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MOF-based spectrophotometric sensors for cholesterol detection: current trends and challenges

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Cholesterol detection is essential for early diagnosis and monitoring of cholesterol-related diseases, such as atherosclerosis, hypercholesterolemia, and liver diseases. A variety of nanomaterials have been designed and synthesized for cholesterol detection *via* electrochemical and spectrophotometric techniques. Metal organic frameworks (MOFs) have emerged as promising detector materials for cholesterol sensing. Recent research explores MOFs as spectrophotometric cholesterol sensors with remarkable performance in terms of limit of detection and selectivity. Given the growing interest in cholesterol sensing, and limited reviews on recent advances in the field, this review critically examines recent advances in MOF-based spectrophotometric cholesterol sensors, outlining the different mechanistic roles of MOFs in cholesterol detection. The review also highlights significant improvements, current challenges, and potential applications of MOF-based sensors for cholesterol detection in point-of-care devices and medical diagnostics.

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1. Introduction

Cholesterol plays a vital role in human health. It is involved in several roles including hormone synthesis, cell membrane integrity and fluidity, vitamin D production, bone health, among others.¹ However, excess cholesterol levels (hypercholesterolemia) can cause atherosclerosis and increase the risk of cardiovascular diseases, strokes and obesity.^{2–5} In fact, hypercholesterolemia is considered the third most common cardiovascular risk factor worldwide while also being the least monitored factor.^{5,6} In addition to atherosclerosis, high cholesterol levels can lead to toxic accumulation in the liver, pancreatic islets, kidneys, and other organs, subsequently increasing the incidence of nonalcoholic fatty liver disease, nonalcoholic steatohepatitis, islet B-cell dysfunction, renal dysfunctions, testosterone deficiencies, osteoporosis and osteoarthritis, among others.⁷ Hence, continuous monitoring of cholesterol levels is crucial in patients with atherosclerosis and other cholesterol-related diseases, for early detection and maintenance of diseases to maintain patient health.⁸

Cholesterol exists in blood serum in two forms: free cholesterol and cholesteryl esters, the latter being the most common.^{9,10} Due to cholesterol's insolubility in aqueous

solutions, they are transported in the bloodstream *via* lipoproteins.¹⁰ Lipoproteins consist of a hydrophobic core carrying triglycerides and cholesteryl esters and a hydrophilic membrane made of phospholipids, free cholesterol, and apolipoproteins.¹⁰ There are many types of lipoproteins, including chylomicrons, high-density lipoproteins (HDLs), low-density lipoproteins (LDLs) and very low-density lipoprotein (VLDLs), with LDLs carrying the majority of cholesterol in the blood.^{9,10} LDLs carry cholesterol from the liver to cells for storage, while HDLs transport cholesterol from cells to the liver, where it is broken down or excreted, as illustrated in Fig. 1.^{1,8–10} Furthermore, LDLs are commonly known as “bad cholesterol,” as elevated levels are correlated with increased arterial wall build up, increasing risks of atherosclerosis.^{7,10,11} Meanwhile, HDLs are commonly coined “good cholesterol,” because of its anti-inflammatory and anti-arteriosclerotic action by removing free cholesterol from cells, a process known as reverse cholesterol transport.¹¹ Due to the complexity of cholesterol availability in the bloodstream, various cholesterol detection methods have been developed.

Cholesterol detection has evolved greatly over the past years. Early methods of detection were simple, quantifying total cholesterol.¹² Later techniques evolved to be more specific, quantifying HDLs, LDLs, triglycerides, free cholesterol, and total cholesterol.^{13–15} The current, standardized method used in hospitals and clinics is the cholesterol oxidase/peroxidase/4-aminophenazone (CHOD-POD-PAP) method, a colorimetric, enzyme-based method.¹⁴ Although widely effective, the assay requires the use of several enzymes, which have low stability, limiting pH, temperature, and operation and storage conditions

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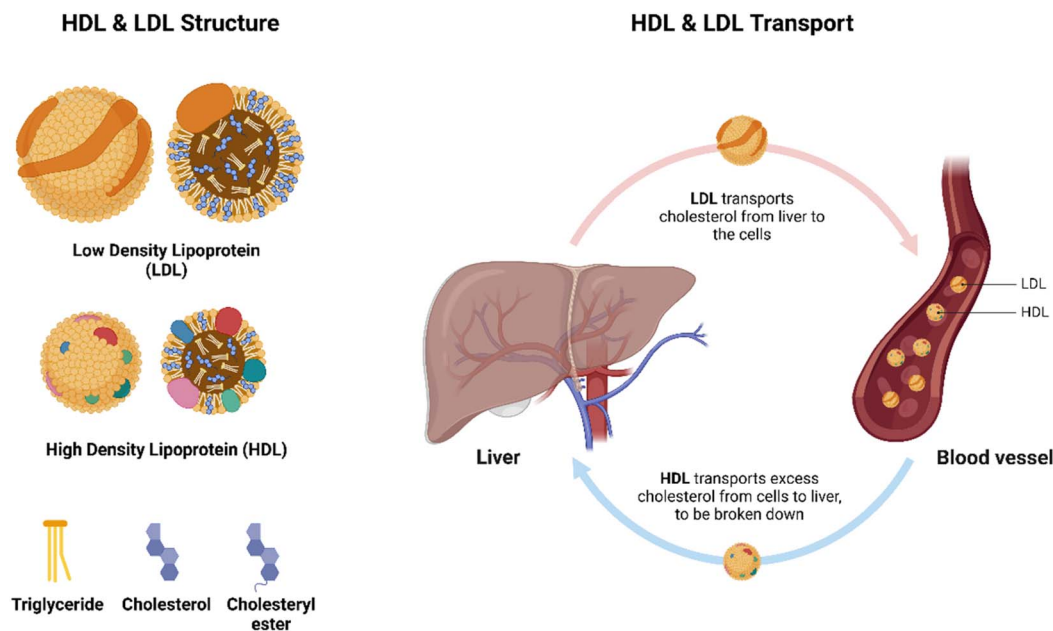



Fig. 1 Structure and metabolic pathway of HDL and LDL.

thus also introducing extra costs.^{13,16–18} Recent studies have shown the high potential of nanomaterials for the development of cholesterol sensors, for rapid and accurate cholesterol quantification.¹⁹ Such nanomaterials include metal nanoparticles, metal oxide nanoparticles, carbonaceous materials, and other composite materials.^{8,19} These have facilitated cholesterol biosensing *via* electrochemical, colorimetric and photoluminescent techniques.^{8,19–22} More recently, metal-organic-frameworks (MOFs) have emerged as superior nanomaterials for cholesterol biosensing, especially for spectrophotometric-based detection (including photoluminescent and colorimetric). This is attributed to the remarkable tunability of the MOF's structures and their large surface areas.^{18,23–31} Furthermore, MOFs exhibit enhanced selectivity, and stability in the immobilization of enzymes.^{16,32,33} Hence, MOFs have great potential for future applications in point-of-care (PoC) cholesterol biosensors.

Research on cholesterol biosensors has grown in popularity over the past two decades. Especially after 2020, a spike in research articles with cholesterol sensing or detection in the title alone is observed, as illustrated in Fig. 2; this reflects the growing interest in cholesterol sensing. Comprehensive reviews have explored electrochemical biosensors for cholesterol detection^{19–22} along with other cholesterol biosensors.⁸ MOFs as photoluminescent sensors³³ and nanozymes^{16,32,34} have also been reviewed extensively. However, to the best of our knowledge, there is a gap in existing reviews as limited focus has been dedicated to reviewing MOFs as spectrophotometric sensors specifically for cholesterol. An increasing interest in MOF-based biosensors for spectrophotometric detection of cholesterol has been also noticed over recent years, growing from 2–4 to 11 publications per year, most of which are journal articles, as illustrated in Fig. 3. Due to this increased research activity, this

paper aims to review and evaluate current research on spectrophotometric MOFs for cholesterol biosensing and address current challenges, propose potential solutions, and outline future research directions. This review exclusively concentrates on spectrophotometric MOF biosensors, which include colorimetric, fluorescent, and chemiluminescent techniques, as electrochemical biosensors have been extensively covered in previous reviews.^{19–22,35} Moreover, this review adds value to the literature by categorizing and analyzing all spectrophotometric, MOF-based cholesterol sensing mechanisms, providing a comprehensive evaluation that has not been previously undertaken.

The outline of this review is as follows: first, traditional and nanomaterial-based spectrophotometric cholesterol detection methods are reviewed (Section 2). Then, the basics of MOFs and their promising properties for applications in sensing are explored (Section 3). Next, a more detailed analysis of MOF-based spectrophotometric cholesterol sensors is conducted, by discussing the roles of MOF in sensing mechanisms and performance (Section 4). Potential integration of these MOFs into sensing platforms is then assessed (Section 5), and the paper concludes with current challenges associated with MOF stability, multi-biomarker detection, continuous, non-invasive sensing and environmental concerns and future directions (Section 5).

2. Cholesterol detection methods

Early methods of cholesterol detection utilized different reagents to enhance cholesterol's chromophore properties for quantification by measuring absorbance of the produced color, a colorimetric technique.^{12,36,37} Later studies utilized enzyme-based colorimetric assays for more accurate and rapid results,



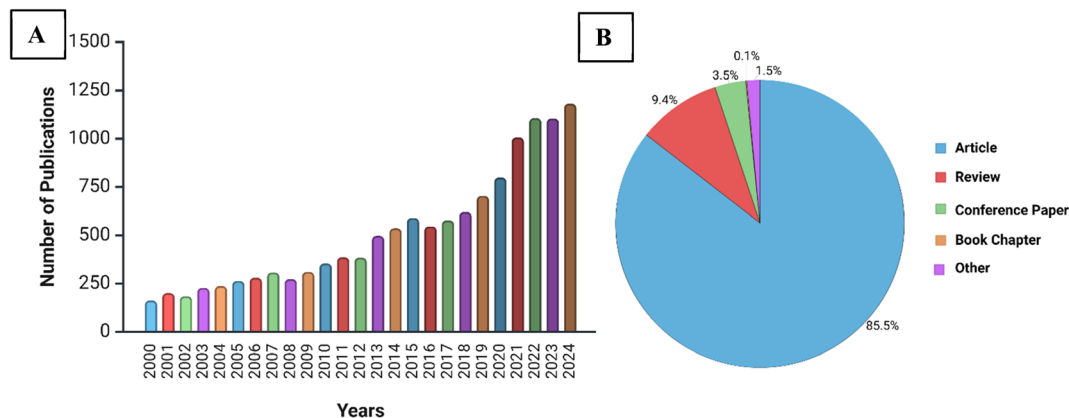


Fig. 2 (A) Growth in and (B) type of Scopus publications from 2000–2024 exploring general cholesterol detection and sensing. Data obtained from Scopus.com, using “cholesterol” and “detection” or “sensor” as the search terms in the article title, keywords, or abstract. Retrieved on Nov 20, 2024.

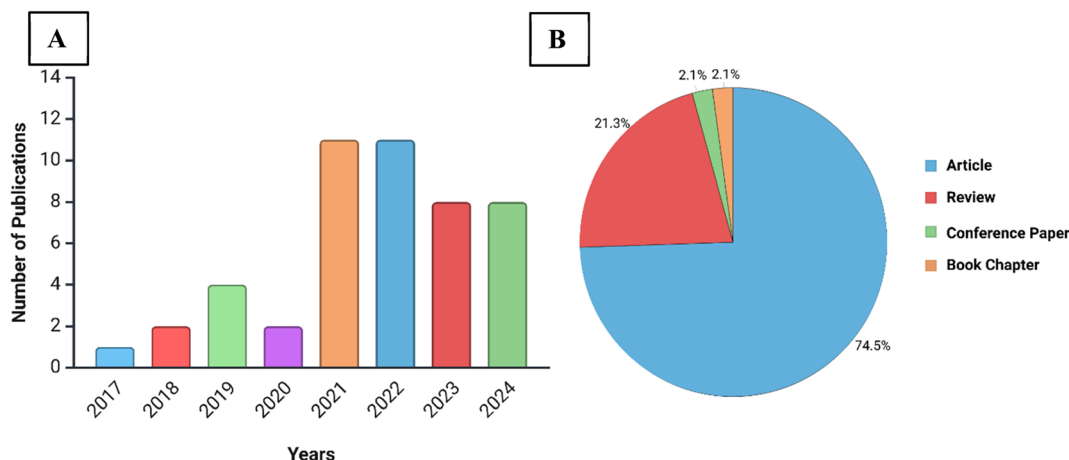


Fig. 3 (A) Growth in and (B) type of Scopus publications from 2017–2024 exploring MOF-based cholesterol detection and sensing. Data obtained from <https://www.scopus.com/>, using “metal organic framework” and “cholesterol” and “detection” or “sensor” as the search terms in the article title, keywords, and/or abstract, retrieved on Nov 20, 2024.

with methods dating back to 1972.^{13,38} Current standardized clinical methods for cholesterol detection also utilize enzymes for a colorimetric assay.^{13–15} However, with advances in research and nanotechnology over the past two decades, various new methods have emerged for cholesterol detection and quantification using different nanomaterials, including electrochemical,^{19,21} colorimetric,^{39–43} and photoluminescent techniques.^{1,44–47} The latter two methods fall under the general umbrella of spectrophotometric techniques. This section herein presents the traditional cholesterol detection method then explores colorimetric and photoluminescent methods for cholesterol detection.

2.1 Traditional, conventional methods

The conventional methods for cholesterol detection vary depending on the medium in which it is being detected. Cholesterol detection is conducted using two methods: clinical, colorimetric assays and reflectance photometric techniques,

often adopted in hand-held devices; both of which are enzyme-based.^{13–15} Electrochemical methods were also introduced, but have since evolved into more recent, improved sensors.^{19,35} Meanwhile, cholesterol quantification in food industry and in research laboratories traditionally adopt chromatographic and spectrometric techniques, such as gas-chromatography-mass spectrometry (GCMS) and high-performance liquid chromatography (HPLC), among others.⁴⁸ This paper herein focuses on medical applications of cholesterol detection, excluding chromatographic and spectrometric detection methods.

2.1.1 Traditional clinical detection. The standardized, traditional clinical cholesterol quantification technique uses an enzyme-based colorimetric bioassay, known as the cholesterol oxidase/peroxidase/4-aminophenazone (CHOD-POD-PAP) method.^{13–15} The analysis involves three main steps. First, blood serum is treated with cholesteryl esterase enzyme, to convert cholesteryl esters into cholesterol. Next, cholesterol oxidase (ChOx) enzyme is used to selectively react with



cholesterol and produce cholest-4-ene and hydrogen peroxide (H_2O_2). Finally, 4-aminoantipyrine, a chemical reagent, is added to react with the produced hydrogen peroxide, in the presence of horseradish peroxidase (HRP) enzyme as a catalyst, to produce a colored quinoneimine dye. The intensity of the produced color is measured using a spectrophotometer and used to determine the concentration of cholesterol.^{13,14} An illustration of these reactions is shown in Fig. 4 below.

The CHOD-POD-PAP method is suitable for the detection of total cholesterol, as well as HDLs, LDLs, and triglycerides. The latter three techniques require pretreatment before analysis.¹⁴ This enzymatic, colorimetric bioassay is widely used as a reference and in clinicals, and is advantageous in its accuracy, selectivity, and the absence of corrosive chemicals.⁵² The method also has a wide linear range of detection, detecting up to 800 mg dL^{-1} and even 1000 mg dL^{-1} of cholesterol (equivalent to 20.7 mM and 25.9 mM , respectively), depending on the commercial kit used.^{14,15,53} Typical levels of cholesterol range around 200 mg dL^{-1} ; however, some kits can detect at lower concentrations of 10 mg dL^{-1} , which is useful for diluted samples.¹⁵ Furthermore, detection is often quick, with incubation times ranging between 5 to 10 minutes, and incubation temperatures varying between room temperature ($20\text{--}25 \text{ }^\circ\text{C}$) or physiological temperature ($37 \text{ }^\circ\text{C}$).^{14,15,53} Although overall advantageous, this traditional method of cholesterol detection is heavily dependent on enzymes, which require specific pH and temperature conditions for storage and operation, and are known for their low stability, short lifetime, and lack of reusability.^{13,16–18} Natural HRP is especially recognized for its low stability to heat and chemical changes, making the method costly and time-consuming.^{17,18} Hence, while advantageous, the CHOD-POD-PAP traditional cholesterol detection method can be significantly optimized.

2.1.2 Reflectance photometric methods. Reflectance photometry for cholesterol detection is among the earliest techniques introduced. Through an enzymatic assay similar to

the CHOD-POD-PAP assay, hydrogen peroxide is produced and interacts with a dye to produce a color change. The color intensity is quantified by measuring the light reflectance off the colored strip.^{54,55} The amount of reflected light is correlated with the hydrogen peroxide concentration and can be subsequently used for cholesterol quantification.^{54,55} This technique dates back to 1980–1990's and was greatly effective during that time for cholesterol detection and other biomarkers, including glucose and hemoglobin.^{54–56} Furthermore, this method has been traditionally incorporated in handheld, commercial devices for cholesterol detection, such as Accutrend,⁵⁷ STANDARD LipidoCare⁵⁸ and CardioChek,⁵⁹ and is advantageous in its rapid nature. However, the accuracy of such devices varies significantly, with many of which do not meet current standards.⁶⁰ Accutrend Plus reported the highest sensitivity (92%) and specificity (89%).⁶⁰ Nonetheless, the varying accuracy poses a challenging limitation, especially when used in commercial, PoC cholesterol detection devices.

2.2 Nanomaterial-based methods

To overcome the limitations of the conventional methods, recent studies have shifted to using different nanomaterials for cholesterol detection. These nanomaterials include metal and metal oxide nanoparticles, carbonaceous materials, including carbon nanotubes, carbon dots, and graphene, and composite materials including two or more types.^{8,19,61,62} Some use nanomaterials to immobilize HRP increasing its stability and prolonging its lifetime, reducing costs.^{44,63} For example, platinum nanoclusters,⁶³ silicon nanoparticles,⁴⁴ and poly(thionine)-modified glassy carbon electrodes⁶⁴ all proved effective in immobilizing HRP for accurate and sensitive cholesterol detection. Others use nanoparticles with peroxidase-like activity, to replace HRP altogether, overcoming its stability challenges.⁶⁵ Examples include nitrogen-doped carbon nanosheets with Fe–Ni entrapped inside,⁶⁶ Cu–salt–Fe composites,⁶⁷

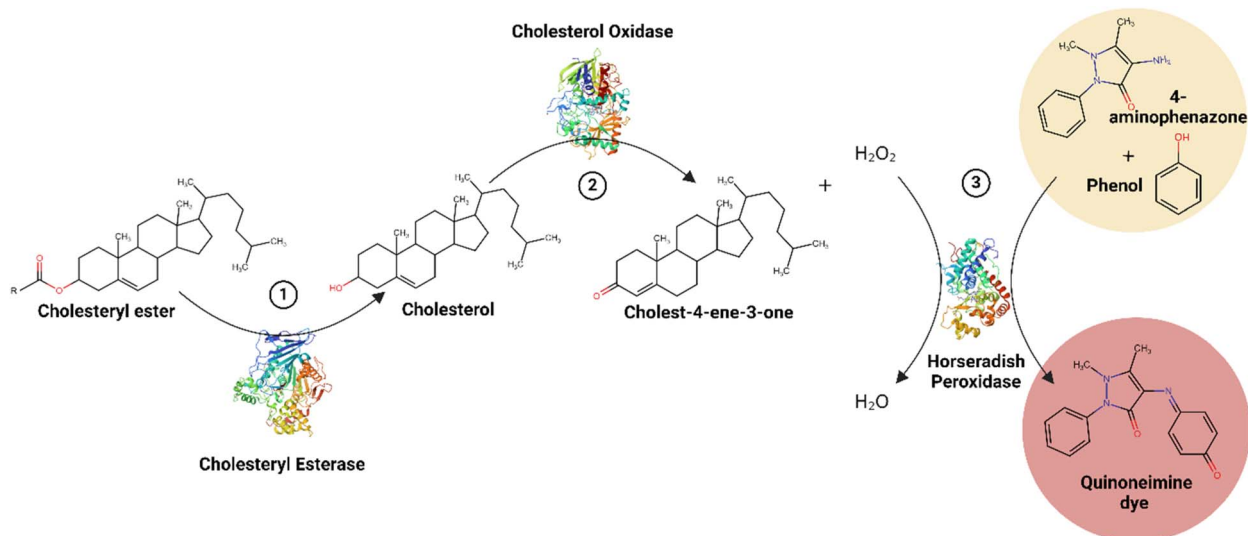


Fig. 4 Traditional, clinical CHOD-POD-PAP method summary.^{49–51}



copper-doped cerium dioxide nanosheets,⁶⁸ all of which exhibited great sensitivity and selectivity to cholesterol. These sensors utilized spectrophotometric^{39–47,69–74} and electrochemical³⁵ measurements for cholesterol quantification. Electrochemical-based cholesterol sensors determine cholesterol concentration by measuring changes in current due to a redox reaction, in potential between a reference and working electrode, or in impedance.^{19,35} Electrochemical sensing methods are advantageous due to their relatively lower cost, portability, and high sensitivity; comprehensive reviews on electrochemical-based cholesterol sensors can be found in ref. 19, 21 and 35. In addition, plasmonic-based, optical techniques of cholesterol detection have also emerged.^{75–78} For instance, bimetallic silver-shelled gold nanorods proved effective in detecting both triglycerides and cholesterol *via* their plasmonic properties, allowing for detection through surface enhanced Raman spectroscopy.⁷⁶ Moreover, graphene oxide coupled with silver nanowires have enabled optical detection of cholesterol at low concentrations.⁷⁵ Comprehensive reviews on plasmonic-based biosensors can be found in ref. 77 and 78.

More recently, spectrophotometric-based sensors have emerged as sensitive and accurate cholesterol sensors.^{39–47,69–74} Spectrophotometric detection methods include colorimetric, fluorescent, and chemiluminescent techniques, illustrated in Fig. 5. A variety of nanomaterials have been designed for spectrophotometric-based cholesterol sensing. These will be herein discussed. An overview of these nanomaterials, the cholesterol sensing operating principle, the limits of detection and linear ranges achieved are summarized in Table 1.

2.2.1 Colorimetric. Colorimetric-based sensors are optical sensors that operate by detecting a change in color upon

a chemical reaction with the analyte. More specifically, this color change occurs within the visible light wavelength region (400–800 nm), which is often detectable by the naked eye, and can be quantified by detecting change in absorbance at a fixed wavelength.⁷⁹ Colorimetric methods of detection are simple, less costly, rapid, and sensitive, with relatively low background interferences.^{40,43} The traditional, clinical cholesterol detection method, explained earlier in Section 2.1, employs colorimetric-based cholesterol detection. However, a significant limitation is the heavy dependence on enzymes, especially HRP, which is known for its instability.^{17,18}

Recent colorimetric sensors integrate nanomaterials with inherent peroxidase-like activity, to replace HRP enzyme for more efficient sensing and longer operation.^{39–43} For example, polypyrrole nanoparticles (PPy NPs), an organic, semi-conducting NP, demonstrated good peroxidase-like activity, and allowed for complete replacement of HRP. In combination with the dye 3,3',5,5'-tetramethylbenzidine (TMB) and ChOx, the PPy NPs facilitated cholesterol detection, with a limit of detection of 3.5 μM .⁴³ Other works explored the use of nanocomposites, combining a minimum of two nanomaterials for improved peroxidase activity and cholesterol sensing.^{39–43} Examples include gold NP-coated Fe_3O_4 magnetic particles (GoldMag),⁴¹ MXene- Ti_3C_2 nanosheets combined with CuS NPs,⁴² and Pd-NPs stabilized with gum kondagogu biopolymer,³⁹ all demonstrating inherent peroxidase activity and combined with a dye (TMB or 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid (ABTS))), for effective sensing of cholesterol in μM ranges. Lower detection limits were achievable in a ZnO/carbon nanotube (ZnO/CNT) nanocomposite, with a linear range of 0.0005–0.5 μM and an LOD as low as 0.0002 μM .⁴⁰ Thus, nanomaterials

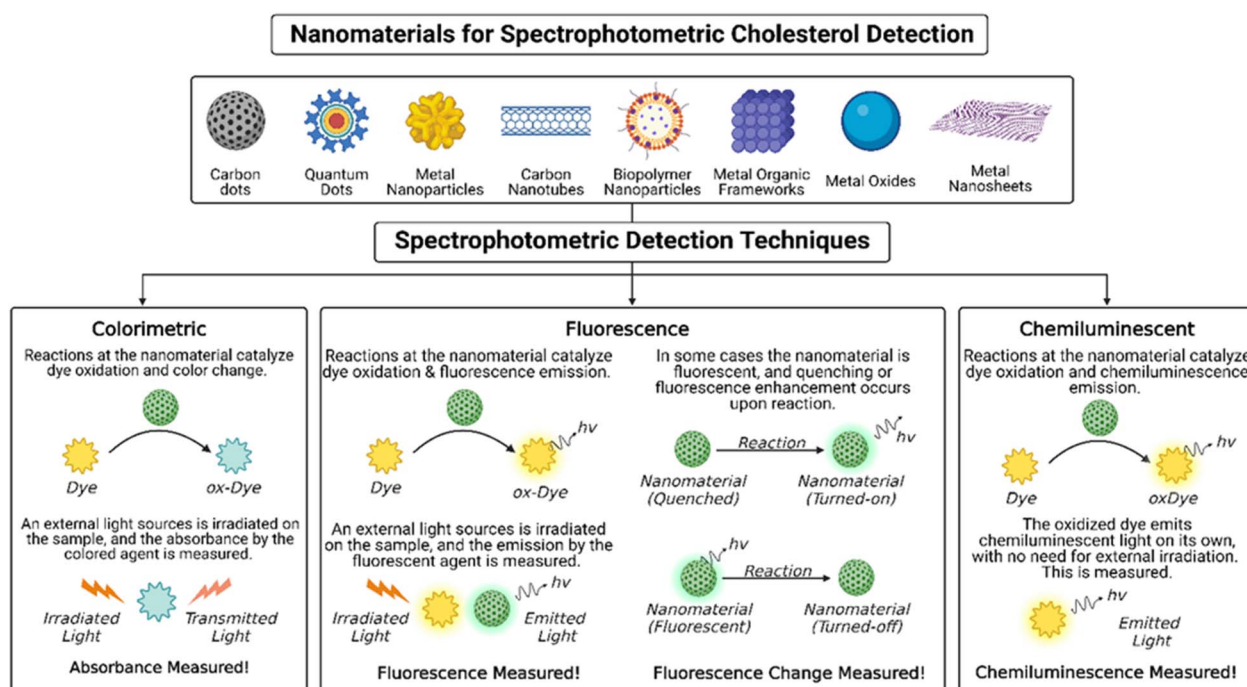


Fig. 5 Overall mechanisms of nanomaterial-based spectrophotometric cholesterol sensors.





Table 1 Spectrophotometric-based cholesterol sensors, excluding MOFs

Nanoparticles used	Detection method	Cholesterol sensing operating principle	Limit of detection (LOD) (μM)	Linear range (μM)	Ref.
Gum kondagogu (a biopolymer) – reduced/stabilized palladium NPs (GK-Pd NPs)	Colorimetric	Sample is treated with ChOx, then GK-Pd NPs has intrinsic peroxidase properties, catalyzing hydrogen peroxide (H_2O_2) reduction and TMB dye oxidation, causing a color change	3.7	5–100	39
ZnO NPs on carbon nanotube (CNT) surface	Colorimetric	After oxidation by free ChOx, peroxidase-like ZnO/CNT catalyzes H_2O_2 reduction and 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) (ABTS) dye oxidation, producing a change in color	0.0002	0.0005–0.5	40
Au@Ag core-shell nanoparticles (Au@Ag NPs)	Colorimetric	After reaction with free ChOx, produced H_2O_2 oxidizes Ag surrounding AuNPs to Ag^+ , leading to a visible color change	0.15	0.3–300	70
Fe_3O_4 particles coated with gold nanoparticles (GoldMag)	Colorimetric	After treatment with ChOx, GoldMag acts as a peroxidase mimic, catalyzing H_2O_2 reduction and ABTS oxidation, resulting in color change	7.8	259–19397	41
MXene- Ti_3C_2 nanosheets and CuS nanoparticles (MXene- Ti_3C_2 /CuS) nanocomposites	Colorimetric	After treatment with ChOx, MXene- Ti_3C_2 /CuS acts as a peroxidase mimic, catalyzing H_2O_2 reduction and TMB oxidation, resulting in color change	1.9	10–100	42
Polypyrrole nanoparticles (PPy NPs)	Colorimetric	Cholesterol is oxidized by free ChOx and PPy NPs act as peroxidase mimic, catalyzing H_2O_2 reduction and TMB oxidation, resulting in color change	3.5	10–100	43
Cholesterol oxidase immobilized gold nanoparticles (ChOx@GNP)	Dual colorimetric and fluorescence	Cholesterol is oxidized by immobilized ChOx and produced H_2O_2 quenches the GNP, leading to decreased fluorescence and a color change	490	650–7760	45
Gold-carbon dot nanocorjugates (Au@CDs)	Dual colorimetric and fluorescence	Cholesterol directly interacts with Au@CDs, destabilizing surface and leading to particle agglomeration. This causes a change in color and fluorescence quenching. No ChOx or HRP needed	2.5	1000–6250	73
Silicon NP (SiNP)	Fluorescence	Sample is pre-treated with ChOx then HRP in the presence of <i>p</i> -phenylenediamine (PPD), producing PPDOx. Produce PPDOx quenches SiNPs by the inner filter effect (IFE), and this quenching is correlated with cholesterol concentration	0.018	0.025–10	44
Poly(vinylpyrrolidone)-protected gold nanoparticles (PVP-AuNPs) (peroxidase substitute)	Fluorescence	After ChOx treatment, produced H_2O_2 enlarges PVP-AuNPs, which then quenches bovine serum albumin-protected gold nanoclusters (BSA-AuNCs). Quenching is measured	1.4	1–100	46

Table 1 (Contd.)

Nanoparticles used	Detection method	Cholesterol sensing operating principle	Limit of detection (LOD) (μM)	Linear range (μM)	Ref.
β -Cyclodextrin carbon dots (β -CDs)	Fluorescence	Cholesterol directly interacts with β -CD cavity, quenching its fluorescence. No ChOx or HRP needed	0.204	0.5–40	71
Carbon dot/hemoglobin complex (CD/Hb)	Fluorescence	Cholesterol directly binds with Hb, disturbing CD quenching and restoring fluorescence. No ChOx or HRP needed	56	0–800	72
Garlic-capped Ag NPs (G-AgNPs)	Fluorescence	Cholesterol directly interacts with G-AgNPs, <i>via</i> ionic or hydrogen bonds, causing fluorescence enhancement	186	400–5170	74
ChOx-immobilized CdSe/ZnS quantum dots	Fluorescent	Cholesterol is oxidized by the immobilized ChOx, then produced H_2O_2 directly quenches CdSe/ZnS quantum dots	10	10–9110	47
Cupric oxide NP (CuO NP)	Chemiluminescence	After ChOx treatment, cupric oxide NPs act as peroxidase mimic, catalyzing luminol conversion to excited state, which later relaxes. Chemiluminescence here is measured	0.17	0.625–12.5	69

with peroxidase-like activity worked as accurate and sensitive colorimetric cholesterol sensors when combined with a dye and ChOx.

However, the need for an external dye increases costs and materials required for the sensor design. Later studies include nanomaterials that demonstrated a color change upon exposure to hydrogen peroxide produced during cholesterol oxidation by ChOx, excluding the need for a peroxidase mimic. For instance, Au@Ag core-shell nanoparticles demonstrated a clear, color change when exposed to hydrogen peroxide after cholesterol oxidation.⁷⁰ The produced hydrogen peroxide etched away the Ag shell, producing Ag^+ and causing a gradual color change from orange to wine red.⁷⁰ A wide linear range of cholesterol detection was achieved (0.30–300 μM) and the LOD was 0.15 μM , with good selectivity in the presence of other biomolecules.⁷⁰ Hence, this unique, colorimetric sensing mechanism allowed for selective and sensitive cholesterol sensing while eliminating the need for HRP. Nonetheless, in all these nanomaterial-based colorimetric methods, ChOx enzyme is still needed as a pretreatment step for selective cholesterol sensing.

2.2.2 Fluorescent. Fluorescent analysis methods detect the emission of light by a fluorophore, or a fluorescent dye, after excitation caused by external light irradiation.⁸⁰ The wavelength of the emitted light is often longer than the absorbed light, and the mechanism of sensing often involves observing fluorescence enhancement/recovery or quenching, depending on the nanomaterial used as the fluorophore.⁸⁰ Nanomaterial-based fluorescent sensors have various advantages, including low costs, ultrasensitive detection to picomolar levels, enhanced selectivity, and allows for real-time, continuous monitoring.^{80–82} A variety of nanomaterials have been used for fluorescence-based detection of cholesterol.^{44–47,71–74} Some sensors use pure nanomaterials of one type, such as silicon NPs⁴⁴ and β -cyclodextrin carbon dots.⁷¹ Other sensors use hybrid, nanomaterial composites combining for enhanced fluorescence detection of cholesterol, such as sensors using garlic capped AgNPs,⁷⁴ CdSe/ZnS quantum dots,⁴⁷ and others.

Similar to colorimetric, nanomaterial-based sensors, fluorescent nanomaterial cholesterol sensors have been used in combination with ChOx and HRP⁴⁴ and also as peroxidase-mimics, with the need for ChOx.^{45–47} For example, fluorescent silicon NPs are used in combination with ChOx and HRP enzyme treatment in the presence of *p*-phenylenediamine, which – upon oxidation – quenches the silicon NPs; this quenching is used to determine cholesterol concentrations.⁴⁴ Meanwhile, poly(vinylpyrrolidone)-protected gold nanoparticles (PVP-AuNPs) proved to be efficient peroxidase-mimics, which replaces HRP and relies only on ChOx treatment. In the presence of oxidized cholesterol and hydrogen peroxide, PVP-AuNPs swell, causing quenching of bovine serum albumin-protected gold nanoclusters (BSA-AuNCs) and subsequent cholesterol detection.⁴⁶

However, fluorescent nanomaterial-based cholesterol detectors are unique in their ability to design enzyme-free cholesterol sensors, for direct cholesterol detection without ChOx or HRP.^{71–74} In these sensors, cholesterol directly interacts with active sites in the nanomaterials, causing a change in



fluorescence. For instance, carbon dot/hemoglobin complex demonstrated great selectivity to cholesterol, where fluorescence enhances upon cholesterol–hemoglobin binding; the sensor had a wide dynamic range (0–800 μM), with a low LOD of 56 μM .⁷² β -Cyclodextrin carbon dots⁷¹ and garlic-capped Ag NPs (G-AgNPs)⁷⁴ also exhibited selective cholesterol fluorescent detection, with β -cyclodextrin carbon dots allowing for micromolar range cholesterol detection and the G-AgNPs allowing for millimolar range cholesterol detection. These sensors are advantageous in their stability, due to the elimination of enzymes. Other nanomaterial sensors display dual colorimetric and fluorescent detection, such as ChOx immobilized gold nanoparticles⁴⁵ and gold–carbon dot nanoconjugates,⁷³ the latter being advantageous in its enzyme-free nature. All these fluorescent, nanomaterial-based sensors exhibited selective cholesterol detection and good detection limits.

2.2.3 Chemiluminescent. Chemiluminescence is the emission of light as a result of a chemical reaction; the light intensity is correlated with the reaction rate; thus, it can be used to determine the concentration of the limiting reactant. In sensing applications, luminol is often used as the light-emitting reagent,⁶⁹ *i.e.* the reagent causing chemiluminescence, among other reagents. Chemiluminescence is advantageous in its rapid nature, relatively simple method, and high sensitivity.⁶⁹ Different nanomaterials have been used to catalyze chemiluminescent reactions between produced hydrogen peroxide and a dye, for effective and rapid cholesterol sensing.^{69,83} For example, CuO NPs exhibited excellent peroxidase-like activity, catalyzing the reaction between hydrogen peroxide, produced from ChOx cholesterol oxidation, and luminol, a chemiluminescent dye. The enhancement in chemiluminescence allowed for micromolar detection of cholesterol, while also replacing HRP for a more stable, long-lasting system.⁶⁹ Hence, chemiluminescent nanomaterials for cholesterol detection have proven effective sensing materials. However, unlike fluorescent-based sensors, current chemiluminescent-based sensors still require the use of ChOx enzyme. Future studies can investigate the complete elimination of enzymes in chemiluminescent, nanomaterial-based cholesterol sensing.

3. MOFs as promising sensing materials

Compared to other nanomaterials, MOFs are gaining rapid attention and superiority in sensing applications. This is attributed to its highly tunable structure, surface functionalization, inherent catalytic and fluorescent properties, among other advantages.^{18,23–31} Understanding the basic composition, structure, and type of MOFs is crucial to better design MOF-based sensors and optimize performance. This section herein explores the basic features of MOFs, their promising advantages for sensing applications.

3.1 MOF basics

Metal organic frameworks are highly crystalline materials, with unique compositions and structures. This section herein briefly explores the basic features of MOFs, including their composition, structures, synthesis procedures, and characterization techniques.

3.1.1 Composition. MOFs are made up of two main components: metal ions or clusters and organic linkers, connected *via* coordination bonds.^{84–87} A variety of metal ions can be used, including zinc (Zn),^{88,89} zirconium (Zr),^{90–93} titanium (Ti),^{94,95} chromium (Cr),^{96,97} iron (Fe), and aluminum (Al),⁹⁸ with transition metal ions of high valence and empty d-orbitals being the majority.^{84,99} These metal ions or clusters act as nodes, forming multiple coordination bonds to join organic linkers and form the overall MOF structure. Like metal clusters, a variety of organic linkers can be used, producing a multitude of MOFs with different structures. These organic linkers, also coined ligands, often contain O- or N-donor functional groups, to donate electrons and form the coordination bond.⁸⁷ Examples of common organic linkers include carboxylates, benzenes, crown ethers, sulfonates, imidazoles, and pyridyl compounds, among others.⁸⁷ Together, the different combinations of metal clusters and organic linkers produce a multitude of MOFs with different properties and structures. Within these structures many spaces exist, which can be occupied by solvent molecules,

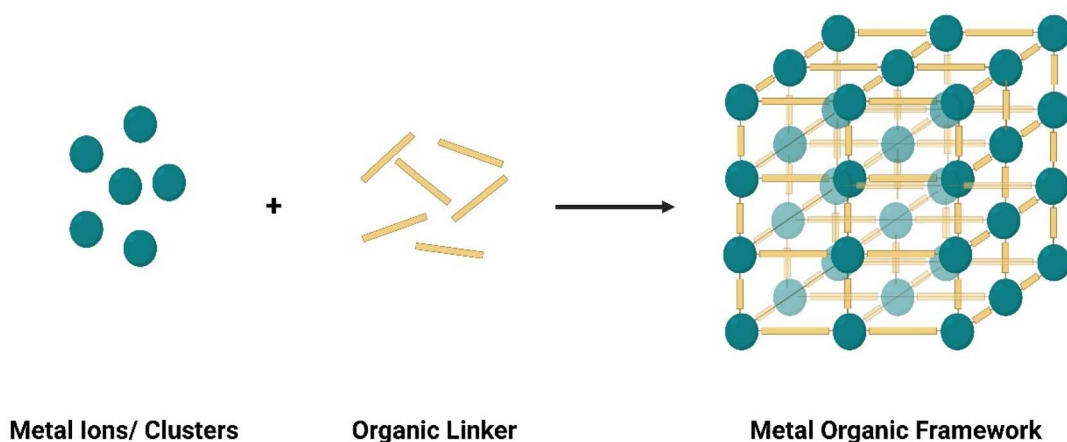


Fig. 6 Generic MOF composition from metal ions and organic linkers.

guest molecules, or others. Fig. 6 below illustrates the general composition of MOFs from its building blocks.

3.1.2 Structures. Different MOF structures or geometries have been established. This variance is dependent on the metal clusters and organic linkers used. More specifically, the coordination number (or ligancy) of the metal clusters, the geometry of the metal ion clusters, the functional group of the organic linker, and the rigidity or flexibility of the organic linker all contribute to the overall MOF structure.^{87,99} For example, metal clusters with only two vacancies often produce linear, one-dimensional (1D) structures, to form 1D MOF nanowires, nanotubes, and other shapes depending on the synthetic protocol. Examples of 1D, linear MOFs include Co-MOF-74,¹⁰⁰ ZIF-8 when synthesized on polycarbonate track-etched (PCTE) template,¹⁰¹ and Mg-MOF-74 when synthesized at a controlled pH,¹⁰² among many others reviewed in ref. 103 and 104. Meanwhile, metal clusters containing three and four vacant sites form more elaborate 3D structures, such as MIL-125(Ti), MIL-101(Cr), and MIL-53(Al), among others.^{99,105,106} Some of the geometries available at the metal cluster–ligand bonding site include linear, trigonal planar, T-shaped, tetrahedral, square planar, pentagonal bipyramidal, octahedral, trigonal prismatic, trigonal bipyramidal, and square pyramidal.⁸⁶ These geometries contribute to the overall MOF topology, which includes cubic, spherical, spindle, and rod structures.^{107–109} The diversity and tunability of MOF structures and shapes is greatly advantageous, allowing for a wide range of applications.

3.1.3 Synthesis & characterization. A multitude of synthetic techniques have been developed for the successful preparation of MOFs. Early methods were rather simple, whereby metal ions and organic linkers were mixed until porous, crystalline MOFs are developed. However, this method requires long synthesis times and can be costly.⁹⁹ With advancements in technology and research, more complex and effective methods emerged. These include microwave-assisted, electrochemical, solvothermal, mechanochemical, sonochemical, diffusion, slow evaporation, and microemulsion methods.^{87,99,107} Each method has its own advantageous and limitations. For instance, slow evaporation and diffusion methods can be performed at room temperature, unlike microwave-assisted methods requiring high temperature and energy costs.⁸⁷ Microemulsion synthesis of MOFs allows for precise control of MOF topology and surface area; however, it is associated with high costs and large production of environmental pollutants.⁸⁷ For large-scale production, continuous flow solvothermal methods are often preferred. More recently, microfluidic methods have emerged for smaller microscale, rapid production of MOFs.¹¹⁰ Hence, cost-benefit analysis is required to choose the optimal synthetic protocol for the desired MOF preparation. For further analysis of MOF preparation methods, readers can refer to ref. 84, 87, 107 and 109.

Characterization techniques are crucial to understand MOF properties and features and confirm accurate preparation. Mechanical and structural properties of MOFs, such as elasticity, microstructure, and surface topography, atomic force microscopy (AFM) can be used. Transmission electron microscope (TEM) and scanning electron microscope (SEM)

techniques can reveal further information about the crystallography properties and defects of MOFs.^{89,111} However, TEM and SEM are more expensive, have risks of beam damage, with TEM being especially sensitive to contaminants.¹¹¹ Meanwhile, Brunauer–Emmett–Teller (BET) has been especially useful for understanding surface area and porosity of MOF structures. Thermal stability can generally be characterized using thermal gravimetric analysis (TGA) while colloidal properties are better characterized using dynamic light scattering (DLS) analyses.¹¹¹ To confirm bonding and understand chemical properties, Fourier-Transform Infrared (FTIR) and Raman spectroscopy are useful.¹¹¹ Many other MOF features can be easily characterized *via* a variety of techniques, which have been extensively reviewed in ref. 111–115.

3.2 Promising properties of MOFs for spectrophotometric biosensing

MOFs demonstrate great promise for spectrophotometric biosensing applications, especially spectrophotometric-based sensing. This is because many MOFs demonstrate inherent fluorescent and/or colorimetric nature, which exhibit changes upon analyte detection.^{33,80,116} Similarly, many MOFs exhibit intrinsic catalytic activity, acting as nanozymes for sensing biological analytes requiring enzyme treatment.^{26,32,34} Moreover, MOFs can easily be hybridized with other nanomaterials for enhanced sensing performance.^{117–119} It may be argued that other nanomaterials, such as metal oxides, quantum dots, and metal NPs, all also demonstrate excellent fluorescence, intrinsic catalytic activity, and potential for hybridization – as shown in Table 1. While this is true, MOFs have additional unique properties that make it stand out as promising nanomaterials for biosensing applications. These unique advantages include high surface area and porosity, tunable structures, and functionalizable surfaces. These unique properties are discussed herein.

3.2.1 High surface area & porosity. MOFs are unique in their high surface area and porosity. MOF surface areas can reach up to 10 400 m² g^{−1},¹²⁰ which is significantly greater than that of graphene (~2000 m² g^{−1}),¹²¹ zeolites (<1100 m² g^{−1}),¹²² and other nanomaterials.¹¹¹ Pore volumes can also extend up to ~3.60 cm³ g^{−1}, which is superior to other nanomaterials.^{111,120} This massive internal surface area and pore volumes facilitate entrapment of more analyte molecules within the MOF framework, thus increasing analyte-MOF collisions and facilitating for more sensitive detection and quantification.^{123,124} Moreover, the large pore volume can facilitate detection of larger biomolecules and analytes, since they can fit into the structure.¹²⁵ Immobilization of larger molecules can also be facilitated in MOF pores, such as enzymes for cholesterol and glucose detection.^{126,127} Table 2 below presents some common MOFs used for developing spectrophotometric based sensing of cholesterol, reporting their composition, surface areas, volumes, and its luminescence properties. While some MOFs do not exhibit inherent fluorescent properties, they can be modified and functionalized to be fluorescent, and can be used as nanozymes to catalyze sensing mechanisms.^{28,126} These



Table 2 Key MOFs used in spectrophotometric cholesterol sensors and their structural properties

MOF	Metal cluster	Organic linker	Morphology	Pore volume (BET, cm ³ g ⁻¹)	BET surface area (m ² g ⁻¹)	Luminescence property	Ref.
MIL-100 (Fe)	Fe-O	Terephthalic acid	Microsphere shell	—	—	Not fluorescent	18
MIL-101 (Cr)	Cr-O	Terephthalic acid	Octahedral crystal	—	—	Not fluorescent	28
UiO-66	Zr-O	Terephthalic acid	Octahedral	0.5453	1094	Weakly fluorescent	27
ZIF-8	Zn-O	2-Methylimidazole	Dodecahedral with rhombus faces	0.617	1275–1664	Not fluorescent	25, 126, 128 and 129
PCN-333 (Al)	Al-O	4'-s-Triazine-2,4,6-triyl-tribenzoic acid (H ₃ TATB)	Octahedral	2.48	2550	Not fluorescent	127
MOF-5	Zn-O	Terephthalic acid	Cubic	—	18.52	Not fluorescent	30
NU-1000	Zr-O	1,3,6,8-Tetra(4-carboxylphenyl)pyrene (H4TBAPy)	Rod	1.31	1872	Fluorescent at 470 nm	31
Co/2Fe bimetallic MOF	Co-O & Fe-O	Trimesic acid	Irregular particles	—	1181	Not fluorescent	24
MIL-101 (Fe)	Fe-O	1,4-Dicarboxybenzene	Octahedral	—	—	Not fluorescent	117
UiO-66-NH ₂	Zr-O	2-Aminoterephthalic acid	Octahedral	—	—	Not fluorescent	130
MoCu-MI	Cu-O & Mo-O	2-Methylimidazole	Nanosheet	—	—	Not fluorescent	29
Cu (PABA)	Cu-O	<i>p</i> -Aminobenzoic acid (PABA)	Ellipsoid	—	—	Not fluorescent	131
CuMOF	Cu-O	Terephthalic acid	Flake-like	—	293.7	Not fluorescent	83
V-MOF	V-O	2-Aminoterephthalic acid	Irregular particles	—	74.74	Not fluorescent	132

MOFs may also be used for enzyme immobilization and stabilization, as discussed later in Section 4.¹²⁷

3.2.2 Tunable structures. MOFs demonstrate great tunability, due to the variety of metal ions and organic linkers that can be used. Conversely, other nanomaterials demonstrate poor tunability due to limited building blocks. For instance, metal NPs are limited in tuning only the metal centers. Similarly, carbon dots and graphene have poor tunability due to limited building blocks (carbon).⁸² Meanwhile, the wide range of organic linkers available for MOF preparation offer unlimited MOF structures and configurations, as different organic linkers can be luminescent at different wavelengths and can interact differently with varying metal nodes. Thus, for spectrophotometric sensing applications, organic linkers and metal clusters can carefully be chosen to target specific analytes, maximizing sensor selectivity.¹²³ This flexibility in tuning nanomaterial luminescent structures to target analytes is unique to MOFs, making them promising candidates for applications in spectrophotometric sensing.

3.2.3 Functionalizable surfaces. Since MOF surfaces majorly consist of organic linkers, surface functionalization is easily facilitated. This subsequently allows for further tuning of MOF sensing properties for more specific and selective analyte detection. The presence of different chemical groups on MOF surfaces act as potential sites for functionalization of specific molecules, which can play a role in attracting analytes (through affinity or guest–host interactions), turning MOF luminescence on or off, altering MOF dispersibility, among others.^{31,116,123,133} Surface functionalization can be through adsorption, coordination bonding, or covalent linkages, depending on the organic linker and molecule being attached.^{31,133} MOFs can also be functionalized with other nanomaterials, to form hybrid nanocomposites with enhanced properties.¹³⁴ Other

nanomaterials are more rigid, with less chances of successful functionalization. Hence, the easily functionalizable surfaces of MOFs can be tuned for specific analyte detection. Combined with MOF's characteristic high surface areas, surface functionalization adds an added advantage for MOFs as promising materials for spectrophotometric sensing applications.

Fig. 7 summarizes the main advantageous properties of MOF that make them promising candidates for spectrophotometric cholesterol detection. These include high surface area and porosity, tunability, easy surface functionalization, and intrinsic luminescence and catalytic activity.

4. MOFs for spectrophotometric cholesterol detection

Recently, MOFs have emerged as promising nanomaterials for spectrophotometric-based cholesterol detection. MOFs can be used alone^{24,28,30,31,127,132} or in hybrid composites with other nanomaterial,^{18,25,27,29,83,117,118,126,128–131} for more effective cholesterol quantification. Generally, MOF-based cholesterol sensors can be categorized into five types, according to the mechanism of sensing adopted, which in turn impacts detector performance. The mechanisms and the performance of these sensors will be discussed below.

4.1 Roles of MOFs in sensing mechanisms

Spectrophotometric sensing of cholesterol through MOFs has been reported widely in the literature. The role of MOFs in cholesterol detection can be categorized into five main types. In some cases, the MOF is used for immobilization of sensing factors,^{16,17,127–130} while other studies primarily adopt mechanisms where the MOF plays an active role in the sensing



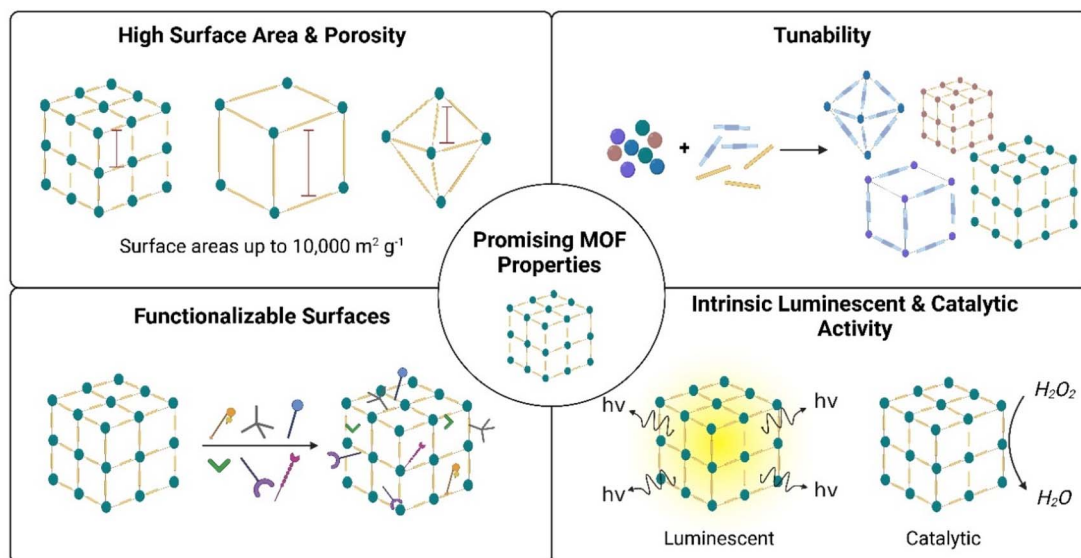


Fig. 7 Summary of promising MOF properties.

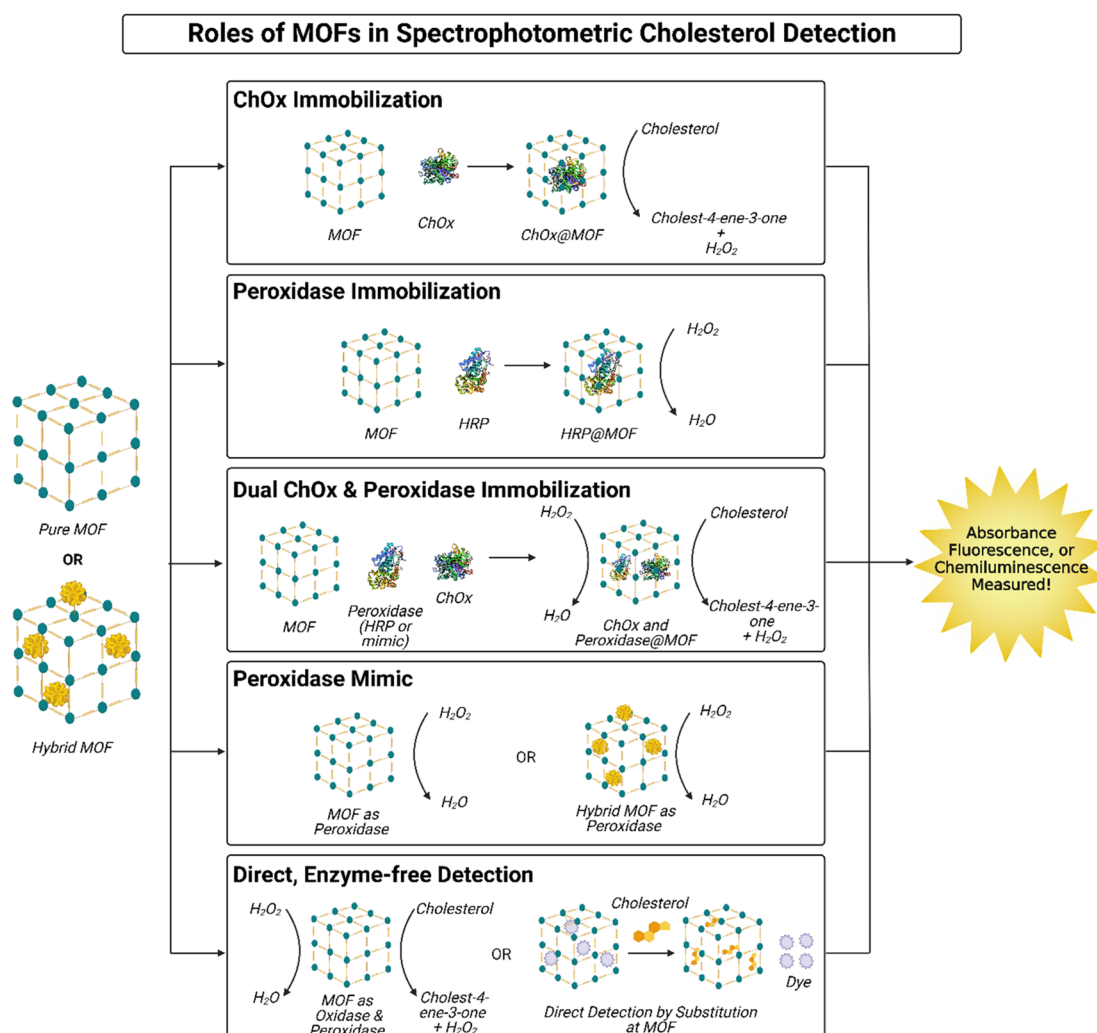


Fig. 8 Summary of MOF roles in spectrophotometric cholesterol detection.



process.^{18,24,27–29,31,83,117,118,126,131,132} Furthermore, these mechanisms implement a variety of spectrophotometric methods, including colorimetry,^{18,27–30,83,127,128,130,131} fluorescence,^{24,25,27,31,117,118,126,131,132} and chemiluminescence.^{83,129} These mechanisms are evaluated and compared in the next few sections. A summary of these mechanisms is illustrated in Fig. 8.

4.1.1 MOF for ChOx immobilization. Enzymes are highly pH- and temperature-dependent, thus requiring great care in use and storage to avoid denaturation.^{16,17} ChOx is a crucial enzyme in cholesterol sensing, as it is needed to catalyze the oxidation of cholesterol to produce hydrogen peroxide, which is subsequently used for cholesterol detection.^{16,25,30} ChOx immobilization and encapsulation in MOFs is an effective method for improving enzyme stability and enhancing its efficiency.^{16,17} However, for effective encapsulation, MOF pore size must be optimal to host the enzyme without extensively impacting its conformation;¹³⁵ small pore dimensions will block out ChOx, preventing immobilization.³²

MOFs have proven to be effective hosts for ChOx immobilization, in turn facilitating cholesterol detection by prolonging ChOx stability.^{25,30} ChOx immobilization in MOFs can also facilitate faster oxidation of cholesterol by the enzyme, through maximizing the reaction surface area between cholesterol and ChOx and concentrating ChOx in a smaller area.^{25,30} For instance, Hassanzadeh *et al.* utilized MOF-5 for ChOx immobilization *via* a post-synthetic method,³⁰ while Ding *et al.* used ZIF-8 (ref. 25) and adopted an *in situ* method, both demonstrating good performance and minimal ChOx leakage. However, it is important to highlight that the MOFs here do not actively interact with cholesterol chemically; they only act as hosts that enable the first step of the cholesterol detection process. Other agents are needed to continue the reaction. Hassanzadeh *et al.* couple ChOx@MOF-5 with an Ag nanocluster decorated MoS₂ nanosheets, the latter acting as a peroxidase mimic to catalyze hydrogen peroxide reduction and in the presence of terephthalic acid, which is oxidized for colorimetric detection of cholesterol (illustrated in Fig. 9A).³⁰ Meanwhile, Ding *et al.* opted for a fluorescent method by encapsulating Au nanoclusters (AuNCs) within the ChOx@ZIF-8 complex and decorating it with polyethyleneimine (PEI) to form a hybrid, MOF nanocomplex. Iron(II) ions (Fe²⁺) are introduced to the system with cholesterol, which react with hydrogen peroxide produced after cholesterol oxidation to produce hydroxyl radicals; this reaction is coined a Fenton reaction. The hydroxyl radicals subsequently oxidize the AuNCs from Au(0) to Au(I), quenching the systems fluorescence. The fluorescence quenching is correlated with cholesterol concentration; thus, cholesterol can easily be quantified.²⁵

The above studies mostly encapsulate ChOx within the MOF pores. Recently, ChOx immobilization on MOF surfaces was explored, rather than inside its pores, *via* a hybrid structure.¹¹⁸ Xu *et al.* designed a nanocomposite of Tb-MOF and MnO₂ nanosheets – which quench Tb-MOF fluorescence; ChOx is combined with copper phosphate for self-assembly of ChOx-copper phosphate nanoflowers on the MOF surface.¹¹⁸ Through this unique design, cholesterol undergoes rapid

cascade reactions, where it is first oxidized by ChOx nanoflowers, then produced hydrogen peroxide reduces MnO₂ sheets, thus allowing for Tb-MOF fluorescence recovery. This fluorescence is measured to determine cholesterol concentration.¹¹⁸ Hence, MOFs have proven useful for ChOx immobilization, in pores or on the surface, and subsequent cholesterol detection *via* fluorescent and colorimetric methods.

4.1.2 MOF for peroxidase immobilization. Like ChOx immobilization, MOFs have also been useful for encapsulating and immobilizing peroxidases, specifically horseradish peroxidase (HRP) – which is traditionally used for in cholesterol detection assays.¹²⁹ HRP is well-known for its low stability and quick degradation.^{17,18} Therefore, HRP immobilization in MOFs can significantly prolong its lifetime and improve cholesterol sensing. ZIF-8 demonstrated excellent encapsulation of HRP, through a one-pot synthesis method combining HRP and ZIF-8 precursors.¹²⁹ However, for a peroxidase@MOF system to effectively detect cholesterol, initial pretreatment with free ChOx is needed to produce hydrogen peroxide, which is later reduced by encapsulated HRP in the presence of a dye for quantification. Xu *et al.* achieved this by using luminol, a chemiluminescent dye when reacted with hydrogen peroxide, in the presence of HRP@ZIF-8, which initiates a chemiluminescence reaction triggering light emission.¹²⁹ The intensity of light emitted can be used for cholesterol quantification. The entire synthetic scheme and detection mechanism is illustrated in Fig. 9B. As such, MOFs also prove effective for encapsulating peroxidases (ex: HRP) and stabilizing them for cholesterol detection.

4.1.3 MOF for co-immobilization of ChOx & peroxidase. While MOFs have proven successful for encapsulating HRP and ChOx separately,^{25,30,129} co-immobilization of these enzymes in one MOF offers greater potential of faster detection using less materials. Through co-immobilization of peroxidases and ChOx in one MOF, cholesterol is oxidized and the product hydrogen peroxide can immediately be oxidized by peroxidase, in the presence of a dye or fluorescent agent, for cholesterol quantification. Zhao *et al.* achieved this by co-encapsulating HRP and ChOx in PCN-333(Al) MOF, with a high loading capacity and good stability.¹²⁷ First ChOx was encapsulated, followed by HRP, as illustrated in Fig. 9C. Peroxidase mimics may also be co-immobilized with ChOx, such as hemin¹²⁸ and AuNPs.¹³⁰ To effectively secure hemin molecules in ZIF-8, hemin was conjugated to poly(1-vinylimidazole) (PVI) then co-encapsulated with ChOx in ZIF- through a one-pot synthesis.¹²⁸ Meanwhile, Cao *et al.* effectively immobilized AuNPs on UiO-66-NH₂ MOF surface and then used molecular imprinting to secure ChOx.¹³⁰ Combined with ABTS^{127,128} or TMB¹³⁰ dye, colorimetric detection of cholesterol was successfully achieved by the MOF systems. For effective cascade detection of cholesterol *via* enzyme co-immobilization, the MOF used should have an optimal pore size and surface area to fit both enzymes while also allowing space for cholesterol to enter and interact with these enzymes. Interfering interactions may also be a concern, especially due to the close proximity of the two enzymes. Nonetheless, this approach has been proven an effective mechanism for cholesterol detection.



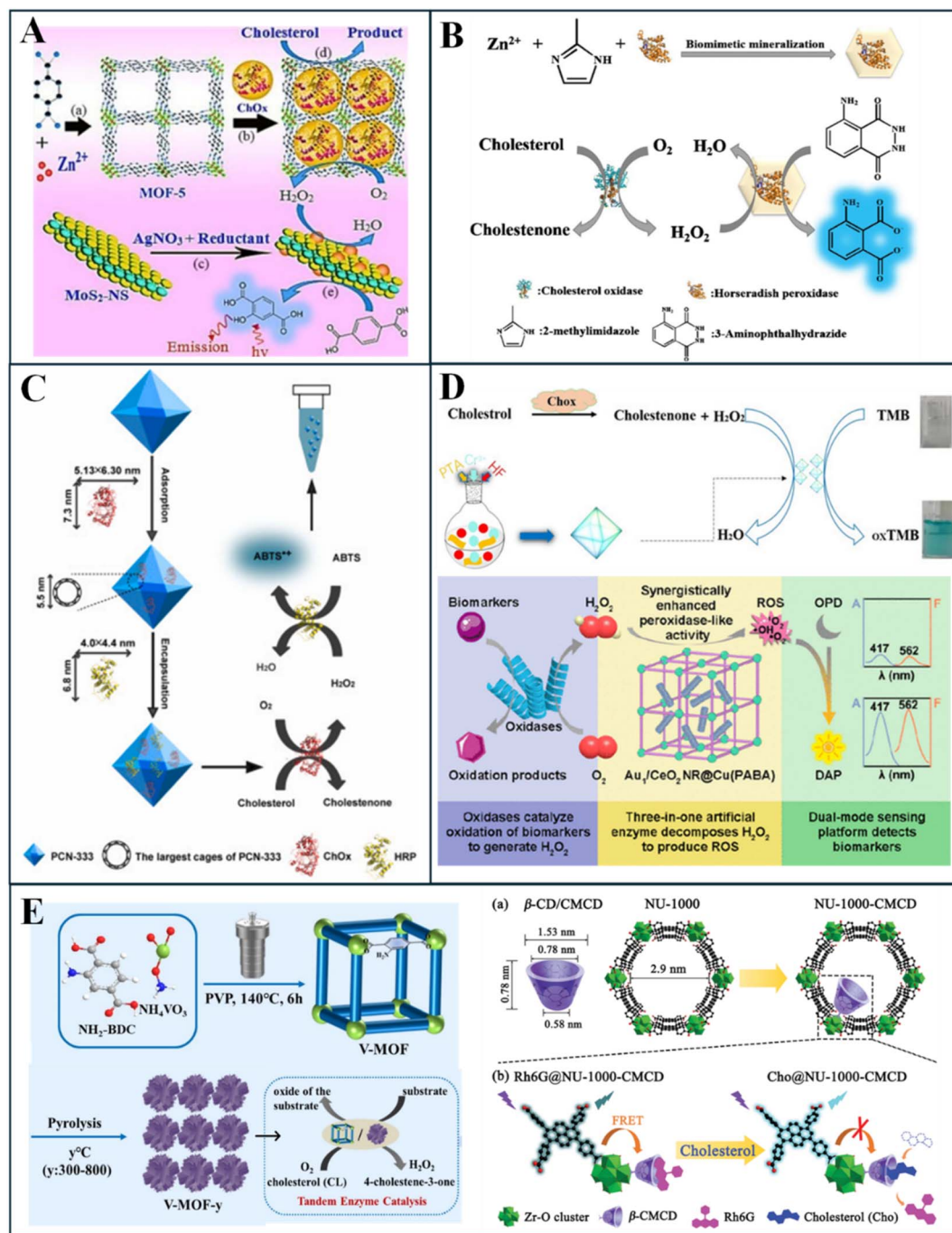


Fig. 9 Roles of MOFs in spectrophotometric cholesterol sensing, including (A) ChOx immobilization, (B) peroxidase immobilization, (C) co-immobilization of ChOx and peroxidase, (D) pure (Top) and hybrid (Bottom) MOFs as peroxidase mimics, and (E) MOFs for direct, enzyme free detection of cholesterol, by dual peroxidase and oxidase activity (left) or direct substitution (right). Reprinted with permission from ref. 28, 30, 31, 67, 129, 131 and 132.

4.1.4 MOF as peroxidase mimic. MOFs have also demonstrated active roles in cholesterol detection, unlike the passive role as an enzyme host discussed earlier. MOFs as peroxidase mimics is the most widely studied mechanism for cholesterol detection in the literature.^{18,24,27–29,83,117,118,126,131} In this approach, the MOF has intrinsic peroxidase activity that catalyzes the reduction of hydrogen peroxide; often time, this is done in the presence of TMB dye, which is oxidized for colorimetric

quantification of cholesterol.^{18,27–29} The absorbance of these colors is correlated with the cholesterol concentration. Pure MOFs can be used for this approach, such as MIL-101(Cr).²⁸ As shown in Fig. 9D (top), MIL-101(Cr) acts as a peroxidase by reducing hydrogen peroxide to H₂O, and subsequently oxidizing TMB to produce a color change for colorimetric detection.²⁸ However, most studies adopt hybridized MOF systems as peroxidase mimics. These include Mo-doped Cu(2-



methylimidazole) (Cu-2MI) MOF,²⁹ MIL-100(Fe) coated Fe₃O₄ magnetic NPs,¹⁸ and graphene quantum dots/zirconium-based MOF (GQDs/UiO-66) nanocomposite.²⁷

Meanwhile, other studies considered fluorescent dyes at oxidation, such as terephthalic acid,²⁷ *o*-phenylenediamine (OPD),¹³¹ and thiamine,¹²⁶ to detect emitted light rather than absorbance. These MOF systems are also often hybridized, with Pd nanoclusters,¹²⁶ Au-loaded CeO₂ nanorods,¹³¹ or graphene quantum dots.²⁷ For example, Cu(PABA) MOF hybridized with Au-loaded nanorods demonstrated more effective peroxidase activity; combined with OPD fluorescent dye, Au-loaded CeO₂ nanorods@Cu(PABA) MOF composite effectively oxidized the dye in the presence of cholesterol, producing a fluorescent change. This mechanism is illustrated further in Fig. 9D (bottom). Peroxidase-like MOFs have also facilitated cholesterol detection by ratiometric fluorescent measurements,^{24,117} whereby one area of the MOF-system is quenched while the other emits fluorescence; the ratio of these intensities can be used to determine cholesterol concentration. Yang *et al.* adapted this through a Eu³⁺-doped MIL-53(Fe) MOF, where hydrogen peroxide exposure enhances MIL-53(Fe) fluorescence and quenches Eu centers.²⁴ MIL-53(Fe) acts as a peroxidase here, catalyzing internal linkages in the MOF thus amplifying its intensity.²⁴ Ye *et al.* followed a similar approach but through a hybrid MOF-QD system, using bifunctional carbon quantum dots (CQDs)@ MIL-101(Fe). Here, MIL-101(Fe) exhibited peroxidase ability and the CQDs are fluorescent; in the presence of OPD dye, hydrogen peroxide catalyzes the oxidation of OPD into highly fluorescent 2,3-diaminophenazine (DAP), which subsequently quenches CQD fluorescence; hence, the ratio between quenching and fluorescence intensity can be used for cholesterol quantification.¹¹⁷ Chemiluminescence detection using peroxidase-like MOFs has also been demonstrated using Co₃O₄@CuMOF hybrid system in combination with the chemiluminescent rhodamine B dye.⁸³ The MOF here catalyzes the chemiluminescent, redox reaction between hydrogen peroxide and rhodamine B dye, in turn emitting light and causing a color change.⁸³ In all cases, ChOx pretreatment of cholesterol is required to produce hydrogen peroxide. This is often done using free ChOx, which degrades rapidly.^{18,24,27–29,83,117,118,126,131} Nonetheless, cholesterol sensors adopting MOFs as peroxidase mimics for the main sensing mechanism have demonstrated excellent promise.

4.1.5 MOF for direct, enzyme-free detection. The final mechanism of cholesterol detection is based on deploying MOFs for direct cholesterol sensing, eliminating the need for any enzymes.^{31,132} In this case, cholesterol directly interacts with active sites on the MOF, leading to a quantifiable spectrophotometric change. Enzyme-free detection of cholesterol is highly advantageous, since it eliminates the risk of enzyme degradation and related uncertainties. Direct, enzyme-free detection of cholesterol has been done through two main mechanisms: using a MOF as a dual oxidase and peroxidase agent¹³² and using a MOF for substitution-based detection.³¹ For example, Wu *et al.* designed a pyrolyzed V-MOF which oxidized cholesterol, mimicking ChOx, and catalyzed the oxidation of OPD in the presence of hydrogen peroxide, mimicking HRP. The

tandem enzyme activity in this MOF facilitated rapid and accurate cholesterol detection, while also possibly saving costs.¹³² The synthetic protocol and subsequent tandem enzyme activity (oxidase and peroxidase) is illustrated in Fig. 9E (left). Meanwhile, the second enzyme-free cholesterol detection approach involves functionalizing a MOF using a ligand with high affinity to cholesterol, such as β -cyclodextrin.³¹ Gong *et al.* functionalized NU-1000 with carboxymethyl- β -cyclodextrin, then loaded the β -cyclodextrin ligands with rhodamine 6G dye, which quenches the MOF's initial fluorescence. Introducing cholesterol substitutes rhodamine 6G in the MOF, subsequently recovering MOF fluorescence and allowing for cholesterol quantitation, as depicted in Fig. 9E (right).³¹ Through this mechanism, cholesterol detection is done directly in one-step, without the need for sequential oxidation and peroxide reactions, minimizing possible interferences.³¹ Hence, enzyme-free cholesterol detection mechanisms are preferred due to their lower costs and enzyme elimination.

Fig. 9 illustrates the roles of MOFs in spectrophotometric-based cholesterol sensors, as reported in the literature and discussed above. These include MOFs for ChOx immobilization (Fig. 9A), peroxidase immobilization (Fig. 9B), co-immobilization of ChOx and peroxidase (Fig. 9C), peroxidase mimics (Fig. 9D), and direct, enzyme-free detection of cholesterol (Fig. 9E).

4.2 MOF sensor performance

Different cholesterol sensing mechanisms impact sensor accuracy, sensitivity, selectivity, LOD, and overall performance. It is crucial to understand the relationship between the sensor performance and the associated detection mechanism to achieve superior performance in terms of response time, selectivity, and accuracy. Herein, the performance of a variety of MOF-based spectrophotometric sensors is explored, based on their mechanism of action. Table 3 summarizes all spectrophotometric MOF-based sensors, reporting the composition, mechanism of sensing, detection limits, and linear ranges.

4.2.1 Sensors using MOFs for ChOx immobilization. Immobilization of ChOx in MOF structures and nanocomposites can significantly improve catalytic activity and stability, thus facilitating improved cholesterol sensing performance.^{25,30,118} For instance, Ding *et al.*²⁵ adopted an *in situ* technique to synthesize a AuNCs/ChOx@ZIF-8/PEI nanocomposite, which facilitated tandem reactions for cholesterol detection with an LOD of 0.073 μ M. Optimal sensor performance was achieved at neutral pH and 37 °C, with a detection time of 15 min. While the results are obtained relatively quickly when compared to traditional cholesterol lab tests, the detection time is quite long for use in point-of-care devices, where cholesterol detection can be achieved in less than five minutes.¹³⁶ Furthermore, the narrow linear dynamic range (0.1–2.4 μ M) limits applications for real life samples, since it poses the need for 1000 time dilutions, which can be challenging and material-consuming.²⁵ An earlier nanocomposite MOF system of ChOx/MOF-5 and AgNC@MoS₂ NSs, in combination with TMB dye, achieved a lower LOD of 0.03 μ M and a wider linear



Table 3 Summary of spectrophotometric MOFs used for cholesterol detection

MOF used	Hybrid or pure MOF	Sensing (light) mechanism used	MOF mechanism	LOD (μM)	Linear range (μM)	Ref.
MOF-5	Pure	Colorimetric	MOF for ChOx immobilization	0.03	0.06–15	30
AuNCs/ChOx@ZIF-8/PEI nanocomplexes	Hybrid	Fluorescence	MOF for ChOx immobilization	0.073	0.1–2.4	25
MnO ₂ nanosheets on the surface Tb-MOFs. Coupled with hybrid nanoflowers are self-assembled by ChOx and Cu ₃ (PO ₄) ₂ ·3H ₂ O	Hybrid	Fluorescence	MOF for ChOx immobilization	1.57	10–360	118
HRP@ZIF-8 nanocomposite	Hybrid	Chemiluminescence	MOF for peroxidase immobilization	0.04	0.1–100.0	129
ChOx & PVI-hemin@ZIF-8	Hybrid	Colorimetric	MOF for co-immobilization of ChOx & peroxidase	NR	5–50	128
PCN-333(Al)/ChOx & HRP	Pure	Colorimetric	MOF for co-immobilization of ChOx & peroxidase	0.6	0.0–40.0	127
UiO-66-NH ₂ @AuNPs-ChOx@MIPs	Hybrid	Colorimetric	MOF for co-immobilization of ChOx & peroxidase	2400	2900–6700	130
Fe ₃ O ₄ magnetic NPs that are coated by MIL-100(Fe)	Hybrid	Colorimetric	MOF as peroxidase mimic	0.8	2–50	18
Mo-doped Cu-2MI (2-methylimidazole) [MoCu-2MI]	Hybrid	Colorimetric	MOF as peroxidase mimic	1.2	2–140	29
MIL-101(Cr)	Pure	Colorimetric	MOF as peroxidase mimic	0.86	1.1–22.9	28
Graphene quantum dots/zirconium-based metal-organic framework (GQDs/UiO-66)	Hybrid	Dual colorimetric & fluorescence	MOF as peroxidase mimic	0.01	0.04–1.60	27
Bimetallic Eu/Fe-MOFs [Eu ³⁺ -doped MIL-53(Fe)]	Pure	Ratiometric fluorescence	MOF as peroxidase mimic	10.5	2000–8000	24
Au-loaded CeO ₂ nanorods (NR) in Cu-based metal-organic frameworks (Cu(PABA))	Hybrid	Dual colorimetric & Fluorometric	MOF as peroxidase mimic	Colori: 0.162 Fluor: 0.178	0.05–40	131
Bifunctional carbon quantum dots (CQDs)@ MIL-101(Fe)	Hybrid	Ratiometric fluorescence	MOF as peroxidase mimic	4.55	5–100 & 100–1000	117
Co ₃ O ₄ @CuMOF NPs	Hybrid	Dual chemiluminescence & colorimetric	MOF as peroxidase mimic	0.026	0.1–300	83
Pd nanoclusters in ZIF-8 (Pd@ZIF-8)	Hybrid	Fluorescence	MOF as peroxidase mimic	0.92	5–1000	126
V-MOF-700 (pyrolyzed at 700 °C in N ₂ atm)	Pure	Fluorescence	MOF for direct, enzyme-free detection	0.38	2–160	132
NU-1000-CMCD	Pure	Fluorescent	MOF for direct, enzyme-free detection	0.40	0–180	31

range of 0.06–15 μM .³⁰ This can be attributed to the high ChOx encapsulation efficiency (25.3%), and greater, preserved catalytic activity due to post-synthetic loading.³⁰ Fig. 10(A1) demonstrates the increase in fluorescence with cholesterol addition, while the linear cholesterol calibration curves and hydrogen peroxide production rates with time of (a) MOF-5 alone, (b) ChOx alone, (c) ChOx@MOF grafted post-synthesis, and (d) ChOx@MOF synthesized *in situ* are all illustrated in Fig. 10(A2 and A3), respectively. As shown, MOF-5 alone does not have any peroxidase activity, while the best sensitivity to cholesterol was achieved by post-synthetically grafted ChOx@MOF, as depicted by higher hydrogen peroxide production (Fig. 10(A3)) and greater slope of calibration curve (Fig. 10(A2)).

Similar to the MOF earlier,²⁵ optimal detection was achieved at neutral pH and physiological temperature (37 °C), but a lower detection time of 10 min. An additional advantage is the recyclability of the sensing material, where 8 cycles of detection

maintained sensing efficiency of 90%, indicating great potential for use in reusable PoC devices.³⁰ Both MOF nanocomposites proved effective for free cholesterol and total cholesterol detection applications, the latter requiring an additional pretreatment step with cholesteryl esterase.^{25,30} Meanwhile, surface functionalization nanoflower-ChOx structures on Tb-MOFs demonstrated a higher LOD of 1.57 μM , but with a wider dynamic range of 10–360 μM , thus showing potential for blood-cholesterol detection applications.¹¹⁸ As such, ChOx immobilization in hybrid MOF sensors have demonstrated great cholesterol sensing performance and potential for real-life applications.

4.2.2 Sensors using MOFs for peroxidase immobilization. Immobilization of HRP, the peroxidase used for traditional cholesterol detection, can improve cholesterol detection through HRP stabilization.¹²⁹ Through a one-pot synthetic protocol, HRP was immobilized in ZIF-8 pores, which



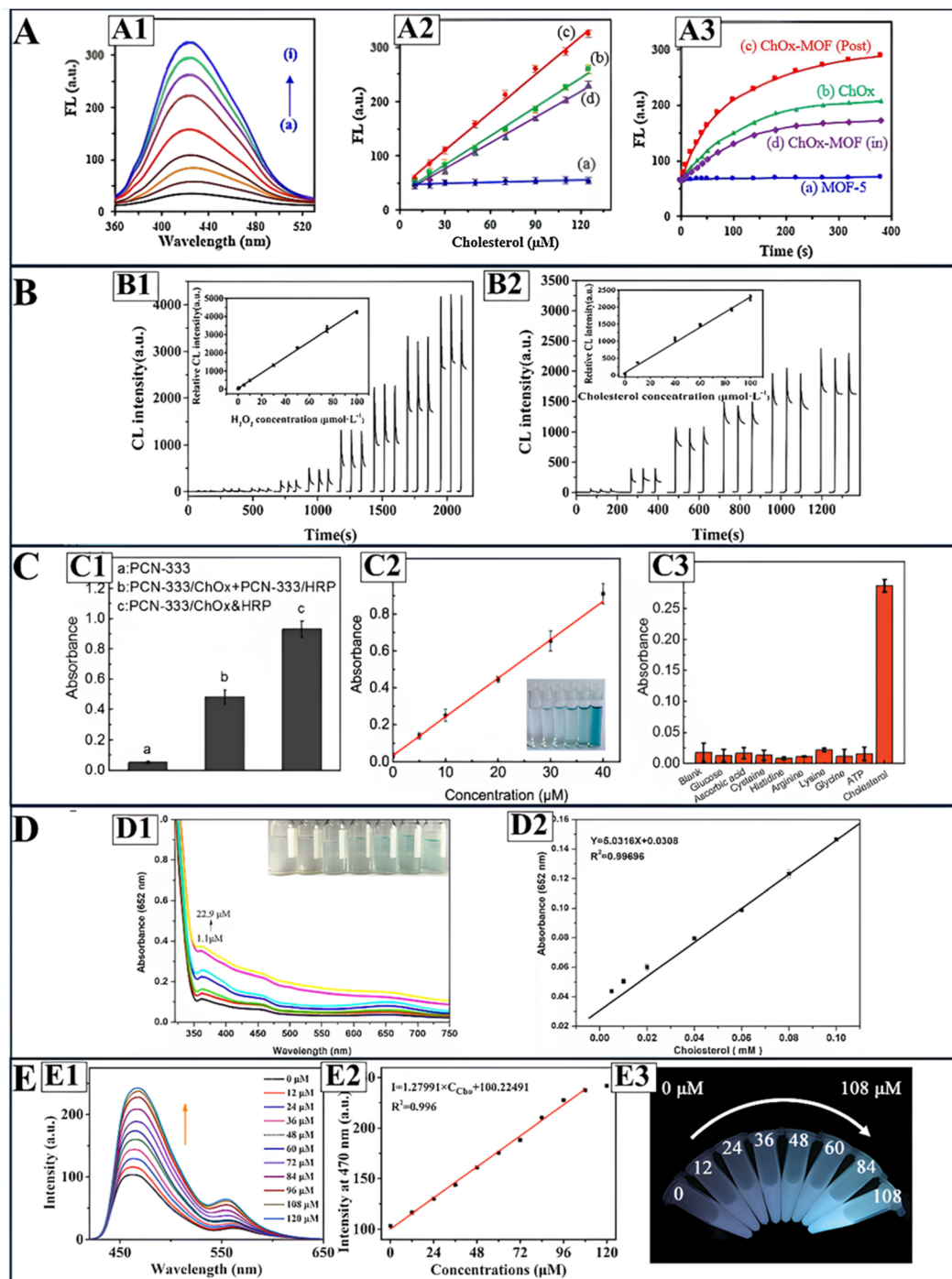


Fig. 10 Sensing performance of MOF-based, spectrophotometric cholesterol sensors. (A) Depicts the performance of ChOx@MOF sensors, where (A1) illustrates turn on fluorescence in presence of cholesterol, (A2) depicts cholesterol calibration curves of (a) post synthetically immobilized ChOx@MOF, (b) free ChOx, (c) *in situ* ChOx@MOF immobilization, and (d) MOF-5, and (A3) illustrates H_2O_2 production with ChOx activity over time. (B) Illustrates the chemiluminescence signals attained using peroxidase@MOF sensors when detecting (B1) H_2O_2 and (B2) cholesterol; inserts represent the calibration curves. (C) Illustrates the (C1) catalytic activity, (C2) cholesterol calibration curve, and (C3) selectivity of co-immobilized ChOx and peroxidase MOF sensors. (D) shows the (D1) turn on fluorescence response to cholesterol and (D2) calibration curve of peroxidase-like MOFs. Finally, (E) depicts the (E1) fluorescence recovery and (E2) calibration curve for direct cholesterol by enzyme-free MOF, with (E3) illustrating the real samples fluorescence at different cholesterol concentrations. Reprinted (adapted) with permission from ref. 28, 30, 67 and 129.

significantly enhanced HRP's temperature tolerance and activity at neutral pH.¹²⁹ This enhancement in HRP activity allowed for cholesterol detection, by chemiluminescence

reaction with luminol, at a low LOD of 0.04 μM and a wide linear dynamic range of 0.1–100.0 μM .¹²⁹ The chemiluminescence response is demonstrated in the hydrogen peroxide and



cholesterol chemiluminescence signals (Fig. 10(B1 and B2), respectively). Furthermore, the LOD and wide dynamic range to hydrogen peroxide and cholesterol is shown by the calibration curves (inserts of Fig. 10(B1 and B2), respectively). This LOD is comparable with results of MOFs with immobilized ChOx,^{25,30} as well as other nanomaterial detection systems utilizing chemiluminescence mechanisms.⁶⁹ The HRP@ZIF-8 – luminol sensor performance was validated using real blood samples; however, the main drawback is the high incubation times required for ChOx pretreatment (30 min at 37 °C). Nonetheless, the system proved very precise (relative standard deviations (RSD) below 2.5%), showing promise for future real-life applications.¹²⁹

4.2.3 Sensors using MOFs for co-immobilization of ChOx & peroxidase. Co-immobilization of ChOx and peroxidases in one MOF system is expected to improve sensor performance due to more rapid, cascade reactions of cholesterol. Many studies tested this through developing a variety of dual ChOx & peroxidase MOF nanocomposites.^{127,128,130} For instance, Zhao *et al.*¹²⁷ co-encapsulated HRP and ChOx in a water-stable PCN-333(Al) MOF while Zhang *et al.*¹²⁸ adopted ChOx and hemin, a peroxidase mimic, in a ZIF-8 MOF. Combined with ABTS dye, colorimetric measurements were used for cholesterol quantitation.^{127,128} Contrary to expectations, both these sensors demonstrated LODs (0.40 & 0.60 μM) and linear ranges (<50 μM) similar to MOF-nanocomposites encapsulating one enzyme type.^{30,129} For PCN-333(Al) in particular, the greatest catalytic activity was achieved when co-encapsulating ChOx and HRP in PCN-333, as compared to mixing single encapsulated ChOx@PCN-333 and HRP@PCN-33 or PCN-33 alone (Fig. 10(C1)). In contrast, the stability of co-immobilized ChOx and peroxidase MOF sensors is greatly improved, exhibiting longer shelf-life and high tolerances to pH and temperature changes.^{127,128} Moreover, the dual-encapsulated ChOx&HRP@PCN-333 MOF demonstrated excellent selectivity in the presence of interfering analytes (Fig. 10(C3)), and the linear range calibration curve and color change with cholesterol addition is shown in Fig. 10(C2). Meanwhile, a more recent study adapted a novel technique for grafting both ChOx and AuNPs (peroxidase mimics) on a UiO-66-NH₂ MOF, through molecular imprinting.¹³⁰ While encapsulation was successful, the colorimetric sensor demonstrated a relatively higher LOD in the milli-molar ranges (2.4 mM), and a narrow dynamic range of 2900–6700 μM .¹³⁰ Since normal blood cholesterol concentrations generally do not exceed 5200 μM ,¹³⁷ the sensor does prove useful for real-life, clinical or PoC applications. However, applications in patients with hypocholesterolemia or hypercholesterolemia may be limited due to the narrow range.^{130,137} This is further supported by the long shelf-life stability and good reusability of the MOF sensing system, where only 4% of catalytic activity is lost after 25 days and sensor performance only drops by 2.5% after eight cycles of reuse.¹³⁰ Thus, co-encapsulation of enzymes in MOFs have showing promising results for cholesterol detection, but further enhancements are needed to optimize performance.

4.2.4 Sensors using MOFs as peroxidase mimic. The need for enzyme immobilization and stabilization, as in earlier

studies, can limit the overall stability and effectiveness of the MOF system, and limit temperature and pH conditions. Therefore, many studies developed MOF-based composites as peroxidase mimics, also known as nanozymes, to minimize enzyme use and maximize cholesterol sensor performance.^{18,24,27–29,83,117,118,126,131} Through colorimetric,^{18,27–29,83,131} fluorescent,^{24,27,117,126,131} and/or chemiluminescent⁸³ mechanisms, these MOFs have demonstrated great performance in sensing cholesterol. These will be further discussed within.

Colorimetric MOF nanocomposites coupled with TMB have facilitated cholesterol detection at micromolar levels, with LODs as low as 0.80 μM .¹⁸ Liu *et al.* used pure MIL-101(Cr) as a peroxidase mimic, achieving an LOD of 0.86 μM and a linear dynamic range of 1.1–22.9 μM .²⁸ While the LOD is low, the linear range is quite narrow, requiring careful dilution of real samples to fit this range. The calibration curve is depicted in Fig. 10(D2) while the increase in color intensity and absorbance with cholesterol addition is shown in Fig. 10(D1). Furthermore, large incubation times (45 min) and higher temperature (60 °C) are required to achieve optimal results.²⁸ Wu *et al.* overcame these limitations through a hybrid nanocomposite of MIL-100(Fe) coated Fe₃O₄ magnetic NPs.¹⁸ In addition to using iron, which is cheaper than chromium, the system achieved a slightly lower LOD (0.8 μM) and a wider linear range (2–50 μM) at a lower incubation time (15 min) and temperature (50 °C). The MOF-Fe₃O₄ NP hybrid system also demonstrated great recyclability, retaining over 90% of its effectiveness after 5 cycles of use.¹⁸ Meanwhile, a Mo-doped Cu-2MI MOF achieved an even wider linear range (2–140 μM) in 15 min at a lower incubation temperature of 37 °C. While the LOD is slightly higher (1.2 μM), this system is unique in its one-step sensing process, where ChOx, MOF peroxidase mimic, TMB, and cholesterol are all incubated in one step,²⁹ unlike other systems requiring sequential treatment with ChOx then the MOF, prolonging detection.^{18,28} Despite the differences in sensor performance, the colorimetric mechanism adapted in these sensors is advantageous because a clear color change can be observed, allowing for applications in simple PoC devices.

MOF-based peroxidase mimics have also been used for fluorescent-based cholesterol detection.¹²⁶ For example, ZIF8 MOFs were decorated with Pd nanoclusters and coupled with thiamine dye, which upon oxidation converts to thiochrome, a fluorescent agent.¹²⁶ The hybrid nanocomposite demonstrated excellent peroxidase activity at alkaline conditions (pH = 10.5), and cholesterol detection was achieved after incubation for 20 min at 45 °C. Detection limits were as low as 0.92 μM , with cholesterol detection spanning over a 5 to 1000 μM , which is among the widest ranges achieved in the literature.¹²⁶ However, fluorescent sensors depending on single emission sources are more susceptible to interferences from external sources, thus increasing chances of errors.^{117,138,139} Therefore, peroxidase-like MOF nanozyme sensors have been designed to sense cholesterol through ratiometric fluorescence,^{24,117} whereby the ratio of two fluorescent signals at different wavelengths is used to determine analyte concentration.^{138,139} Ratiometric fluorescence-based sensors demonstrate lower signal-to-noise



ratios due to the elimination of background interferences.^{138,139} Yang *et al.*²⁴ developed an Eu-doped MIL-53(Fe) MOF for cholesterol quantitation by ratiometric fluorescence, where Eu centers are quenched and MIL-53(Fe) fluorescence increases with hydrogen peroxide; MIL-53(Fe) simultaneously acts as a peroxidase mimic to catalyze hydrogen peroxide reduction. The ratio of the quenched and enhanced fluorescent signals allowed for cholesterol detection at millimolar levels, with an LOD of 0.0105 mM and a linear range of 2000–8000 μM .²⁴ The LOD is much higher than the single-emission fluorescent sensor in ref. 126 and the use of Eu can add costs due to it being a rare metal.¹⁴⁰ A later ratiometric fluorescent cholesterol sensor by Ye *et al.*¹¹⁷ demonstrated better cholesterol detection with an LOD of 4.55 μM and an advantageously wide linear range of 5–1000 μM . Rather than metal-doping, Ye *et al.* functionalized MIL-101(Fe) MOF with carbon quantum dots, which facilitated more sensitive cholesterol detection at lower concentrations.¹¹⁷ Therefore, MOFs as peroxidase mimics can also facilitate fluorescent-based cholesterol detection.

Recent studies designed peroxidase MOF-based systems with dual fluorescent and colorimetric detection of cholesterol.^{27,131} Chen *et al.*¹³¹ achieved this through a nanocomposite of CeO₂ NRs and Cu(PABA) MOF peroxidase mimic. In combination with OPD dye, cholesterol introduction and hydrogen peroxide peroxidase reaction catalyzes the oxidation of OPD to DAP, a fluorescent, yellow dye. Colorimetric measurements of the yellow dye absorption allowed for a cholesterol detection limit of 0.162 μM , while fluorescence intensity of DAP allowed a slightly higher cholesterol detection limit of 0.178 μM . Meanwhile, both colorimetric and fluorescent measurements allowed for micromolar cholesterol quantitation in the linear range of 0.05–40 μM .¹³¹ The peroxidase activity of the MOF-nanocomposite was enhanced by the incorporation of CeO₂ NRs, and the sensor demonstrated excellent stability in extreme pH and thermal conditions, as well as a long shelf-life extending to 30 days.¹³¹ A QDs/UiO-66 nanocomposite dual mode colorimetric and fluorescent cholesterol sensor demonstrated a lower cholesterol LOD of 0.01 μM but a narrower dynamic range (0.04–1.60 μM) at room temperature, constraining the dilution needed for real samples.²⁷ Meanwhile, Amirzehni *et al.*⁸³ explored a novel, dual chemiluminescent and colorimetric detection technique, rather than fluorescence, using the chemiluminescent dye rhodamine B.⁸³ Here, the Co₃O₄@-CuMOF hybrid nanocomposite mimics peroxidases by catalyzing the reaction between rhodamine B and hydrogen peroxide, consequently causing a color change and producing a chemiluminescent intensity.⁸³ Quantification of cholesterol through these signals facilitated micromolar cholesterol detection over a wide range (0.1–300 μM) with an LOD of 0.026 μM .⁸³ Compared to fluorescence, chemiluminescence-based detectors often demonstrate greater sensitivity due to lower background noise as no external light source is needed, thus greater overall sensor performance.^{141–144} Nonetheless, further improvements are needed to widen the linear range. However, these dual-mode sensors demonstrate great potential for use in medical laboratories or in PoC devices for rapid cholesterol detection.

While the need for HRP is minimized using MOFs as peroxidase mimics, most cases require pretreatment with ChOx for cholesterol oxidation and hydrogen peroxide production to detect free cholesterol.^{18,24,27–29,83,117,118,126,131} Moreover, for total cholesterol quantitation, further pretreatment with cholesteryl esterase is needed,^{13,14} which all add to the overall sensor performance time and may require different conditions. However, this versatility also opens windows for using these peroxidase MOF nanozymes for sensing any biomolecules that can produce hydrogen peroxide, such as glucose, galactose, uric acid, lactic, and others, as proven in many studies.^{117,126,128,131} Multi-biomarker applications of MOF-based sensors are of great potential, and future directions can focus on further improving these systems for accurate detection and clinical, real-life applications.

4.2.5 Sensors using MOFs for direct, enzyme-free detection. Despite the great success of enzyme-based MOF sensors, current trends are shifting towards enzyme-free sensing systems, to minimize instabilities and prolong shelf-lives and operation times. Enzyme-free MOF-based cholesterol sensors are scarce, with only two systems reported in the literature. Wu *et al.*¹³² designed a V-MOF, pyrolyzed at 700 °C, to achieve dual oxidase and peroxidase behavior. Coupled with OPD, cholesterol can directly be oxidized by the MOF, then the produced hydrogen peroxide catalyzes conversion of OPD to DAP, resulting in fluorescence. Cholesterol detection was achieved at a LOD of 0.38 μM and over a linear range of 2–160 μM .¹³² In addition to applications in cholesterol detection, the MOF proved useful for *in vitro* quantifications of cholesterol in cell membranes, a unique application with various biological impacts.¹³² Gong *et al.*³¹ designed an enzyme-free MOF system with similar performance (LOD 0.4 μM), using β -CMCD-decorated NU-1000 MOFs. However, their design was based on a substitution reaction between cholesterol and rhodamine 6G dye at the β -CMCD pores, causing recovery of MOF fluorescence (Fig. 10(E1)). The dynamic range is shown in the calibration curve in Fig. 10(E2), while the fluorescence recovery can be visually seen when irradiated with UV-light in Fig. 10(E3).³¹ Nevertheless, both MOF-based sensors demonstrated great selectivity and stability, eliminating the need for enzymes and expanding sensor applications to *in vitro* assays.^{31,132}

Fig. 10 highlights the key performance of MOF-based spectrophotometric cholesterol sensors, depending on the role of the MOF: for ChOx immobilization (Fig. 10A), peroxidase immobilization (Fig. 10B), co-immobilization of ChOx and peroxidase (Fig. 10C), peroxidase mimics (Fig. 10D), and direct, enzyme-free cholesterol detection (Fig. 10E). Performance metrics illustrated include cholesterol detection fluorescent spectra, calibration curves, and selectivity studies, among others.

5. Integration into biosensing platforms

With the high costs and labor needed for laboratory testing, portable sensing platforms have grown in popularity, with



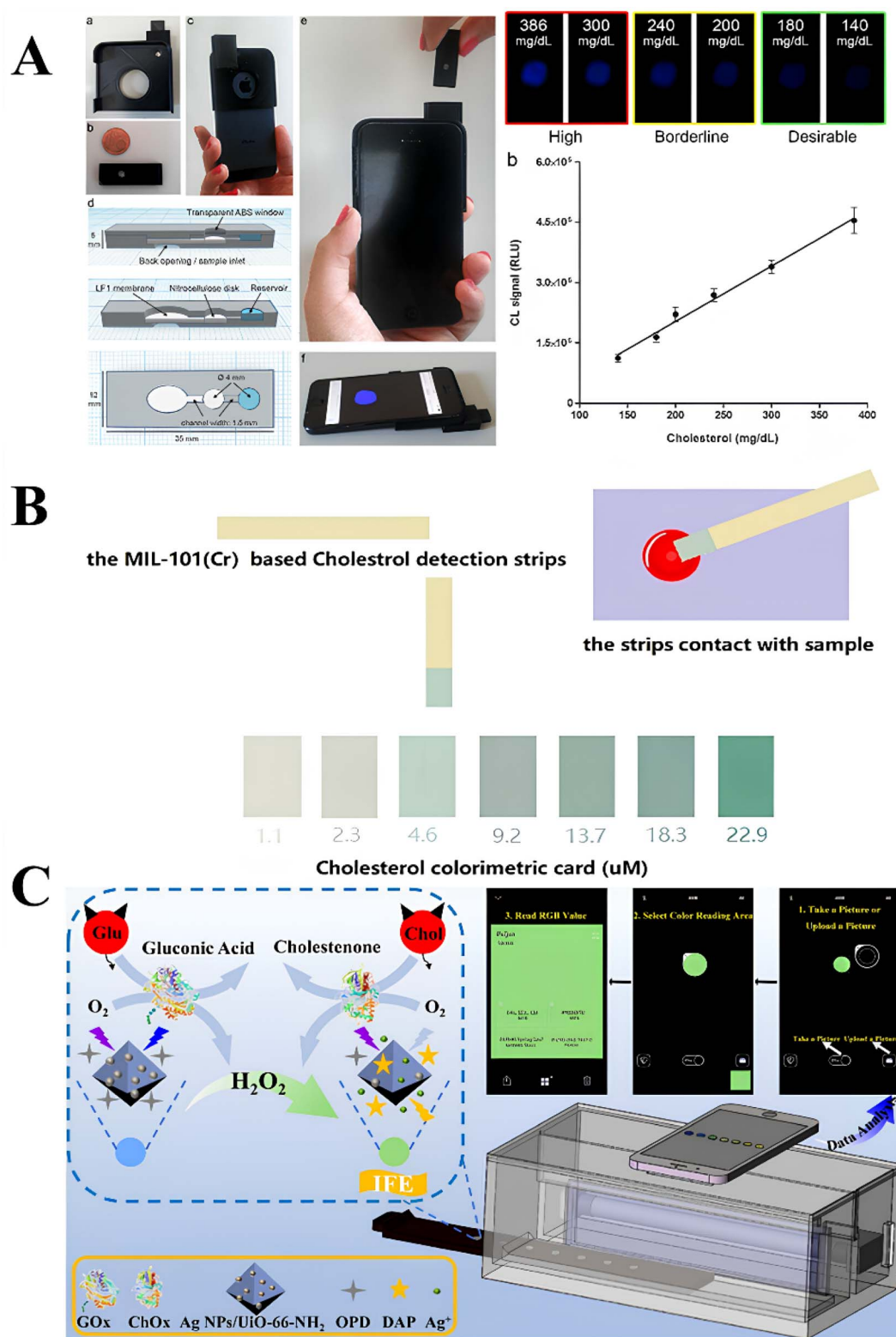


Fig. 11 Illustration of different PoC devices for spectrophotometric cholesterol detection. (A) 3D printed micartridge containing inlet for sample and reservoir for reagents is attached to smartphone, where chemiluminescence emission is detected for a range of cholesterol concentrations. Enzyme-based reagents are used. Reprinted (adapted) with permission from Roda *et al.*¹⁴⁷ Copyright 2024 American Chemical Society. (B) MIL-100(Cr) based test strips for colorimetric detection of cholesterol, where different color intensities represent the cholesterol concentration. Reprinted (adapted) with permission from Liu *et al.*²⁸ Copyright (2024) John and Wiley Sons. (C) A MOF-base fluorescence-based PoC device for cholesterol detection, where MOF and enzymes are immobilized in a box, and fluorescence measurements taken using smartphone camera. RGB analysis used to determine cholesterol concentration. Reprinted (adapted) with permission from Guo *et al.*¹⁵⁰ Copyright 2024 American Chemical Society.



research focusing on developing rapid, accurate, and portable point-of-care (PoC) sensing devices for biomarker and metabolite quantification.^{144,145} Few, spectrophotometric sensing platforms based on smartphone devices have been developed and tested in the literature.^{146–148} For instance, disposable test strips impregnated with cholesteryl esterase, cholesterol oxidase, 4-aminoantipyrine (4-AAP) and aniline sodium salt were combined with a cloud-enabled smartphone and a photochemical dongle to for colorimetric-based detection of cholesterol *via* red green blue (RGB) analysis.¹⁴⁹ With only 30 μL of blood, the smartphone-based sensing platform can quantify cholesterol concentrations within 2 minutes. Oncescu *et al.*¹⁴⁶ also used standard cholesterol strips for smartphone-based detection, whereby the smartphone's flash was used to illuminate the strip and measure colorimetric changes. Two ranges of cholesterol detection were achieved: low cholesterol concentrations (below 2600 μM) and high cholesterol concentrations (above 10 300 μM).¹⁴⁶ However, since normal cholesterol ranges lie around 5.2 mM, the sensor's application is quite limited.^{14,15,36} Furthermore, an important factor to consider when designing enzymatic strips is the method of immobilization, because weakly immobilized enzymes can lead to leaching and inaccurate analyte quantification.¹⁴⁹ Some immobilization techniques often used for test strips include adsorption, encapsulation, entrapment, and covalent linkages.¹⁴⁸

Roda *et al.* developed a similar smartphone-based PoC while using a chemiluminescent-based system with cholesteryl esterase, ChOx, HRP and luminol.¹⁴⁷ A 3D printed, miniature cartridge is used (shown in Fig. 11A), replacing paper strips used previously.^{146,148,149} Through the cartridge accessory contains a back opening where blood samples are introduced. Through a membrane, the sample travels to the reservoir where it mixes with the reagents; here the reaction starts. Since it is a chemiluminescent system, no external light is needed; therefore, the smartphone camera is used to simply capture emitted light from the sample (chemiluminescence), which is displayed on the phone screen.^{142,147} Only 15 μL of blood is needed for sensing, which can be easily obtained through a finger prick. The smartphone-based platform and its accessory is shown in Fig. 11A. Moreover, the sensing platform had a linear range between 20 mg dL^{-1} and 386 mg dL^{-1} , equivalent to 520–10 000 μM , which lies within the typical cholesterol ranges observed in patients.¹⁴⁷ The calibration curve and detection range, along with the expected signal shown in the smartphone at different concentrations is depicted in Fig. 11A. Although advantageous, these systems are heavily enzyme-dependent, reducing long term stability of test strips or solutions.^{146–149}

However, as explored earlier (Section 3.2), MOF-based spectrophotometric sensors can achieve lower LODs in rapid times with high sensitivity and selectivity. As such, future sensing platforms can replace these test strips and enzymatic solutions with MOF-based nanocomposite sensors for better sensor performance and stability, with lower costs. Guo *et al.*¹⁵⁰ achieved this by creating test strips with Ag NPs/UiO-66- NH_2 and OPD. Paired with a portable device, where the sample test strip is placed, and a smartphone, the system can be used to measure

colorimetric intensities, determine RGB intensities, and subsequently calculate cholesterol concentrations by taking a photograph.¹⁵⁰ The MOF sensing mechanism, the portable device, and the RGB pictures and analysis are all depicted in Fig. 11C. A much lower LOD of 10 μM was obtained, with high selectivity to cholesterol.¹⁵⁰ However, larger response times were needed for accurate measurements, which is a drawback not observed in many PoC devices. Other MOF-based cholesterol sensors have also demonstrated potential for use in PoC devices, such as MIL-101(Cr), which was successfully developed into simple test strips.²⁸ Once in contact with blood samples, the test strip changes color, and the resulting color is compared to a standard test strip to determine cholesterol concentration range.²⁸ The color changes correlated with different cholesterol concentrations are shown in Fig. 11B.

Research on integration spectrophotometric-MOFs into PoC sensing device is relatively recent. Future studies can design PoC devices that use these strips for accurate cholesterol quantification by measuring RGB intensities. MOF-assays may also be developed into miniature cartridges, especially chemiluminescent systems, as inspired by earlier studies.¹⁴⁷ The field of PoC devices for cholesterol detection using MOFs is still young, with many gaps for research and advancement of current technologies and developing new technologies. Fig. 11 illustrates current PoC devices for spectrophotometric cholesterol detection, using enzymes or MOFs.

6. Current challenges & future directions

Despite the variety of advantages offered by MOF-based spectrophotometric sensors for cholesterol detection, many challenges are faced, requiring further research and development to enhance current sensors for optimal results.

6.1 Stability of sensing elements

Cholesterol has very poor solubility in aqueous mediums, and more optimal solubility in organic solvents.⁹ Meanwhile, if enzymes like ChOx and HRP are used, an aqueous medium mimicking physiological condition is needed for optimal catalytic activity. Hence, MOF-based sensors must exhibit good stability in both aqueous and organic solvents, and over a range of pHs and temperatures, for best sensing applications.³³ This challenge is further emphasized when introducing hybrid nanocomposites of different nanomaterials, requiring exhaustive stability tests. While most reported MOF-based sensing materials have been reported to have good performance in different mediums,^{130,131} one expects the process to developing these systems to be lengthy and difficult. Moreover, as MOF-nanocomposite complexity increases, challenges in identifying the most suitable buffer and pH also arises, introducing new challenges. Future research can look into tackling this issue through high throughput testing applications, to rapidly understand MOF stability in different solvents at one. Furthermore, artificial intelligence and machine learning tools can be used to predict preliminary results, which can save time and



steer research towards the correct direction. Enzymes also contribute greatly to the stability of the sensing platform, due to their high sensitivity to changes in temperature, pH, solvent, *etc.*^{151–153} Hence, future studies should aim to develop enzyme-free systems with high sensing accuracy and performance for enhanced stability and expanded lifetime of sensing platform.^{73,151,152} Nonetheless, MOF-based sensors are still advantageous and show great promise for real-life, PoC applications.

6.2 Multi-biomarker detection

While accurate sensing of cholesterol alone is crucial, practical application of MOF-based sensors for multiple biomarker detection are of greater importance and value.^{131,149} However, detection of multiple biomolecules using one MOF-system is quite challenging, introducing limitations in the sensor design, input areas, materials, concentrations, and much more. This is especially highlighted in PoC applications, where buying multiple sensors for each biomolecule can be costly. Although challenging, multiplex biomarker detection has been implemented successfully in ref. 128, 131 and 149, for dual detection of glucose and cholesterol¹²⁸ or for detection of glucose, cholesterol, and uric acid.^{131,149} However, these MOF-sensors can be time consuming. Future studies can focus on improving multiplex cholesterol, MOF-based sensors for more rapid detection at higher sensitivity, perhaps through exploring microfluidics as a promising option since it has been proven useful for electrochemical-based cholesterol detection.¹⁵⁴

6.3 Continuous, non-invasive sensing

Patients with hypercholesterolemia, hyperlipidemia, and high risks of cardiovascular diseases require continuous monitoring of cholesterol levels for rapid treatment when spikes occur.¹⁵⁵ While multiple, small blood samples can be taken (through finger pricks) and tested using MOF-based spectrophotometric sensors,¹⁴⁹ the process is very invasive, and patients may resist multiple monitoring stages throughout the day. Adapting spectrophotometric MOF-based nanocomposites into real-time, non-invasive sensors is a significant challenge, and a promising future direction of research. To tackle non-invasiveness, MOF-based cholesterol sensors can be adapted for detecting cholesterol in saliva¹⁵⁶ or tear¹⁵⁵ samples, as done in other electrochemical sensors. Designing wearable devices with these MOFs can allow for real-time monitoring.^{155,157} These fields of research are still in infancy, with MOFs showing great potential in advancing the field of spectrophotometric cholesterol sensing.

6.4 Integration into wearable devices

While MOF-based spectrophotometric has demonstrated high accuracy in cholesterol detection at low concentrations, their integration into wearable devices for continuous cholesterol monitoring remains a challenge. Smart wearable devices for quantification of metabolic biomarkers, including glucose, cholesterol, *etc.*, require miniature sizes and high accuracy and sensitivity to target analyte in the presence of various interferences.^{158,159} Most MOF-based sensors reported in the literature require specific temperature, pH, and environmental

conditions for optimal operation, and deviations from optimized conditions can lead to poor accuracies and sensitivities.^{158,160} This poses a significant challenge because wearable devices are subjected to various interferences, including skin temperature and pH variations, contamination with dust, sweat, hair interference, among other factors.^{158,160} This can alter the performance of the wearable biosensor, leading to inaccurate readings and impeding commercialization and clinical integration. While a study succeeded in creating wearable contact lenses for cholesterol monitoring, further clinical trials are needed for full evaluation of their safety and efficacy.¹⁶¹ Thus, future studies should focus on developing more robust, stable sensing systems, using MOFs or other nanomaterials, for more versatile applications in wearable sensors for early detection of diseases and continuous monitoring.

6.5 Scalability & commercialization

Scaling up lab biosensing and chemo-sensing system into point-of-care (PoC) devices, with potential for commercialization and public use, is an emerging field or of high importance in healthcare.^{162–167} Although numerous studies have developed accurate cholesterol sensing systems, scaling up such systems into PoC devices for commercialization remains a persistent challenge faced in sensing platform development. Various electrochemical-based cholesterol sensors have been commercialized and scaled-up.²¹ However, more recent spectrophotometric-based cholesterol detection systems remain restricted to research labs. This can be attributed to lack of reliable methods for scaling up production of MOFs and MOF-based composites,¹⁶⁸ and the costs associated with spectrophotometric detection components.¹⁶⁹ Integration of these nanomaterial-based sensors into microfluidic systems may improve scalability and speed up commercialization, due to wide range of studies on scaling up production of microfluidic systems.¹⁶⁶ Hence, future research should aim to investigate methods for improving scalability while maintaining accuracy and reliability of the sensing systems – to allow for their integration into PoC devices and future commercialization at low costs.

6.6 Environmental concerns

The high stability of MOFs, although useful in sensing applications, poses environmental concerns relating to persistence and degradability.^{170,171} This challenge is significantly highlighted when MOF-based sensors are implemented in PoC devices, since proper disposal becomes more challenging and control over the process is limited.¹⁷⁰ If the MOFs used contain toxic metals or organic linkers, which is common in many MOFs, public and environmental health, including aquatic life, become threatened.^{171,172} Moreover, preparation of MOFs often require harsh acidic conditions and reagents, further increasing environmental concerns relating to these materials.^{170,171} To minimize environmental impacts concerning synthetic protocols, new synthesis pathways are being developed using less materials.^{173,174} Furthermore, circular economy principles show great applications in mitigating environmental impacts of MOF



synthesis, whereby materials are reused, recycled, and recovered to minimize waste.¹⁷⁰ Moreover, MOFs are now being designed using biodegradable linkers, to eliminate both environmental and biological toxicities.^{173,175} Still, further research is needed to design effective, non-toxic MOFs specific to cholesterol detection, through eco-friendly synthetic approaches.

7. Conclusions

Overall, research on cholesterol sensors continues to grow, with recent interests in spectrophotometric, MOF-based sensors for cholesterol detection. In comparison to other nanomaterials, MOFs have high surface areas, large pore sizes, and highly tunable structures, allowing for flexibility in designing sensitive cholesterol sensors. Spectrophotometric MOFs for cholesterol sensing have specifically grown in popularity over the last decade, as illustrated in Fig. 2. With these increased research interests and the absence of a comprehensive review of these sensors, this paper aims to analyze and review current research on spectrophotometric MOFs for cholesterol sensing. MOFs are valued over other nanomaterials due to their unique high surface area and porosity, flexible tunability, and easy surface functionalization. Furthermore, in contrast to the traditional colorimetric approaches, MOF-based cholesterol sensors follow five different mechanisms of sensing, where MOFs are used for ChOx immobilization peroxidase immobilization, dual ChOx and peroxidase immobilization, peroxidase mimics, and direct, enzyme-free detection of cholesterol. To the best of our knowledge, this paper is the first to categorize these MOF roles and analyze them. Furthermore, spectrophotometric-based MOFs have great potential in PoC device applications, especially smartphone-based devices.

However, some challenges regarding MOF stability and environmental impacts arise. Future research can help overcome these challenges by:

- Developing spectrophotometric MOFs for multi-biomarker detection, including cholesterol, glucose, uric acid, triglycerides, and insulin, *etc.*
- Exploring development of non-invasive and real-time monitoring cholesterol sensors. This may be done through integrating microfluidics in the sensor platform, for miniaturization and continuous detection.
- Integrating artificial intelligence (AI), where it can be used for optimizing MOF design, predicting sensor performance, and monitoring changes into cholesterol concentrations to predict patient health.

Overall, although the field is quite recent, future applications of spectrophotometric MOFs for cholesterol sensing appear highly promising.

This review paper offers a comprehensive analysis of applications of MOFs in spectrophotometric-based cholesterol sensing, which has not been previously reviewed to the best of the authors' knowledge. The review enables to categorize the various roles that MOFs play for cholesterol sensing and explains the associated attachment mechanisms. Furthermore, this review is informative given its inclusion of the recent

literature and advances, as well as real-life applications of MOF-based spectrophotometric sensors in PoC devices. On the other hand, we note that the present review is greatly focused on spectrophotometric MOF detection and thus excludes many studies dealing with cholesterol sensors adapting other techniques, such as plasmonic-based detection techniques.^{65,77,78} Moreover, the paper focused solely on cholesterol detection, disregarding literature detecting other important biomarkers, like glucose, uric acid, *etc.* Thus, the review serves as a focused evaluation and analysis of the current literature on MOF-based spectrophotometric cholesterol sensors, which may not be adequate for readers interested in exploring broader sensing platforms including multiple biomarkers or nanomaterials. Nonetheless, this review remains important and timely for researchers interested in developing or improving MOF-based spectrophotometric sensors and integrating them into PoC sensing devices.

Data availability

Data available on request from the authors.

Conflicts of interest

The authors declare no conflict of interest.

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