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Development of the design and synthesis of metal–organic frameworks (MOFs) – from large scale attempts, functional oriented modifications, to artificial intelligence (AI) predictions

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Owing to the exceptional porous properties of metal–organic frameworks (MOFs), there has recently been a surge of interest, evidenced by a plethora of research into their design, synthesis, properties, and applications. This expanding research landscape has driven significant advancements in the precise regulation of MOF design and synthesis. Initially dominated by large-scale synthesis approaches, this field has evolved towards more targeted functional modifications. Recently, the integration of computational science, particularly through artificial intelligence predictions, has ushered in a new era of innovation, enabling more precise and efficient MOF design and synthesis methodologies. The objective of this review is to provide readers with an extensive overview of the development process of MOF design and synthesis, and to present visions for future developments.

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1. Introduction

Metal–organic frameworks (MOFs) are a class of crystalline materials characterized by their unique porous structures and

exceptional properties.^{1–3} Composed of metal ions or clusters coordinated to organic ligands, MOFs feature intricate frameworks with distinct pore environments. This architecture grants MOFs an extraordinary surface area, often surpassing that of traditional porous materials, making them highly effective for a variety of applications.^{4–10} The versatility of MOFs arises from the vast combinatorial possibilities of metal nodes and organic linkers, allowing for tailored synthesis to meet specific needs. This adaptability has propelled MOFs to the forefront of research and industrial applications.^{11–15}

In theoretical research, the development of MOFs has significantly expanded the fields of coordination chemistry

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Zhou's group

The Zhou Research Group focuses on interdisciplinary research spanning Chemistry, Biochemistry, and Materials Science and Engineering with an emphasis on porous materials such as metal–organic frameworks (MOFs), porous polymer networks (PPNs), and covalent and coordination cages. By exploiting the extreme tunability of MOFs, PPNS and other porous materials, the group seeks to understand the role that material architecture and functionality has on its chemical and physical properties and how modifying these properties can lead to the development of MOFs and PPNS for applications related to gas and energy storage, separations, catalysis, enzyme immobilization and drug delivery.



and topology, deepening the understanding of coordination geometries and topological structures.^{16–20}

In practical applications, the potential of MOFs is amplified by their stability and recyclability, positioning them as sustainable alternatives to conventional materials. As advancements in MOF design and synthesis continue to emerge, these materials hold promise for addressing some of the most pressing challenges in energy, environment, and health. The ongoing exploration of MOFs represents a vibrant and rapidly evolving frontier in materials science.^{21–25}

Traditional design principles and synthesis methods heavily rely on high-throughput attempts within a continuous “trial and error” process to combine different metals and ligands, leading to the creation of various novel structures. Building on the solid foundations of primary research, functional-oriented design and modification methods have been developed, reducing the time and cost associated with numerous attempts. These advanced methods allow for the introduction and modification of target coordination fragments and functional groups at specific positions to achieve desired properties and functions. However, the vast number of possible combinations of metal nodes and organic linkers makes the experimental synthesis and modification of MOFs time-consuming and resource-intensive processes. Recently, with the evolution of computer science and technology, significant advancements in machine learning (ML) and artificial intelligence (AI) have further accelerated the design and modification of new structures for targeted applications. AI-assisted MOF synthesis can address these challenges by leveraging ML algorithms and computational models to predict optimal synthetic routes and material properties. Recent research progress has focused on developing AI models that can rapidly screen large chemical spaces, predict MOF stability and performance, and even suggest novel MOF structures with desired characteristics. These AI tools can analyse vast amounts of experimental data, helping to identify patterns and correlations that are not easily discernible by human researchers. The necessity of AI in MOF synthesis arises from the sheer complexity of the design space and the desire to accelerate the discovery and development of novel MOFs for emerging applications, reducing both the cost and time of experimentation while enhancing the precision and scalability of the synthesis process. These technologies enhance the efficiency and precision of identifying optimal combinations and configurations, thereby revolutionizing this field and opening new frontiers in materials science (Fig. 1).

1.1 Origin and development of MOFs

The origin and development of MOFs represent a remarkable journey in the field of porous science, beginning in the late 20th century. Initially rooted in coordination chemistry, the early concepts of MOFs evolved from the synthesis of coordination polymers.^{26–28} The pivotal moment came in the 1990s with the pioneering work of researchers, such as Prof. O. M. Yaghi, who demonstrated that combining metal salts with organic linkers could create highly porous, crystalline structures with vast surface areas and customizable properties.^{29–31} This innovation provided

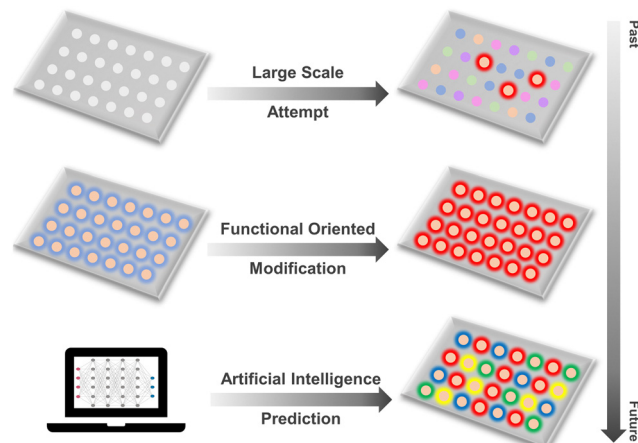


Fig. 1 Schematic representation of the development of the design and synthesis of MOFs.

a new class of materials with significant potential for various applications based on their unique pore properties.

The subsequent decades saw rapid advancements in MOF research, driven by improvements in synthesis techniques and computational modelling (Fig. 2). These advancements enabled the precise design and functionalization of MOFs, significantly broadening their applicability. The ability to tailor the pore size, shape, and functionality of MOFs has led to their exploration in numerous fields, including environmental remediation, energy storage, and drug delivery.^{32–36}

Nowadays, MOFs continue to be a vibrant area of research, with ongoing efforts focused on discovering new frameworks, enhancing their stability, and developing scalable production methods. The continuous innovation in this field promises to address some of the most pressing challenges in technology and industry, positioning MOFs as a cornerstone of future scientific and industrial advancements.^{37–40}

1.2 Machine learning and artificial intelligence

The evolution of computer science and technology has catalyzed significant advancements in the fields of ML and AI. AI technology emulates the human intelligence processes, enabling machines to perform complex tasks such as natural language understanding, vision perception, and decision-making. ML, a primary state of AI, focuses on developing algorithms that allow computers to learn from data and make accurate predictions (Fig. 3).^{41–44}

With the surging development of computational power and data storage capacities, modern computers are endowed with the ability to employ ML and AI models to process and analyse vast datasets, uncovering patterns and insights beyond human reach. The iterative learning capabilities of ML and AI—through methods such as supervised, unsupervised, and reinforcement learning—have been enhanced by advancements in hardware, software, and data acquisition techniques, which drives innovation and efficiency in healthcare, finance, transportation, and beyond. In healthcare, these technologies enable breakthroughs in disease detection and personalized medicine. In



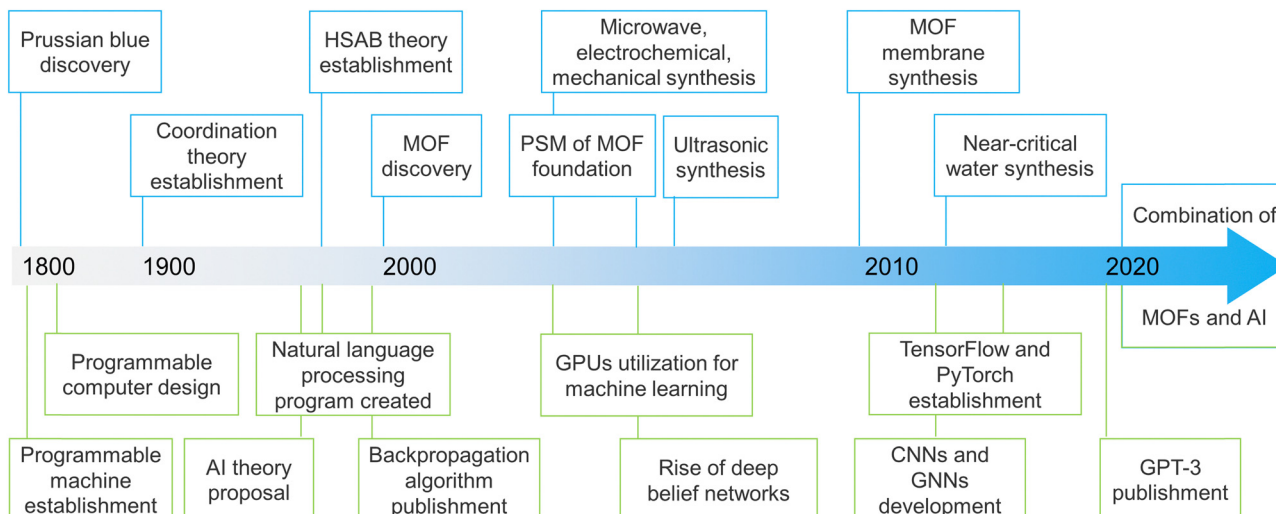


Fig. 2 Simplified development timeline of MOFs and AI. HSAB stands for hard–soft acid–base, while PSM stands for post-synthetic modification. CNN stand for convolutional neural network, while GNN stands for graph neural network.

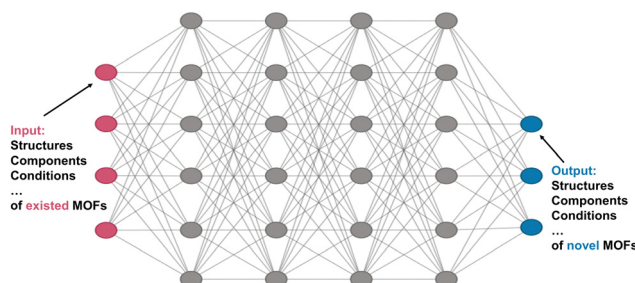


Fig. 3 Schematic representation of the ML and AI-guided MOF design and synthesis.

finance, they improve fraud detection and algorithmic trading. Autonomous vehicles, intelligent virtual assistants, and personalized recommendation systems illustrate their broad impact on daily life.^{45–47}

Ongoing research aims to refine ML and AI methodologies, emphasizing model interpretability, robustness, and ethical considerations. The continuous interplay between technological development and ML/AI advancements promises to address complex global challenges, augment human abilities, and foster unprecedented opportunities for innovation. This dynamic field remains at the forefront of the digital transformation, representing a pivotal area of growth in modern technology.^{48–50}

2. Traditional design principle and synthesis method

As a class of crystalline porous materials constructed through the coordination interactions between metal ions and organic ligands, the structures of MOFs depend heavily on the selection of metals, ligands, solvents, and other reaction conditions, like

temperature and pressure. Based on this, the design principles and synthesis methods of MOFs are pivotal in determining their properties and functions. Traditionally, researchers have employed high-throughput screening under a continuous “trial and error” process to construct novel MOFs with specific components and spatial structures, employing various synthesis methods. While this approach is straightforward in design, it is both time and cost consuming.

2.1 Traditional design principle

The design of MOFs revolves around the deliberate selection and combination of metal nodes and organic linkers to create frameworks with specific structures, properties, and functionalities. The key principles primarily involve the careful matching of metal salts, ligands, solvents, and reaction conditions (Fig. 4).

In general, the synthesis of MOFs is a complex process that typically involves three key stages: the formation of coordination bonds, nucleation, and crystal growth. Initially, metal ions

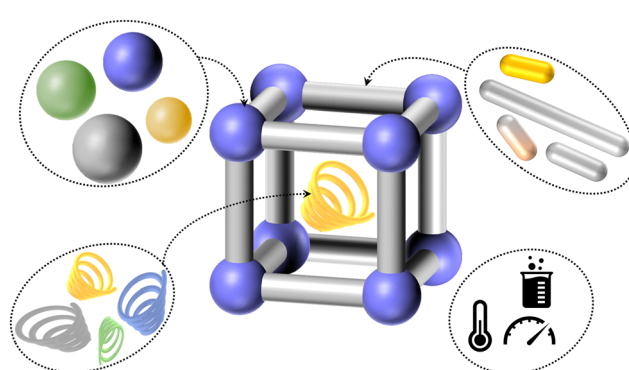


Fig. 4 Design principle of MOFs through the selection of metal salts, ligands, solvents, and reaction conditions.

or clusters interact with organic ligands to form coordination bonds, establishing the foundational framework of the material. This step defines the basic connectivity and geometry of the MOF structure. Once coordination bonds are formed, nucleation begins, where small, stable clusters—known as crystal nuclei—emerge. The conditions at this stage, such as temperature, solvent, and concentration, significantly influence the size and number of these nuclei, which in turn affects the final morphology of the MOF. After nucleation, the crystal growth phase follows, during which the nuclei expand into well-defined, extended structures as more metal–ligand units are added. The rate of crystal growth and the resulting size and quality of the MOF crystals can be controlled through various synthesis parameters, such as reaction time, temperature, and solvent selection. Together, these stages govern the overall properties of the MOF, including the porosity, stability, and surface area, which are critical for its performance in applications like catalysis, gas storage, and sensing.

The hard and soft acids and bases (HSAB) theory plays a significant role in guiding the design and synthesis of MOFs by providing a theoretical framework for understanding and predicting the interactions between metal ions and organic linkers. In essence, HSAB theory helps researchers optimize the choice of metal centers and organic ligands based on their “hard” or “soft” characteristics, which in turn affects the stability, functionality, and overall properties of the resulting MOF. For applications requiring hard or soft metals, corresponding hard or soft ligands will be selected to connect the metal centers and form stable frameworks. Similarly, for applications requiring hard or soft ligands, matching hard or soft metals will be chosen to ensure optimal compatibility and stability. The HSAB theory serves as a powerful guide for MOF design and synthesis by enabling the rational selection of metal nodes and organic linkers to optimize interactions based on their hard or soft characteristics.

The synthesis of MOFs is also governed by a delicate balance between dynamic processes and thermodynamic equilibrium. The dynamic of MOF synthesis involves kinetic processes such as nucleation, crystal growth, and reaction rates, while thermodynamic equilibrium dictates the stability of the final product and its structural phases. The challenge in MOF synthesis lies in balancing the dynamics and thermodynamics to achieve the desired structures and properties. In some cases, the kinetic pathway dominates, leading to metastable MOF phases that are kinetically trapped and may possess unique properties, such as open frameworks with high porosity but lower stability. In other cases, allowing the system to reach thermodynamic equilibrium results in more stable and crystalline MOFs with fewer structural defects. By adjusting synthesis parameters, researchers can manipulate both the dynamic and thermodynamic aspects of MOF formation.

2.1.1 Metal salt selection. The choice of metal nodes is a critical factor in the design and synthesis of MOFs (Fig. 5). Metal nodes, which can be metal ions or clusters, serve as the central coordination points in MOFs, linking organic ligands into extended and porous structures. The selection of

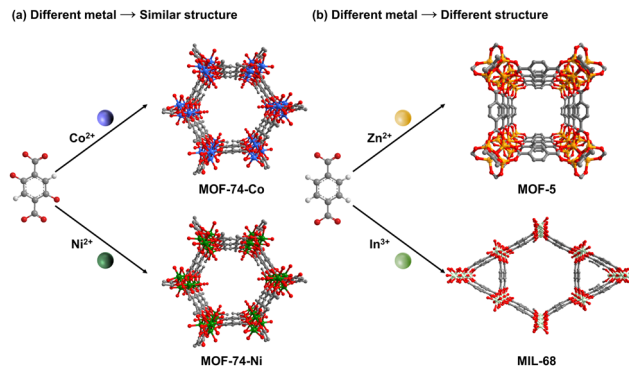


Fig. 5 Metal selection influence on the structures of MOFs under similar conditions. Atom code: C, grey; O, red; H, white; Co, blue; Ni, green; Zn, yellow; In, light green.

appropriate metal nodes significantly influences the structural, physical, chemical, and biological properties of MOFs, thereby determining their suitability for various applications. Main group metals (especially Mg, Ca, Al, In, *etc.*),^{51–54} fourth-period transition metals (particularly Ti, Cr, Fe, Co, Ni, Cu, Zn, *etc.*),^{55–61} fifth-period transition metals (notably Zr, Cd, *etc.*),^{62,63} and lanthanide metals^{64–66} are frequently employed as metal nodes.

Benefiting from the diverse functions of these metals, different metals are chosen based on desired properties:

- Structural property: metals like Zr, Hf, lanthanides, *etc.* are often selected to synthesize cluster-based MOFs.
- Physical stability: metals such as Zr, Al, Cr, Fe, *etc.* are usually used to construct highly stable frameworks.
- Chemical reactivity: metals like Fe, Co, Ni, Cu, *etc.* are frequently employed for catalytic applications.
- Biocompatibility: metals such as Mg, Ca, Fe, Zn, *etc.* are generally chosen for biocompatible MOFs.

Additionally, the combination of metal salts and ligands should adhere to the HSAB theory to optimize coordination and stability.^{67,68} As for the selection of the anions, the departure ability of the anions and the solubility in certain solvents are often regarded as the primary consideration.^{69,70}

2.1.2 Ligand selection. The choice of ligands is a pivotal aspect in the design and synthesis of MOFs (Fig. 6). Ligands can connect to the metal nodes to form the periodically extended network structures. The selection of appropriate ligands profoundly influences the framework's architecture, stability, porosity, and functionality, making it crucial for tailoring MOFs to target applications such as gas storage, catalysis, drug delivery, and sensing. Various ligands are often employed, mainly including the carboxylates, imidazolates, phosphonates, pyrazoles, triazoles, *etc.*^{71–75}

Benefiting from the diverse choice of these ligands, different ligands are chosen based on desired properties:

- Architecture: ligands determine the connectivity and geometry of MOFs, mainly including the length, rigidity, number of joints, and functional groups of the ligands.
- Framework stability: the properties of the ligands affect the stability of MOFs. For example, ligands with robust



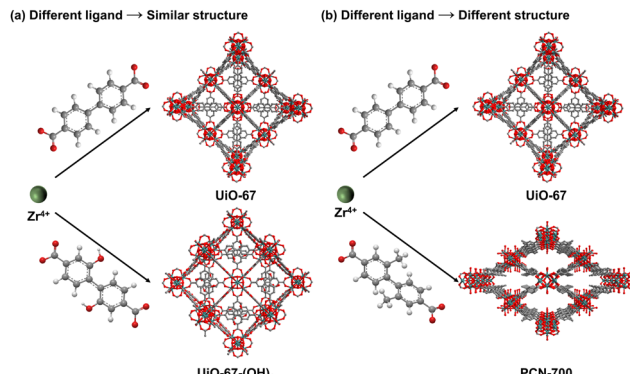


Fig. 6 Ligand selection influence on the structures of MOFs under similar conditions. Atom code: C, grey; O, red; H, white; Zr, dark green.

chemical bonds can enhance the framework's resistance to hydrolysis, thermal degradation, and chemical attack.

- **Variable porosity:** the size and shape of ligands can be varied to tune the porosity of the MOFs, affecting its capacity for gas adsorption, separation, and storage.

- **Functional property:** ligands impart specific functionalities to MOFs with their target functional groups, which is beneficial to form excellent host–guest interactions in applications, for example, hydrogen bond, halogen bond, $\pi \cdots \pi$ interaction, *etc.*

Also, the choice of ligands should adhere to the HSAB theory to optimize the coordination interaction and enhance the stability. Furthermore, in some cases, the organic linkers will be transformed to their corresponding metal salts for better solubilities in certain solutions.^{76,77}

2.1.3 Solvent selection. The choice of solvent plays a crucial role in the synthesis of MOFs (Fig. 7). Solvents do not only act as the medium in which the reactants dissolve and interact but also influence the crystallization process, morphology, and properties of the MOFs. The selection of the appropriate solvent is essential for optimizing reaction conditions, achieving high-quality crystals, and tailoring the properties of the resulting MOFs for specific applications. Some commercial chemicals with great solubilities were used, including water, methanol/ethanol, *N,N*-dimethylformamide (DMF), *N,N*-dimethylacetamide (DMA), chloroform, *etc.*^{78–83} Also, in many cases, the extra additions of acid or base can be beneficial

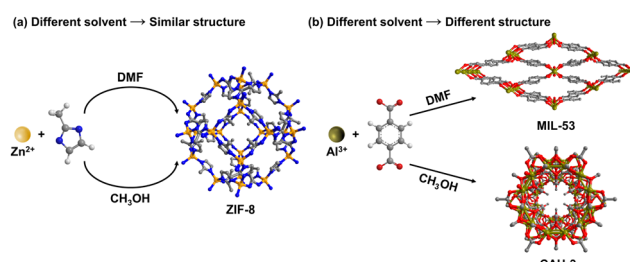


Fig. 7 Solvent selection influence on the structures of MOFs under similar conditions. Atom code: C, grey; O, red; H, white; Zn, yellow; Al, dark yellow.

for generating higher quality crystals or higher crystallization rates.^{84,85}

From the various solvents, there are some basic conditions for consideration:

- **Solubility:** solvents determine the solubility of metal salts and organic linkers, affecting the availability of reactants and the overall reaction kinetics. Suitable solvents can enhance the reactivity and facilitate the formation of desired coordination bonds.

- **Saturation vapor pressure:** the saturation vapor pressure of the solvent affects the pressure in the reactor, which further influences the reaction rate and the sizes and shapes of the products.

- **Chemical compatibility:** the solvent must be chemically compatible with the reactants and products, avoiding unwanted side reactions or decomposition.

- **Environmental and safety issues:** solvent toxicity, flammability, and environmental impact are also crucial factors. Green solvents such as water and ethanol are preferred for their lower environmental impact and safer handling properties.

2.1.4 Reaction condition selection. The choice of reaction conditions can also influence a lot in MOF synthesis, mainly including the reaction temperature, reaction time, cooling steps, the synthesis methods, *etc* (Fig. 8).^{86–90} These parameters significantly influence the nucleation, crystal growth, and overall quality of the MOFs. Optimizing these conditions is essential for tailoring the structural, physical, and chemical properties of MOFs to meet specific application needs.

From the various reaction conditions, there are some basic conditions for consideration:

- **Reaction temperature:** the synthesis temperature controls the kinetic energy of the reactants, influencing the rates of nucleation and crystal growth. Higher temperatures generally accelerate reaction rates and promote the formation of larger crystals, but they must be balanced to prevent decomposition of sensitive ligands or coordination structures.

- **Reaction time:** the duration of the reaction affects the extent of nucleation and crystal growth. Insufficient reaction time can lead to incomplete reactions and poor crystallinity, while excessive reaction time may cause overgrowth or degradation of the framework.

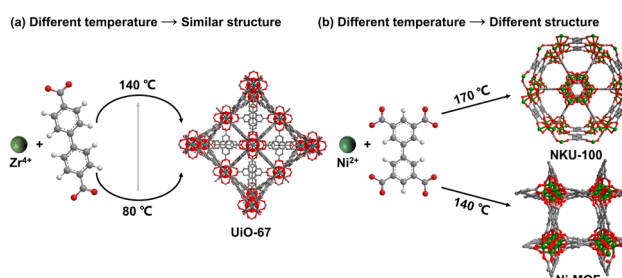


Fig. 8 Temperature selection influence on the structures of MOFs under similar conditions. Atom code: C, grey; O, red; H, white; Zr, dark green; Ni, green.

- Cooling steps: the rate at which the reaction mixture is cooled affects the crystallization process. Slow cooling allows for the formation of well-defined, high-quality crystals, whereas rapid cooling often results in smaller, less ordered structures.

- Synthesis methods: the synthesis methods generally include solvothermal method, microwave method, electrochemical synthesis, mechanical grinding, ultrasonic synthesis, and near-critical water synthesis. These methods can greatly affect the reaction time through the different energy intervention forms, which will be discussed in detail below.

- Activation methods: removing solvent residues from the pores of as-synthesized MOFs are crucial step for applications such as adsorption and catalysis. The solvent residues are typically trapped in the MOF's pores after synthesis, and removing them often involves solvent exchange and thermal activation. In solvent exchange, generally a more volatile solvent replaces the initial solvent within the pores, while thermal activation is typically performed by heating under vacuum. In some cases, supercritical CO_2 drying is used to remove solvents without causing structural collapse, which usually happens with conventional drying methods. This process of activation is critical to achieving a fully open and accessible pore structure, maximizing the surface area and allowing the material to function effectively in its intended applications.

In summary, traditional design principles necessitate a deep understanding of coordination chemistry and crystallography, along with numerous pairing attempts and adjustments. This approach has led to the development of many novel coordination structures and porous frameworks.

2.2 Synthesis method

The synthesis of MOFs involves a variety of methods, each offering control over the framework's structure, property, and scalability. The primary synthesis methods mainly include solvothermal method, microwave method, electrochemical synthesis, mechanical grinding, ultrasonic synthesis, and near-critical water synthesis. The choice of synthesis methods for MOFs has profound implications for their design and application potential, as different methods directly influence the structural property, functionality, and scalability of the resulting materials. MOF design is intrinsically linked to the desired application, and selecting the appropriate synthesis approach is critical to tailoring specific properties like porosity, stability, and catalytic activity.

For instance, solvothermal method is widely used for producing highly crystalline MOFs with uniform pore structures, making them ideal for gas storage, separation, and catalysis. However, this method often requires high temperatures, long reaction time, and toxic solvents, posing challenges for scalability and environmental sustainability. Rapid synthesis techniques like microwave method, and ultrasonic synthesis have emerged as promising methods for producing MOFs quickly. These methods offer great potential for applications where time efficiency is critical, such as in commercial-scale production or the development of MOF-based sensors. However, controlling the morphology and ensuring the homogeneity of the materials

synthesized through these faster methods can be difficult, which may affect their performance in applications that require precise structural control. Greener synthesis technique, such as mechanical grinding, offer more sustainable alternatives by reducing solvent usage and energy consumption, which is well-suited for applications where environmental impact and cost efficiency are critical, such as large-scale industrial processes and environmental remediation. Yet, achieving the same level of structural precision and crystallinity as traditional methods can be challenging, which may limit their application in areas requiring highly ordered frameworks, like selective catalysis or drug delivery.

Ultimately, the synthesis method must be carefully chosen to align with the desired application, as it affects not only the material's physical and chemical properties but also its feasibility for real-world use. Meanwhile, for energy-related applications, such as hydrogen storage or battery technologies, the synthesis process must prioritize high surface area, stability, and ease of mass production. Balancing these factors—efficiency, scalability, sustainability, and functional precision—remains one of the central challenges in optimizing MOF design for diverse applications.

2.2.1 Solvothermal method. Solvothermal synthesis is the most common method for producing MOFs, where metal salts and organic linkers are dissolved in a solvent and heated in a reaction vessel to promote crystal growth. The particle size and shape can be regulated and controlled by adjusting the reaction temperature, reaction time, and cooling steps. Generally, solvothermal synthesis allows for high crystallinity and purity.^{91–94}

Various reaction vessels are frequently used in experimental operations, mainly including glass vials, glass flasks, pressure-resistant bottles, Schlenk bottles, and Teflon autoclaves (Table 1). The primary advantage of glass vessels over Teflon autoclaves is their ability to facilitate real-time observation of crystal growth processes.

Glass vials with plastic lids are inexpensive but can hardly withstand temperatures over 100 °C and are prone to air leakage, resulting in low pressure. In contrast, vials with metal lids are more expensive but can endure higher temperatures.

Compared to other vessels, glass flasks are suitable for larger-scale synthesis, though they generally cannot be sealed for safety reasons, which highly limits the application range for the atmospheric pressure condition.

Pressure-resistant bottles offer a higher sealing capability and can withstand higher pressures, though the internal pressure must be estimated beforehand for safety, which requires a low liquid level height, high solvent boiling temperature, and low reaction temperature.

Schlenk bottles are used for some special reactions requiring low humidity or an oxygen-free environment, mainly involving some unstable metals or ligands. Also, this type of vessels mainly provides the atmospheric pressure condition.

Teflon autoclaves are among the most widely used vessels due to their excellent resistance to heat, acids, and alkalis, allowing them to withstand much higher temperatures and pressures. Besides, autoclaves are much safer due to the



Table 1 Main pros and cons of different reaction vessels for MOF synthesis

Vessels	Pros	Cons
Glass vial	<ul style="list-style-type: none"> • Real-time observation • Cheap and convenient 	<ul style="list-style-type: none"> • Low temperature and pressure
Glass flask	<ul style="list-style-type: none"> • Real-time observation • Large-scale synthesis 	<ul style="list-style-type: none"> • Normally atmospheric pressure
Pressure-resistant bottle	<ul style="list-style-type: none"> • Real-time observation • Withstand high pressure 	<ul style="list-style-type: none"> • Safety issue
Schlenk bottle	<ul style="list-style-type: none"> • Real-time observation • Low humidity or oxygen-free environment 	<ul style="list-style-type: none"> • Normally atmospheric pressure
Teflon autoclave	<ul style="list-style-type: none"> • Withstand high pressure • High safety 	<ul style="list-style-type: none"> • Unobservable reaction process

durable metal shell following standard operations. However, the device is relatively expensive and inconvenient.

2.2.2 Microwave method. Microwave heating operates on the principle of dielectric heating, where the energy is absorbed by molecules in the reaction mixture, causing them to oscillate and generate heat through molecular friction. This process results in extremely fast and uniform heating, which is particularly beneficial for reactions that require precise temperature control. Microwave-assisted synthesis has emerged as a powerful and versatile technique in the field of chemical synthesis, particularly for the fabrication of MOFs. This method leverages microwave radiation to heat the reaction mixtures, offering distinct advantages over conventional heating methods. The rapid and uniform heating provided by microwaves can significantly shorten the reaction time from several days to several hours, improve product yields, and allow for better control over the physical properties of the synthesized materials, like the sizes and shapes. This technique enables the rapid screening of different metal–ligand–solvent combinations, accelerating the discovery of new MOFs with target properties and functionalities. Additionally, the ability to precisely control reaction conditions makes it possible to fine-tune the porosity, surface area, and stability of the resulting frameworks.^{95–97}

2.2.3 Electrochemical synthesis. Electrochemical synthesis of MOFs operates on the principle of electrolysis, where an electric current is applied to an electrolyte solution to drive the assembly of metal ions and organic ligands. This method allows for precise control over the concentration of metal ions, reaction kinetics, and deposition rates, leading to high-quality MOF samples. Electrochemical synthesis has gained prominence as an innovative and efficient method for the fabrication of MOFs, which utilizes an electrochemical cell to drive their formation, offering distinct advantages in terms of reaction control, purity of products, and environmental sustainability. During the synthesis process, metal ions are continuously supplied from the anode, enabling the controlled growth of MOFs with certain sizes and morphologies. Electrochemical synthesis has been successfully employed to produce a diverse range of MOFs with tailored properties for specific applications. This method is particularly advantageous for the synthesis of thin films and coatings of MOFs on various substrates, which are important for applications in catalysis, sensing, and energy storage. The ability to precisely control the deposition

process makes it possible to engineer MOFs with desired morphologies and functionalities.^{98–100}

2.2.4 Mechanical grinding. Mechanical grinding has emerged as an innovative and sustainable alternative in MOF synthesis. This technique, a branch of mechanochemistry, involves the use of mechanical force to induce chemical reactions between solid reactants, typically using a ball mill or mortar and pestle. Mechanical grinding offers several advantages over conventional methods, such as reduced solvent usage, lower energy requirements, and shorter reaction times. Additionally, it often results in high product yields and can facilitate the synthesis of MOFs with unique properties unattainable through traditional routes. The mechanochemical approach for MOF synthesis is not only environmentally friendly but also opens new avenues for the design and discovery of novel materials. By leveraging the principles of green chemistry, mechanical grinding represents a paradigm shift towards more sustainable and efficient production of MOFs, aligning with the growing demand for eco-friendly and cost-effective manufacturing processes in materials science.^{101–103}

2.2.5 Ultrasonic synthesis. Ultrasonic synthesis also serves as a promising alternative technique for MOF fabrication, leveraging the power of ultrasonic waves to drive chemical reactions. This method, also referred to as sonochemistry, utilizes high-frequency sound waves to generate localized high-energy conditions through acoustic cavitation—the formation, growth, and implosive collapse of bubbles in a liquid medium. The extreme temperatures and pressures generated by this process facilitate rapid and efficient chemical reactions, often leading to the formation of MOFs in significantly shorter times and under milder conditions compared to solvothermal method. The cavitation effect not only accelerates the reaction kinetics but also enhances the uniformity and crystallinity of the resulting MOF structures.^{104–106}

2.2.6 Near-critical water synthesis. Near-critical water synthesis is a newly developed method. High-temperature water is being studied as a medium for processes such as organic reaction, waste destruction, and nanoparticle formation. This field arises because properties of water change dramatically as it approaches the critical point. For example, the dielectric constant decreases to values more typical of non-polar solvents, making organic compounds, such as the ligands used for the synthesis of MOFs, soluble. Additionally, the ionic



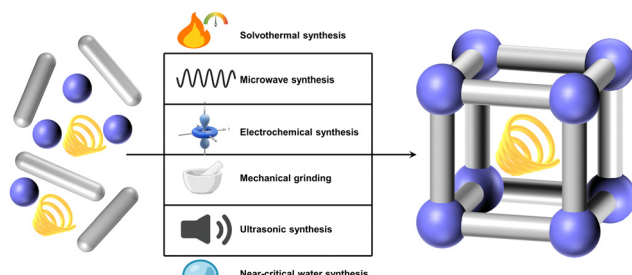


Fig. 9 Classical synthesis methods of MOFs.

product of the reaction solution increases by up to three orders of magnitude, reaching a maximum at around 280 °C, and creating a solvent with simultaneously raised concentrations of H⁺ and OH⁻ ions. Furthermore, the water can be reused in the reaction without purification. If necessary, processes such as ion exchange can be employed to remove unreacted carboxylic acids and metal ions.^{107–110}

In summary, the synthesis methods have effectively utilized various forms of energy intervention to achieve highly efficient energy usage, resulting in more efficient, cost-effective, and environmentally friendly synthesis processes (Fig. 9).

3. Functional oriented design and modification method

Functional oriented design involves the selection of appropriate metal nodes, organic linkers, and solvents for the MOF's structures, guided by the desired properties and applications, particularly in areas such as luminescence, chirality, catalysis, separation, storage and delivery, and electrochemistry. Modifications can be achieved through post-synthetic treatments, focusing primarily on the modification of metals (metal exchange and coordination modification), linkers (linker functionalization, linker scissoring, and linker exchange), and guests (guest exchange) (Fig. 10). These modifications enhance

the MOF's target performance. By employing a functional-oriented design and modification strategy, researchers can optimize MOFs for specific tasks, creating bespoke materials that address complex challenges in various technological domains. This methodology not only broadens the scope of MOF applications but also paves the way for innovative solutions to contemporary scientific and industrial needs.

The HSAB theory and dynamics and thermodynamic equilibrium also play a significant role in the modification of MOFs, offering valuable insights into how different reagents interact with the metal centers and ligands within a MOF structure. By applying HSAB principles, researchers can design more post-modifications to tailor MOF properties for specific applications, such as catalysis, gas storage, and sensing. This understanding helps ensure that modifications are not only efficient but also result in stable, functional materials that perform well under the desired operating conditions. While for the dynamics and thermodynamic equilibrium, unlike the initial MOF synthesis, where the formation of coordination bonds and crystal growth are dictated by the primary synthesis conditions, the functional modification often involves chemical reactions or exchanges that occur within the pre-established framework. This requires careful control over both kinetic factors (reaction rates, diffusion of reactants) and thermodynamic principles (stability, equilibrium of products). The ability to control reaction rates, optimize conditions for achieving equilibrium, and ensure the long-term stability of modified MOFs opens new possibilities in tailoring these materials for complex tasks, such as catalysis, gas storage, sensing, and drug delivery.

3.1 Functional oriented design

Based on the accumulated experience from numerous syntheses attempts and studies of the relationships between the structures and the properties of MOFs, the functional oriented design of MOFs with targeted functions, or the modification of MOFs to achieve specific functionalities, has attracted great attention. This approach significantly expands the application prospects and reduces the time needed for exploration and development.

3.1.1 Luminescence. The luminescent properties of MOFs make them highly attractive for applications in sensing, imaging, light-emitting devices, and photonics.^{111–115} These materials can exhibit diverse photophysical behaviours such as fluorescence, phosphorescence, and delayed fluorescence, depending on their composition and structure. By engineering the electronic environments of the metal nodes, the conjugation pathways of the organic linkers, and the specific properties of functional guests, researchers can create MOFs with targeted emission wavelengths and intensities tailored to certain needs.^{116–120}

Additionally, luminescent MOFs offer unique advantages in terms of stability, porosity, and functionality compared to traditional luminescent materials. Their porous nature allows for guest encapsulation, which can lead to enhanced luminescent properties or novel sensing mechanisms. The modular design of MOFs also enables the incorporation of multiple

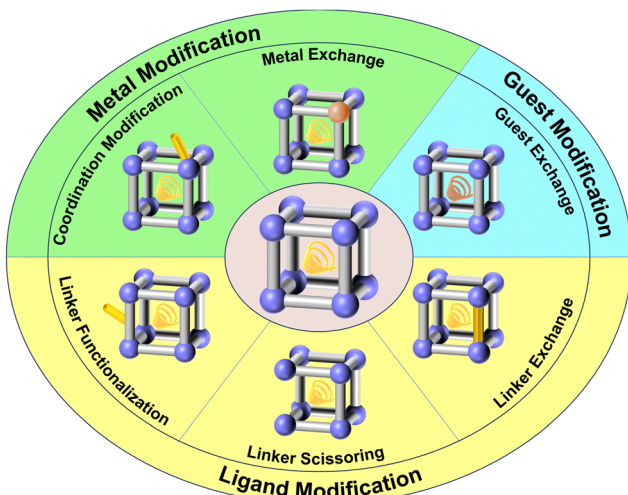


Fig. 10 Classical modification methods for MOFs.



luminescent centers within a single framework, facilitating multifunctional applications.^{121–125}

Due to the abundant combinations of metals, ligands, and guests, MOFs can possess various luminescence sources and multiple luminescence centers, simultaneously. Metals such as Cu(I), Ag(I), Eu(III), and Tb(III) are frequently employed as luminescent centers, while other metals that do not exhibit strong competitive absorption of excitation light, like Mg(II), Zn(II), Al(III), and Zr(IV), are used as metal nodes to link luminescent ligands. Ligands with highly conjugated structures or functional groups, such as pyrene rings and amino groups, are also often used as luminescent sources. As for the guests, metal ions like Eu(III) and Tb(III), coordination units such as $[\text{Ir}(\text{ppy})_3(\text{bpy})]^+$, and luminescent dyes can serve as additional luminescent centers within MOFs (Fig. 11).^{126–130} These diverse components contribute to the versatile luminescence properties of MOFs, enabling their use in a wide range of applications. Besides the direct synthesis to introduce luminescence centers into MOFs, nearly all the classical modification methods can bring the luminescence centers into MOFs, which enriches the luminescence sources and makes it easy for MOFs to possess multiple luminescence centers.

3.1.2 Chirality. Chirality, a concept widely recognized in organic chemistry, extends its intriguing characteristics to MOFs, creating a fascinating intersection of geometry, symmetry, and functionality. The chirality in MOFs can arise from several sources, including the inherent chirality of the organic linkers, the chiral arrangement of the framework itself, or the incorporation of chiral guests within the pores. This structural handedness imparts unique properties to chiral MOFs, such as the ability to differentiate between enantiomers of a compound, a feature highly desirable in pharmaceuticals, pesticides, and fine chemicals.^{131–135}

Understanding and harnessing the chirality of MOFs involves a multidisciplinary approach, integrating principles from coordination chemistry, crystallography, and materials science. Researchers are continually developing new synthetic strategies to design and construct chiral MOFs with tailored properties, aiming to exploit their chiral nature for practical applications, such as the enantioselective recognition, separation, asymmetric catalysis, and nonlinear optics. This field is not only rich in fundamental scientific inquiry but also holds promise for innovative solutions to complex industrial challenges.^{136–138}

The introduction of chirality into MOFs is a vibrant area of research that combines the precision of chemistry with the ingenuity of materials science. While several methodologies are employed to introduce chirality into MOFs, each leveraging different aspects of chemical design and synthesis. The most common approach is the use of pure chiral ligands during the MOF synthesis, while the organic molecules that are inherently chiral. These chiral ligands can transfer their chirality to the entire framework, resulting in chiral MOF structures. Another method involves the incorporation of achiral ligands in a way that induces a chiral arrangement within MOFs. This can be achieved through the choice of specific metal centers and reaction conditions that favour the formation of chiral frameworks. Post-synthetic modification is another powerful strategy, where an initially achiral MOF is functionalized with chiral molecules, thereby imparting chirality to the framework, which mainly includes the coordination modification to the metal nodes, the chiral linker functionalization and exchange to the ligands, and the ion/molecule exchange to the guests. Additionally, templating methods, where chiral molecules or ions are used as templates during the synthesis, can also guide the formation of chiral MOFs by influencing the spatial arrangement of the building blocks (Fig. 12).^{139–145}

The precise control over the chirality of MOFs through these methods not only enhances our understanding of chiral materials but also creates new avenues for practical applications. By tailoring the synthesis process, researchers can design chiral MOFs with specific pore sizes, shapes, and functionalities, enabling their use in a wide range of advanced technological applications.

3.1.3 Catalysis. MOFs have emerged as a versatile class of materials in the field of catalysis, offering a unique platform for facilitating a wide range of chemical reactions, which exhibit many attributes that are particularly advantageous for catalysis. The large surface area and porosity allow for the accommodation and diffusion of reactant molecules, enhancing the contact between reactants and active sites. The abundant potential open metal sites and switchable dynamic metal sites also give rise to the sufficient opportunities for contact. The modular nature of MOFs indicates that their components can be precisely designed and synthesized to incorporate specific catalytic sites, such as metal ions, metal clusters, or functionalized organic linkers. This design flexibility enables the creation of MOFs tailored for specific catalytic processes, ranging from

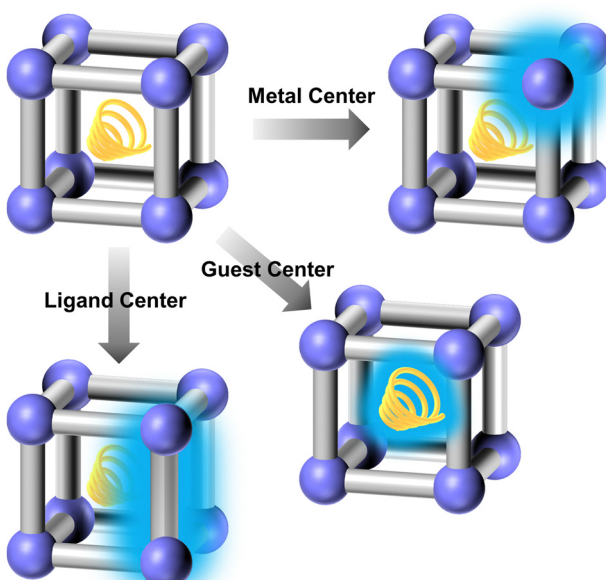


Fig. 11 Various luminescent centers in MOFs.



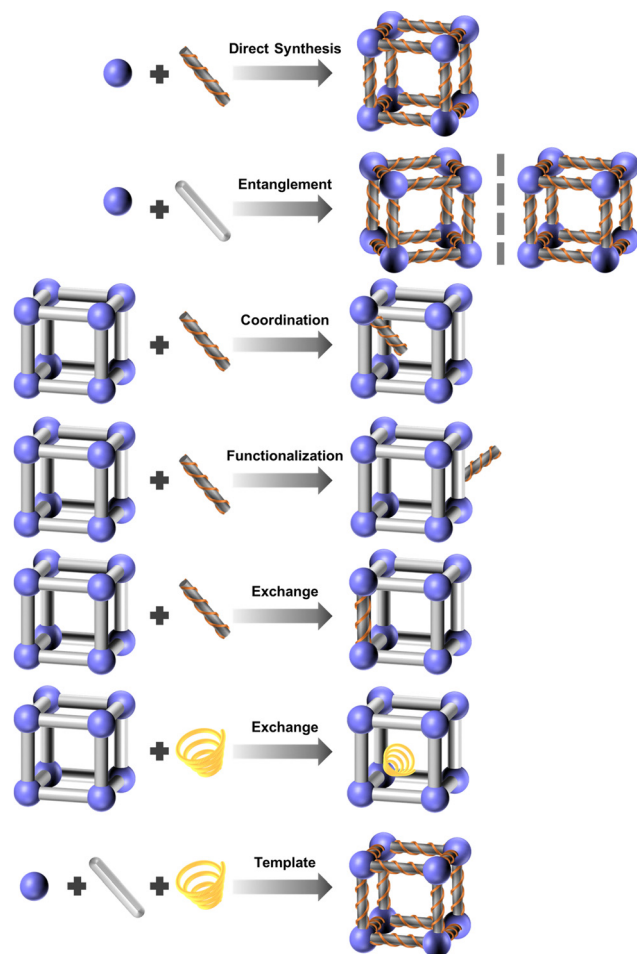


Fig. 12 Classical construction methods for chiral MOFs. The original guests in the framework have been omitted for clarity.

organic transformations and gas adsorption to photochemical and electrochemical reactions.^{146–150}

Besides, the MOF catalysts possess the potential for heterogeneity and recyclability. Unlike many traditional homogeneous catalysts, MOFs can often be recovered and reused without significant loss of activity, offering economic and environmental benefits. Additionally, the stability of MOFs under various reaction conditions, including high temperatures and acidic or basic environments, further extends their applicability in industrial processes. Researches into MOF-based catalysis are continuously evolving, with innovative strategies being developed to enhance the performance and scope of these materials. The confluence of these advancements positions MOFs as a promising frontier in catalysis, with the potential to revolutionize processes in chemical manufacturing, environmental remediation, and sustainable energy solutions.^{151–154}

Many functional metals and groups have been widely introduced into MOFs both through direct synthesis and the post-synthetic modification, mainly through the coordination modification and the guest exchange methods, especially for the transition metal elements Fe, Co, Ni, Cu, Ag and functional

linkers porphyrin and phthalocyanine. The abundant and highly adjustable metal sites for MOFs give rise to the efficient constructions of novel catalysts for certain usage.^{155–160}

3.1.4 Separation. By exploiting precise control through direct synthesis or modification of pore dimensions, pore shapes, and surface functionalities, MOFs can selectively capture and separate specific ions or molecules from complex mixtures in both liquid and gaseous states, which highly relays on the pore size and inner chemical environments. This capability offers significant advancements in applications ranging from environmental remediation, clean energy production to industrial purification and biochemical processing.^{161–165}

There are two main research directions in this field: energy utilization and environmental protection. For energy utilization, MOFs have shown great potential in separating petroleum products, including gaseous products like low-chain alkanes, olefins, and alkynes, as well as liquid products such as C₆, C₇, and C₈ mixtures, and downstream products like xylene (Table 2). Additionally, crucial metal resources, such as Li⁺, Co²⁺, Ni²⁺, and Ln³⁺, can also be efficiently separated using MOFs.^{166–170}

For environmental protection, MOFs have demonstrated excellent separation effects for a series of toxic gases, ions, and persistent organic pollutants. Examples include SO₂ and NO₂ in automobile exhaust (Table 3), heavy metal ions in water environments, and perfluorinated compounds and dyes in natural environments. As the field continues to evolve, the integration of MOFs into separation technologies promises to enhance efficiency, reduce energy consumption, and contribute to more sustainable practices across diverse industries.^{173–177}

3.1.5 Storage and delivery. MOFs have garnered significant attention in the fields of storage and delivery due to their unique structural and pore properties, which make MOFs ideal

Table 2 Xylene separation abilities of selective MOFs in vapor phase

MOF	Selectivity	Ref.
MAF-X8	<i>p</i> - > <i>m</i> - > <i>o</i> - > eb	166
Mg-CUK-1	<i>p</i> - > <i>o</i> - > <i>m</i> -	167
sql-1-Co-NCS	<i>o</i> - > <i>p</i> - > <i>m</i> - > eb	168
MIL-101	<i>o</i> - > eb > <i>m</i> - > <i>p</i> -	169
Co ₂ (bdc)	<i>o</i> - > eb > <i>m</i> - > <i>p</i> -	170
ZU-61	<i>o</i> - > <i>m</i> - > <i>p</i> -	171
MFM-300	<i>m</i> - > <i>o</i> - > <i>p</i> -	172

Table 3 SO₂ and NO₂ adsorption capacities of selective MOFs at 1 bar and 298 K

MOF	Capacity (mmol g ⁻¹)	Ref.
[Ir]@NU-1000	SO ₂ 10.9	178
Ni(bdc)(ted) _{0.5}	SO ₂ 10.0	164
SIFSIX-1-Cu	SO ₂ 11.0	176
MFM-601	SO ₂ 12.3	179
MIL-101(Cr)	SO ₂ 18.4	180
MFM-170	SO ₂ 17.5	181
MFM-300-Sc	SO ₂ 9.4	182
ECUT-111	SO ₂ 11.6	183
MFM-300-Al	NO ₂ 14.1	184
MFM-300(V)	NO ₂ 13.0	185



candidates for the efficient storage and delivery of a wide range of substances, including gases, liquids, and certain therapeutic agents, through guest exchange processes,^{186–190} which highly relays on the pore volume and inner chemical environments.

In the context of gas storage, MOFs offer solutions for storing and delivering hydrogen, methane, and other gases with high volumetric and gravimetric capacities, presenting promising advancements for clean energy applications. Additionally, MOFs demonstrate superior performance in storing and releasing volatile organic compounds, which is crucial for environmental monitoring and industrial processes. MOFs' ability to encapsulate and release drugs in a controlled manner also holds significant potential for targeted drug delivery systems, improving the efficacy and reducing side effects of various treatments.^{191–195}

As research progresses, the integration of MOFs into storage and delivery systems promises to revolutionize various industries by enhancing efficiency, increasing capacity, and providing more sustainable and controlled methods of substance management. The versatility and adaptability of MOFs underscore their potential to address contemporary challenges in energy storage, medical treatments, and environmental protection, heralding a new era of advanced materials science.

3.1.6 Electrochemistry. MOFs have emerged as a highly promising class of materials in the field of electrochemistry, offering transformative potential for a range of applications, including energy conversion, energy storage, and energy applications.

In energy conversion, MOFs offer significant potential to enhance the efficiency, selectivity, and sustainability of various electrochemical reactions, especially in oxygen evolution reaction (OER), hydrogen evolution reaction (HER), oxygen reduction reaction (ORR), and carbon dioxide reduction reaction (CO₂RR), which convert the electrical energy into chemical energy. Target metals are frequently employed as the catalyst sites, including Fe, Co, Ni, Cu, and In.^{196–200}

In energy storage, MOFs have shown great promise in the development of advanced batteries and supercapacitors. Their high porosity and customizable surfaces enable the efficient storage and rapid transfer of ions, leading to improved charge/discharge rates and increased capacity. MOFs are also being explored for use in fuel cells and electrolyzers, where their structural versatility allows for the optimization of catalytic processes, enhancing the overall efficiency of energy conversion and storage.^{201–205}

As for the energy applications, MOFs are making significant strides in the field of electrochemical sensing and environmental monitoring. The precise control over their chemical functionalities and pore environments allows for the selective detection of various analytes, including gases, ions, and organic compounds, with high sensitivity and specificity.^{206–210}

As researches continue to advance, the integration of MOFs into electrochemical applications promises to drive significant improvements in performance, sustainability, and functionality. This integration not only opens new avenues for energy storage and conversion but also enhances capabilities in

environmental sensing and beyond, marking a pivotal step forward in the application of advanced materials in electrochemistry.

3.2 Modification method

Post-synthetic modification involves the deliberate alteration of the MOF's structure and functionality based on its initial synthesis, which allows for the exchange of metal centers, introduction of new functional groups, or the alteration of the guests within the MOF framework without disrupting its overall architecture. Target-guided modification provides a versatile toolkit for tailoring the physical and chemical environment within the pores, thus optimizing MOFs for specific applications. Post-synthetic modification allows researchers to introduce new functionalities, improve stability, or adapt the structural features of a pre-synthesized MOF without altering its underlying framework. This capability is especially valuable for applications in gas storage, catalysis, drug delivery, and sensing, where specific properties such as pore size, chemical reactivity, or surface hydrophilicity are critical. For example, modifying the metal center can enhance catalytic efficiency or improve sensing capabilities, while the modification towards the linker can alter the physical and chemical environment of the framework and pores, affecting gas storage, catalysis, and sensing performance. Additionally, the modification to the guest molecules can also influence the pore environment, leading to changes in storage capacity and delivery efficiency.

3.2.1 Metal exchange. Metal exchange on the framework of MOFs involves replacing the metal ions within the pre-formed framework with different metal species, which mainly involves fourth row transition metals and lanthanide metals, respectively (Fig. 13). This process normally can be performed under relatively mild conditions, preserving the integrity of the MOF structure while enabling the selective substitution of metal ions, which can significantly alter the structural, electronic, and catalytic properties of MOFs, thereby opening new possibilities for customization and application-specific optimization. Metal exchange is a post-synthetic modification strategy that enables researchers to fine-tune the functionality of MOFs without the need to synthesize entirely new frameworks from scratch.^{211–215}

Metal exchange greatly expands MOFs with various properties and functions. It allows for the introduction of metals which may be challenging to incorporate during the initial synthesis due to incompatibility with the organic ligands or

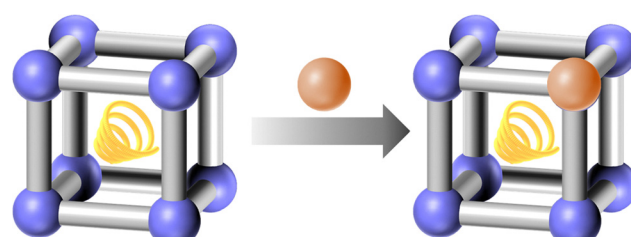


Fig. 13 Metal exchange process for MOFs.



reaction conditions. Additionally, metal exchange can be used to create multi-metallic MOFs, which can exhibit synergistic properties that are not attainable with single-metal frameworks. This flexibility is particularly beneficial for catalytic applications, where the nature of the metal center plays a crucial role in determining the activity and selectivity of the catalyst.

3.2.2 Coordination modification. Coordination modification involves altering the coordination environment around the metal nodes or clusters within MOFs with potential open metal sites after its initial synthesis (Fig. 14), which can significantly impact the physical and chemical properties of MOFs, allowing for fine-tuning of its functionalities to meet specific application requirements. This process can also be performed under relatively mild conditions, preserving the integrity of the MOF structure. By modifying the coordination sphere, researchers can introduce new reactive sites, adjust the electronic properties, and improve the stabilities of MOFs.^{216–220}

The advantages of coordination modification are numerous. It offers a versatile approach to introduce functional groups that may not withstand the conditions of direct MOF synthesis. This method enables the precise control of the metal-ligand interactions and the creation of coordinatively unsaturated sites, which are essential for catalytic activities and enhanced adsorption properties. Coordination modification also facilitates the incorporation of multiple types of ligands, leading to multifunctional MOFs with synergistic properties.

Traditional coordination modification often involves one-end coordination to the metal nodes. Recently, the sequential linker installation method, which aims at dual-end coordination, has attracted great attention due to their highly defined position and the great stability of the installed linkers. By enabling the precise adjustment of the organic linkers within the framework, this method allows for the creation of highly

specialized MOFs tailored to meet the demands of advanced technological applications.^{221–225}

3.2.3 Linker functionalization. Linker functionalization involves the post-synthetic modification of the linkers within the MOF structure. This process allows for the introduction of various functional groups into the framework, tailoring the physical and chemical environment within the pores without altering the overall architecture of the MOF. By modifying the linkers, researchers can significantly regulate the properties of the MOF, including its reactivity, stability, and interaction with guest molecules. Generally, there are three main pathways for linker functionalization: organic reactions,^{226–230} coordination interactions,^{231–233} and partial exchanges^{234,235} (Fig. 15).

For MOFs with certain functional groups, such as aldehyde group, amino group, carboxyl group, double bond, and triple bond, organic reactions can introduce molecules with complementary functional groups to enhance specific properties and functions. In MOFs that possess ligands with potential coordination sites, like carboxyl groups, hydroxyl groups, and nitrogen-containing heterocyclic units (pyridine, pyrazine, piperazine, *etc.*), metals can be attached to the ligands through coordination interactions. Additionally, for MOFs whose ligands contain coordination units, such as metal-porphyrin and metal-phthalocyanine, the metals within these ligands can be partially exchanged with other metals.

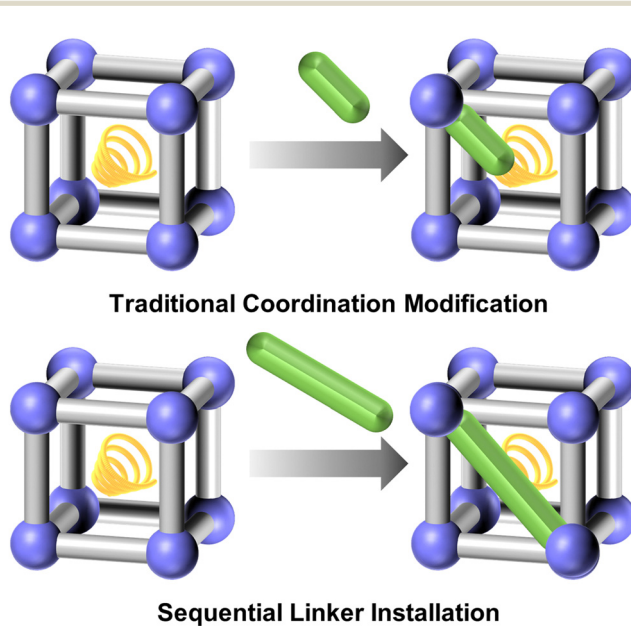


Fig. 14 Coordination modification process for MOFs.

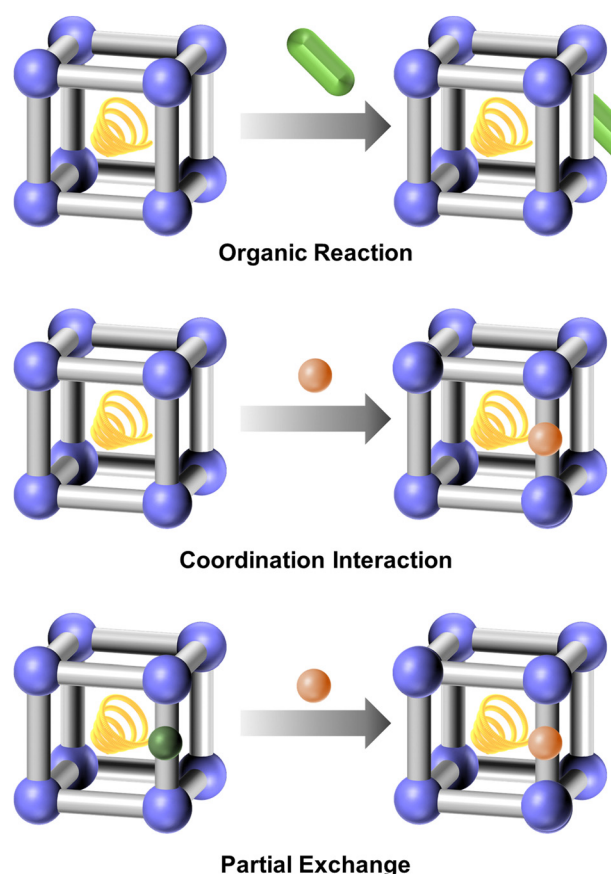


Fig. 15 Linker functionalization process for MOFs.



The advantages of linker functionalization are numerous. This technique enables the precise incorporation of functional groups that may not survive the harsh conditions of direct MOF synthesis, thus expanding the range of achievable modifications. Functionalized linkers can enhance the MOF's ability to interact with specific molecules, improving its performance in applications such as catalysis, where active sites are crucial, or in gas storage, where selectivity and capacity are paramount. Moreover, functionalization can be used to introduce responsive or stimuli-sensitive groups, creating smart materials that react to environmental changes. This functionalization strategy enables precise control over the physical and chemical environment within MOFs, facilitating the development of materials with tailored properties for advanced applications.

3.2.4 Linker scissoring. Linker scissoring involves the post-synthetic cleavage of the organic linkers within the MOF structures (Fig. 16), which strategically breaks the linkers, altering the framework's connectivity and porosity without destroying the MOF architecture completely. By selectively scissoring the linkers, researchers can introduce novel properties and functionalities, create porous defects, or modify the pore structures, significantly influencing the material's properties and enhancing its performance in various applications.^{236–240}

Pathways for linker scissoring mainly include chemical cleavage, where specific chemical reagents are used to break the linkers; photochemical scissoring, which uses light to induce cleavage; and mechanical methods, such as applying pressure or shear forces. Each method offers unique advantages and can be tailored to achieve the desired modifications while maintaining the overall integrity of the MOF structure.

The usage of linker scissoring allows for the generation of open metal sites and the creation of hierarchical porosity, which can improve the accessibility of active sites and enhance the material's catalytic activity. This pathway also enables the introduction of functional groups at the scission sites, facilitating further chemical modifications. Additionally, linker scissoring can be used to modulate the mechanical flexibility and stability of MOFs, making them more suitable for specific applications.

3.2.5 Linker exchange. Linker exchange involves the post-synthetic substitution of the organic linkers within a MOF structure with different linkers (Fig. 17), which allows for the introduction of new properties and functionalities without the need to synthesize new structures. The pathway

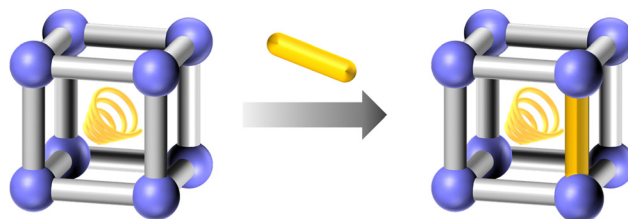


Fig. 17 Linker exchange process for MOFs.

of implementing this route is simple and facile, usually achieved by heating and soaking in the solvent or vapor phase of target chemicals, which provides specific advantages depending on the desired modification and the structural stabilities of MOFs.^{241–245}

The advantages of linker exchange are obvious. It offers a flexible approach to modify the properties and functions of MOFs, enabling the incorporation of functional groups that may not withstand the conditions of direct synthesis. This technique can also improve the material's stability and robustness, enhance its adsorption capacity, and introduce reactive sites for catalysis. Additionally, the linker exchange method allows for the development of multifunctional MOFs with synergistic properties by combining different types of linkers within the same framework.

3.2.6 Guest exchange. Guest exchange involves the post-synthetic replacement of ions or molecules within the pores of MOFs with different guest species (Fig. 18). This process can be performed under relatively mild conditions which allows for the dynamic modification of the MOF's properties without altering its fundamental structure. By exchanging guest molecules, researchers can fine-tune the MOF's pore environment, optimize its performance for specific applications, and introduce new functionalities.^{246–250}

Guest exchange provides a flexible approach to modify the internal environment of MOFs, enabling the introduction of various functional groups or active species that can be incompatible with direct synthesis. Guest exchange can generally enhance the material's stability, reactivity, and selectivity, making it more effective for specific applications such as molecular sensing and catalysis, where the presence of target guest molecules can significantly influence catalytic activity. Additionally, guest exchange can be used to create responsive MOFs that can adapt to different environmental conditions or stimuli.

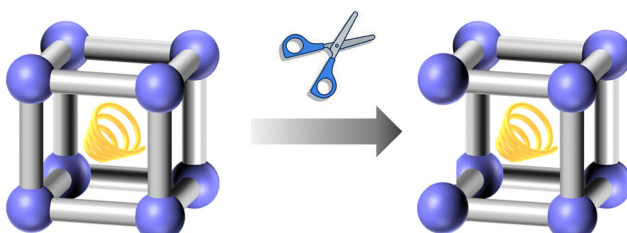


Fig. 16 Linker scissoring process for MOFs.

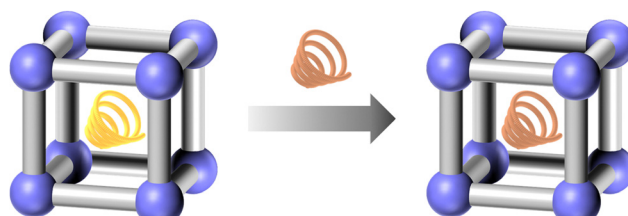


Fig. 18 Guest exchange process for MOFs.



4. AI methodologies and AI-guided synthesis

The recent technological advancements in ML and AI have sparked a proliferation of new systems and methodologies across various fields. From conversational agents like ChatGPT to breakthroughs in sports analytics,²⁵¹ medicine,²⁵² and beyond, AI has become an indispensable tool for predictions, information gathering, complex calculations, and other critical applications. In chemical research, ML models harness the power of AI and vast datasets to develop increasingly accurate predictions. These predictions can be experimentally validated, with the resulting data fed back into the models to further refine their accuracy. By identifying patterns in massive datasets that would be impractical for humans to detect within a similar timeframe, ML and AI allow scientists to focus their efforts on laboratory testing and validation of model-generated insights, thus accelerating the pace of discovery and innovation.²⁵³

4.1 AI methodologies

Before diving into how ML and AI can be used to help chemists perform more efficient research, it is crucial to understand how the ML models are developed, and how they use established databases to grow their algorithm. In this section, a general overview of database establishment for ML models, as well as training ML models to develop algorithms and perform better calculations and predictions will be presented in the context of MOF research.

4.1.1 Database establishment. To build a ML model, the system must first be fed up with data that is specific to the task which the user requires the model to perform. Instead of just explicitly coding a set knowledge base, ML allows for more fluid logic from a system by enabling the system to learn relationships through observing the data it is fed up with and finding patterns in it. This means that as more relevant data is given, not only successful trials but also failed trials, the model is able to offer better predictions.²⁵⁴ Database establishment is a crucial step in the process of making a functional ML model, and greatly helps the training processes. Once the model has been trained, it will then be validated to ensure accuracy before being used to solve tasks.

The four types of data that a database can be composed of are structured, unstructured, semi-structured, and metadata (Fig. 19).²⁵⁵ As the name suggests, structured data is highly organized. Think of an excel spreadsheet of addresses, with each column having one piece of the address. This type of data is considered structured and easy to handle by a ML model. Unstructured data does not have a predefined format, and therefore is much more difficult for the model to consume. Most normal internet media like images, portable document formats (PDFs), and word documents are considered unstructured. Semi-structured databases are not as structured as a structured database, but do have some organization pattern, such as hyper text markup language (HTML) data. Metadata is not a type of raw data, but instead is the extracted key points

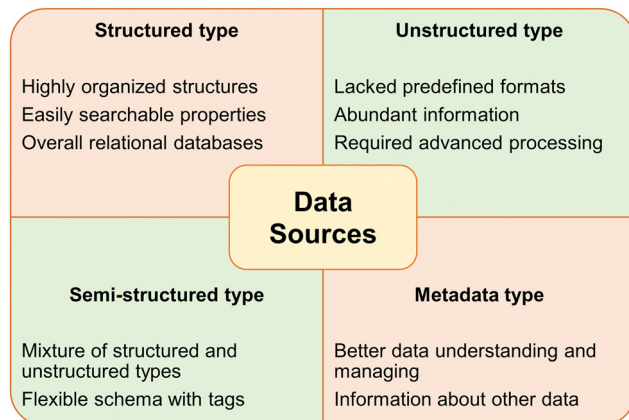


Fig. 19 Four types of data for machine learning.

from a full data set. Metadata presents the most important information for the user, without diluting it with the full data set.²⁵⁵ Framing these types of data into the context of MOF research and database establishment, if the ML model retrieves its data from PDFs of journal manuscripts or raw experimental data, it would be considered as an unstructured database, but if it retrieves the data from the Cambridge Structural Database (CSD) or other ordered database, it would be considered metadata. Once the data has been selected, it is classified as either training data or testing data. As the names suggest, training data is used to develop and optimize the ML models, while testing data is then used to evaluate whether the model can correctly assess a problem and produce a reasonable outcome.²⁵⁶

Due to the nature of chemistry, the information is often not in a format that is consumable by the ML model. For this reason, the data must first be refined to ensure it is in a language that the model can understand. Many different works may synthesize the same MOF structure, but there could be inconsistencies, missing data, or errors in some of the works. For this reason, all of the data for a single MOF must be compiled and refined into a single dataset, which can help make the model more accurate.²⁵⁷ Furthermore, each MOF structure must be simplified into a Simplified Molecular Input Line Entry System (SMILES), which is a language that can be understood by a ML model, and all other forms of media is translated in a similar way.²⁵⁸

There is an array of currently available MOF databases to be used as training and testing data for a ML model, including CoRE MOF, SynMOF, and CSD. CSD is the largest and most popular structural database available, containing not only MOF and other material structures, but also small molecules and other types of experimentally determined three-dimensional structural information, numbering over 1.4 million structures and including about 70 000 MOF structures.^{259,260} Many journals even require a submission of any crystallographic information file (CIF) to the CSD before publishing a manuscript, making the database a rich source for MOF data, and there are multiple examples of MOF databases built from CSD



data.^{261,262} While CSD is a rich resource for crystal structures, it is not exclusive to MOFs and so it would be counterintuitive to train a model using the full database. By filtering the CSD down while only MOF data is included, ML models are able to efficiently gather the data needed to run whatever task is required. There is even a subset data package within the CSD specifically for MOF data.²⁶³

CoRE MOF is a large database with over 14 000 MOF structures.^{258,264} In some structures, bound solvent molecules can affect the structure or can be a desired use for some materials. For this reason, the database is separated into free solvent removed (FSR) and all solvent removed (ASR), allowing a ML model to draw from whichever data type it requires.²⁵⁸ This database also includes searches for different types of organic linkers, possible open metal sites, as well as calculations for the structures. The CoRE MOF database draws from the CSD and supplements any missing structures by using Web of Science and manually imported CIFs. The data is then refined to be ready for computations, allowing it to be directly used as a source for a ML model. This makes CoRE MOF a ready-to-use database for ML and has been used for MOF screening purposes.^{265–267} CoRE MOF has even been used as a training database for a ChatGPT-type program that utilizes large language models, called ChatMOF.²⁶⁸ CoRE MOF has also been used to establish specific database for synthesis predictions, namely SynMOF. By extracting the synthesis information (metal, linker, time, temperature, solvent, and modulator) for the MOFs contained in the CoRE MOF database and combining it with the structural information contained in the CSDs, SynMOF is able to predict the synthesis conditions required for theoretical MOF structures.²⁶⁹ By utilizing SynMOF, it is possible to go from a predicted structure to the synthesis conditions required to generate it, enabling chemists to better determine the synthesis conditions, and saving time and chemicals. Another popular database that utilizes CoRE MOF data is MOFX-DB.²⁷⁰ A unique feature of MOFX-DB compared to the previous databases is that it includes both experimentally discovered and hypothetical structures retrieved from ML models for both MOFs and zeolites. MOFX-DB combines data from CoRE MOF,²⁵⁸ hypothetical MOF (hMOF),²⁷¹ and ToBaCCo²⁷² for MOF data, and IZA²⁷³ and PCOD-syn²⁷⁴ for zeolite structures. MOFX-DB also includes the calculated adsorption data for the MOFs, totaling over 160 000 MOFs, as well as all of the necessary data to reproduce the calculated data. This makes MOFX-DB a rich source for ML training, especially for finding ideal adsorbents.

4.1.2 Algorithm and model. Once the database has been established, the data can be used to train a ML model. There are four main ways to train a ML model: supervised, unsupervised, semi-supervised, and reinforcement learning, based on whether the data received by the system is labeled or not (Fig. 20).²⁵⁵ In supervised learning, the output data is labeled by a supervisor, usually a human, which is an explanation of the input data. The combination of the input and its corresponding labeled output explanation forms a training example, and a collection of these examples constitutes the training

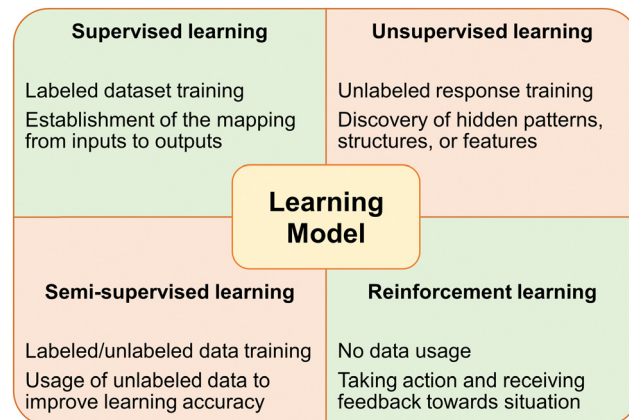


Fig. 20 Four main ways to train machine learning model.

dataset. The presence of the output explanation can be crucial for advancing the logic of the ML model, and since it is labeled by a human, the model gets a head start in understanding the logic it should apply. The two main types of supervised learning models are regression models and classification models.²⁷⁵

Unsupervised learning is defined if the output data is not manually tagged before being used for training. A lot of times, the volume of data is too large to be tagged manually, or the data cannot be tagged due to the type of data it is. When a ML model encounters this type of data, the goal is to extract trends or an underlying structure from the database. Due to the volume of most contemporary databases, ML relies heavily on unlabeled data and the trends the system can gain from it. It is possible for deep learning algorithms to be produced from unsupervised data, allowing for training from a large database, and increasing the potency of the deep learning algorithm.²⁷⁶

Sometimes, however, a supervisor is able to add a small sample of labeled to the system, giving the system a better understanding of how to classify the data it receives. When this method is applied, it is called semi-supervised learning, which has advantages over the previous two types. Semi-supervised learning systems are able to take unlabeled data, and by using the classification logic established by the labeled data, can classify this data in a way that the supervised system cannot. Furthermore, semi-supervised systems require less supervision, because they can become increasingly independent as the system is fed up with more unlabeled data and refines its logic.^{255,275}

The final type of ML technique is reinforcement learning, which does not rely on existing data. Similar to classical reinforcement learning, the system observes an input state and decides to perform a certain action based on the observation. If the action is correct, the system is 'rewarded', while if the action is incorrect, the system is 'reinforced'. The information about the relationship between the state and correctness of action is then stored and used as learning for the next state. As more input states are observed, the system is able to fine-tune its decision making due to the reward/reinforcement based on the action the system performs.^{255,275}

There are many different types of computational models that can be used for the designs and predictions of MOFs, such as Grand Canonical Monte Carlo (GCMC), regression model, convolutional neural network, and graph neural network for studies, creating a toolbox for a variety of purposes. GCMC is a model for studying systems in equilibrium and analyzing related properties. It is widely employed in MOF chemistry to simulate the adsorption properties of materials. By calculating the energy exchange and particle exchange between different states of a material during adsorption, it is possible to determine which molecule gets adsorbed best, as well as the possible mechanisms of adsorption, by comparing the energy strain of each adsorption state of the material. Furthermore, the heat of adsorptions for different molecules can be calculated within a specific MOF structure.²⁷⁷ Two types of potential energies are usually calculated for GCMC adsorption calculations, one is based on guest-guest interactions and the other is based on host-guest interactions using the force field of atoms in the MOF. A force field provides an approximate representation of the interactions between atoms, serving as a key metric for describing the potential energies of these interactions.²⁷⁸ By pairing GCMC calculations with ML algorithms, it is possible to quickly screen a large sample of MOF structures, theoretical and experimental, to determine which has the best simulated adsorption capabilities for a certain substance or mixture.²⁷⁹⁻²⁸¹

Another computational model that is commonly used for adsorption studies is the Gaussian Process Regression (GPR) model. GPR tends to require less data for training compared to other methods. Usually for adsorption data, temperature and pressure of the system, combined with the pore volume and surface area of the MOF are the parameters set for regression modeling.²⁸² Other parameters can be used for other types of predictions, such as catalysis, where the pH and reactant and catalyst concentrations can be used as parameters to determine which catalyst works best for a certain pollutant. GPR is especially helpful when a non-linear relationship between input and output is present, due to the ability to have multiple kernel functions within the algorithm allowing for increased flexibility, especially when resolving uncertainties in the data. Kernel functions are multidimensional functions that quantify similarities between two pieces of data in a certain abstract data space. With a set number of parameters, no matter the size of the data space, the kernel functions can be compared to see which works best for a certain data set.²⁸³ These qualities allow GPR models to be powerful and versatile modeling tools.

With the information resulted from previous mentioned models, deep learning neural networks algorithms can be further utilized, specifically convolutional neural networks (CNNs) and graph neural networks (GNNs). By taking the calculations from GCMC or other models, and using crystal graphs, which is a representation of crystal materials that the network can understand, CNNs are able to be trained to have optimized parameters, allowing them to automatically describe the properties of crystalline materials. CNN doesn't necessarily require fine granularity in the input data, which can increase

the computational efficiency and improve the model generalization by identifying broader, more generalizable patterns in the data. Using crystal graphs, CNNs are able to predict material properties using the data they were trained on, giving CNNs flexibility similar to GPR models. By discerning the training data for a specific purpose, the researchers are able to make CNN models predict a vast array of MOFs' performance for specific tasks, such as gas storage or separation.²⁸⁴⁻²⁸⁶ CNNs have even been shown to be able to characterize MOF structures from experimental X-ray diffraction patterns after being trained on mostly theoretical data.²⁸⁷ The other main type of neural network is the GNN. CNNs are mostly used for image-based data, including crystal structure data, while GNNs are better for graph related tasks, such as atomistic material representation.²⁸⁸ In order to train a GNN, the MOF structure must be portrayed in a graphical representation by turning the atoms into nodes and the bonds as edges. The nodes are labeled as a certain element by assigning input node features such as electronegativity, location on the periodic table, radius, *etc.* This atomistic graph is then converted into a line graph and then the training data is run with a goal of minimizing the error, usually mean squared error. Once again, based on the parameters of the training data, the GNN can be tuned to look for a specific purpose in the MOF structures, such as gas adsorption based on surface area and pore size, or catalytic efficiency based on band gaps and other desired properties. It has also been shown that traditional GCMC calculations are able to be used as a comparison to the GNN predictions, ensuring accuracy of the trained model.²⁸⁹

There are also examples of large language models being combined with ML algorithms to develop smart assistants for MOF discovery and optimization.^{290,291} These smart assistants are able to effectively gather information and screen large quantities of data to answer an input task much quicker than just using conventional computational methods such as GCMC. By combining conventional computational methods with ML models, chemists are able to quickly screen possibilities, allowing them to spend more time in the lab testing the output from the models.

4.2 AI-guided synthesis

The structure and application of MOFs can be finely adjusted by altering the metal nodes, organic linkers and the reaction conditions. This high degree of tunability poses a challenge for researchers seeking the most suitable MOF for specific applications under optimal synthesis conditions. AI offers a promising solution to streamline this process by accelerating the screening of optimal MOFs based on structure and functionality. This chapter explores AI-guided enhancements in synthesis conditions and the design of functional structures, as well as AI-guided approaches to innovate new MOF designs and synthesis methodologies.

4.2.1 AI-assisted synthesis optimization. The synthesis of MOFs involves the self-assembly of metal nodes and organic linkers. Various synthesis methods have been developed, while multiple parameters within the synthesis conditions can be



tuned, significantly influencing the crystallinity, morphology, and surface area of the final MOF products. These parameters include the choice of solvents, pH value of the solution, concentrations of metals and organic linkers, type and amount of additives, reaction time, temperature, pressure, and more. Screening optimal conditions can be a time-consuming, labor-intensive, and costly process. ML and AI offer a promising solution to reduce the effort in condition screening and facilitate the novel synthesis of pristine MOFs.

ML model can be trained using the SynMOF database, which was generated through automatic data extraction from the CoRE MOF database and literatures, recording six parameters, including metal source, linker, solvent, additive, synthesis time, and temperature, for the recorded MOFs. A web tool was developed based on this model, which, when given the CIF file of a MOF structure, can predict synthesis conditions including temperature, reaction time, solvent, and additives. This model demonstrates better performance compared to experts' general intuition. Although the model provides only basic predictions of the synthesis conditions without detailed information such as reactant concentrations and additive amounts, it highlights the potential of this tool to accelerate the optimization of the MOF synthesis process.²⁶⁹ Genetic algorithm (GA) can be used to optimize the experimental synthesis conditions of HKUST-1, while nine different synthesis parameters, including volume of water, dimethylformamide, methanol, ethanol, isopropanol, ratio of reactants, reaction temperature, microwave strength and reaction time, were studied. With over 120 conditions tested, the importance between the nine parameters can be determined and a better weighted calibration can be introduced to enhance the algorithm. The algorithm was also applied to Zn-HKUST-1, which didn't yield crystalline MOF with previously reported condition for HKUST-1. Out of 20 algorithm-generated conditions, two conditions yielded Zn-HKUST-1 crystals.²⁹² Synthetic Conditions Finder (SyCoFinder) can also be used for the optimization of the synthesis of Al-PMOF, while the original reported solvothermal reaction for Al-PMOF had a low 40% yield and long reaction time as 16 hours. Five parameters (type of solvent, concentration, temperature, microwave strength, and reaction time) were optimized by the GA algorithms, and an alternative condition was generated for the synthesis of highly crystalline Al-PMOF. Under the new condition, the reaction time was shortened to 50 minutes, with around 80% yield achieved.²⁹³ SyCoFinder has also been proven for the synthesis optimization of HKUST-1, while five parameters including the amount of metal and linker, amount of modulator, ultrasonic cleaning time, and spray cleaning time, were optimized for the synthesis of HKUST-1 surface anchored MOFs. In two generation and a total of 30 trials, HKUST-1 SURMOFs could be synthesized with 84% crystallinity and a perfect 100% [111]-orientation. One of the challenges of synthesizing SURMOFs is the layer-by-layer growth of the MOF film. With the larger number of deposition cycles, the influence from underlying self-assembled monolayer decreased, and the uniformity of the MOF orientation is lost. However, with the algorithm-optimized condition, HKUST-1

SURMOFs retain high 92% in [111] direction with 95% crystallinity even after 80 cycles.²⁹⁴

Besides, ML and AI can also support the development of new process for MOF synthesis. SyCoFinder can be utilized to optimize the synthesis condition of ZIF-8 and ZIF-67 with microdroplet-based spray method, a newly developed method for efficient continuous synthesis of MOF particles. Four parameters, including the furnace temperature, operating pressure, water/methanol ratio, and linker/metal ratio, were optimized. 100 parameter sets for ZIF-8 synthesis were generated by SyCoFinder and 21 of the most diverse sets were tested. Within the experiment set of the second generation, highly crystalline ZIF-8 with the surface area of $1748 \text{ m}^2 \text{ g}^{-1}$ was synthesized, which is among the highest reported values and predominated by MOF micropores rather than mesopores from interparticle voids. The same condition was applied for the synthesis of ZIF-67, while ZIF-67 with high crystallinity and surface area was also obtained.²⁹⁵

Overall, the integration of ML and AI in MOF synthesis not only enhances the efficiency and efficacy of existing methods but also paves the way for innovative synthesis techniques, thereby advancing the field of MOF research and applications.

4.2.2 Functional oriented property predictions and structure optimization. The applications and functionalities of MOFs are intricately linked to their structural properties. Traditionally, screening datasets to uncover MOFs with desired characteristics involves high-throughput computational screening (HTCS) utilizing methods like density functional theory (DFT) or GCMC.²⁹⁶ However, this process is often time-consuming, expensive, and inefficient due to the vast amount of data involved. The effectiveness of a ML model hinges on various factors including dataset scope and selection, choice of ML algorithm, feature selection and weighting, and thorough model evaluation. The selection of appropriate metrics and models is contingent upon the specific objectives of the data analysis.

Leveraging their experience in AI-assisted synthesis optimization and the in-depth study of the relationships between MOFs' structures and functions, researchers have developed an auxiliary tool for MOF design and synthesis. This tool focuses on functional-oriented property predictions and structure optimizations, significantly enhancing the efficiency of designing MOFs for specific functions. Key applications including water treatment, gas adsorption, gas separation, energy transfer, and sensing will be discussed in detail below (Fig. 21).

For water treatment, artificial neural networks (ANNs) are excellent models due to their ability to handle many parameters with complex nonlinear relationships. In toxic and harmful ion adsorption of water treatment, the influencing parameters are complex and interrelated, which makes ANNs particularly suitable for predicting and optimizing the performance of MOFs in ion adsorption. The research on predicting the performance of MOF thin-film nanocomposite membranes through an ANN model by collecting parameters such as MOF size, loading, pore size, polyamide thickness, pressure, and salt concentration, can indicate that thickness and pore size are the



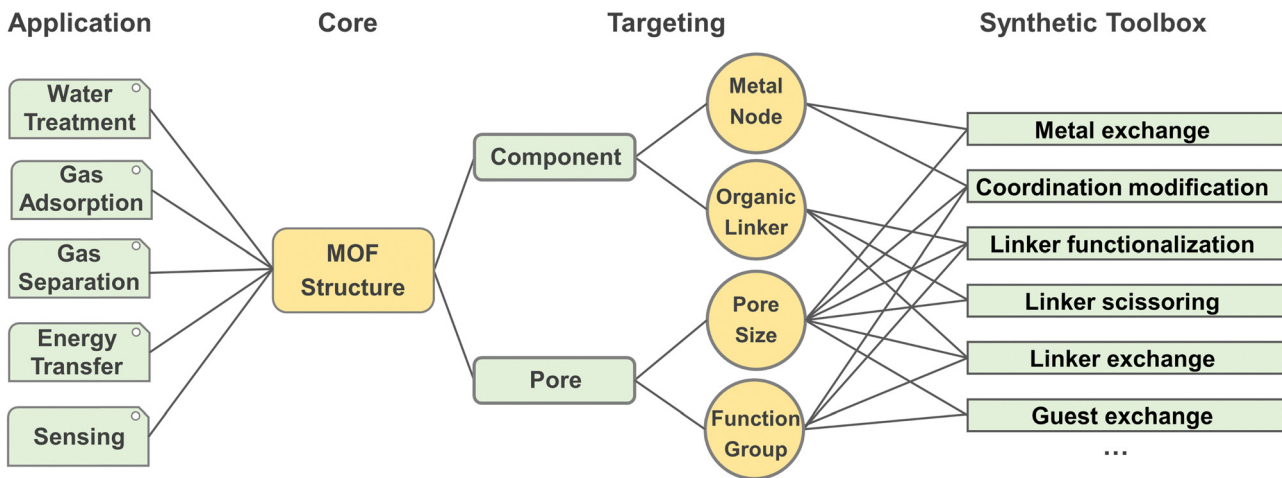


Fig. 21 Simplified functional oriented property predictions and structure optimization processes.

most critical factors affecting performance.²⁹⁷ ANN model can also be employed to predict the performance in phosphate adsorption for water treatment. Several factors affecting MOF adsorption performance were analysed, including MOF types, synthesis methods, modification techniques, and operational conditions (initial concentration, adsorbent dose, pH, contact time, and temperature). The construction and evaluation of the ANN model indicate that initial phosphate concentration and modulating agents significantly impacted adsorption performance prediction.²⁹⁸

For gas adsorption, several energy and exhausted gases, are widely studied. For energy gases, such as H₂, a dataset of approximately 98 000 real and hypothetical MOFs were used to screen their gas adsorption abilities by the Bayesian Optimization (BO) method, aiming to identify structures with high H₂ storage capacity. By using this method, it is possible to find the optimal candidate MOFs by screening less than 0.027% of the database. Despite the high complexity of training, BO models have the following advantages: requiring fewer samples, identifying the global optimum rather than the local, and being suitable for black-box optimization.²⁹⁹ Besides, ANN model, least squares support vector machine (LS-SVM) model, and generalized regression neural network (GRNN) model can also be employed to study the H₂ storage capacity, indicating that surface area and adsorption enthalpy are the most critical variables.^{300,301} For CH₄, random forest (RF) method was trained on a dataset of over 130 000 MOFs from the hMOF database, while the percentage of metal was introduced as a variable to represent open metal sites, achieving an *R*² value of 98%.³⁰² Also, the practical application was further considered, aiming to prevent water-induced degradation of MOF structures through RF method.³⁰³ GCMC simulations and ANN regression models can also be employed to study the CH₄ adsorption ability of 2224 MOFs.³⁰⁴ For CO₂, DFT calculations and GCMC simulations can be used to study the CO₂ adsorption performance of MOFs, suggesting that the -NH₂ functional group has the strongest binding affinity with CO₂.³⁰⁵ GCMC

simulations also confirm that molecular selectivity is a crucial indicator for CO₂ capture.³⁰⁶ Other gases like N₂ can also be predicted, while RF method based on CoRE MOF database and DFT calculations based on ARC-MOF database can be employed for the gas adsorption performance study.^{307,308}

For gas separation, through the comparisons of 2434 MOFs from the CoRE MOF and CSD databases, the selection of the database notably impact the gas adsorption and ideal selectivity calculations under low-pressure conditions in higher H₂/CH₄ Henry's coefficients.²⁶⁵ Multi-component-GCMC method to screen the hMOF database can be employed to identify the best materials for C₂H₆ and C₂H₄ separation, discovering that the porosity is the key factor.³⁰⁹ GCMC simulations with an XGBoost (eXtreme Gradient Boosting) ML model can be combined and employed for screening MOF materials for high Xe/Kr separation performance, while the porosity, density, pore volume, and pore limiting diameter are the key features influencing adsorption performance.³¹⁰ 5249 MOF membranes and 31 494 MOF/polymer mixed-matrix membranes (MMMs) were used for the evaluation in the separation of six gas pairs (He/H₂, He/N₂, He/CH₄, H₂/N₂, H₂/CH₄, N₂/CH₄), which indicates that physical properties such as porosity have a stronger correlation with the gas adsorption and diffusion properties of MOFs, rather than chemical descriptors like atomic types and degree of unsaturation.³¹¹

For energy transfer, several highly conductive MOFs were developed through optimization. Potential conductive MOF structures can be identified by transfer learning.²⁶⁷ Crystal graph convolutional neural network (CGCNN) was employed to address bandgap prediction, which generates approximate crystal graphs and requires only 7 minutes to compute the properties of 13 058 MOFs, whereas traditional DFT calculations would take 1.5 million hours.³¹²

For sensing, ANN model was used to predict glucose concentration in Huangshui, demonstrating excellent fitting and predictive performance with low root mean squared error and high residual predictive deviation, which showed superior detection



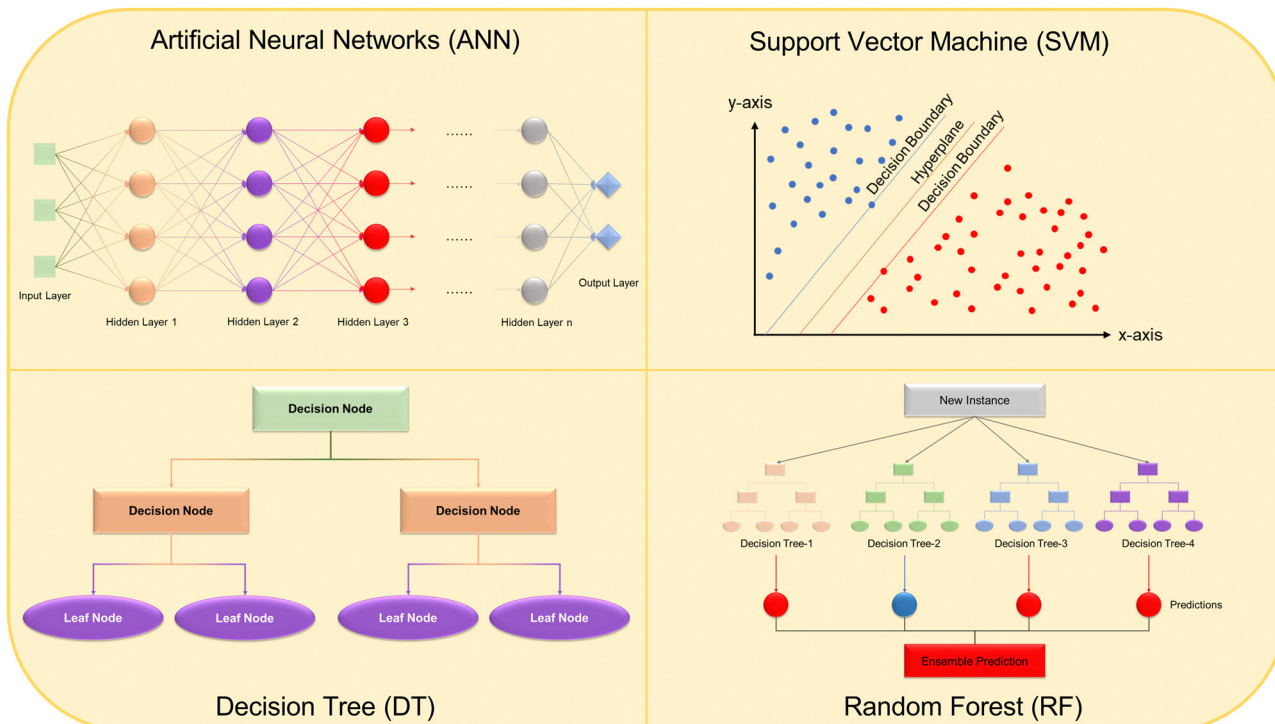


Fig. 22 Classical logics for the models of machine learning.

performance and extended the sensor's linear detection range to 11 mM (Fig. 22).³¹³

4.2.3 AI-guided new MOF design and synthesis. The AI-driven development of novel MOFs can be achieved mainly in three processes: (i) screening potential MOFs from hMOF databases; (ii) utilizing structure–activity relationship knowledge to assist structural design; (iii) employing algorithms to generate novel MOFs.

For the screening process when constructing hMOF database, the comprised MOF structures were identified by high-throughput screenings which involves simulating the performance of all candidates in the database for comparison. The construction of hMOF databases can be traced back to 2000 for the development of a method named “automated assembly of secondary building units (AASBU)” to generate MOF structures. Secondary building units (SBUs) from atoms were built by using Lennard-Jones potential to parameterize the interaction energy between atoms. Then structures were generated by calculating the interactions between SBUs with a given number of SBUs per unit cell and space group. The simulated annealing Monte Carlo algorithm was used to optimize the energy of the final structure by allowing building units to rearrange and adjusting cell size and distance between steps so that the resulting structure is stable enough to be synthesized.³¹⁴ Later, geometric methods were developed as an alternative approach to minimize the energy, which can be classified as “bottom-up” and “top-down”. The “bottom-up” approach begins with the SBUs, through sequential connection to form a periodic crystal structure.³¹⁵ While the “top-down” approach starts with a given network or topology and then maps the appropriate building

blocks to the network to generate the structure.^{316,317} In this process, AI can accelerate the screening by introducing advanced algorithms to reduce the number of operation objects like performing a preselecting process to filter un-promising MOFs before running property simulations. The related algorithms can be mainly divided into three types, tree-based (*i.e.* decision trees, random forests), kernel-based (*i.e.* support vector machines), and artificial neural network,^{318–321} which provide the foundation for the subsequent structural design.

For the structure–activity relationship knowledge assistant, structure–activity relationship study is a cornerstone of modern chemistry which provides a systematic approach to understand how molecular structure influences properties, and thus enable to tailor materials for various applications. Employing AI to study structure–activity relationship can quickly identify the key features of high performance among a huge MOF database, which will be used to further generate promising MOF structures. Collectively, relative studies underscore the transformative impact of ML and AI methodologies in the rational design and synthesis of MOFs with various structures and properties.^{322–326}

For the employment of algorithms to generate novel MOFs, introducing an evolutionary algorithm (EA) is a valuable approach. EA is one kind of ML algorithm that can start from a small set of MOFs and achieve high-performance structure discovery through continuous modification evolution. In EA, a linker is randomly chosen from the population to undergo one of several evolution operations (like linker modifications). The produced linker can be inserted back into the population only if it passes all the filters such as the torsions number < 8 ,



Table 4 Typical examples for AI-assisted MOFs

Method types	Application	R^2
XGBoost ³³⁵	As adsorption	0.93
LightGBM ³³⁶	As adsorption	0.996
ADA-GPR ³³⁷	Hg/Ni removal	0.998
ANN ²⁹⁸	Phosphate adsorption	0.73–0.96
ANN ²⁹⁷	Water purification	0.9062
RF ³³⁸	Water adsorption	0.941
RF ³³⁹	Methane adsorption	0.98
RF ³⁰³	HCHO capture	0.867
XGBoost ³⁴⁰	CO ₂ /N ₂ /CH ₄ adsorption	0.9945/0.9948/0.9911
GBDT ³⁴¹	H ₂ /CH ₄ /CO ₂ adsorption	0.864
GPR ³⁴²	CO ₂ adsorption	0.98
General regression neural network ³⁴³	H ₂ storage	0.9986
GNN ³⁴⁴	CO ₂ adsorption	0.95
CNN ³⁴⁵	Methane adsorption	0.947
ANN ³³⁸	Water adsorption	0.982
ANN ³⁰⁰	H ₂ storage	0.93
RF ³⁰⁹	Ethane/ethylene separation	0.89
XGBoost ³⁴⁶	N ₂ /O ₂ separation	0.93
XGBoost ³¹⁰	Xe/Kr separation	0.973
Tree-based pipeline optimization tool ³¹¹	He/H ₂ /CH ₄ /N ₂ separation	0.99/0.80/0.73/0.84
ANN ³⁴⁷	Xe/Kr separation	0.972
ANN ³⁰⁵	CO ₂ /N ₂ separation	0.905
Categorical boosting ³⁴⁸	Ibuprofen adsorption	0.76
ANN ³⁴⁹	Glucose detection	0.9996
RF ²⁶⁷	Conductivity	0.97
CNN ³¹²	Bandgap prediction	0.876

GBM for gradient-boosting machine; ADA for adaptive; GPR for Gaussian process regression; GBDT for Gradient boosting decision tree.

and sites number = 2 to make sure it can form a MOF while its capacity should also be greater than the lowest one in the current population so that it has research value. As a result, the linkers possessing high capacity would evolve and the population could enlarge while excluding the low-performance ones.^{327–334} The integration of EAs with ML/AI techniques has significantly advanced the design and discovery of high-performing MOFs.

AI-assisted new MOF design is not limited to the development and optimization of the algorithm and model to increase the accuracy and speed. It is also trying to reduce the knowledge requirement to use the algorithm, which can predict MOF properties and generate MOFs with specified properties from natural language inputs. Some of the results reported and their applications have been summarized and listed in Table 4. Coefficient of determination (R^2) reflects how closely the predicted values align with the ideal line, which is easy to interpret and allows for the comparison across different models and objectives. While R^2 alone sometimes cannot fully capture the model's accuracy and can be misleading when comparing models with different numbers of variables. In addition, overfitting is a common issue in predictive models, which must be carefully addressed. It occurs when a model captures noise or random fluctuations in the training data instead of learning the underlying patterns. This phenomenon leads to poor performance on new, unseen data and undermines generalization of the model. To prevent overfitting, it is essential to split the available data into two parts: training set and testing set. The training set is used to build and tune the model, while the testing set, which the model has not seen during training, is

employed to evaluate its generalization ability. Only no significant drop in performance on the testing set ensures that the model is robust and can accurately predict outcomes for new data.

Although there have been many works applying AI to the design of new MOFs, AI itself is also a growing discipline. Therefore, the application of more advanced AI algorithms for MOF design is necessary. In addition, interdisciplinary cooperation also needs to be improved. Many works only show the potential of different AI models, but there are few examples of reaching the laboratory to achieve synthesizing new MOF structures with performance breakthroughs, leading to a potential risk of the optimal structure not being able to be synthesized due to its unstable nature. Furthermore, the automated integration platform combining synthetic robots and ML algorithms for novel MOF synthesis has not been achieved yet. Such a platform will largely accelerate the synthesis of novel MOFs with the liberation of human labour.

5. Conclusion and outlook

In summary, MOFs have experienced remarkable growth and innovation over the past decades, establishing themselves as versatile materials with a wide range of applications. Advancements in their design and synthesis have been driven by a combination of large-scale synthesis attempts, functional-oriented modifications, and the integration of ML and AI for predictive modelling. Large-scale synthesis efforts have successfully constructed numerous novel structures, providing



valuable experience in the design and synthesis of these materials. Functional-oriented modifications have further expanded the utility of MOFs by tailoring their properties for specific applications, leading to the development of MOFs with enhanced stability, activity, and other properties that meet the demands of complex industrial processes and environmental sustainability. The integration of ML and AI in MOF research represents a significant leap forward, with AI-driven predictions accelerating the discovery of novel MOF structures, optimizing synthesis conditions, and predicting material properties with high accuracy, which has significantly reduced the time and resources required for experimental trial and error.

In the coming years, the field of MOF design and synthesis is expected to evolve through several transformative trends, but it will also face critical challenges that must be addressed for the full potential of these materials to be realized. One major trend is the deeper integration of ML and AI with computational chemistry, which will continue to accelerate the discovery of novel MOF structures with highly specialized properties, particularly for emerging applications in energy storage, carbon capture, catalysis, and biomedicine. However, a key challenge will be the development of accurate, large-scale datasets needed to train AI models, as well as addressing the complexity of predicting MOF behavior in real-world conditions where multiple variables interact, which may need plenty of work for AI and researchers to explore collaboratively. Another emerging trend is the push towards sustainable and green synthesis methods, which are essential for scaling MOF production while minimizing environmental impact. Developing energy-efficient, low-waste, and solvent-free methods remains a significant challenge, especially when transitioning from laboratory-scale synthesis to large-scale industrial processes. Additionally, MOFs' long-term stability under harsh operational conditions, such as in industrial catalysis or environmental remediation, presents another challenge. Addressing these challenges will require interdisciplinary collaboration, innovative research methodologies, and significant investment in both fundamental and applied sciences.

The future of MOF design and synthesis is poised to be even more dynamic and impactful, with several promising directions on the horizon:

- Deeper integration with computational techniques: the continued advancement of ML and AI will further refine the predictive capabilities for MOF design. Enhanced computational power and more sophisticated models will enable the discovery of novel MOFs with unprecedented properties, tailored for emerging applications in energy, environment, and biomedicine.

- Sustainable and environmentally friendly synthesis: as environmental concerns become more pressing, the development of sustainable and green synthesis methods for MOFs will gain prominence. Efforts to utilize renewable resources, reduce energy consumption, and minimize waste will be critical in MOF production, especially large-scale synthesis, more environmentally friendly.

- *In situ* characterization and real-time monitoring: advances in *in situ* characterization techniques will provide deeper insights into the dynamic behavior of MOFs during synthesis. Real-time monitoring of MOF synthesis will enhance the understanding of the formation and growth processes, allowing for better control of synthesis conditions and more efficient production of MOFs.

Overall, the design and synthesis of MOFs stand at the cusp of a new era, driven by innovative methodologies and interdisciplinary efforts. The convergence of large-scale attempts, functional modifications, and AI predictions will continue to propel the field forward, offering transformative solutions to global challenges and ushering in a new age of advanced materials.

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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