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Application and Development Prospect of Nanoscale Iron Based Metal-Organic Frameworks in Biomedicine

Xiujuan Peng¹, Li Xu¹, Min Zeng², Hao Dang¹

¹Department of Clinical Laboratory, The Third Hospital of Mianyang (Sichuan Mental Health Center), Mianyang, Sichuan, 621000, People's Republic of China; ²School of Materials and Chemistry, Southwest University of Science and Technology, Mianyang, Sichuan, 621010, People's Republic of China

Correspondence: Min Zeng; Xiujuan Peng, Email zengmin@swust.edu.cn; 34472941@qq.com

Abstract: Metal-organic frameworks (MOFs) are coordination polymers that comprise metal ions/clusters and organic ligands. MOFs have been extensively employed in different fields (eg, gas adsorption, energy storage, chemical separation, catalysis, and sensing) for their versatility, high porosity, and adjustable geometry. To be specific, Fe^{2+}/Fe^{3+} exhibits unique redox chemistry, photochemical and electrical properties, as well as catalytic activity. Fe-based MOFs have been widely investigated in numerous biomedical fields over the past few years. In this study, the key index requirements of Fe-MOF materials in the biomedical field are summarized, and a conclusion is drawn in terms of the latest application progress, development prospects, and future challenges of Fe-based MOFs as drug delivery systems, antibacterial therapeutics, biocatalysts, imaging agents, and biosensors in the biomedical field.

Keywords: Fe-based MOFs, bio-application, nanocomposites, nanomedicine, green synthesis

Introduction

Metal-organic frameworks (MOFs) refer to porous coordination polymers (PCPs) formed through the self-assembly of metal ions (eg, Zn, Cu, Fe, Cr, Co, and Ni) and organic ligands (eg, phosphonates, carboxylates, sulphonates, and imidazolates).^{1,2} Due to the different reaction conditions (eg, the type of solvent, reaction temperature, reaction time, and the ratio of ligand to metal), MOFs are capable of self-assembling to form one-dimensional (1D),³ two-dimensional (2D),^{4,5} or three-dimensional (3D)⁶ skeleton topologies. Almost unlimited reasonable combinations are generated with the continuous expansion of metal clusters, organic ligands, and topological structures.⁷ In this context, MOF materials exhibit a wide range of properties, including high porosity (from ultra-micropores to mesopores), high surface area, large pore volume for packaging, chemical and thermal stability, and so forth.^{8,9} Because of the above-described characteristics, MOFs have been extensively employed in a wide variety of research fields (eg, gas storage, photochemistry, catalysis, separation process, diagnosis, and drug delivery).^{10–13} Besides, they are considered the most promising materials in biomedicine and nanomedicine.^{9,14}

In the past few years, nanoscale MOFs (NMOFs) has played a crucial role in biomedical engineering. (i) NMOFs are the most fascinating nanocarriers in drug delivery, thus becoming better candidates for carrying anticancer drugs. NMOFs are more flexible and have better biodegradability compared to other delivery systems based on nanoparticles (eg, nano-silica).¹⁴ (ii) The selection of internal components of NMOFs (eg, encapsulated fluorescent agents or organic dyes) can be adopted in the fields of biosensors and biological imaging.¹⁰ (iii) The customized components of NMOFs can also be employed as therapeutic agents for a wide variety of treatments (eg, using drugs as organic ligands of MOFs, or using active metals as central metal ions of MOFs), which can be applied in the fields (eg, biocatalysis, antibacterial, and bacteriostatic).¹³

Fe is one of the most abundant transition metal ions that are identified in the human body and other animals, and it serves as a vital component of hemoglobin. It serves as an excellent metal ion candidate for building coordination polymers because of its clear biological function and biological safety.¹⁵ Accordingly, Fe-MOFs (eg, MIL-100,¹⁶ MIL-88,¹⁷ MIL-53,¹⁸ MOF-n,¹⁹

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and PCN^{20,21}) have aroused extensive attention in different biomedical applications. Its advantages are presented as follows: (i) It exhibits an open structural framework and a large specific surface area, such that the reduction of the framework can be induced to form a coordinated unsaturated iron site; it is selected for application in several fields (eg, biocatalysis and antibacterial). The open metal sites exhibit antibacterial activity.^{16,19} (ii) For its excellent chemical stability and high porosity, it can be combined with other organic and inorganic materials, and it is selected for several fields (eg, sensing and fluorescence detection).²¹ (iii) With good biocompatibility and biodegradability, it can load drugs and metal particles in the frame, and it is selected for drug transportation, diagnosis, and treatment, and other fields.¹⁷ (iv) It is easy to synthesize, and it can be produced through green synthesis and optimal scale.^{15,16} In general, Fe-MOFs become a powerful platform for achieving different biomedical goals for the above-mentioned biological characteristics of iron.

To date, there have been rare comprehensive summaries of Fe-MOFs and their biomedical progress despite a considerable number of comments on MOFs and their biomedical applications. Iron-based MOFs, as a subclass of the huge MOFs family, have made great progress in intelligent design and application strategies of various nano iron-based MOFs materials over the past 5 years. Hence, Fe MOFs and their biomedical progress are comprehensively summarized and systematically analyzed. In this study, the key index requirements for Fe-MOF materials in the biomedical field are first analyzed. Subsequently, the latest progress of Fe-MOFs in biomedical applications is systematically presented (eg, drug delivery systems, biosensors, biocatalysts, image guidance, and antibacterial), as shown in Figure 1. Lastly, the development prospects and future challenges of Fe-MOFs in biomedical applications are summarized.



Figure 1 Application of Fe-based MOFs nanomaterials in biomedicine. Adapted from these studies.^{36,80,89,104,115,116,138,139,162,166.}

Key Indicators of Fe-MOFs Materials in Biomedical Applications

In general, Fe-MOF materials show the biomedical potential for their high porosity, strong adsorption capacity, good biocompatibility, as well as extensive fictionalizations advantages.¹ Notably, for the design of Fe-MOFs in biomedical applications, no matter the routes (host MOF or BIOMOF) or application (drug delivery, imaging, or combination of both), several vital indicators should be followed.²²

Accurate Particle Size Control

Fe-MOF materials employed in the biomedical field must have accurate particle size control and size optimization.²³ Figure 2 shows the application of MOF materials in biomedicine and particle size dependence relationships. As indicated by the results, (i) the reduction of particle size will affect its specific surface volume and increase its reactivity. (ii) The penetration ability of particles is enhanced with the reduction of size. (iii) Larger particles can be retained in the cell tissue, and they can lead to the increased stability of buffer solution and the reduced aggregation of particles. (iv) The different mechanisms of endocytosis (eg, phagocytosis, megacytosis, fossil-mediated endocytosis, or reticulin-mediated endocytosis) are particle size-dependent.¹ Thus, the size, charge, shape, and density of MOFs will significantly affect the cell uptake, circulation and retention time, biological distribution, and internalization mechanism.²⁴

In general, MOFs materials with micron or even millimeter size are usually not suitable for biological applications.^{25,26} For instance, in the case of injection, the particle size should not exceed 200 nm; the size range between 50 and 300 nm can provide the best-circulating half-life for parenteral administration (Figure 2). This is also why the synthesis of nanoscale drug delivery systems has become such a vital strategy.²⁷ Many literatures have adopted different strategies to control the particle size of MOF. For instance, crystal growth kinetics can be controlled by adjusting reaction parameters (eg, the type and quantity of solvent, reagent concentration, temperature, and pH) or introducing auxiliary additives (coordination modifiers, surfactant-mediated synthesis, etc.).^{23–27} Accordingly, the study of a simple synthetic process for accurately controlling the particle size, morphology and crystallinity of Fe-MOFs has become a vital research direction in the application of this material in the biomedical field.²⁷

Simple Synthesis Process Route

Different synthesis processes can affect the particle size, morphology, and crystallinity of Fe-MOFs for their advantages and disadvantages. A comprehensive schematic diagram of important synthesis technologies of Fe-MOFs materials is



Figure 2 Application of Fe-MOF nanomaterials in biomedicine and particle size dependence relationship.



Figure 3 Synthetic methods of Fe-MOFs and their composites.

presented in Figure 3. Hydrothermal or solvothermal methods are the most common methods to synthesize Fe-MOFs materials.^{28–32} The above method usually forms porous metal-organic framework crystals through self-assembly nucleation and crystal growth in a sealed autoclave at a temperature higher than or close to the boiling point of the solvent, as well as through a long time of heating. The above method requires high time and energy consumption, which limits the feasibility of its large-scale synthesis and practical applications. The microwave radiation-based synthesis method is a highly promising synthesis technology of metal-organic skeleton for its several advantages (eg, rapid reaction and environmental friendliness).³³ Compared with conventional hydrothermal synthesis, microwaves can directly interact with mobile charge in solvent and solid, such that it can save energy and reaction time.

Ultrasonic-assisted synthesis refers to a type of energy-saving method for chemical reactions with high product selectivity and high speed at ambient temperature.^{34,35} The above method is to promote the chemical reaction in the solution containing the precursor through the interaction with the liquid through the cyclic mechanical vibration of the ultrasonic wave. The power, type, and ultrasonic time of the ultrasonic generator significantly affect the nucleation rate and crystal growth, and it is easy to form an amorphous MOF structure. Mechanochemical synthesis shows the potential of industrial scale for the advantages of zero waste generation and high output.^{36,37} In this synthetic route, mechanical energy causes chemical bond breaking and chemical reactions rapidly. Despite the improvements in reaction kinetics and rapid nucleation rate and the capability of rapidly synthesizing nano-sized particles, the above method is easy to form defective MOFs crystal structures.

The spray drying method has the advantages of simplicity, rapidity, less waste generation, and high solvent recovery.³⁸ In this method, through simple evaporation steps, the combined effect of evaporation of atomized droplets and local heating may lead to new chemical reactions. The advantage of the above method is that it can form dry spherical powder continuously and rapidly in a short time with minimum manufacturing cost, thus boosting the industrial production and large-scale application of MOFs. Flow chemistry refers to a mature technology for large-scale synthesis of functional nano-MOF materials.³⁹ The above method is to rapidly form MOF microcrystals through the continuous conversion of mixed reagents in a flow reactor at the reaction temperature. It has the advantages of high reproducibility and precise control of reaction parameters and can realize continuous production by adding reactant solution. Compared with the structure formed based on the conventional batch method, the obtained MOFs structure exhibits better crystal-linity and porosity.

The existing synthesis methods have their advantages and disadvantages, and their practical applications are still limited (Table 1). It can be predicted that a great deal of research work in the future will be devoted to designing accurate, controllable, simple, clean, and sustainable synthesis routes for Fe-MOF materials while reducing the impact of their preparation on the environment.

Synthesis Methods	Fe-MOFs	Organic Linker/LD50	Size (nm)	Morphology	Application	Refs.
Hydrothermal	MIL-100	Trimesic acid (H3BDC)/(8.4 g/kg)	sic acid (H3BDC)/(8.4 g/kg) ~5µm Irregular Elec		Electrochemical sensor	[28]
	MIL-88B	Terephthalic acid (H2BDC)/(5 g/kg)	600–800nm	Nano-column	Drug release Scaffold Implantation	[29]
	MIL-127	3,3′,5,5′- azobenzenetetracarboxylic acid (H4-TazBz)	~lµm	Micrometric cubic	Drug delivery	[30]
Solvothermal	MIL-88A	Fumaric acid/ (endogenous ligands)	~50nm	Nanorods	Drug delivery	[31]
	MIL-53	2-aminoterephthalic acid (2-NH2- H2BDC)	~5µm	Hexagonal prism	Adsorption	[32]
	Fe-(AIP)	5-aminoisophthalic acid (5-AIP)/ (1.6 g/kg)	-	-	Catalytic	[6]
	Fe-2MI	2-methylimidazole/(1.4 g/kg)	I~2µm	Spherical shape	Photocatalytic	[13]
	Fe-NDC	2,6 napthalenedicar boxylic acid/ (5 g/kg)	300–500nm	Rod-like shape	Photothermal materials	[14]
Microwave-assisted	MIL-101	I,4-benzenedicarboxylic acid/ (5 g/kg)	150–250 nm	Octahedral	Octahedral Tetracycline	
	NH2-MIL-101	2-Aminoter-ephthalic acid	cid 100–200 nm Octahedral adsorption		adsorption	
Ultrasound-assisted	MIL-88A	Fumaric acid/ (endogenous ligands) 55–88nm Prismatic Ethyler adsort 3		Ethylene adsorbents	[35]	
Mechanochemical	MIL-100	Trimesic acid (H3BDC)/(8.4 g/kg)	100nm-10μm	Irregular	Drug delivery	[36]
	MIL-88A	Fumaric acid/ (endogenous ligands)	~5µm	Irregular		[37]
Spray-Drying Continuous-flow	MIL-100	Trimesic acid (H3BDC)/(8.4 g/kg)	~2µm	Sphere		[38]

Table I Summarizes Achievements Related to Fe-MOFs, Solvents Used in Their Preparation, and Their Applications

Biocompatibility and Non-Toxicity

Biocompatibility and non-toxicity are the necessary conditions for the application of Fe-MOF materials in biomedicine.⁴⁰ Thus, a complete toxicity evaluation (eg, cytotoxicity and in vivo research) takes on great significance, determining the applicability of MOF materials in vivo, and it is also one of the serious obstacles limiting the application of a wide variety of MOFs materials in biomedicine.⁴¹ In general, biocompatibility research includes blood reaction, immune reaction, and tissue reaction. The evaluation of biological toxicity mainly comprises (i) the initial toxicity evaluation of the cations and ligands that make up the MOFs materials using LD50 (half lethal dose parameter),⁴² (ii) the test of the cytotoxicity of MOFs to gain insights into the cytotoxicity at the cell level,⁴³ (iii) toxicity research in vivo, which is the critical test since it presents the most direct and valuable information.⁴⁴

Non-toxic metal cations should be employed to control the toxicity of MOFs, and the toxicity of cations is also significantly correlated with their ability to accumulate in vivo. Iron refers to a metal ion used by almost all living things, and is employed in biological processes (eg, DNA biosynthesis, oxygen transport, and cell energy generation).⁴¹ Although Fe(III) is neutral to tissues and can be safely transported in the non-active state of oxidation and reduction, its high reactivity enhances its destructive power.⁴⁵ Figure 4A presents the key players in iron metabolism. When iron is excessive, it is also termed hemochromatosis, which can result in cell damage, cell death, and organ failure, mainly affecting the liver, heart, pancreas, thyroid, and central nervous system. Accordingly, Fe(III) should be adopted to



Figure 4 Biocompatibility evaluation of Fe-MOFs nanomaterials: (A) Key players in iron metabolism. Reprinted from Vogt A-CS, Arsiwala T, Mohsen M, et al. On Iron Metabolism and Its Regulation. *Int J Mol Sci.* 2021;22:4591. Creative Commons.⁴⁵ (B) In vitro and in vivo models employed to assess nano materials' toxicity. Reprinted from Loret T, Rogerieux F, Trouiller B, et al. Predicting the in vivo pulmonary toxicity induced by acute exposure to poorly soluble nanomaterials by using advanced in vitro methods. *Part Fibre Toxicol.* 2018;15:25. Creative Commons.⁴⁶ (C) Evaluation of MIL-89(Fe) toxicity on embryonic zebrafish development. Reprinted from Al-Ansari DE, Al-Badr M, Zakaria ZZ, et al. Evaluation of Metal-Organic Framework MIL-89 nanoparticles toxicity on embryonic zebrafish development. *Toxicology Reports.* 2022;9:51–960. Creative Common.⁴⁴ (D) Effects of MIL-100(Fe) on human normal liver cells (HL-7702) cells' viability and morphology. Reprinted from Chen G, Leng X, Luo J, et al. In Vitro Toxicity Study of a Porous Iron(III) Metal-Organic Framework. *Molecules.* 2019;24(7):1211. Creative Commons.⁴⁷

synthesize Fe-MOF materials that can be employed in biomedicine, and the intake, distribution, and utilization of iron should be strictly controlled.⁴⁵

The toxicity of MOF materials also depends on the type of organic ligand.⁴⁸ Figure 4B presents the in vitro and in vivo models employed to assess MOFs' toxicity. Both exogenous and endogenous substances can serve as organic ligands. There are many types of exogenous ligands, and functional substituents can be added to change the ADME (eg, absorption, distribution, metabolism, and excretion) process, and affect the drug–carrier interaction, making the delivery process more controllable. On the other hand, endogenous ligands (eg, fumaric acid) can be metabolized by organisms after MOF degradation. However, only a few MOFs based on endogenous ligands show appropriate porosity and structural stability.^{47,49,50} Table 1 also lists some LD50 example data of typical organic ligand toxicity, suggesting that their toxicity is at an acceptable level for biological applications.

Given the complete toxicity evaluation of Fe-MOF, Al-Ansari et al and Chen et al studied the toxicity of MIL-89 and MIL-100(Fe) in vivo (Figure 4C and D).^{44,47} As indicated by all the parameters studied (eg, serum, enzymology, and

histology), MIL-89 had no acute or subacute toxicity symptoms, and its degradation and excretion mechanism was believed to be that nano-Fe-MOFs particles were rapidly isolated by liver and spleen, and then further biodegradable and directly eliminated in urine or stool, without metabolic and substantive toxicity.⁴⁴ Within a safe dose of 80 µg/mL, MIL-100(Fe) particles had excellent biocompatibility with human normal liver cells (HL-7702) and hepatocellular carcinoma cells (HepG2), low cytotoxicity, and allowed a high cell survival rate.⁴⁷

The existing Fe-MOF materials still lack systematic biocompatibility and biological toxicity research, which is also one of the possible reasons for the delayed clinical application of Fe-MOF materials.⁵¹ Metal ions and organic connectors significantly affect the biocompatibility and toxicity evaluation of MOF, while the solvent used in the synthesis of MOF and the crystal size of MOF will be the key research direction of the biocompatibility of Fe-MOF materials.

Chemical Stability and Controlled Degradation

The stability, reusability and controllable degradation of materials are also important conditions for the application of Fe-MOF materials in biomedicine.^{40,52} The stability of evaluation materials mainly includes mechanical stability, chemical stability, thermal stability, stability under acid and alkali conditions, and stability under physiological conditions. Different application environments may have very different requirements for the stability of materials. Industrial applications usually require materials that are very stable in thermal, chemical, and mechanical aspects. During drug transportation, it is necessary to be very stable in water, heat, pressure, and harsh chemical environment. Thus, moderately stable and reusable Fe-MOF materials can be recognized as an advantage in biomedical applications.

The stability of Fe-MOFs particles is primarily dependent on several parameters (eg, its components,⁵² ligands, and metal ions, medium, temperature,²⁹ pH value,^{32,53} and physiological conditions⁵⁴). To be specific, it is the potential representing the degree of electrostatic repulsion between adjacent particles in dispersion ζ . Moreover, it is one of the vital parameters to characterize the stability of components. In general, the greater the potential ζ , the greater the charge repulsion force between particles will be, and the less likely the particles will be to agglomerate. The thermal stability of materials can be monitored through thermo-gravimetric analysis (TGA) and differential scanning calorimetry (DSC). When the temperature reaches over 400 °C, the MOF structure collapses and decomposes to form Fe₂O₃, suggesting that Fe-MOFs show great thermal invariance in the physiological environment.^{54,55}

The stability of numerous MOF structures, when exposed to water, is a vital issue. Accordingly, the systematic determination of the stability of MOF under water and acid–base conditions is crucial for its application in biomedicine. Bezverkhyy et al studied the stability of MIL-101 and MIL-53 in liquid water.⁵⁵ They found that: (i) dispersion of Fe-MOFs solid in water will lead to pH reduction, (ii) Due to the function of solid acid catalysts, MIL-101 can extend the effective pH range up to 10.2, and (iii) MIL-101 exhibited good reusability and stability in liquid water. However, there has been scarce research evaluating the stability of MOF in acidic, neutral, and alkaline solutions.

Besides, the stability of Fe-MOF in the biological medium is a crucial parameter.⁴⁶ If the material is non-biodegradable, the toxicity may also increase due to accumulation in the body.^{56–58} Iron serves as a major participant in biological metabolism (eg, DNA biosynthesis, oxygen transport, and cell energy production). Under physiological conditions, iron ions are absorbed by intestinal cells in the duodenum and then loaded onto transferrin, such that they can be transported throughout the body to the places where high iron requires (eg, bone marrow, where red blood cells are produced). Aged red blood cells are recognized and swallowed by macrophages and degraded in cells.⁴⁵ In general, MOF materials are used in the form of nanoparticles. When the particle size is within the range of 20–30nm, they mainly undergo kidney elimination, while larger particles (30–300nm) can be rapidly absorbed by mononuclear phagocytic system cells, and mainly exist in the liver, spleen, and bone marrow. However, there have been few studies on the stability of Fe-MOF materials under physiological conditions and the in vivo controlled degradation thus far, and more in vivo research should be conducted.

Overall analysis can suggest that the above-described key indicators of Fe-MOF materials in biomedical applications have interrelated characteristics.^{59,60} On the one hand, the organic connector in MOFs, the solvent employed in synthesis and the crystal size significantly affect the biocompatibility and toxicity evaluation. On the other hand, the structural stability and degradation tendency of MOF are dependent on the type of metal and ligand, the diameter of nanoparticles, as well as physiological conditions. Thus, to meet the clinical application needs of Fe-MOF materials, basic research on

material synthesis, biocompatibility, and controllable degradation performance continues to be its main direction, covering the development of more organic ligands, nanocomposites, etc.

Application Status of Fe-MOFs Materials in Medical Treatment

Fe-MOF materials have become one of the most favorable candidates in biomedical applications for their high surface area and porosity, strong adsorption capacity and extensive functionalization advantages.^{7,61} The latest research of Fe-based MOFs in the fields of drug delivery, biosensors, biological imaging, biocatalysis, antibacterial and bacteriostasis has made great progress over the past few years, and its development prospects and future challenges should be discussed comprehensively.⁶²

Drug Delivery Systems

The components of numerous active drugs (API) are almost insoluble or unstable in the organism, and they are easy to rapidly and widely metabolize, and their distribution in the organism is usually non-selective.⁶³ This will lead to the generation of low therapeutic effect and may cause damage to healthy cells and tissues and serious adverse side effects. Conventional drug delivery systems (DDS) materials (inorganic materials, including silica,⁶⁴ porous carbon,⁶⁵ or zeolite⁶⁶) are prone to cause the so-called "explosive effect" arising from insufficient adsorption capacity and no way to delay or control the release of drugs. The interaction between organic ligands and metals in MOF materials contributes to controlling the gradual release and degradation of API, including release under specific conditions (eg, pH, temperature, osmotic pressure, or through enzyme activity); it can also target specific cells through recognizing ligands (eg, folate and peptide), or using the magnetic field to reach specific target or position, and so forth.^{67–72} Accordingly, the Fe-MOFs drug packaging and delivery system has been proposed and widely investigated for its high porosity, regular structure, excellent adsorption capacity, and high biocompatibility, as well as its wide range of functional advantages.^{72–83} Table 2 shows the current research status of the application of Fe MOF in the transportation of some typical drugs.

In general, there are four common ways to synthesize Fe-based MOFs drug delivery systems, as shown in Figures 5 and 6, which are elucidated as follows: (i) Adsorption synthesis strategy. First, Fe-MOFs with specific properties are synthesized and activated, and drug molecules are added to the outer surface of the MOF carrier through chemical adsorption (covalent bonding with structure).⁶⁹ (ii) Indirect packaging strategy. MOF materials are synthesized first, and then bioactive compounds are blended into the channels, pores, or cavities of MOF through physical action (non-covalent bond) by impregnation or high

Drug	Fe-MOFs	Solvent	Loading Capacity	Time-Release (h)	Release Test Medium	Refs.
Aspirin	MIL-100	Ringer's solution	I I.87wt %	24	PBS, SGF	[36]
Tetracycline	MIL-88	Water	~10 wt %	12	PBS (pH=7.4)	[69]
Ibuprofen	MIL-88B	Hexane	7.88 wt %	148	NS	[71]
	MIL-88B	Hexane (0.5mg/mL)	6.0μg/cm2	24	PBS+Tween 20	[72]
Naproxen	MIL-101	Water	-	24	SGF (pH=2) SIS (pH=7.4)	[74]
Theophylline	MIL-100	HCI (0.03M)	32%	48	Gamble's solution	[75]
5-Fluorouracil	Fe-MOF	Water	77 wt %	24	PBS	[77]
Celecoxib	MIL-100	Water (20 mg/L)	~20 mg/g	312	PBS (pH=7.4)	[78]
Curcumin	Fe-MOF	Ethanol (2.7 mg/mL)	-	24	PBS+ Tween 80	[41]
Vancomycin	MOF-53	Deionized water (25 mg/mL)	20 wt %	-	PBS	[82]
Gentamicin	MIL-100	Dichloromethane Distilled water	600µg/mg 200µg/mg	12	PBS, SGF, SIS	[83]

Table 2 Summary of the Drug, the MOF Type, Loading Capacity, and In Vitro Release Studies in This Review

Abbreviations: PBS, phosphate buffer solution; SGF, Simulated gastric fluid; NS, Physiological saline; SIS, Simulated intravenous solution.



Figure 5 Different strategies for incorporating biomedical related reagents into Fe-MOFs nanomaterials.

pressure.^{36,67} (iii) Direct encapsulation strategy. Metal ions, organic ligands, and drug molecules are co-dissolved, and drug molecules are directly encapsulated into the MOFs carrier during the formation of the MOFs structure.^{73,75} (iv) Direct assembly strategy. A wide variety of biological molecules (eg, amino acids,⁷⁹ peptides,⁸⁰ and nitrogen-containing bases) or drug molecules are employed as ligands, and the biological metal-organic framework (BioMOF) is directly assembled and synthesized through the coordination bond between them and the metal Fe node.^{41,81–83}

The basic methods of drug packaging exhibit special characteristics and unique defects. The major disadvantage of adsorption synthesis and indirect packaging methods, ie, the most used synthesis strategy for drug packaging systems thus far, refers to the heterogeneous distribution of active drugs in the MOFs structure. In this heterogeneous distribution, drug molecules generally tend to concentrate on the outer surface of MOFs, which makes delivery dynamics difficult to control while reducing drug utilization and efficacy (Figure 6A). The direct packaging strategy can help to improve the heterogeneous distribution. However, it is very important to evaluate the affinity of solvents and drugs to obtain better packaging efficiency, as shown in Figure 6B. Thus, the selectivity of metal ions, organic ligands, drug molecules, and solvents should be high, as shown in Figure 6C–E. The strategy of direct assembly and synthesis of BioMOFs employs biomolecules as organic ligands, which can make drugs uniformly distributed while showing the advantages of low toxicity and high biocompatibility.⁸⁴ However, the low symmetry of biological ligands may cause an unpredictable self-assembly process, and the chemical stability and thermal stability of BioMOFs cannot conform to the requirements of conventional biological materials.

In brief, the conventional adsorption synthesis and indirect packaging strategies remain the focus of future research for their simple methods and diverse structures. The direct packaging method has been scarcely investigated, whereas it has broad application prospects. BioMOFs materials will serve as an excellent drug delivery platform and a superstar in the field of smart materials for their beautiful structure, rich supramolecular chemistry, and unique bionic characteristics.



Figure 6 The practical application of Fe-MOFs in DDS: (A) Application of MIL-100 (Fe) in Guest Encapsulation. Reprinted from Souza BE, Möslein AF, Titov K, et al. Green reconstruction of MIL-100 (Fe) in water for high crystallinity and enhanced guest encapsulation. ACS Sustainable Chem Eng. 2020;8(22):8247–8255. Creative Commons.³⁶ (B) The combined formulation of the NH2-MIL-101(Fe)/d-pen and NH2-MIL-101(Fe)/CPT-11 and the anticancer mechanism. Reprinted from Ji HB, Kim CR, Min CH, et al. Fe-containing metal-organic framework with D-penicillamine for cancer-specific hydrogen peroxide generation and enhanced chemodynamic therapy. *Bioeng Transl Med.* 2023;8: e10477. Creative Commons.⁷⁷ (C) Green hydrothermal synthesis of iron based MOFs drug carriers,⁷⁴ (D) MOF-53 (Fe) nanoparticle embeds vancomycin drug. Lin S, Liu X, Tan L, et al. Porous Iron-Carboxylate Metal–Organic Framework: a Novel Bioplatform with Sustained Antibacterial Efficacy and Nontoxicity. ACS Appl Mater Interfaces. 2017;9 (22):19248–19257. Copyright (2017), American Chemical Society.⁸² (E) DOX@Fe-MOF nanocrystals for overcoming cancer resistance/metastasis. Reprinted from Yao XX, Chen DY, Zhao B, et al. Acid-Degradable Hydrogen-Generating Metal-Organic Framework for Overcoming Cancer Resistance/Metastasis and Off-Target Side Effects. Adv Sci. 2022;9:2101965. Creative Commons.⁸⁰

Biosensor

Biosensors are devices that accurately detect biochemical reactions with simple biological molecules, amino acids, proteins, enzymes, tissues, nucleic acids, cells, or microorganisms involved.⁸⁵ In the field of health care and medicine, accurate and rapid diagnosis and monitoring using developing biosensors that are capable of recording physical, chemical or biological changes and converting them into measurable signals takes on critical significance.⁸⁶ The development of biosensors primarily aims to develop efficient sensing materials. Fe-MOFs are also considered promising candidates for biosensors for their high porosity, functional sites, and high mass transfer capacity. Moreover, Fe-MOFs, as a vital material for biosensors, have numerous advantages (eg, versatility, toxicity, biosafety, and biodegradability).⁸⁷

From Figure 7 and Table 3, we can see that the core role of Fe-MOFs in biosensors mainly includes:

(i) Carrier of sensing elements, porous Fe-MOFs exhibit strong surface area and adsorption capacity, thus making Fe-MOFs easy to be combined with various functional materials (eg, DNA molecules,⁸⁸ enzymes,⁸⁹ antibodies,⁹⁰ nanoions,⁹¹ and dyes⁹²), and they have been extensively employed as carriers of sensitive elements in a wide variety of biosensors, as shown in Figure 8A–C.



Figure 7 Application Prospects and Challenges of Fe-MOFs in the Field of Biosensors.

- (ii) Enzyme-mimic elements, Fe-MOFs show catalytic activity (also known as nanoenzymes). The above-mentioned nano-enzymes have the same or even stronger catalytic performance as natural biological enzymes, and can maintain stability under various biological unfriendly conditions.¹⁰⁶
- (iii) Electrochemical signal transduction and electrochemical sensors. Fe-MOFs as electrochemical sensors also arouse wide attention for their strong redox activity and many catalytic active sites.¹⁰⁷ However, the original characteristics of MOF (eg, slow diffusion and low conductivity) significantly hinder its efficient electrochemical application, as shown in Figure 8D. The modification of electroactive ligands, other electroactive

Functions	Sensitive Materials	Fe-MOFs	Sensing Methods	Analytes: Limit of Detection	Refs.
(i) Carrier of sensing elements	Glucose oxidase	MIL-88B- NH ₂	Colorimetric	Glucose/(0.487 μM)	[89]
	Au	MIL-53	UV-vis spectrometer	Chloramphenicol/(8.1 ng/mL)	[90]
	MXene	Fe-MOF	Electrochemical	As(III)/(0.58 ng L-1)	[93]
	CD63-aptamer	Fe-MIL-88	Visual determination	Exosomes/(5.2×104 particles/µL)	[94]
	Ag-NPs	MIL-101(Fe)	Colorimetric	H ₂ O ₂ / (0.23 μM)	[95]
	Co	TCPP(Fe)	Chemiluminescence	Glucose/(10.667µg/L)	[96]

 Table 3 Summary of Biosensing Applications of Fe-MOFs in This Review

(Continued)

Functions	Sensitive Materials	Fe-MOFs	Sensing Methods	Analytes: Limit of Detection	Refs.
(ii) Enzyme-mimic elements	Eu3+	MIL-53(Fe)	Fluorescence intensity	Cholesterol/(10.5 μM)	[97]
	Co	MIL-101(Fe)	Fluorescence Colorimetric	Vanillin/(104 nM)	[98]
	Fe3+	MIL-53(Fe)	Colorimetric	Biothiols (Cys)/(120 nM)	[99]
	Hemin	MIL-88-NH ₂	Colorimetric	H ₂ O ₂ /(0.06 μM)	[100]
(iii) Electrochemical sensors	Fe	NporMOF	Electrochemical	NO/(I.3 μM) H ₂ O ₂ /(I.I μM)	[101]
	Ferrocene	PCN-222(Fe)	Electrochemical	Dissolved oxygen (0.3 mg mL-1)	[102]
	Fe ₃ O ₄ /PPy	ZIF-8	Electrochemical	Glucose/(0.33 μM)	[103]
(iv) Optical sensors	Fe3+	PCN-224	UV-vis spectrometer	H ₂ O ₂ (22 μM) Glucose(0.3 mM)	[104]
	Fe2+	Fe(II)-MOF	Photoluminescence	Hg(II)/(1.14 nM)	[105]

Table 3 (Continued).

materials, or metal nanoparticles is a vital research direction to increase the electrochemical activity of sensors.^{93,101,102,108}

(iv) Optical sensors, Fe-MOF have good oxidase-like catalytic performance, can be applied to colorimetric detection of color detection, and have the advantages of simple, fast, cheap, and direct visual readout.^{94,97–99,104,109} It has been widely used in the detection of enzymes,^{98,106} DNA,⁹⁴ hydrogen peroxide,^{95,100,109} glucose,^{89,96,103} and so on. Besides, using the specific coordination of hydroxyl or amino groups in Fe-MOF with heavy metals, based on colorimetric and PL chemosensory, can provide highly sensitive and selective detection of arsenic⁹³ or mercury.¹⁰⁵

As depicted in Figure 7, the vital problems of clinical application of Fe-MOFs in biosensors mainly include:

- (i) In situ application or direct injection of Fe-MOFs biosensor is the vital factor for clinical application. Notably, in-situ
 detection with real-time diagnostic analysis takes on critical significance in detecting factors for toxicity and
 biosafety.⁸⁷
- (ii) At different stages of the sensor based on MOF, the comparative study of selectivity and repeatability detection takes on crucial significance. Most of the published MOF-based biosensors use low-selectivity physical adsorption or chemical processes to identify target analytes, which are not suitable for analysis and detection in complex matrices.¹⁰⁸ Accordingly, selective detection of biomarkers at different stages of various diseases is helpful to develop rapid diagnostic clinical therapies; Reusable MOF materials are also very important in large-scale applications.¹⁰⁹
- (iii) Rapid, accurate, and sensitive detection of MOFs biosensors is also recognized as a critical condition for clinical application accurate detection. If the specific recognition of biological probes (eg, antibodies or aptamers) can be directly coupled with those highly stable MOFs, the above bottleneck problem can be effectively solved. At present, although many technologies and methods have been developed to synthesize Fe-MOFs and their derivatives, the target optimization and controllable modification required for synthesis and detection, as well as Fe-based MOF materials with controllable active surfaces, is still a huge challenge.¹¹⁰

Fe-MOFs show great application prospects in the field of biosensors. The synthesis and preparation place a major focus on biocompatible materials with specific pore microenvironments, stable structures and morphology, as well as controllable surface activity. Performance research focuses on high sensitivity in situ detection and wearable detection combining



Figure 8 The practical application of Fe-MOFs in Biosensors: (A) Integrated with Glucose oxidase as a biomimetic Glucose biosensor. Reprinted from Weiqing X, Jiao L, Yan H, et al. Glucose oxidase-integrated metal-organic framework hybrids as biomimetic cascade nanozymes for ultrasensitive glucose biosensing. ACS Appl Mater Interfaces. 2019;11:25, 22096–22101. Copyright (2019) American Chemical Society.⁸⁹ (B) Efficient Biocatalytic System for Biosensing by Combining MIL-53(Fe) MOF-Based Nanozymes and G-Quadruplex (G4)-DNAzymes. Reprinted from Mao XX, He FN, Qiu D, et al. Efficient Biocatalytic System for Biosensing by Combining Metal- Organic Framework (MOF)-Based Nanozymes and G-Quadruplex (G4)-DNAzymes. Anal Chem. 2022;94(20):7295–7302. Copyright (2022) American Chemical Society.⁹⁰ (C) PCN-224(Fe) hybridized gold nanoparticles as a bifunctional nanozyme for glucose sensing. Tong PH, Wang JJ, Hu X-L, et al. Metal-organic framework (MOF) hybridized gold nanoparticles as a bifunctional nanozyme for glucose sensing. Chem Sci. 2023;14:7762–7769. Creative Commons.⁹¹ (D) Fe-MOF-525 Enables Benchmark Electrochemical Biosensing. Small. 2023. 2301933. Creative Commons.⁹³ (E) MIL-88A-Derived Fe3O4@C Hierarchical Nanocomposites for Electrochemical Sensing. Wang Y, Li X, et al. The MIL-88A-Derived Fe3O4-Carbon Hierarchical Nanocomposites for Electrochemical Sensing. ¹⁰⁴

conventional biological insight with digital instruments; the unique advantages of Fe-MOFs in versatility, biosafety and biodegradability are bound to become the focus of research in the field of biosensors.

Biocatalysis

Biocatalysis refers to the process of using enzymes as catalysts to simulate complex biological transformation and then chemical transformation.¹¹¹ Enzymes, ie, the most efficient catalyst, exhibit unique characteristics (eg, high substrate affinity and specificity). However, the practical application of natural enzymes is hindered by high cost, highly restricted catalytic conditions, and fragility for their complex and fine structure, and natural enzymes should be protected by carrier materials.^{112,113} To solve the above-described problems in the practical application of enzymes, enzyme immobilization technology and the synthesis of efficient biomimetic catalysts have become the research focus in the field of biocatalysis.¹¹⁴



a) Porous structure, b) Metal nodes, c) Organic linkers

Figure 9 Applications of Fe-MOFs nanomaterials in the Field of Biocatalysis.

Metal-organic frameworks (MOFs) have excellent tunability, good biocompatibility and significant surface modification in structural design.^{113,115} As candidates in the field of biocatalysis, they have become increasingly popular.

It can be seen from Figures 9 and 10 that the core applications of Fe-MOFs in the field of biocatalysis mainly include:

- (i) Natural enzyme immobilization matrix: Immobilizing the host enzyme on the porous organic framework carrier can protect the enzyme from the effect of harsh industrial conditions while promoting the selective diffusion of guest molecules through the carrier, and facilitating its recovery and recycling together with the carrier. MOF, a natural enzyme immobilization matrix, primarily comprises surface adhesion, covalent bond, pore interception and coprecipitation.⁵⁴ This requires the preparation of MOF materials with adjustable porosity, adjustable morphology and large surface area, and the immobilization of enzymes to stable MOFs with enzyme activity retention through appropriate strategies, as shown in Figure 10A and B. This is the critical and challenging research direction in this field.^{115,118–120}
- (ii) Biomimetic enzyme mimics: To address the problems in the practical application of natural enzymes and inspired by the catalytic mechanism of natural enzymes, bionic schemes that mimic some key characteristics of enzymes have become effective in developing efficient biomimetic catalysts.¹²¹ Biomimetic Fe-MOFs catalysts can satisfy some enzymatic characteristics (eg, the structural characteristics of enzymes, active sites, microenvironment, co-catalytic sites, and transmission channels) while showing the advantages of wider application conditions and stronger stability (Figure 10C). Besides, they will exhibit catalytic properties that natural enzymes do not exhibit, which are elucidated as follows: (1) homogeneous separation of biomimetic MOF catalytic sites to avoid self-accumulation and deactivation; (2) appropriate hydrophilic and hydrophobic pore properties to promote the recognition and accumulation of reactant molecules; (3) synergetic microenvironment to improve the reactivity between active sites and substrates; (4) restricted pore size, shape and function to improve chemistry, regionally, stereology and selectivity; (5) auxiliary active sites to improve catalytic efficiency and reaction rate; (6) simple reuse and high stability. Compared with enzymes, the applicable pH range, reaction temperature and solvent of biomimetic MOFs are significantly broadened.^{89,96,116,122-124} With the significant progress in the latest characterization technology to reveal the structure and catalytic mechanism of enzymes, the important synergy of microenvironments and co-catalytic sites has been recognized progressively, such that considerable scientific and technological efforts have been attracted for the development of efficient biomimetic MOFs catalysts.
- (iii) Biomimetic enzyme/natural enzyme hybridization: Due to the comprehensive advantages of the selectivity of natural enzymes and the controllable catalytic activity of nanoenzyme, nanoenzyme/natural enzyme hybridization takes on critical significance in biosensor, treatments, and catalysis, as shown in Figure 10D. Extensive research should be conducted to properly address the interaction mechanism and biocompatibility of biomimetic enzyme/natural enzyme.^{117,125}



Figure 10 The practical application of Fe-MOFs in biocatalysis: (A) Cu/Au/Pt TNP-modified TCPP-(Fe) nanozyme. Reprinted from Wu P, Gong F, Feng X, et al. Multimetallic nanoparticles decorated metal-organic framework for boosting peroxidase-like catalytic activity and its application in point-of-care testing. *J Nanobiotechnol.* 2023;21:185. Creative Commons.¹¹² (B) MIL-101(Cr/Fe) used as a platform for MP-8 enzyme immobilization. Reprinted from Kesse X, Sicard C, Steunou N, et al. Encapsulation of Microperoxidase-8 into MIL-101(Cr/Fe) Nanoparticles: a New Biocatalyst for the Epoxidation of Styrene. *Eur J Inorg Chem.* 2023;26:e202300040. Creative Commons.¹¹⁵ (C) Multi-compartmental MOF microreactors for chemo-enzymatic cascade catalysis. Reprinted from Tian D, Hao R, Zhang X, et al. Multi-compartmental MOF microreactors for chemo-enzymatic cascade catalysis. Nat Commun. 2023;14:3226. Creative Commons.¹¹⁶ (D) Fe-MOF based bio-/enzyme-mimics used for cancer treatment and anti-tumor principle. Reprinted from Xiang X, Pang H, Ma T, et al. Ultrasound targeted microbubble destruction combined with Fe-MOF based bio-/enzyme-mimics nanoparticles for treating of cancer. *J Nanobiotechnol.* 2021;19:92. Creative Commons.¹¹⁷

In general, Fe-MOFs improve the stability and reusability of encapsulated biocatalysts while expanding their application scope in the field of biocatalysis.¹²⁶ The research focus will be placed on the development of simple and mild enzyme immobilization technology, the design and synthesis of novel MOF with a mesoporous structure to facilitate the transport of enzyme materials, and gaining insights into more detailed interaction mechanism between enzyme and Fe-MOF to more effectively boost industrialization. The integration of biocatalysts and Fe-MOFs is highly promising for industrial biocatalysis in the future.

Bioimaging and Phototherapy

The current cancer treatment methods mainly include surgery, chemotherapy, radiotherapy, endocrine therapy, and targeted therapy. To increase the therapeutic effect of chemotherapy, the image-guided chemical-photothermal combined therapy strategy combining chemotherapy and phototherapy has aroused wide interest.^{127,128} The above-mentioned method primarily refers to a treatment method that employs targeted recognition imaging or development technology to transform light energy into heat energy in the vicinity of tumor tissue by injecting materials with high light and heat conversion efficiency, and kill cancer cells locally. The above-described technology makes surgical treatment less invasive and more accurate, and it is capable of shortening hospital stays and reducing repeated operations.¹²⁹ At

present, common imaging and photothermal agents comprise metal nanostructures, carbon-based materials, metal oxides and chalcogenides, metal-organic frameworks (MOFs), Mxenes, and so forth. To be specific, chemically and thermally stable Fe-MOFs are employed in drug delivery, biosensors, biocatalysts, and other fields, and they exhibit excellent optical and photothermal properties. Besides, they have been commonly adopted as imaging agents for magnetic resonance imaging (MRI)¹³⁰ and photoacoustic imaging (PAI).¹³¹ Furthermore, they serve as a photothermal agent and carrier for photothermal therapy (PTT)¹³² and photodynamic therapy (PDT).¹³³

(i) Imaging agent, developer, and carrier: Magnetic resonance imaging (MRI), a non-invasive method for imaging and tissue characterization of living organisms, has been the most widely applied and effective imaging technology.¹³⁴ Because Fe³⁺ is a paramagnetic metal ion, Fe-MOF has become an excellent candidate for MRI imaging, as shown in Figure 11.¹³⁰ The existence of numerous paramagnetic metal centers makes Fe-MOFs tend to exhibit higher relaxation than small molecular contrast agents (CA). In general, a higher relaxation value will help to improve the sensitivity and specificity of contrast media. However, most of the single Fe-based MOF materials have only moderate relaxation (relaxation), which leads to the lack of sensitivity of MRI imaging.^{135–137}

Photoacoustic imaging (PAI) and photothermal imaging (PTI) are novel non-invasive soft tissues medical imaging modes under photoacoustic (PA) and photothermal effects.¹³⁹ They refer to the generation of sound waves by local tissues irradiated by pulsed laser and the reception of local thermal expansion signals by ultrasonic detectors. Both PAI and PTI have the characteristics of high resolution, rich optical contrast, and tissue penetration depth, and show broad application prospects in visualizing tissue structure and function. Fe-MOF materials have less research in the application of PAI, which may be related to the weak optical absorption of the above-mentioned materials.^{140,141}

(ii) Therapeutic agent and carrier for photothermal therapy and photodynamic therapy: Photothermal therapy (PTT) refers to a treatment method that uses materials with high photothermal conversion efficiency to kill cancer cells locally by converting light energy into heat energy.^{132,142} Photodynamic therapy (PDT) has been confirmed as



Figure II (A) Schematic illustration of imaging-guided therapy. (B) Photodynamic tumor therapy (PDT) by reversing multiple resistances. Reprinted from Liu P, Zhou Y, Shi X, et al. A cyclic nano-reactor achieving enhanced photodynamic tumor therapy by reversing multiple resistances. *J Nanobiotechnol.* 2021;19:149. Creative Commons.¹³⁸ (C) MRI imaging of orthotopic pancreatic murine tumors. Reprinted from Rojas JD, Joiner JB, Velasco B, et al. Validation of a combined ultrasound and bioluminescence imaging system with magnetic resonance imaging in orthotopic pancreatic murine tumors. *Sci Rep.* 2022;12:102. Creative Commons.¹³⁰ (D) PAI images and (E) Photothermal photographs of doxorubicin@MIL-100 compound. Reprinted from Zhiming H, Caina X, Liang Y, et al. Multifunctional drug delivery nanoparticles based on MIL-100 (Fe) for photoacoustic imaging-guided synergistic chemodynamic/chemo/photothermal breast cancer therapy. *Mater Design.* 2022;223:111132. Creative Commons..¹³¹

a novel technology that exploits the photodynamic effect to diagnose and treat diseases.^{133,138,143} Its mechanism of action is to induce photosensitizer to produce reactive oxygen species (ROS) by laser of specific wavelength, and produce cytotoxicity by oxidizing reaction with adjacent biological macromolecules, triggering tumor cell damage and even death. PTT and PDT are characterized by minimally invasive, long-term, high-precision, and controllable treatment process in the treatment of malignant tumors, and they have aroused wide attention by scientific researchers. In terms of PTT or PDT, non-toxic or low-toxic and biodegradable iron (III) based MOF serves as an ideal carrier of photosensitized Fe-soc-MOF nanoparticles combined with ICG.¹⁴⁴ It will be its main research direction in facilitating the near-infrared (NIR) absorption of Fe-MOFs and increasing the light stability, especially in the multi-functional therapeutic agent combining MOFs with other functional materials.^{145–149}

Fe-MOFs exhibit high research value and broad application prospects in image-guided chemical-photothermal therapy strategy. First, given the accumulation, retention, and metabolism of residues in the body, the degradation mechanism and biological stability of existing Fe-MOFs should be investigated in depth. Second, it is important to synthesize Fe-MOF materials with highly repeatable physical properties and systematically evaluate the toxicity in vivo. Besides, in PAI and PTT technology, the use of nanoparticles (NPs) with unique magnetism and excellent biocompatibility combined with appropriate Fe-MOFs to synthesize nanocomposites that can provide effective treatment and diagnosis will be its main research direction.

Antimicrobial MOFs

With the increasing resistance of pathogens to antibiotics (AMR) over the past few years, infections arising from microorganisms (eg, bacteria, fungi, viruses, and parasites) pose increasingly serious harm to public health.^{150,151} Thus, the development of antibacterial agents with high bactericidal activity (eg, antibacterial peptides, bacteriophages, and antibodies) has attracted great attention.¹⁵² Although the conventional phytochemical antibacterial agents have high biocompatibility, their effect on treating pathogenic microorganisms is poor, and their stability and bioavailability in the water environment have also been limiting factors.^{153,154} The scientific progress of nanotechnology and material science has endowed nanoparticles (NPs) with higher antibacterial activity, including metals and their compounds (eg, metal oxides, metal salts, and metal hydroxides), inorganic-organic hybrid nanoparticles (NPs) and polymers, promising alternative methods for the treatment of AMR-associated infections.^{155–159} However, their application in clinical research has triggered several problems (eg, cytotoxicity and non-targeted effects).

MOF materials and MOF matrix composites are considered a type of potential antibacterial materials, though their antibacterial effect research remains in its infancy.^{160,161} Fe-MOFs show several advantages (eg, low-cost composition, easy preparation, water solubility, and long-term stability), and they significantly apply to medical sterilization and bacterial eradication, as well as the growth inhibition of antibacterial agents.^{162–165} Figure 12 presents the active site and main applications of antibacterial Fe-MOFs and their composites. The main research directions of Fe-MOFs in antimicrobials are presented as follows.

- (i) MOFs as metal ions reserve: Fe-MOFs can serve as sites of active metal ions, ie, original antibacterial. For instance, existing research has suggested that MIL-101 (Fe) exhibits high-free radical scavenging activity, and the MIC values of Escherichia coli (ATCC 512), Enterobacter planus (ATCC 256), Lactobacillus pneumophilus subspecies (ATCC 4), Bacillus cereus, Staphylococcus aureus (ATCC 256), Pseudomonas aeruginosa (ATCC 32), and Candida albicans reach 16, 256, 10536, 10541, 33152, 6538, and 9027, respectively.⁴³ As indicated by the above analysis, Fe-MOFs molecule exhibit high antibacterial activity against microorganisms, fungus, and yeast, as shown in Figure 13A and B.^{162,163}
- (ii) MOFs containing bioactive links: For instance, it is obtained through the self-assembly of Fe³⁺ ion and doxorubicin hydrochloride (DOX) molecule Fe-DOX@Gd-MOF, showing higher stability and controllable DOX release, as shown in Figure 13C.¹³⁹ Nevertheless, there has been scarce research in this category.
- (iii) For antimicrobial control delivery carriers: For instance, silver ion was loaded into Fe-MOFs as a typical antibacterial agent to form an antibacterial system, showing excellent antibacterial behavior against Staphylococcus aureus and



Figure 12 Active site and main applications of antibacterial Fe-MOFs nanomaterials and Their Composites.

Escherichia coli.^{167,168} Gentamicin (GM) can also be encapsulated with Fe-MIL-100 through a simple permeation procedure. Antibacterial tests on Staphylococcus aureus, Streptococcus epidermidis and Pseudomonas aeruginosa also confirmed that the released transgenic antibiotic activity was retained.⁸³ Fe-MOF-112 prepared in the form of nanoparticles has been used as a nano-carrier for the precise transport of 3-azido-d-alanine (d-AzAla).¹⁶⁰ After intravenous injection, 112 NPs have been found to accumulate in the infected tissue and release d-AzAla in the inflammatory environment with high H₂O₂, which is selectively incorporated into the cell wall of MRSA bacteria.¹⁶⁰ Figure 13D and E shows the antibacterial mechanism of La³⁺, Gd³⁺ metal ion doping¹⁶⁵ and Carvacrol encapsulated in Fe-based MOFs.¹⁶⁶ On the other hand, Fe-MOFs can also appear in the form of various composites, including fiber, polymer, gel or nanocomposite.^{169–172}

In general, biosafety Fe-MOF materials seem to be very suitable for medical sterilization, bacterial eradication or as carriers of antiviral agents, because they combine the advantages of two different fields, namely inorganic antibacterial agents with long sterilization time and broad-spectrum sterilization.^{173,174} However, at present, there are few research results on antibacterial Fe-MOFs. We can foresee that research in this field will increase significantly under the context of the recent COVID-19 pandemic and the rise of potential risks in the microbial world.

Discussion and Future Outlook

This study focuses on the key index requirements of Fe-MOFs materials in biomedical applications, as well as their extensive research in biomedical applications (eg, drug delivery, biosensors, photothermal therapy and antibacterial activity), suggesting the huge scientific and technological potential of the above-described materials. Fe-MOFs material serves as an ideal candidate in the biomedical field for its numerous advantages. For instance, its good biocompatibility and biodegradability can be applied to drug delivery; large specific surface area and high porosity provide a solid basis for the use of biosensors and biocatalysts; the open structural framework and active metal sites expand the ability of photothermal therapy and antibacterial; finally, its simple green synthesis process is also conducive to large-scale production. However, the research of Fe-MOF materials for biomedical applications is in the early stage of exploration, many long-term obstacles and several challenges remain.

(i) The size optimization, shape control and structural stability of Fe-MOFs take on critical significance in drug delivery and biosensors. The uniform size and stable morphology are conducive to expediting the distribution and reuse of Fe-MOFs in the organism. However, it is difficult for the self-assembly synthesis process to apply to the



Figure 13 (A) Surface-Anchored MOFs-Cotton Material with antibacterial properties. Reprinted from Rubin HN, Neufeld BH, Reynolds MM. Surface-anchored metal-organic framework-cotton material for tunable antibacterial copper delivery. ACS Appl Mater Interfaces. 2018;10(17):15189–15199. Copyright (2018) American Chemical Society.¹⁶² (B) The antibacterial mechanisms of Fe(III)-MOF towards different types of microorganisms, fungus, and yeast. Reprinted from Sheta SM, Salem SR, El-Sheikh SM. A novel Iron (III)-based MOF: synthesis, characterization, biological, and antimicrobial activity study. *J Mater Res.* 2022;37(14):2357–2367. Creative Commons.¹⁶³ (C) Synthesis of a mesoporous MIL-100(Fe) bacteria exoskeleton Reprinted from Permyakova A, Kakar A, Bachir J, et al. In Situ Synthesis of a Mesoporous MIL-100(Fe) Bacteria Exoskeleton. *ACS Materials Lett.* 2023;5(1):79–84. Copyright (2023) American Chemical Society.¹³⁹ (D) Smart MIL-88B(Fe) coating as a host matrix for the antibiofilm compound. Reprinted from Claes B, Boudewijns T, Muchez L, et al. Smart metal-organic framework coatings: triggered antibiofilm compound release. *ACS Appl Mater Interfaces.* 2017;9(5):4440–4449. Copyright (2017) American Chemical Society.¹⁶⁵ (E) Antibacterial properties of Carvacrol encapsulated in MIL-100 (Fe) nanoparticles. Reprinted from Caamaño K, Heras-Mozos R, Calbo J, et al. Exploiting the Redox Activity. *ACS Appl Mater Interfaces.* 2022;14:10758–10768. Creative Commons.¹⁶⁶

synthesis of Bio-MOFs due to the low symmetry of many biological ligands. Moreover, the size, shape, and porosity of Fe-MOFs can hardly be accurately controlled. Thus, improving the structural stability, accurate particle size control, and simple synthetic process route are crucial in widening their biological application. This will serve as crucial content to guide the design and application of Fe-MOF materials in the future.

- (ii) A considerable number of physiological mechanisms of Fe-MOFs (eg, biodegradation mechanism, pharmacokinetics, antibacterial and bacteriostasis mechanism) remain unclear. Although some Fe-MOFs materials or their nanocomposites have been extensively explored in vitro and in vivo, there is little research on the metabolic mechanism of Bio-MOF materials. Further exploring the biological logic and molecular level theoretical research of absorption, distribution, metabolism, and excretion (ADME) in vivo under complex substrate conditions (eg, the effect of pH, temperature, or other active molecules) is also a vital research direction of Fe-MOFs.
- (iii) The application of single-structure Fe-MOF materials is also restricted by biocompatibility and cytological toxicity. Their biosafety and biodistribution in vitro and/or in vivo must approach the requirements of practical biomedical applications. Future research will focus on the development of mixed metal MOF materials based on highly biocompatible metal ions (eg, Ca and Zn), functional MOF materials combined with bioactive ligands, and graded MOF materials with surface modification, so as to significantly promote the expansion of the application range of Fe-MOF materials.

In brief, early reports on Fe-MOF materials in the biomedical field have revealed their extraordinary and unique properties, while some challenges should be explored in future research. With the latest development of MOF-based nanocomposites, their application in biomedicine provides exciting new avenues for the clinical development of nanotechnology. Hence, technologies based on Fe-MOF materials have brighter prospects in the biomedical field.

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Disclosure

The authors declare no conflicts of interest in this work.

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493 I