

REVIEW

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Design, analysis, and application of metal–organic framework derived carbons

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Metal–organic framework-derived carbons (MOF_dCs) have emerged as a rapidly growing class of porous materials over the past 15 years. Inspired by the principles of templated synthesis used in organic chemistry and nanomaterials, MOF_dCs combine the highly tunable features of metal–organic frameworks with the versatility and stability of traditional porous carbon materials. Although many advances in MOF_dC applications have been driven by an exploratory approach, there is a growing body of systematic, hypothesis-driven studies of the underlying chemistry and materials science of MOF_dC structure, design, and structure–property relationships. This review summarizes developments in the systematic design and characterization of MOF_dCs and strategies developed to control the structure of these materials. The role of variables such as pyrolysis temperature, gas environment, and the atomic- and nanoscale structure of the template MOF are described, with particular emphasis on their relationship to MOF_dC structure and applications. By focusing on the fundamental principles underpinning MOF_dC design, this work will lead to more rationally designed MOF_dC materials and greater efficiencies in the development of MOF_dCs for a host of applications.

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Introduction

Porous materials are of growing importance in our society, as we seek highly efficient catalysts and adsorbents for energy storage and environmental remediation. Charcoal, a porous carbon material, has been described in sources from cultures

across the world from as early as 1500 BC and has been used for medicinal applications and water purification for centuries.^{1–4} In the late 20th and early 21st centuries, chemists such as Robson, Yaghi, and Kitagawa introduced a new type of porous material, known as porous coordination polymers or metal–organic frameworks (MOFs).^{5–10} These materials exhibited both permanent porosity and highly tunable structures and represented a significant advance in obtaining a high level of control over the structure of porous materials. Research over the last 25–30 years has expanded greatly on these early reports to produce MOFs that are more stable^{11–17} and have

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various functionalities apportioned throughout the framework structure.^{8,17–22} However, despite the sophistication of these materials, their stability is often poor compared to more established porous materials, such as porous carbon or zeolites, thus limiting their applicability.^{23–27} Consequently, the commercialization of MOFs has been limited to a small number of niche applications and the need for a highly controllable porous material remains.^{26–28}

Templated synthesis is a key strategy by which structural control has been achieved in fields such as organic chemistry, biochemistry, and nanomaterials.^{29–37} As their cost has decreased and their ease of synthesis has improved, MOFs have begun to be used as templates for the design and synthesis of new MOF-derived materials. One example of this is the advent of MOF-derived carbon (MOFdc) materials, which are porous carbons derived from the thermal decomposition of MOFs.^{37–40} Some MOFdc are also decorated with metal species derived from the inorganic components of the MOFs. Compared to their template MOFs, MOFdc are typically much more thermally and chemically robust,^{37,38} which provides greater opportunities for their application under working conditions for electrocatalysis and other industrial chemical processes.^{41–44} Most importantly, the structure of the template MOF can be tuned or altered to impart specific properties into the MOFdc. By encapsulating guests inside the framework, doping the structure with additional metal ions to form bimetallic MOFs, or altering the connectivity or composition of the inorganic or organic building units, the structure of MOFdc can be tuned in a fashion that is unusually controllable for an amorphous material.

Early MOFdc development

Many early MOFdc required the incorporation of additional carbon sources into the template framework to circumvent excessive loss of carbon through combustion. Xu and co-

workers reported one of the first MOFdc, which was synthesized *via* carbonization of a polyfurfuryl alcohol@MOF-5 composite at 1000 °C under an argon atmosphere. Under these conditions, the organic linkers and polyfurfuryl alcohol were transformed into a porous carbon material with a high surface area.³⁹ Due to the high carbonization temperature and the low boiling point of zinc, no metal was retained in the MOFdc structure, although zinc oxide could be observed at lower carbonization temperatures. Notably, the carbon structure contained micropores, mesopores, and macropores. Later research has shown that the micropores in MOFdc structures are typically derived directly from the micropores of the template MOF.⁴⁵ Xu later extended this research to test the application of these materials as electrodes for supercapacitors.⁴⁶

Following this seminal work, other researchers applied similar strategies to generate MOFdc from other template MOFs. Furfuryl alcohol remained a popular additive and was later applied in the carbonization of zeolitic imidazolate frameworks (ZIFs) by several researchers (Fig. 1a).^{47,48} Gao and coworkers studied the incorporation of various different carbon additives, including a phenolic resin, carbon tetrachloride, and ethylenediamine, finding that carbonization of additive-free MOF-5 produced a MOFdc with high porosity, while carbonization of MOF-5 with all three additives produced a more graphitic carbon.⁴⁹

Eventually, MOF carbonization strategies that did not require the inclusion of an additional carbon source were developed (Fig. 1b). Around the time that Xu and coworkers published their MOFdc study, other researchers reported the synthesis of MOF-derived multiwalled carbon nanotubes,



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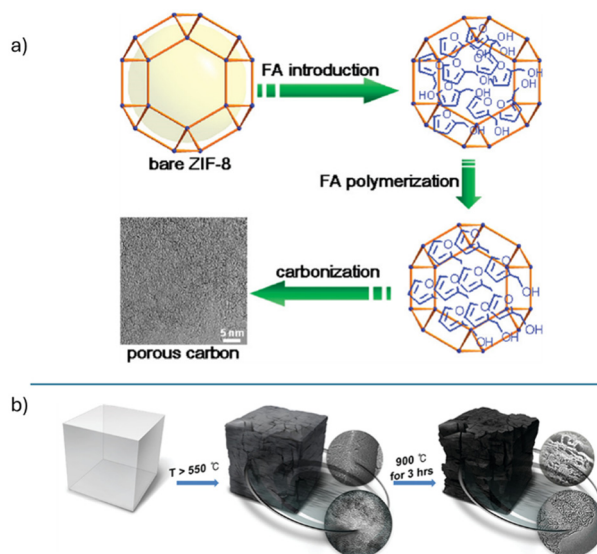


Fig. 1 (a) Incorporation of furfuryl alcohol prior to carbonization increases carbon content in the final MOFdc material. Reprinted with permission from ref. 47. Copyright 2011 American Chemical Society; (b) direct carbonization of MOF crystals can also be achieved. Reprinted with permission from ref. 54. Copyright 2012 American Chemical Society.



where the metal center of the MOF was used to catalyze the formation of the carbon structure.⁵⁰ However, examples of direct MOF carbonizations where the porous structure of the template MOF was retained did not arise until the early 2010s.

One of the earliest examples was the direct carbonization of an aluminium-based MOF by Yamauchi and coworkers.⁵¹ The MOF in question had previously been carbonized with the addition of furfuryl alcohol,⁵² however Yamauchi and coworkers found that the carbon content of the 1,4-naphthalene dicarboxylic acid linker was in fact sufficient to produce a carbon matrix with high porosity after the material was treated to remove the alumina nanoparticles derived from the inorganic building units (IBUs).⁵¹ Notably, the carbonization temperature was a key variable that affected the porosity of the MOFdc, with carbonization at 800 °C producing the MOFdc with the highest surface area. At temperatures above 900 °C, the MOFdc exhibited a drastic reduction in porosity due to graphitization and collapse of the carbon pore structure.

In contrast to the Al-MOF template described above, many MOFdc are formed from Zn-MOFs, as the low boiling point of zinc metal leads to the removal of the metal species without any subsequent processing. Banerjee and Kurungot directly carbonized several zinc-based MOFs and coordination polymers to produce porous carbons that required no post-synthetic treatment due to the evaporation of the zinc. In an effort to maximize carbon content and surface area, this study explored a range of different linkers with varied lengths and aromaticity.⁵³ Park and coworkers also reported the direct carbonization of IRMOFs 1, 3, and 8 (IRMOF = isorecticular MOF). They were able to tune the pore size of the MOFdc and optimize them for hydrogen uptake by altering both the structure of the template MOF and the pyrolysis temperature.⁵⁴

One of the most popular MOF templates for the formation of MOFdc is ZIF-8. This framework is comprised of zinc ions as IBUs and 2-methylimidazolate linkers that provide sufficient carbon content to form highly porous MOFdc without the incorporation of additional carbon sources. Furthermore, ZIF-8 is both inexpensive and synthetically simple to produce in the quantities required for carbonization, as cost is a key limitation to the commercialization of MOFs and MOFdc.^{47,55–59}

As the field of MOFdc has begun to mature, more complex MOFdc structures are being generated. While some MOFdc research remains focused on simple porous carbon structures, many researchers have begun to synthesize MOFdc with N- or S-doped carbons or containing metal species derived from the template MOF.^{47,60} These more complex structures represent a significant advance in the versatility of MOFdc, as they provide several additional structural variables that can be tailored to form materials with specific structures or functionalities.

Structural features of MOFdc

There is a great deal of diversity in the structure of MOFdc, however there are several features that are common to many MOFdc (Fig. 2). First is the presence of a porous carbon

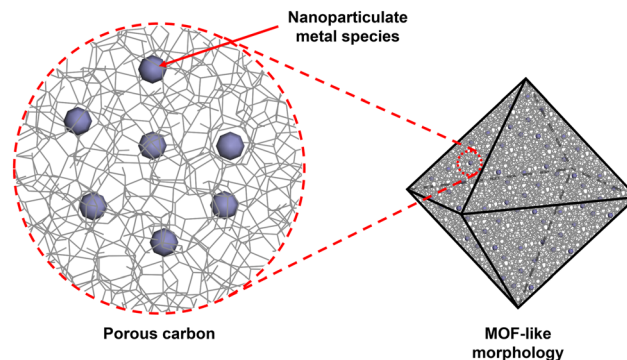


Fig. 2 Schematic of MOFdc structural features showing MOF-like morphology, porous carbon, and nanoparticulate metal species.

matrix, which is a universal feature in these materials.^{46,49,52,61–63} Originally derived from a combination of the organic linker and additional carbon-containing guests, and later from the organic linkers alone, the structure of this porous carbon is strongly related to the structure of the template MOF. The carbon typically contains a mixture of micropores and mesopores, the sizes of which are derived from the pores of the original MOF template.^{45,49,52,64} Pore size distributions of MOFdc obtained *via* gas sorption analysis show that the micropores in the carbon are of the same size as the micropores in the MOF. The mesopores are generated by a combination of combustion of the organic components and *in situ* defect formation leading to partial structural collapse during the carbonization.^{45,64–66} The relative concentrations of micro- *versus* mesopores can be tuned based on the pyrolysis conditions. Harsher carbonization conditions, such as high temperatures or oxidizing gas environments, lead to the formation of greater mesoporosity, as the harsh conditions generate more defects in the MOF during the pre-carbonization heating and encourage a greater degree of combustion of the organic components. Depending on the composition of the organic linker or organic additives, the carbon matrix can also have additional elements doped into the non-graphitic carbon structure. Most common is nitrogen-doping, which is achieved by either including a nitrogen-rich guest like urea in the MOF prior to carbonization^{60,67} or by designing a MOF structure with nitrogen-rich linkers, such as the ZIF-8/ZIF-67 system.⁴⁷ Carbonization of these nitrogen-rich templates produces N-doped carbon, which enhances the energy storage properties and the electrochemical activity of the material.^{40,47,61,62,68} MOFdc with sulfur-doped and phosphorus-doped carbons have also been reported, however these are less common.⁶⁹ MOF-derived materials that do not contain a carbon matrix can also be generated,^{70–72} however these materials are outside the scope of this review.

On a macro scale, MOFdc also inherit structural features from the template MOFs. As highly ordered crystalline materials, MOFs generally grow in well-defined morphologies, such as octahedra, cubes, or rods. Upon carbonization, these morphologies are well-maintained, with MOFdc usually exhibiting the same morphology as the template MOF



crystals.^{40,73,74} Due to the loss of material through partial combustion or evaporation of the metal species, the faces of octahedral or cubic MOFdc are often sunken in.⁷⁵ Furthermore, due to the loss of relatively strong coordination bonds in three dimensions and the generation of additional grain boundaries, MOFdc are generally more brittle than the template MOFs.⁷⁶

Another feature that is common, though not universal, in MOFdc is the presence of inorganic components. While some MOFdc research focuses on the structure of the carbon components and will deliberately remove the inorganic components,^{51,52} the field of metal-containing MOFdc is growing rapidly.^{37,68,77–79} Inorganic components typically exhibit nanosized domains, as the temperatures of MOF pyrolysis are often not high enough to melt or sinter the metal species into larger domains.^{80,81} In some cases, the inorganic species are single atoms supported on the carbon,^{62,77,82} however this is often difficult to achieve due to the high temperatures of the carbonization and the close proximity of the metal ions in the template MOF structure. A variety of metal species can be formed in a MOFdc, including metals, metal oxides, metal nitrides, metal carbides, and metal sulfides.^{45,79,83–87} In cases where more than one metal ion is contained within the IBU or hosted in the pores of the MOF, multimetallic inorganic species or alloys are often formed.^{88–92} Due to the small size of the domains and inorganic node distortions that occur upon heating, metal species in MOFdc are often poorly ordered, which makes traditional bulk scale or average structure analysis challenging or poorly representative of the true structure of the species.^{83,84,93–95} Consequently, many researchers in the field have relied on local structure analysis techniques, such as X-ray or neutron total scattering or extended X-ray absorption fine structure (EXAFS) to complement average structure analysis.^{45,83,84,94–96}

Effect of carbonization conditions on MOFdc structure

More recently, research into the influence of carbonization conditions on the MOFdc structure and composition has begun to be studied. An early work in this area was performed by Banerjee, Poddar, and coworkers, who compared carbonization of a range of MOFs in both air and nitrogen environments. They found that the inorganic species produced from carbonization in air exclusively formed metal oxides, regardless of the identity of the metal ($M = \text{Mg}, \text{Mn}, \text{Co}, \text{Cu}, \text{Zn}, \text{Cd}$). However, carbonization in nitrogen produced metallic nanoparticles of cobalt and copper, while the other metals were not reduced under these conditions due to their lower oxidation potential.⁸⁷

In another such study, Zhou and coworkers conducted a detailed investigation of the iron MOF PCN-250 (PCN = porous coordination network) and the effects of pyrolysis temperature and gas environment on the structure of the resulting iron species and carbon matrix in the MOFdc (Fig. 3).⁴⁵ They found that pyrolysis in oxidizing gases produced MOFdc with



Fig. 3 Influence of MOF pyrolysis conditions on the structure of PCN-250-dC.

low surface areas and low carbon content, suggesting that the oxidative conditions caused combustion of the linker before it could be carbonized. In contrast, non-oxidizing atmospheres typically produced much higher surface areas and higher carbon content, suggesting that a non-oxidizing environment is key for retaining the porous structure of the MOF after carbonization. Chapman and coworkers similarly investigated the pyrolysis of PCN-250 under reducing atmospheres, corroborating the results reported by Zhou and expanding the scope of the research to consider the structure of metal oxide nanoparticles derived from multimetallic PCN-250.⁹⁴ Zhou and coworkers also demonstrated that the linker decomposition process involves an oxidative decarboxylation, which in turn reduces the iron of the MOF cluster to iron(0) before later being oxidized, while Chapman and coworkers identified metal node distortions prior to structural collapse, suggesting that both node and linker dynamics contribute to the MOF decomposition.^{45,65,66,94} This process also shed light on differences in the iron oxide phase present following pyrolysis in different gas environments. While carbonization in oxidizing gases produced a single phase of iron oxide due to uniform oxidation of the iron(0) intermediate during the carbonization, carbonization in non-oxidizing gases produced multiple iron oxide phases due to the CO_2 produced during the carbonization and post-carbonization oxidation upon exposure to ambient air.⁴⁵ Carbonization in more strongly reducing gas environments produced iron(II) oxide, perhaps by inhibiting the oxidative decarboxylation of the linker.⁹⁴ Notably, this mechanistic understanding of the carbonization process has subsequently been applied to target the formation of MOFdc with specific structural features.⁹⁷ Cheetham, Wang, and coworkers also observed the effect of gas environment on the metal species of a MOFdc. Carbonization of a manganese triazolate framework under rigorously oxygen-free conditions produced nanoparticulate manganese nitride that was air-stable, a property not found in bulk manganese nitrides. In contrast, carbonization in air produced manganese oxide.⁸³ As the nitrogen of the manganese nitride was derived directly from the triazolate linker, this suggests that different reaction mechanisms may be at play depending on the gas environment.



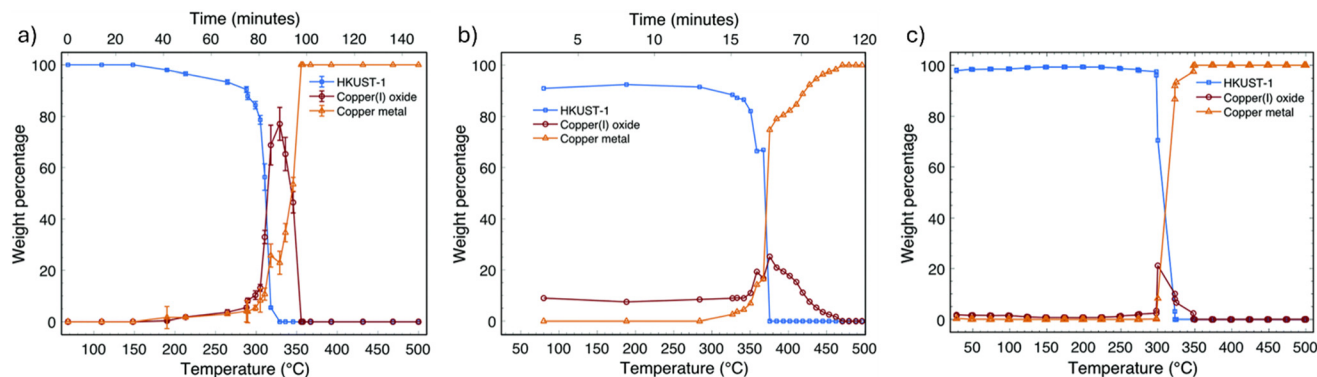


Fig. 4 Evolution of MOF, Cu(I) oxide, and Cu(0) phases during carbonization of HKUST-1, as measured by XAS (a), X-ray PDF (b), and PXRD (c). Reproduced from ref. 96 with permission from the Royal Society of Chemistry.

Zhou and coworkers also noted some significant differences in MOF_dC structure based on pyrolysis temperature. When Zn-MOF-74 was carbonized below 600 °C, ZnO nanoparticles were produced, while carbonization above 600 °C caused sintering of the ZnO into a bulk domain.⁸⁴ This sintering causes an increase in mesoporosity as the ZnO sinters to the exterior of the MOF_dC, leaving mesoporous voids on the interior of the material. While this specific sintering behavior had not been previously observed in MOF_dCs, other temperature dependent effects have been described. The first MOF_dC reported was calcined at 1000 °C to ensure evaporation of the zinc species derived from the IBU, however it was also possible to obtain zinc oxide-containing MOF_dCs by carbonization at lower temperatures, albeit at the expense of high porosity.³⁹ The rate of heating can also influence this sintering behavior. More rapid heating ramps typically produce larger, more crystalline metal species, as the activation energy for grain growth can more readily be overcome.^{98,99} Notably, the effect of heating ramp is generally negligible on the decomposition pathways or the identity of the final metal species, as demonstrated by pyrolysis of HKUST-1 (HKUST = Hong Kong University of Science and Technology) at heating ramps from 2–5 °C min⁻¹ (Fig. 4).⁹⁶ By conducting *in situ* powder X-ray diffraction (PXRD), X-ray absorption spectroscopy (XAS), and X-ray total scattering pair distribution function (PDF) experiments, the authors observed that the MOF decomposed at 300–325 °C into a mixture of Cu(0) and Cu(I) oxide, the latter of which was an intermediate that was subsequently reduced to Cu(0) at higher temperatures (above 350 °C). While the Cu(I) oxide intermediate was not observed upon rapid heating (over 2500 °C min⁻¹), this difference was ascribed to the limited time/temperature resolution that could be achieved in the analysis and does not necessarily indicate a change to the mechanism of Cu(0) formation.⁹⁶

Effect of MOF template structure on MOF_dC structure

In addition to the carbonization conditions, features of the MOF template structure such as dimensionality, IBU structure,

and connectivity also play a role in determining the structure of the MOF_dC. When comparing the carbonization products of copper MOFs HKUST-1 and F-MOF-4 (F-MOF = fluorinated metal–organic framework), the 3-dimensional HKUST-1 produces larger copper particles than the 2-dimensional F-MOF-4.⁸⁷ This enhanced growth was hypothesized to be a consequence of the dimensionality, as the HKUST-1-derived copper particles are agglomerated from all three dimensions. This hypothesis also held for the comparison between the 2-dimensional cobalt MOF MOF-CJ4 and the 3-dimensional cobalt MOF Co-HFMOF-D (HFMOF = partially fluorinated metal–organic framework).⁸⁷

The effect of linker structure on the pore structure of the carbon matrix of the MOF_dC has also been investigated. In another study, several zinc MOFs with a range of linkers were carbonized to produce metal-free MOF_dCs of varied structure. Carbonization of MOF templates with more rigid and aromatic linkers produced porous carbons that maintained the morphology and pore structure of the template MOF, while more flexible linkers produced carbons with much less well-ordered pore structures.⁵³

The influence of MOF structure on the resulting MOF_dC was further explored by Zhou and coworkers. When comparing pyrolysis of Zn-MOF-74 and UiO-66 (UiO = University of Oslo), they found that carbonization of Zn-MOF-74 and its higher connected and more thermally stable linker produced MOF_dCs with higher surface area by preventing pre-carbonization pore collapse.⁸⁴ Furthermore, they found that carbonization of UiO-66 produced primarily cubic zirconia due to the templating effect of the cubic Zr₆-oxo clusters, despite this phase typically being considered unstable at particle sizes above 2 nm.¹⁰⁰

Cheetham, Wang and coworkers note that while oxygen-free carbonization of a manganese triazolate framework produces manganese nitride, carbonization of the analogous iron and cobalt triazolates under the same conditions produce iron carbide and cobalt metal respectively.⁸³ While this was not explored in depth, it further emphasizes that different inorganic components in the MOF can result in different reac-



tion pathways and different inorganic products. Likewise, Zhou and Chapman have both demonstrated that incorporation of a heterometal into the cluster of PCN-250 can produce different metal phases following carbonization, with PCN-250-Fe₂Ni in particular favoring the formation of fcc-Ni due to the favorability of FeNi alloy formation and selective Ni extraction from the cluster during the pyrolysis process.^{94,97}

One of the most significant advances in MOFDCs in the last decade is the development of the metal ion “fence” strategy to form MOF-derived single atom catalysts (SACs) supported on a porous carbon matrix (Fig. 5). Following reports of Co SACs formed from bimetallic ZIF structures,^{101,102} this strategy was more systematically developed by Wu, Li, and coworkers, who carbonized a mixed cobalt/zinc ZIF to produce single cobalt atoms anchored to a N-doped carbon matrix.⁷⁷ By incorporating zinc ions in the ZIF structure as a “fence” between cobalt ions, the cobalt ions were too spatially separated to undergo significant aggregation and thus remained embedded as single atoms in the carbon with a porphyrinic planar structure. Wang and coworkers further investigated the relative activity of MOF-derived Co-SACs and MOF-derived cobalt nanoparticles for nitroarene hydrogenation, showing that the SACs were significantly more active.¹⁰³ This fencing strategy has since been used with other metal dopants, including iron,^{62,104–107} manganese,¹⁰⁴ nickel,^{107–110} and copper.^{110,111}

An alternative strategy that relies on the same principle is that described by Zhou, Li and Fischer, in which the porphyrin center of a porphyrinic zirconium-based MOF was partially metalated with nickel and several other transition metal ions.⁸² As with the fence strategy, the incomplete metalation of the porphyrin center resulted in significant distance between the nickel ions and prevented agglomeration or sintering into larger particles upon carbonization of the MOF, instead producing SACs on a porous carbon support. However, this strategy is not universally applicable. When a zinc-porphyrin MOF metalated with palladium is used in place of the zirconium-porphyrin MOF metalated with nickel, the final material con-

tained a palladium-zinc alloy instead of single palladium atoms dispersed on the carbon.¹¹²

More complex metal species can be produced by designing MOF templates with more than one metal ion in the IBU. Prussian blue, a coordination polymer known since the early 1700s, can be calcined to form porous multimetallic iron oxide structures, with partial substitution of the iron atoms by heterometals such as manganese or cobalt.^{90,91} Another popular MOF template for the formation of alloys is M-btc (H₃btc = benzene 1,3,5-tricarboxylic acid), as multimetallic MOFs of this structure can be readily synthesized in one-pot reactions simply by varying the concentration of each respective metal ion in the reaction mixture.^{88,89}

It is also possible to incorporate secondary metals by impregnating the MOF with the metal or metal precursor prior to carbonization. This strategy has commonly been performed with the ZIF-8/ZIF-67 system, as techniques for impregnation of metal nanoparticles in this MOF are well-established. This is a particularly desirable strategy when the metal of interest is too expensive to be used as the backbone of the MOF, for example noble metals such as gold or platinum.^{113,114} In many cases, this strategy is used to form alloys or multimetallic species that are more active than the individual metals alone.^{92,114–117}

Structure–function relationships in MOFDCs

Gas storage

As with most materials, the functionality of MOFDCs is dictated by their structure. Inspired by the promise of MOFs in gas storage applications, early work with MOFDCs similarly focused heavily on the gas storage properties of the materials. The first report of a MOFDC involved a discussion of the hydrogen uptake properties of the material,³⁹ which remained a popular application for MOFDCs throughout the early 2010s.^{47,54,87} Crucially, MOFDCs outperformed porous carbons produced by other methods due to the high concentration of ultramicropores.^{54,118} Furthermore, some highly graphitic MOFDCs are excellent materials for sensing aromatic compounds due to their high surface area and the affinity of the graphitic sp²-hybridized carbons for aromatic molecules.⁵¹ MOFDCs have also been used as sorbents for liquids and some dyes due to the hydrophobicity of the carbon and their high porosity.^{75,119–122} In addition to their high uptake of oil and dyes, the presence of magnetic cobalt- or iron-based nanoparticles derived from the MOF IBUs can significantly improve recovery and recyclability of the sorbent.^{75,119}

Catalysis

MOF-derived bimetallic metal species have also been used as Fenton catalysts, where peroxide and hydroxide radicals are generated to degrade organic pollutants.¹²³ Similarly, MOF-derived SACs, often formed using the fencing strategy, have also been applied as Fenton or Fenton-like catalysts,^{106,124} as

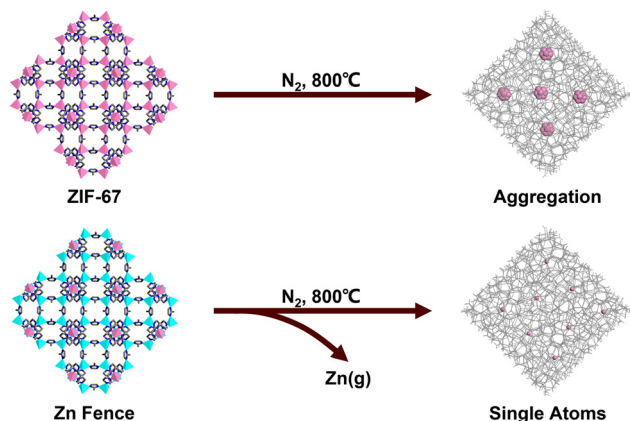


Fig. 5 “Fencing” strategy for the generation of ZIF-derived Co-SACs. The procedure is generalizable to other metals.



have MOF-derived cobalt nanoparticles¹²⁵ and partially carbonized Ce-TCPP nanocomposites (H₄TCPP = tetrakis(4-carboxyphenyl)porphyrin).¹²⁶ Even metal-free MOFDCs have been applied as catalysts for peroxydisulfate activation due to the high activity of nitrogen-doped carbon structure.^{55,56}

MOFDCs are also strong candidates for large-scale industrial catalysis at elevated temperatures and pressures.^{41,42,127,128} Fischer–Tropsch synthesis of hydrocarbon chains from syngas is an important industrial reaction which is typically performed at high temperatures and pressures.⁴¹ While MOFs may not be stable under these conditions, MOFDCs are significantly more resistant to high temperatures.³⁷ A variety of iron- and cobalt-based MOFs have been carbonized and applied for Fischer–Tropsch catalysis, with either ultrasmall metal nanoparticles or SACs as the active species. Gascon and coworkers have produced several iron-based MOFDCs that equal or outperform commercial Fischer–Tropsch catalysts.^{42,127}

Electrocatalysis

Due to the high concentration and good dispersion of metal species and the conductivity of the porous carbon matrix, MOFDCs have also shown significant promise as electrocatalysts for a variety of reactions. This review will briefly summarize several key reactions for which MOFDCs have been used as catalysts and the types of MOFDCs that have been applied in this way. Readers are directed to other recent reviews for a more detailed discussion of the design and application of MOFDCs for electrocatalysis.^{61,68,129–132}

Electrochemical nitrogen reduction is an important reaction for energy storage as it presents a key avenue for hydrogen fixation and ammonia production. MOFDCs have recently been applied in this process.^{132,133} In one such example, Cu-btc was calcined at several different temperatures to produce either a Cu-SAC or copper nanoparticles. The Cu-SACs were highly effective at promoting nitrate reduction while suppressing the accumulation of toxic nitrite ions.¹³⁴ Li and coworkers similarly found cobalt-doped iron nanoparticles derived from MOF-74 were effective nitrate reduction catalysts.¹³⁵ Nitrate is not the only substrate that has been converted to ammonia by MOF-derived catalysts. A bismuth-based MOFDC was used to fix nitrogen under ambient conditions with very high faradaic efficiency.¹³⁶

Carbon dioxide utilization is another key process for which MOFDCs have been applied as electrocatalysts. The ZIF-8 system is a popular template for this reaction due to the propensity to form SACs upon carbonization. Both monometallic Ni-SACs and tandem systems that include Ni-SACs and either Cu- or Fe-SACs within the same porous carbon have been generated.^{107,108,110} Other nickel-based MOFs have also been carbonized to produce Ni-SACs with promising results.^{88,137} Similarly, HKUST-1-derived Cu–C composites have been applied for CO₂ electroreductions to both methanol and ethanol with high faradaic efficiencies and very low overpotentials (Fig. 6).¹³⁸ Titanium-based MOFs have also been applied as templates for the generation of carbon dioxide reduction

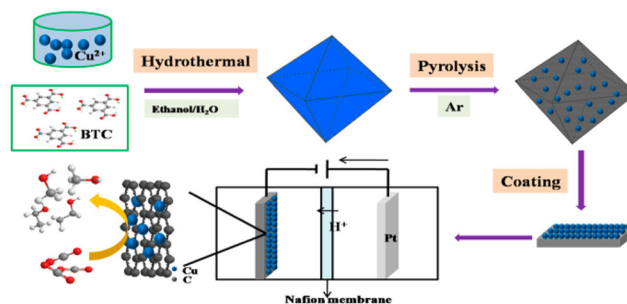


Fig. 6 Synthesis and utilization of HKUST-1-derived Cu–C composites for CO₂ electroreduction to C₁ and C₂ alcohols. Reprinted with permission from ref. 138. Copyright 2017 American Chemical Society.

reaction (CO₂-RR) catalysts. In one example, MIL-125-derived Mg-doped titania (MIL = Materials Institute Lavoisier) was used to photocatalytically reduce carbon dioxide, with the magnesium dopant promoting the generation of surface Ti³⁺ ions to improve charge separation. Furthermore, the Mg-doping increased the surface area of the titania by discouraging sintering.¹³⁹ In another, a mixed titanium-iron MOF was used to generate titanomaghemite nanoparticles with a high Fe/Ti ratio that cannot be achieved *via* traditional inorganic synthetic methods. This MOFDC was also an effective electrocatalyst for CO₂-RR.¹⁴⁰

Electrochemical water splitting reactions are also an important target reaction for MOFDC catalysts. Oxygen reduction reactions (ORR) are one of the most common applications in the literature for metal-containing MOFDCs.^{44,61} While early work in this area focused on the generation of iron nanoparticles,¹⁴¹ the ZIF-8/ZIF-67 system later became a popular template for the synthesis of these catalysts due to its propensity to form SACs, a consequence of the high nitrogen content in the carbon matrix and the ability to “fence” the Co ions with zinc in a bimetallic framework.^{77,105,142–144} Cobalt is generally the most active SAC for these reactions, however iron and copper sites have also been employed.^{105,145} Other MOFs have also been used as templates for generating SACs as ORR catalysts, such as Fe-doped NH₂-MIL-101(Al).¹⁴⁶ These materials have been found to exhibit higher ORR activity than commercial Pt/C catalysts, representing an important step towards reducing society’s reliance on noble metals and increasing the sustainability of ORR.^{105,147}

In addition to ORR, MOFDCs have been employed as multifunctional catalysts that are active towards both ORR and oxygen evolution reactions (OER), the latter of which is significantly more challenging due to its slow kinetics.^{43,44,68,148} A diverse range of template MOFs have been used to generate such catalysts. In one example, a Zn-MOF was used as a host for copper or platinum nanoparticles and was carbonized to form a nanoparticle@porous carbon material without the inclusion of IBU-derived nanoparticles.¹⁴⁹ More common, however, is the generation of IBU-derived nanoparticles, such as through the carbonization of thiourea@Fe-btc to form an FeS/Fe₃C@C composite or the carbonization of a cobalt-



vanadium mixed-metal MOF to form cobalt and vanadium nitride nanoparticles.^{85,86}

Energy storage

Finally, energy storage has recently emerged as an application for MOFDCs due to their high specific capacity, high surface area, and robustness towards volume expansion of guest materials.^{79,150,151} Importantly, the porous carbon of MOFDCs is often graphitic in nature, thus the materials can be engineered to exhibit high capacities and enhanced ion transport compared to non-porous graphite.^{84,152,153} These properties mean that MOFDCs often excel as anode materials for lithium, potassium, and sodium ion batteries.^{151–157}

The conductive nature of the carbon also allows MOFDCs to be used as matrices for other high-capacity anode materials.

Silicon anode lithium ion batteries are highly desired due to the extremely high theoretical capacity (4200 mA h g^{-1}), however challenges such as volumetric expansion upon lithiation limit the amount of silicon that can be incorporated without experiencing severe capacity fade.^{158–160} Several studies have demonstrated the utility of MOFDCs as a porous carbon host for nanoparticulate silicon (Fig. 7).^{150,160–162} The high porosity of the carbon structure enables highly efficient lithium ion diffusion, leading to high specific capacity and high capacity retention.¹⁶¹ Furthermore, the mechanically flexible porous carbon structure of the MOFDC mitigates the detrimental effects of silicon expansion, due to the porous carbon acting as a buffer to reduce strain or induce *in situ* formation of ultrasmall silicon nanodots.^{150,160}

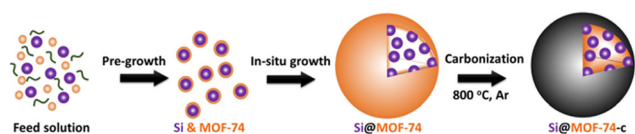


Fig. 7 Synthesis of Si@MOFDC composite anode materials. Reprinted with permission from ref. 150. Copyright 2023 American Chemical Society.

Conclusions and outlook

MOFDCs are an attractive alternative to MOFs due to their enhanced stability and conductivity, and an improvement upon traditional porous carbons due to their microporosity and tunability. As demonstrated in this review, a vast array of MOFDC structures can be selectively synthesized through strategies such as judicious choice of carbonization conditions,

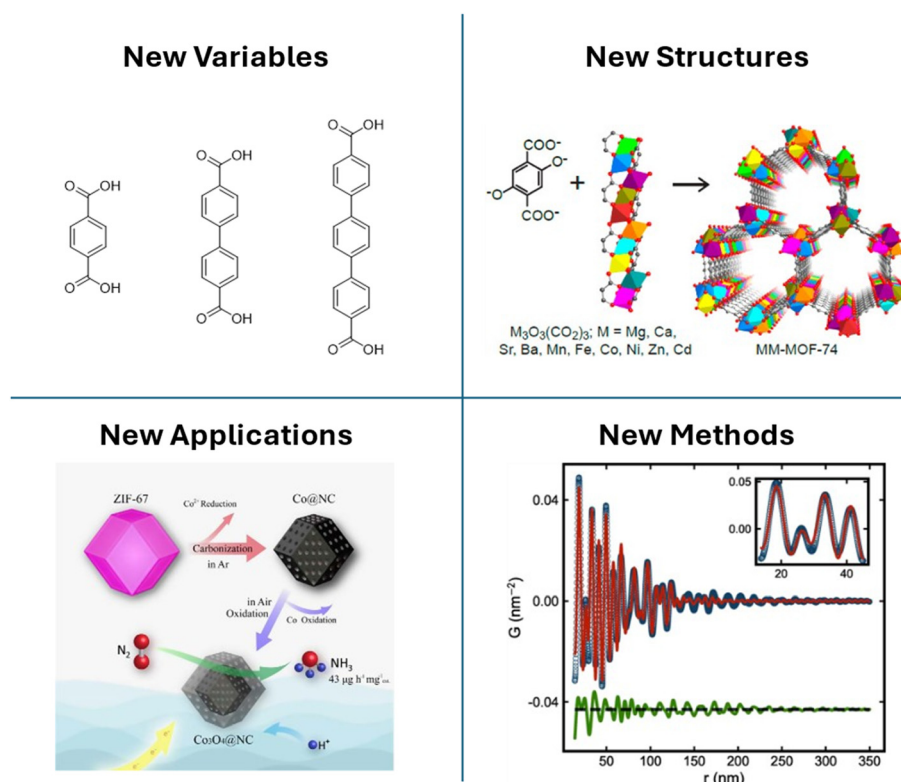


Fig. 8 Future directions for MOFDC research include investigation of previously unstudied structural variables (e.g. linker length, top left), targeted design of multimetallic inorganic structures using multimetallic MOF platforms (top right, reprinted with permission from ref. 167. Copyright American Chemical Society 2014), targeted design of electrocatalysts at commercial scale (bottom left, reprinted with permission from ref. 133. Copyright American Chemical Society 2019) and use as a model system for developing advanced analytical techniques (e.g. sasPDF, bottom right, reproduced from ref. 169 with permission from the International Union of Crystallography).



tuning the MOF structure, and incorporating guests in the MOF prior to carbonization. As the field of MOFDCs continues to grow, there are a number of areas ripe for future research (Fig. 8).

While the literature to date has focused primarily on the templating effect of the IBU and framework structure and the impact of pyrolysis conditions, there are many other variables that have yet to be considered in the design of MOFDCs. Variables such as linker aromaticity and rigidity have been studied for their influence on the carbon structure,⁵³ however their impact on the structure of the metal species has not been considered. Other factors, such as linker length and connectivity, have not yet been systematically studied for closely related MOF templates, such as isorecticular expansions of a template MOF.^{11,163} The inclusion of additional functional groups on linkers also provides a pathway for altering the structure of the metal species in the MOFDC, as they may interact differently with various chemical moieties throughout the carbonization process.^{93,164}

MOFDCs also represent a step forward in the applicability of MOF-based materials due to their improved stability over the template MOFs. To this end, future research should consider the design of MOFDCs tailored to specific applications by using these principles. As electrocatalytic applications are widely investigated, future studies should employ systematic approaches to develop highly active, but inexpensive MOF-derived electrocatalysts, for example using bimetallic MIL-100(Fe) as a template instead of the more costly PCN-250.^{44,97,165,166} Additional developments, such as the design of trimetallic (or more) MOFs as precursors for new and unusual inorganic materials, should also be pursued.^{140,167}

Finally, MOFDCs provide a fascinating platform for the development of new analytical techniques and the advancement and extension of existing techniques. The nature of MOFDCs as poorly ordered solid materials that may contain regions of greater structuring, such as metal nanoparticles, challenges our existing conceptions on how to study the structure of materials. A swathe of advanced complementary techniques is required to fully describe the structure of MOFDCs and new techniques may be able to extract information from unexpected places. Given the often small size of the inorganic nanoparticles in MOFDCs, the combination of small- and wide-angle X-ray scattering (SAXS and WAXS) would provide an avenue to study the coevolution of X-ray scattering and Bragg diffraction peaks *in situ*, thus supporting models derived from other advanced *in situ* techniques.⁹⁶ Due to the differences in intensity of the scattering and diffraction features and the challenges of simultaneous detection across wide q-ranges, this type of analysis will inspire novel approaches in detector design and data collection methods that can be similarly applied in characterization of other advanced materials. Other cutting-edge techniques, particularly in the realm of local structure analysis, can also be further developed using MOFDCs as a model system. Due to their poor crystallinity, proper characterization of MOFDCs often requires the use of local structure analysis techniques, such as PDF or far-infrared

(terahertz) spectroscopy.^{45,94,168} While it is possible to collect this type of data using conventional instrumentation, the cost and time required to obtain high quality data typically necessitates the use of synchrotron radiation. As the field of MOFDCs continues to grow, it will spur further advances in high-intensity X-ray sources and high-resolution detectors to enable greater accessibility, as well as the development of other advanced local structure analysis techniques, such as small angle scattering PDF (sas-PDF) and single-atom-sensitive electron energy loss spectroscopy (EELS).^{169,170}

Author contributions

J. A. P. and H.-C. Z. conceptualized the manuscript, J. A. P. and Y. Y. prepared the manuscript, H.-C. Z. conducted supervision.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

Conflicts of interest

There are no conflicts to declare.

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