational design strategies, the actual synthetic process still requires substantial efforts. Ultimately, the main goal regards the ability to precisely construct the desired material for the intended purpose; in perspective, the propensity for vast advancements in materials chemistry based on zeolite-like MOFs is indubitably highly valuable and holds great promise for novel materials and applications.

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Notes and references

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