

Part Five

Physical Applications

12

Luminescent Metal–Organic Frameworks

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12.1

Introduction

Luminescence is a commonly reported property of metal–organic frameworks (MOFs), in large part because these structures typically contain aromatic or conjugated linker groups that readily fluoresce, or incorporate luminescent lanthanide ions. Because the moieties are largely immobilized by the crystal structure, the lifetimes and quantum efficiencies may be higher than those exhibited by the isolated luminophore in solution. MOFs are not completely rigid, however; their well-known structural flexibility can lead to dramatic changes in the local coordination environment and corresponding variations in emissive properties. The nanoporosity of these structures also creates new opportunities to impart light-emitting properties that are not available to traditional inorganic complexes. In particular, infiltrated guest molecules that are themselves emitters or sensitizers can be immobilized in close proximity to luminescent centers. This can influence the emission properties of the neat MOF in the form of wavelength shifts, intensity changes, or new emission resulting from excimer or exciplex formation. Since fluorescence spectrometers are readily available, many investigators report the excitation and emission spectra of newly synthesized MOF structures. As a result, more than 300 reports of luminescent MOFs are now available in the literature. Relatively few studies delve into the underlying spectroscopy or use luminescence as a tool for probing electronic structure and energy transfer. Nevertheless, it is apparent that luminescence holds considerable potential not only to serve as a diagnostic tool for probing fundamental aspects of MOF behavior and structure, but also to show potential for a variety of applications. As might be expected, chemical sensing is a logical possibility, but as will be seen later in this chapter, the potential extends well beyond this to include radiation detection, nonlinear optical effects, and solid-state lighting.

The purpose of this chapter is first to introduce the reader to the basic concepts of luminescence relevant to much of the current MOF literature. We then discuss luminescence associated with individual components of MOFs, each of which displays characteristic features. Examples from the recent literature are provided that illustrate the multidimensional character of MOF luminescence. However, a

comprehensive review is not presented, as this was recently done [1]. As will be seen, a detailed understanding of luminescence phenomena in MOFs is beginning to emerge. Although still in its early stages, the results demonstrate that the unique characteristics of these supramolecular materials provide a platform for understanding light emission with both unparalleled synthetic flexibility and a highly ordered, inherently quantifiable, structure. From a practical point of view, it is also becoming clear that luminescence spectroscopy is a superb tool for assessing the structural integrity and purity of MOFs, with sensitivity far exceeding that of powder X-ray diffraction (PXRD). The chapter concludes with an overview of recent developments concerning applications of luminescent MOFs.

12.2

Luminescence Theory

In general, luminescence is the result of radiative decay (i.e., emission of light) by electronically excited molecules following the absorption of energy. The absorbed energy can originate from a variety of sources, which leads to different sub-classifications of luminescence. When the energy is absorbed from electromagnetic radiation in the form of photons (light), the subsequent emission of light is defined as *photoluminescence*. The nonthermal production of light by a chemical reaction is termed *chemiluminescence* [2–4], with the special case involving living organisms being referred to as *bioluminescence* [5–7]. In some cases, the presence of an electric field can cause electron–hole recombination in a material, producing *electroluminescence* [8, 9]. If the energy absorbed by the molecules is due to the material being bombarded with high-energy ionizing radiation (>10 eV), the emission of light is referred to as *radioluminescence*, while a beam of high-energy electrons imparted upon a material may lead to *cathodoluminescence*. Additionally, mechanical forces [10–12] acting on a material may also lead to the emission of light in the form of *triboluminescence* (mechanical stress, scratching, or fracture) or *piezoluminescence* (pressure). Although there appear to be several methods to excite a molecule or group of molecules in a MOF electronically to yield emission of light, to date only a few have been investigated, the most prevalent of which has been photoluminescence (PL).

12.2.1

Photoluminescence

Although many excellent texts are available concerning the theory of luminescence in general [13–15] and PL [16, 17] in particular, a general, albeit simplified, overview of the concepts is warranted. For the sake of simplicity in describing the photophysical processes in question, let us consider only the organic molecule, which acts as a linker or strut in the formation of the MOF. Before a molecule is excited by the absorption of light, the molecule resides in its lowest electronic energy level or ground state. Typically, an organic molecule has a singlet ground state (S_0), but in rare cases a triplet ground state (T_0) is possible. Higher energy electronic states exist (S_n or

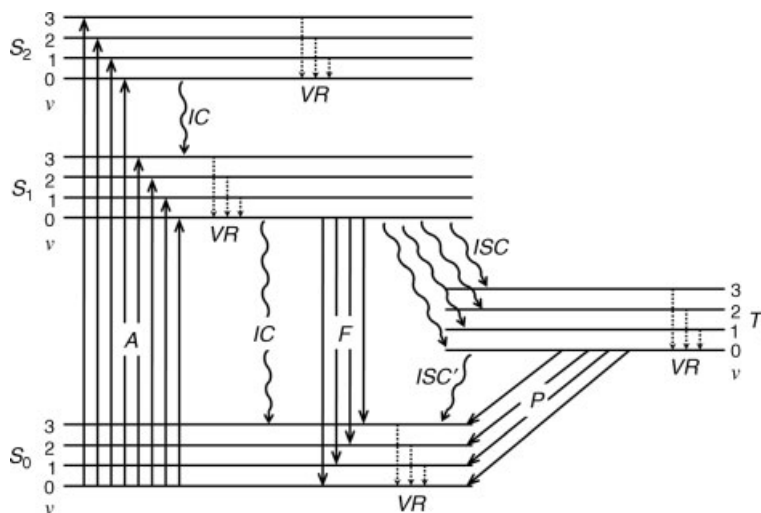


Figure 12.1 Jablonski diagram depicting the photophysical processes of a conjugated organic molecule. Absorption (*A*) of energy is followed by fast, non-radiative processes such as vibrational relaxation (*VR*) and internal conversion (*IC*) down to the lowest vibration level of the first excited state ($S_{1,0}$). From here there are three processes for the deactivation of $S_{1,0}$: the non-radiative processes of internal

conversion (*IC*) and singlet-triplet intersystem crossing (*ISC*), as well as the radiative process of fluorescence (*F*). If the deactivation process leads to singlet-triplet *ISC*, further relaxation from the lowest excited triplet state ($T_{1,0}$) can proceed through either radiative (Phosphorescence, *P*) or non-radiative (triplet-singlet intersystem crossing, *ISC'*) deactivation to the ground state.

T_n ; $n \geq 1$), and it is these elevated states to which an electron is promoted upon the absorption of energy. Additionally, each electronic state is associated with several vibrational energy levels and together the electronic and vibrational levels constitute a vibronic state ($S_{n,v}$; $v = 0, 1, 2, 3, \dots$). A Jablonski diagram (Figure 12.1) is a simplified graphical representation of the vibronic states for a given molecule wherein higher energy levels are stacked vertically over lower ones. Jablonski diagrams are useful for depicting the various photophysical processes involved in the absorption and eventual loss of excess energy by the molecule, but do not account for geometric distortions. The vertical nature of the transitions is reflective of the fact that absorption occurs in a time frame that is too short for significant relocation of nuclei (Franck–Condon principle) and therefore rearrangement of electronic energy states is negligible.

In the ground state, nearly all of the molecules exist in the lowest totally symmetric (a_1) vibrational level ($S_{0,0}$) at room temperature, and so it is from this vibronic state that absorption of energy leads to excitation to elevated energy levels. In order to energize a molecule sufficiently and promote it to a higher electronic state, the use of ultraviolet (UV) or visible light is required. Light of lower energy (and longer wavelength) such as infrared (IR) is typically not energetic enough to cause promotion to higher electronic states, but can be useful for the excitation of higher vibrational levels of the ground electronic state. Upon excitation, the molecule begins rapidly to lose excess vibrational and electronic energy to the surrounding medium

(when in a condensed phase) in a nonradiative thermal deactivation process. In the first phase, the molecule loses excess vibrational energy through vibrational relaxation (VR), in which the molecule undergoes a radiationless transition from a higher to a lower vibrational level within the same electronic state. This is followed by a radiationless transition between electronic states of the same multiplicity called internal conversion (IC) and possibly further VR within the new electronic state. Taken together, VR and IC constitute thermal deactivation of the molecule carrying it rapidly to the lowest vibrational level of the lowest excited state (10^{-13} – 10^{-12} s). From the $S_{1,0}$ excited state, the molecule can undergo deactivation via one of three first-order processes. For many molecules it is possible to become deactivated by nonradiative internal conversion (rate constant k_{ic}). However, for some molecules containing relatively large gaps in energy between the lowest excited and ground states, this process is unfavorable, and the majority of these surrender their excess energy via the spin-allowed emission of light known as fluorescence (F, k_f). The third possibility is the formally spin-forbidden intersystem crossing (ISC, k_{isc}) to the lowest excited triplet state T_1 . The excited triplet state must also undergo deactivation and can do so through either spin-forbidden radiative emission known as phosphorescence (P, k_p) or through nonradiative spin-forbidden ISC ($k_{isc'}$). As there are several competing pathways for the deactivation of the lowest excited singlet state, the number of molecules that fluoresce is only a fraction of the total population of excited molecules. This ratio between the number of molecules which fluoresce and the total excited population is the fluorescence efficiency or *quantum yield of fluorescence* (Φ_f). The quantum yield is a value between 0 (no fluorescence) and 1 (all excited molecules fluoresce) and is related to the first-order rate constants for the deactivation pathways (Equation 12.1). Similarly, the ratio of the number of molecules to undergo phosphorescence to the number of molecules to be deactivated in any of the other processes is given by the related Equation 12.2.

$$\Phi_f = \frac{k_f}{k_{ic} + k_f + k_{isc}} \quad (12.1)$$

$$\Phi_p = \frac{k_p k_{isc}}{(k_{isc'} + k_p)(k_{ic} + k_f + k_{isc})} \quad (12.2)$$

Another useful property of PL, especially in the comparison of fluorescence and phosphorescence, is the concept of the lifetime (τ) of a given excited state. The lifetime is defined as the average time the molecule exists in the excited state before the excited population is reduced by a factor of e (exponential decay). The lifetime of an excited state is therefore the inverse of the summation of the rate constants for the potential decay pathways deactivating that state and is given by the following equations:

$$\tau(S_1) = \frac{1}{k_{ic} + k_f + k_{isc}} \quad (12.3)$$

$$\tau(T_1) = \frac{1}{k_{isc'} + k_p} \quad (12.4)$$

The *natural* lifetime (τ_n) is the intrinsic lifetime of the luminophore in the absence of nonradiative decay and can be calculated from the ratio of τ to Φ . As the processes involved in deactivation of S_1 through the T_1 excited state are all spin-forbidden, the lifetimes are typically much longer for phosphorescence (10^{-4} – 10^0 s) than that observed for fluorescence (10^{-11} – 10^{-7} s).

12.2.2

Fluorescence Quenching

Although many types of molecule are capable of absorbing energy to become electronically excited, relatively few exhibit the phenomenon of luminescence. This is due in part to *quenching*. Essentially, any process capable of diminishing the intensity of fluorescence or phosphorescence can be considered a quenching mechanism [18], including the rapid nonradiative de-excitation of $S_1 \rightarrow S_0$ via internal conversion. External molecules called quenchers are also capable of de-exciting a molecule; this will either diminish or completely eliminate the ability of the excited molecule to luminesce. Quenching in MOFs is important from an applications aspect as it can be both detrimental, leading to a reduction in the desired property (luminescence), and beneficial in cases where the quenching of a known luminescence signal is used as a sensor mechanism for detecting the presence of the quencher moiety. Luminescence quenching is observed in one of two forms, *dynamic* and *static*. Dynamic quenching is characterized by the collision of a quencher moiety (acceptor) with an excited-state luminophore (donor), resulting in de-excitation of the donor and concomitant excitation of the acceptor. The newly excited acceptor may or may not be luminescent in its own right, but any resulting emission will always be of lower intensity than what would be observed from the donor. In dynamic quenching, the quencher and potential luminophore must come into contact during the lifetime of the luminophore excited state, and is therefore a time-dependent process. Additionally, dynamic quenching is dependent on the concentration $[Q]$ and diffusion of the quencher molecule and obeys the Stern–Volmer equation (Equation 12.5). In static quenching, interaction between a potential luminophore and a quencher molecule takes place in the ground state, leading to the formation of a non-luminescent complex. Here the efficiency of quenching is governed by the rate of complex formation (k_c) and $[Q]$ (Equation 12.6).

$$\frac{\Phi}{\Phi_0} = \frac{1}{1 + k_q \tau_0 [Q]} \quad (12.5)$$

$$\frac{\Phi}{\Phi_0} = \frac{1}{1 + k_c [Q]} \quad (12.6)$$

12.2.3

Energy Transfer

Another important process that can affect the luminescence of MOFs is energy transfer (ET). It may be possible, during the lifetime of the excited state, to transfer the

excitation energy from an excited molecule (donor) to another molecule in the ground state (acceptor). This is essentially another method of quenching the donor molecule as it will become de-excited and no longer luminescent. The acceptor absorbs the energy and is excited, which may lead to luminescence (albeit at a lower intensity), or it may deactivate via a nonradiative pathway so that as a system the luminescence will be lost. There are two mechanisms for ET distinguished by the nature of the donor–acceptor interaction. In *Förster resonance energy transfer* (FRET; Förster type; dipole mechanism; coulombic mechanism), the donor and acceptor molecules are not in contact and can be separated by as much as 10 nm [13, 18, 19]. For FRET to occur, at least some spectral overlap between the emission of the donor and the excitation of the acceptor must exist and the rate of energy transfer (probability) is dependent on both the amount of overlap and also the distance between the two molecules, with the rate decreasing in relation to $(1/r)^6$. In *excitation energy transfer* (Dexter type; exchange mechanism), the orbitals of the donor and acceptor molecules must be in direct contact. Dexter-type ET is sometimes considered as a double-electron transfer process, as the highest energy electrons from the donor (located in the LUMO) and the acceptor (located in the HOMO) are exchanged. Since the acceptor gains an electron in its LUMO while giving up an electron from its HOMO, the acceptor becomes excited while conversely the donor is deactivated. While Dexter-type ET requires spectral overlap as in FRET, this process is also diffusion controlled and depends on the viscosity of the solvent (i.e., the environment) [19].

12.3

Ligand-Based Luminescence

12.3.1

Solid-State Luminescence of Organic Molecules

The study of luminescence from organic compounds in the solid state is of increasingly high importance. In addition to the interest in fundamental physical studies, implementation of organic materials as active components in solid-state devices has led to increased attention in this area. Relevant examples include organic light-emitting diodes (OLEDs), thin-film sensors, and solid-state lasers. A major challenge for the development of such devices is suitable control over the morphology and interfacial properties of emissive organic species during device fabrication. Factors such as luminophore orientation, intermolecular distance, packing density, and details of the local molecular environment often have a significant impact on the resulting emission (and absorption) wavelength, intensity, and lifetime. Highly fluorescent organic molecules often feature an extended π -conjugated system (although this is not a requirement for emissive behavior). Such molecules share features in common with ligands often used in the field of MOFs, where a rigid, π -conjugated “linker” is often used to form the struts connecting metal ions or clusters. The overall structure of a MOF is mainly determined by geometric

requirements arising from the strong coordination between the metal ions or clusters and the ligand groups, and may be varied by choosing suitable starting materials and/or growth conditions. MOFs, therefore, have a degree of structural predictability and can allow for a controllable platform for arranging luminophores in order to alter the solid-state emission properties of the material. Conversely, the structure of an entirely organic solid-state material (i.e., a molecular solid) is often dictated by a multitude of weaker crystal packing interactions, including π - π stacking and hydrogen bonding, and there is little control over the structure ultimately formed in the solid state. For MOFs, the wide variety of organic ligands and coordination geometries available to afford predictable structures, and also the ability to accommodate guests, potentially allow for many degrees of freedom and, consequently, rational modulation of the emissive properties of the material.

Several possibilities exist for the type of emission displayed by MOFs containing π -conjugated ligands. In some cases, there is little or no electronic interaction between the organic and inorganic moieties, leading to ligand-based luminescence, as will be discussed in Section 12.3.2. These ligands may be isolated from each other in the structure, resulting in emission characteristic of a dilute solution of the free ligand, or the intermolecular distance and/or orientation may be such that significant interaction occurs, leading to ligand-to-ligand charge transfer (LLCT) emission with a typically broadened and red-shifted emission profile. Sections 12.3.3 and 12.3.4 describe materials for which the luminescence properties are determined not only by the organic molecule(s), but also through interaction with the metal unit(s). Two limiting cases are possible, ligand-to-metal charge transfer (LMCT) or metal-to-ligand charge transfer (MLCT). The particular type of emission observed depends on the structure of the MOF (which dictates the spacing and orientation between the ligands and metal units), the electronic configuration of the metal, and the degree of orbital overlap between the organic and inorganic moieties. Several detailed structure–property relationships have been described for luminescent MOFs, using both steady-state and time-resolved measurements, with the number of reports in this area increasing significantly over the past few years. The following sections are intended to exemplify the important features and principles involved for correlating structure with observed luminescence in MOF materials.

12.3.2

Ligand-Based Luminescence in MOFs

Ligand-centered luminescence typically occurs for MOFs in which there is little or no electronic interaction between the organic and inorganic portions of the structure. Typically, this is the case for structures containing metal ions that have no d–d or f–f transitions possible, most commonly for MOFs with d^{10} Cd^{II} , Zn^{II} , and Ag^{I} ions. An excellent example of this is provided by the IRMOF series of Yaghi and co-workers [20], in which the bonding between anionic carboxylate linkers and Zn^{II} ions is essentially ionic and involves very little charge transfer (see the discussion below). However, the presence of a d^{10} metal ion does not necessarily dictate that the structure will display ligand-based emission, as upon light absorption the ligand may

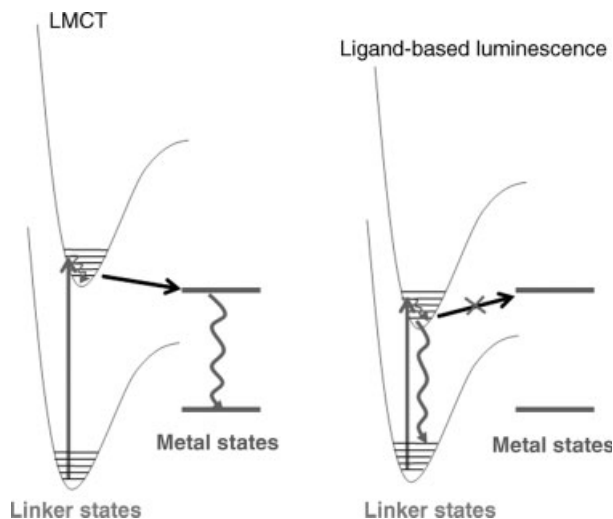


Figure 12.2 Energy level diagram depicting HOMO and LUMO states of the organic ligand, and intersystem crossing (LMCT) to emissive metal states (left). If π -conjugation in the ligand is increased, the HOMO–LUMO gap is decreased, making LMCT inefficient and,

therefore, ligand-based emission is observed (right). In addition, coupling to surrounding solvent or guest molecules may also affect the efficiency of these transitions for porous structures.

facilitate charge transfer to higher energy, emissive metal states if the ligand HOMO–LUMO gap is large. Moreover, as will be seen throughout this chapter, the complexity of MOF structures can frequently produce unexpected luminescent features that defy placement within straightforward categories such as this.

The IRMOFs illustrate the effect of increasing linker conjugation. When this occurs, the π – π^* energy gap decreases [13], making energy transfer to the metal less efficient. The resulting ligand-based emission resembles that of the corresponding ligand in dilute solution (Figure 12.2). A variety of IRMOFs exhibit emission similar to that of their isolated ligands, including IRMOF-11 $[\text{Zn}_4\text{O}(4,5,9,10\text{-tetrahydro-2,7-pyrenedicarboxylate})_3]_n$, IRMOF-13 $[\text{Zn}_4\text{O}(2,7\text{-pyrenedicarboxylate})_3]_n$, and $[\text{Zn}_4\text{O}(4,4'\text{-stilbenedicarboxylate})_3]_n$ [1]. Each of these structures exhibits emission similar to that of the corresponding dilute ligand in solution, indicating that ligand-centered luminescence dominates and there is little influence of the Zn_4O cluster on the electronic structure of the ligand. In addition, the absence of other bands to the red of this emission indicates there is no *significant* LLCT.

A second factor that strongly influences ligand-based MOF emission is the possibility of interpenetration within the structure. The three IRMOF structures just discussed consist of interpenetrated frameworks as a consequence of the increased length of the rigid, linear linker. MOFs with interpenetrated structures consist of two or more frameworks intertwined, and details of the structure, such as distance between the interpenetrated frameworks and dihedral angle of the organic ligands, will lead to variations observed in the emission wavelength, fine structure,

and intensity. Although significant LLCT was ruled out by detailed experimental studies in the $[\text{Zn}_4\text{O}(\text{4,4'}\text{-stilbenedicarboxylate})]_n$ structure [21] (as discussed in more detail below), this may not necessarily be the case for other interpenetrated MOFs, such as the pyrene- and tetrahydropyrene-based IRMOFs, for which careful studies of these two structures have not yet been carried out. However, other IRMOFs that exist as interpenetrated and non-interpenetrated versions have been probed in more detail. Meek *et al.* compared the excitation and emission spectra of the interpenetrated and non-interpenetrated versions of two IRMOFs constructed from the same ligand [22]. IRMOF-9 and -10 are constructed from 4,4'-biphenyl dicarboxylic acid (with IRMOF-9 being interpenetrated), and IRMOF-15 and -16 are constructed from 4,4'-terphenyldicarboxylic acid (IRMOF-15 is interpenetrated). In the terphenyl MOFs, the interpenetrated structure exhibits red-shifted emission compared with the non-interpenetrated variant, suggesting that interpenetration induces greater charge transfer, possibly due to smaller linker-linker and linker-metal distances. In the biphenyl structures, the interpenetrated MOF (IRMOF-9) displays a completely new band at ~ 500 nm, indicating the formation of a new excited state. This emission is broad and unstructured, suggesting that it involves charge transfer between biphenyl linkers in close proximity.

One might ask why this section did not begin with a discussion of IRMOF-1, the simplest of the cubic IRMOF structures, in which the linker is 1,4-bdc (1,4-benzenedicarboxylate) [23]. Here, although the 1,4-bdc linker is fully conjugated, the HOMO-LUMO energy gap is large; the excitation maximum of the isolated linker in dilute DMSO solution occurs at 338 nm. In contrast, a green emission upon UV excitation is observed from IRMOF-1 in some cases. This has been attributed to an LMCT process, whereby energy transfer occurs from the 1,4-bdc singlet excited state to the Zn_4O clusters [23]. Although this mechanism is supported by excitation spectra indicating that light absorption occurs via the benzene unit and by *ab initio* studies [24], recent work has demonstrated that ligand-centered luminescence is actually the dominant process and that the green emission attributed to LMCT is due to synthetic byproducts present in the structure or degradation of the structure itself (framework collapse) [25]. This investigation emphasized that unambiguous knowledge of the structure and constituents (including contaminants) of a material is necessary for accurate interpretation of the physical causes behind observed luminescence properties. Additionally, it demonstrates that changes in framework structure, undetectable by standard characterization methods such as PXRD, can be easily identified by the significantly more sensitive luminescent methods.

The effects on the emission properties of varying the orientation and spacing between ligands have also been demonstrated in the case of Zn stilbene MOFs [21]. Two MOF structures with different topologies were synthesized from the same starting materials via changes in the growth conditions and solvent, yielding both a two-periodic net $[\text{Zn}_3\text{L}_3(\text{DMF})_2]_n$ and a cubic three-periodic framework $[\text{Zn}_4\text{OL}_3]_n$ structure, where $\text{L} = 4,4'\text{-stilbenedicarboxylate}$ (Figure 12.3). Both materials exhibit intense ligand-based luminescence, similar in shape and vibronic structure, signifying little to no influence of the different metal cluster types present in each structure. However, emission from the two-periodic structure is red shifted and

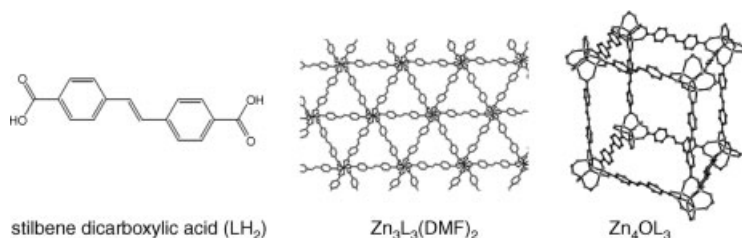


Figure 12.3 Structure of 4,4'-stilbene dicarboxylic acid, the layered 2D Zn stilbene MOF structure, and an individual cubic unit of the catenated 3D Zn stilbene MOF structure.

broadened from that of the cubic three-periodic structure, indicating a greater degree of interaction between luminophores in the two-periodic structure. This is consistent with the 6.0 Å, cofacial spacing between ligands in successive layers observed for this structure. It should be noted that the extent of this intermolecular interaction is small compared with that observed for the ligand itself in the solid state, which exhibits a significantly broader and red-shifted emission. The three-periodic cubic structure consists of anisotropically catenated cubes (edges are closer along the *b*-axis), a likely result of very moderate stilbene–stilbene interactions. The closest linkers are not cofacial in these structures, but rather tilted with a 49° dihedral angle with respect to one another. In this case, the MOF exhibits luminescence similar to a dilute solution of the ligand, a situation where LLCT is minimal.

An additional point of interest, exemplified by the stilbene MOF structures just discussed, is that the MOF materials often display increased emission intensity compared with the ligand itself. In the framework structures, the stilbene units are held tightly through strong metal coordination, and their motion is restricted. This inhibits the nonradiative *cis*–*trans* isomerization process that occurs readily for stilbene in solution upon absorption of light, and results in a concomitant increase in the fluorescence intensity of the rigidified ligand in the MOFs [26, 27]. An increase in the radiative lifetime typically leads to an increase in the fluorescence quantum yield by reducing the efficiency of such nonradiative pathways. Biexponential emission decay was observed for the two-periodic Zn stilbene structure, consistent with two emission pathways for the stilbene (i.e., isolated ligand-based emission and LLCT). However, the lower density cubic Zn stilbene MOF exhibited monoexponential emission decay (consistent with isolated stilbene units), with a 0.50 ns lifetime, approximately five times larger than for stilbene in solution. The increase in lifetime is indicative of the rigidity provided by incorporation into the MOF structure. As minor structural changes can lead to observable changes in luminescence properties, this allows for a natural extension of ligand-based luminescence into the field of sensing (Section 12.6). For instance, upon changing the guest solvent molecules in the cubic Zn stilbene MOF, the emission is found to shift in wavelength. This may be a result of a change in local dielectric environment for the ligand or minor changes in the overall framework structure, such as expansion or compression of the catenated units.

As luminescence can be extremely sensitive to local environment, the study of spectral properties has become a standard method for materials characterization as it is a simple, rapid indicator of structural changes and/or maintenance of structural integrity after MOFs have been subjected to a new environment. For example, recent work has shown that the ethylene bridge of stilbene in the three-periodic cubic Zn MOFs can be partially brominated without collapsing the structure [28]. However, full bromination interrupts the interpenetrated cubic units, leading to partial collapse, as was quickly determined by measurement of the luminescence spectra (revealing a broadened, red-shifted spectrum) and later verified via detailed structural studies. Another interesting aspect, supported by luminescence studies, is the change in stability to evacuation between structures upon bromination. As-prepared stilbene zinc three-periodic MOFs do not maintain their structure after full evacuation, a likely result of a change in the π - π interactions between interpenetrated cubes, which alter the stilbene-stilbene distance and disturbs the structure. This leads to lower than expected surface areas ($560 \text{ m}^2 \text{ g}^{-1}$) and a weakened, red-shifted luminescence with loss of vibronic structure upon evacuation (see Figure 12.4). However, after partial bromination, the inter-cube attractions are reduced and the structure remains intact upon evacuation, with an observed increase in surface area ($1600 \text{ m}^2 \text{ g}^{-1}$) despite the fact that the bromine atoms occupy some of the pore space. The emission spectrum of these partially brominated MOFs is nearly unperturbed from that of the as-prepared three-periodic material before evacuation or bromination, as shown in Figure 12.4 (presumably, the brominated stilbene units are substantially quenched and do not contribute significantly to the spectra).

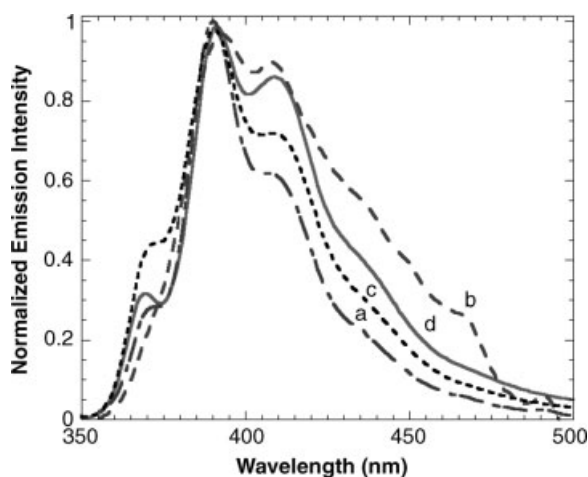


Figure 12.4 Emission spectra of 3D Zn stilbene MOFs with (a) chloroform guest molecules and (b) after full evacuation. The emission spectra of partially brominated Zn stilbene MOFs in (c) a chloroform environment

and (d) after full evacuation reveal the stability of the structure to bromination and subsequent evacuation, as evidenced by their similarity to (a).

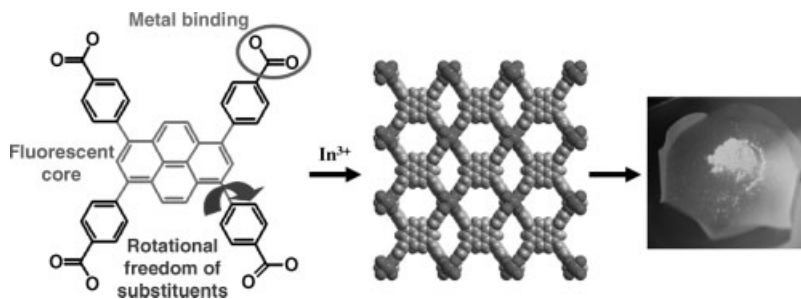


Figure 12.5 $[\text{In}_2(\text{OH})_2(\text{TBAPy})]$ MOFs exhibit bright fluorescence in the extended network. The lifetime and intensity of the emission increases due to strong coordination within the framework structure. Reprinted with permission from [29]. Copyright 2010 American Chemical Society.

A recent study by Rosseinsky and co-workers also clearly demonstrates the direct influence of local ligand environment on the overall emission characteristics for MOF materials based on a luminescent pyrene core, $[\text{In}_2(\text{OH})_2(\text{tbaPy})]_n$ [tbaPy = 1,3,6,8-tetrakis(*p*-benzoic acid)pyrene] [29]. The tbaPyH₄ ligand itself exhibits an emission lifetime of 89 μs, which increases to 110 μs upon incorporation into the MOF, attributed to rigidification. Bright blue luminescence is observed upon coordination of the ligand in the MOF (see Figure 12.5), with a blue shift of 66 nm from that of the free ligand due a reduction of LLCT through isolation of the ligands from one another in the framework. Interestingly, upon evacuation of guest DMF molecules, the emission red shifts, the quantum yield decreases, and the fluorescence lifetime decreases. This is a direct result of the structural distortion which increases the degree of through-space coupling of the pyrene cores, elegantly demonstrated by single-crystal diffraction studies. Upon resolution with DMF, the original structure and luminescence properties are restored. Altering the identity of the guest molecules is also found to cause a change in emission, as will be discussed in Section 12.6.1.1. Other cases of luminescence enhancement due to the rigid environment in MOFs have also been reported [30–33].

12.3.3

Ligand-to-Metal Charge Transfer in MOFs

LMCT-based emission is often reported for Zn^{II} and Cd^{II} MOFs when the sufficiently absorptive ligand has a relatively large HOMO–LUMO gap (as discussed in the previous section), leading to significantly red-shifted emission compared with the ligand itself. Many MOFs displaying LMCT emission contain bdc, benzenetricarboxylate (btc), or both linkers [34–37]. In these cases, green emission is associated with UV excitation and assigned to an LMCT. However, as discussed above, this may be a consequence of an impurity (possibly ZnO). Often, ligand-based luminescence competes with LMCT, and both cases can be observed under certain circumstances. For example, $\{\text{Zn}_3(1,4\text{-bdc})(1,3,5\text{-btc})_2[\text{NH}(\text{CH}_3)_2][\text{NH}_2(\text{CH}_3)_2]_2\}_n$ emits at 430 nm due to LMCT, with a shoulder at 370 nm arising

from emission from the btc ligand [34]. There are also reported examples of LMCT from MOFs containing ligands other than bdc and btc. Subtle variations in the environment can alter the efficiency of energy transfer from the ligand to the metal center, as described by Huang *et al.* for Cd^{II} 2,6-di(4-triazolyl)pyridine-based MOFs [38]. Ligand-based luminescence is observed for the hydrated structure, a result of water in the lattice preventing efficient charge transfer. However, the emission gains significant LMCT character upon dehydration as the charge-transfer process dominates.

The efficiency of LMCT is a function of the orbital overlap between ligand and metal and, therefore, the observation of ligand-based luminescence versus LMCT can also depend on the type of metal center. A recent study of 18 MOF structures containing tetrapyridyl ligands bound to either Zn^{II} , Cd^{II} , or Hg^{II} metal centers demonstrated how subtle variations in the structure can alter the emission characteristics [39]. These materials emit in a multitude of colors, from violet to yellow, depending on the structure type and metal ion. The luminescence intensity for all of the Zn^{II} and Cd^{II} structures was enhanced compared with the free ligand due to the rigidity afforded by coordination in the MOF structure. However, the Hg^{II} structures display reduced emission intensity, attributed to a heavy-metal quenching effect.

In another recent interesting study, the photoluminescence spectra of three structures containing Cd^{II} , 4,4'-bipyridine, and 2-amino-1,4-bdc were found to be characteristic of either a mixture of LMCT and ligand-centered emission, or only LMCT [40]. Upon desolvation and framework distortion, the LMCT character is quenched, while ligand-centered luminescence grows in. Main-group metal complexes may also display LMCT in certain cases, for example $[\text{Pb}_4(1,3\text{-bdc})_3(\mu_4\text{-O})(\text{H}_2\text{O})]_n$ exhibits a strong emission peak at 424 nm originating from LMCT between delocalized π -bonds of the bdc groups and p-orbitals of the Pb^{II} ions [41].

12.3.4

Metal-to-Ligand Charge Transfer in MOFs

MLCT is less commonly reported for MOFs than LMCT, and is typically observed for $\text{d}^{10} \text{Cu}^{\text{I}}$ - and Ag^{I} -based MOFs where there is the possibility of d-electron transfer into low-lying empty ligand states (e.g., for π -acid ligands). For example, *N*-heterocyclic-based Cu^{I} MOFs displaying a multitude of geometries were shown to luminesce with varying degrees of intensity [42] (assigned to a $\text{Cu} \rightarrow \text{CN}$ MLCT process), and $[\text{Cu}(1,2,4\text{-btc})(2,2\text{-bipyridine})]_n$ was found to exhibit intense blue MLCT emission with a lifetime of 13.62 ns [43]. These mixed-valence complexes also exhibit intervalence charge transfer (IVCT).

Excitation of the two-periodic layered MOF $\{\text{Ag}[4\text{-(2-pyrimidylthiomethyl)benzoic acid}]\}_n$ at 370 nm results in an intense green emission with peak maximum at 530 nm, the origin of which was assigned to LMCT and/or MLCT modified by metal-centered (ds/dp) states mediated through Ag–Ag interactions [44]. Similarly, $\{[\text{Ag}(4,4'\text{-bipyridyl})][\text{Ag}(1,2,4\text{-Hbtc})]\}_n$ exhibits an intense emission band with a maximum at 502 nm ($\lambda_{\text{ex}} = 410 \text{ nm}$) due to MLCT [43].

12.4

Metal-Based Luminescence

12.4.1

Metal Luminophores

When discussing metal ions and their salts, only the rare earth elements and some uranyl compounds are typically luminescent. For example, all of the lanthanide trivalent ions (Ln^{III}) except La^{III} and Lu^{III} are luminescent and their emissions span the spectrum from the UV (Gd^{III}) to the visible (Pr^{III} , Sm^{III} , Eu^{III} , Tb^{III} , Dy^{III} , and Tm^{III}) to the near-infrared (NIR) (Pr^{III} , Nd^{III} , Ho^{III} , Er^{III} , and Yb^{III}) region [45]. Some of these elements are fluorescent, whereas others exhibit phosphorescence, although in some instances they can exhibit both. For compounds containing uranium, only the U^{VI} uranyl ion $[\text{UO}_2]^{2+}$ is fluorescent and typically emits in the range 520–620 nm.

12.4.2

Lanthanide Luminescence and the Antenna Effect

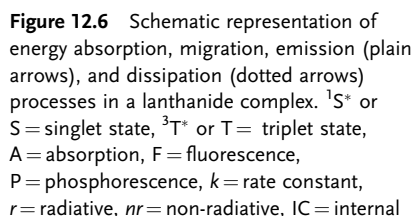
Trivalent lanthanide ions (Ln^{III}) are attractive luminophores because they exhibit spectrally narrow emission with long lifetimes in the solid state and in solution [46]. The 4f orbital is shielded by a filled $5s^25p^6$ sub-shell, thus screening the inner-shell 4f–4f electronic transitions which generate well-defined energy gaps for the electronic levels allowing for the line-like emission. However, electronic transitions for Ln^{III} ions are formally forbidden by parity selection rules (Laporte's rule), typically leading to weak absorbance and, therefore, low brightness. One method of overcoming the weak absorption cross-sections of these ions has been to complex them with strongly absorbing organic molecules. Additionally, complexation often prevents quenching of luminescence by solvent molecules. When efficient vibronic coupling exists between the organic ligand and Ln^{III} moieties, direct energy transfer from the organic molecules' excited state to the metals' elevated energy levels is possible (Figure 12.6). This coupling, first identified by Weissman in 1942, leads to a large increase in luminescence intensity and is widely known as the “antenna effect” or “luminescence sensitization” [45, 47]. For Ln^{III} ions, the electronic dipole transitions are of the same order of magnitude as the magnetic dipole transitions, and therefore both are readily seen in the optical spectra.

12.4.3

Examples of Metal-Based Luminescence

12.4.3.1 Metal-Centered Luminescence

Many isostructural series of Ln^{III} -containing MOFs are known; however, typically, only the luminescence properties of the Eu^{III} and Tb^{III} analogs are reported [48–60]. The likely reason why these two metals predominate in luminescence reports is twofold: they benefit from strong coupling to most organic ligands used for MOF synthesis (antenna effect) and their red and green emission in the visible range is



easier to detect with common laboratory equipment than the longer wavelength emission of several other Ln^{III} ions. In complexes comprised of Eu^{III} , the $^5\text{D}_0 \rightarrow ^7\text{F}_J$ transition series ($J=0-4$) is the most readily observable, typically with the $J=1$ and 2 transitions being both the strongest and most useful for structural determination and sensing. The dominant emission of the $^5\text{D}_0 \rightarrow ^7\text{F}_2$ transition usually occurs around 615 nm, leading to the brilliant red color characteristic of Eu^{III} luminescence. The emissions observed for Tb^{III} come from the $^5\text{D}_4 \rightarrow ^7\text{F}_J$ transitions ($J=3-6$), with the strongest typically being that of $J=5$ and 6 and giving rise to the strong green luminescence in the 540–555 nm range characteristic of Tb^{III} complexes.

An illustrative example of metal-centered luminescence was provided by Li *et al.*, who reported a series of Ln^{III} MOFs based on [Ln₄(OH)₄]⁸⁺ cubane-like secondary building units (SBUs) [61]. The structures, which have the unit formula {[Ln₄(OH)₄(3-sba)₄(H₂O)₄·*n*H₂O]}_{*n*} (Ln = Eu^{III} (**1**), Gd^{III} (**2**), Tb^{III} (**3**); 3-sba = 3-sulfobenzoate), are three-periodic porous frameworks with unique (3,12) topology. The PL of **1** and **3** is characteristic of Ln^{III} metal-centered luminescence. The excitation spectra of **1**, monitored at the strongest Eu^{III} transition (⁵D₀ → ⁷F_{*n*},

614 nm), displays an intense, broad band centered at 286 nm that was attributed to the ligand-centered $\pi \rightarrow \pi^*$ transition, in addition to several much weaker narrow bands attributed to direct f–f transitions within the Eu^{III} metal center ($^7\text{F}_0 \rightarrow ^5\text{H}_4$, 318 nm; $^7\text{F}_0 \rightarrow ^5\text{G}_6$, 362 nm; $^7\text{F}_0 \rightarrow ^5\text{G}_{0-4}$, 377–382 nm; $^7\text{F}_0 \rightarrow ^5\text{L}_6$, 395 nm; $^7\text{F}_0 \rightarrow ^5\text{D}_3$, 416 nm). The emission spectra of **1** ($\lambda_{\text{ex}} = 286$ nm) exhibit several narrow spectral lines assigned as $^5\text{D}_0 \rightarrow ^7\text{F}_j$ ($j = 0-4$) and dominated by the hypersensitive $j = 2$ peak at 614 nm, resulting in bright red luminescence. No emission corresponding to the $\pi^* \rightarrow \pi$ transition of the ligand is observed, indicating that the fluorescence is entirely metal-centered emission. The lifetime of the $^5\text{D}_0$ state was also determined as $\tau = 0.135$ ms, based on the strongest transition. In the case of **3**, the excitation spectrum was monitored at 545 nm ($^5\text{D}_4 \rightarrow ^7\text{F}_5$) and again a strong absorption band at 288 nm was observed and assigned as the $\pi \rightarrow \pi^*$ transition. Additionally, the significantly weaker f–f transitions for Tb^{III} were also present ($^5\text{G}_2$, 351 nm; $^5\text{D}_3$, 359 nm; $^5\text{D}_2$, 369 nm; $^5\text{D}_4$, 377 nm). The emission spectra ($\lambda_{\text{ex}} = 350$ nm) of **3** is dominated by the hypersensitive $^5\text{D}_4 \rightarrow ^7\text{F}_5$ (545 nm with a shoulder at 549 nm), but also displays the remaining transitions from the lowest excited state to the ground-state manifold ($^5\text{D}_4 \rightarrow ^7\text{F}_{6-3}$: $j = 6$, 489 nm; $j = 4$, 528–590 nm; $j = 3$, 622 nm). The lifetime for the $^5\text{D}_4$ state was observed to be $\tau = 1.035$ ms.

In another example of metal-centered emission, Choi *et al.* described electron transfer and size-selective electron oxidation of organic compounds by a MOF based on Eu^{III} and 1,3,5-benzenetribenzoate [62]. When the ligand is selectively excited ($\lambda_{\text{ex}} = 285$ nm), the MOF displays characteristic Eu^{III} luminescence, emitting narrow lineshapes at wavelengths attributed to $^5\text{D}_0 \rightarrow ^7\text{F}_j$ ($j = 1, 2, 3, 4$; 593, 615, 653, 698 nm) with no observable emission from the ligand. This suggests efficient LMCT and metal-centered emission. The authors investigated this material for the ability to facilitate electron transfer from the organic moieties to the Eu^{III} metal ions. Upon irradiation of the MOF, while in the presence of an acetonitrile solution of N,N,N',N' -tetramethyl-*p*-phenylenediamine (TMPD), with strong UV light (365 nm; 50 mW cm^{-2} ; 10 min), the solution turned deep blue, indicative of the formation of the oxidized $\text{TMPD}^{\bullet+}$. This material also showed strong absorption at 565 and 614 nm after irradiation, both in good agreement with the presence of $\text{TMPD}^{\bullet+}$. Furthermore, the authors reported size-selective one-electron oxidation of several other organic compounds together with detailed experimental (time-resolved absorption and emission spectroscopy; confocal microscopy) and theoretical observations (Marcus theory calculations).

An interesting *solvatochromic effect* was reported by Lin *et al.* [63] and provides another example of metal-centered luminescence. They describes a series of porous three-periodic Ln^{III} MOFs, $\{[\text{Ln}_4(\text{bpt})_4(\text{dmf})_2(\text{H}_2\text{O})_8] \cdot (\text{dmf})_5 \cdot (\text{H}_2\text{O})_3\}_n$ [H_3bpt = biphenyl-3,4',5-tricarboxylic acid; $\text{Ln} = \text{Eu}$ (**4**), Gd (**5**), Tb (**6**), Dy (**7**), Ho (**8**), or Er (**9**)]. The ligand H_3bpt was specifically targeted because it was expected to absorb UV light strongly and act as a good sensitizer of Ln^{III} ions, and because other multicarboxylate organic molecules afford extended architectures when coordinated to Ln^{III} ions. Compound **7** displays no observable fluorescence when excited using UV radiation; however, **4** and **6** exhibit characteristic red and turquoise emissions, respectively. The emission of the $\pi^* \rightarrow \pi$ transition seen in the emission spectrum

of the isolated ligand ($\lambda_{\text{ex}} = 300 \text{ nm}$) was not observed for either of the MOF compounds, suggesting efficient sensitization and metal-centered luminescence. Again, the dominant peaks were attributed to the so-called hypersensitive transitions: $^5\text{D}_0 \rightarrow ^7\text{F}_2$ (617 nm) in the case of Eu^{III} and $^5\text{D}_4 \rightarrow ^7\text{F}_5$ (544 nm) for the Tb^{III} compound. Quantum yields were also obtained for these two compounds with $\Phi_4 = 0.1442$ and $\Phi_5 = 0.6087$ obtained for the Eu^{III} and Tb^{III} MOFs, respectively. The high value for the Tb MOF is surprising, particularly given the presence of nine-coordinated water molecules, which one would expect would quench the Ln^{III} luminescence. The compounds were annealed in air at 220°C for 4 h, causing the quantum yields unexpectedly to decrease ($\Phi_{4\text{A}} = 0.052$; $\Phi_{5\text{A}} = 0.326$). The initial quantum yields were only partially recovered upon rehydration ($\Phi_{4\text{B}} = 0.12$; $\Phi_{5\text{B}} = 0.582$), suggesting a possible irreversible change in the compounds upon dehydration and subsequent rehydration. The variation in PL for the as-synthesized, dehydrated, and rehydrated forms of these two Ln MOFs suggests a possible solvatochromic effect in these materials, which is counter to the commonly understood phenomenon of water-quenched luminescence.

The efficiency of energy transfer from absorbing linkers to lanthanide ions within MOFs is not uniform, as shown by a detailed analysis conducted by Soares-Santos *et al.* [64]. They measured the PL of a series of Ln^{III} MOFs, showing that Eu^{III} and Tb^{III} are more efficiently sensitized than their Nd^{III} and Sm^{III} counterparts. In this investigation, 2,3-pyrazinedicarboxylate (2,3-pzdc) and oxalate (ox) combine with Ln^{III} ions to generate two-periodic bilayer structures with unit formula $[\text{Ln}_2(2,3\text{-pzdc})_2(\text{ox})(\text{H}_2\text{O})_2]_n$ (Ln = Ce, Nd, Sm, Eu, Gd, Tb, or Er). In this structure, individual cationic layers interconnect via ox^{2-} . In the Nd^{III} and Sm^{III} cases, the more intense spectral peaks (Nd , $^4\text{F}_{3/2} \rightarrow ^4\text{I}_{11/2}$; Sm , $^4\text{G}_{5/2} \rightarrow ^6\text{H}_{9/2}$) were used to monitor the excitation; several narrow peaks corresponding to transitions from the ground states (Nd , $^4\text{I}_{9/2}$; Sm , $^6\text{H}_{5/2}$) to higher energy levels were observed. Additionally, intense, broad absorption bands from 250 to 400 nm attributed to ligand $\pi \rightarrow \pi^*$ transitions were observed for both MOFs. This ligand absorption is stronger than for the metal transitions, suggesting more efficient sensitization via ligand excited states than for direct excitation of metal transitions. The emission spectra for both the Nd^{III} and Sm^{III} compounds were characteristic of their Ln^{III} ions, with emissions in the NIR region (e.g., for the Nd^{III} compound the strongest observed peak was for $^4\text{F}_{3/2} \rightarrow ^4\text{I}_{11/2}$ at 1064 nm). However, both of these compounds also exhibit large and broad emission bands from 380 to 540 nm, indicative of $\pi^* \rightarrow \pi$ emission from the ligands, suggesting incomplete LMCT. In the case of Gd^{III} , the excited energy levels of the metal ions are higher than that of the S_n or T_n energy levels of the ligand, precluding the possibility of energy transfer from the ligand to the metals. As such, the emission for this compound was attributed solely to the two ligands, and through detailed analysis the authors were able to determine that the emissions are due to the excited triplet states of the two ligands. Additionally, they determined the relative energies for the two triplet states and deduced that 2,3-pzdc is a better sensitizer than ox. In the case of the Tb^{III} and Eu^{III} compounds, PL similar to the other materials described in this section is observed. The excitation spectra exhibit intense, broad bands attributed to $\pi \rightarrow \pi^*$ transitions centered on the ligands and also

weaker and much narrower peaks assigned to transitions from the ground state to higher energy excited states for the Ln^{III} ions, while the emission spectra are characteristic of the respective Ln^{III} ions and show no observable transitions from the ligands. This suggests that either the efficiency of ligand-to-metal energy transfer is higher in the cases of Tb^{III} and Eu^{III} than those of Nd^{III} and Sm^{III} , or that there is some nonradiative deactivation of ligand excited states in the former two cases not observed in the latter.

As a final example of metal-centered luminescence, Li *et al.* described a group of MOF materials in which LMCT can be essentially “switched off” and either ligand-centered or metal-centered emission is observed based on the selection of metal ion [65]. They reported a series of MOFs constructed from the same ligand, 2-(pyridine-4-yl)-1*H*-imidazole-4,5-dicarboxylic acid (H_3pidc), but with lanthanide or non-lanthanide metal ions (Fe^{II} , Cd^{II} , Zn^{II} , Eu^{III} , Tb^{III} , and Y^{III}). In the compounds containing Cd^{II} , Zn^{II} , and Y^{III} , the observed emission is broad and slightly blue shifted relative to the free ligand, suggesting only ligand-centered emission. However, in the compounds containing Tb^{III} and Eu^{III} , no broad ligand-centered emission is observed, while sharp, narrow emissions characteristic of the Ln^{III} ion are detected. In this study, we see that the presence of Ln^{III} ions permits efficient LMCT, preventing the $\pi^* \rightarrow \pi$ intraligand transitions while at the same time generating metal-based excited states, leading to Ln^{III} emission. When the Ln^{III} ions are replaced with d^{10} metals, no LMCT occurs and the PL is attributed solely to ligand-centered emission.

12.4.3.2 Metal-to-Metal Charge Transfer (MMCT)

The potential for energy transfer between metals within a MOF also exists. In one report, two isostructural 4,4'-bipy templated frameworks were prepared using Tb^{III} and 1:1 $\text{Eu}^{\text{III}}\text{--Tb}^{\text{III}}$ [66]. Both of these structures exhibit efficient through-space sensitized luminescence originating from the lanthanide centers. However the compounds containing $\text{Eu}^{\text{III}}\text{--Tb}^{\text{III}}$ show greater sensitization efficiency than the structure comprised solely of Eu^{III} . The quantum yields for the materials were reported to be $\Phi_{\text{Eu}} \approx 0.22$ and $\Phi_{\text{Tb}} \approx 0.32$. Perhaps even more interesting, the mixed-metal MOF exhibits a metal-to-metal antenna effect in which the luminescence expected from the Tb^{III} luminophore is nearly completely quenched, leaving an emission spectrum for the material dominated by the Eu^{III} luminophore. Due to the absence of the Tb^{III} emission, and also the increased quantum yield ($\Phi_{\text{Eu/Tb}} \approx 0.39$ compared with $\Phi_{\text{Tb}} \approx 0.32$), it is likely that energy transfer from the higher energy Tb^{III} $^5\text{D}_4$ state to the Eu^{III} $^5\text{D}_0$ state occurs.

Metal-to-metal charge transfer could also be useful as a structural probe. Soares-Santos *et al.* described a series of mixed-ligand lanthanoid frameworks constructed from 2,5-pyridinedicarboxylate and 1,4-phenylenediacetate along with Eu^{III} , Tb^{III} , and two non-equivalent mixtures of Eu^{III} and Tb^{III} [67]. Each of these MOFs demonstrates sensitized metal-centered luminescence through an antenna effect, while the mixed-metal MOFs exhibit Tb^{III} -to- Eu^{III} MMCT. In the mixed-metal frameworks, this energy transfer can serve as a structural probe by indicating the extent of metal ion dispersal throughout the material. MOFs having single-metal ion

clusters would be expected to exhibit very little of this Tb^{III}-to-Eu^{III} MMCT. Conversely, when the metal ions are homogeneously dispersed throughout the structure, the Tb^{III}-to-Eu^{III} MMCT should be efficient and easily observed.

12.4.4

Lanthanide Luminescence as a Probe of the Metal-Ligand Coordination Sphere

An illustrative example of how lanthanide luminescence can be used as a structural probe of the metal-ligand coordination sphere is that of Eu^{III}. The ratio of relative intensities for the $^5D_0 \rightarrow ^7F_J$ ($J=2:J=1$) transitions is very sensitive to the site symmetry of the Eu^{III} centers. Since the electric dipole transition ($J=2$) is hypersensitive to the ligand environment whereas the magnetic dipole transition ($J=1$) is practically insensitive, it is sometimes possible to elucidate some aspects of symmetry for the material even in the absence of other structural data [68]. This effect is also observed in Ln-based MOFs, particularly an Eu^{III}-doped version of MIL-78 [69], an open-framework structure comprised of one-periodic inorganic columns of eight-coordinate polyhedra connected by bridging trimesate linkers. Luminescence data depict a nearly equivalent intensity for the $^5D_0 \rightarrow ^7F_2$ and $^5D_0 \rightarrow ^7F_1$ transitions ($^7F_2:^7F_1 \approx 1$), suggesting that the Eu^{III} ions lie in an environment of high symmetry relative to the ligand, which was independently confirmed by single-crystal X-ray diffraction (XRD) data. In another example, a MOF structure exhibiting chirality shows the ability of Ln^{III} luminescence to be used as a structural probe. Yue *et al.* synthesized a group of homochiral Ln phosphonate frameworks with the enantiomerically pure ligand *s*-*N*-(phosphonomethyl)proline and several lanthanide ions (Ln = Tb, Dy, Eu, Gd) [59]. The chirality of the MOF arises from the chiral organic ligand, leading to a MOF structure comprised of a series of one-periodic triple-stranded helices. PL spectra for both the Tb^{III} and Eu^{III} analogs exhibit significant splitting in the $^5D_4 \rightarrow ^7F_J$ series ($J=6-4$), indicative of a strong crystal field that results from the asymmetric environment of the metal center. While symmetric systems often have forbidden transitions, the asymmetry inherent in these MOF structures leads to high intensity ratios for the $^7F_2:^7F_1$ transitions, and also 7F_0 transitions that are more intense than even the 7F_1 transition, an unusual occurrence.

12.5

Guest-Induced Luminescence

In the previous two sections, we have discussed the phenomenon of luminescence as it arises directly from the MOF itself, from either the organic linker or the inorganic cluster. However, many MOFs have periodic scaffolds that generate well-defined, essentially permanent, nanopores and large free-volume cavities. These unfilled cavities of controllable size allow the encapsulation of a wide variety of guest molecules. In relation to MOF-based luminescence properties, the ability to encapsulate guest molecules could prove useful in a number of ways. For instance, a non-luminescent MOF could be infiltrated with a luminescent guest molecule,

which lends luminescent properties to the hybrid MOF–guest material, thereby imparting desired luminescent properties into the MOF. It may also be possible that upon infiltration of the MOF, a new luminescent species is formed through MOF–guest interactions (e.g., exciplex) with different luminescence properties from that of either the pristine MOF or guest molecules by themselves. In this case, both the MOF and guest may or may not be luminescent on their own, but upon complexation they generate a new species capable of luminescence. Additionally, the inclusion of a guest molecule could lead to modulation of the inherent luminescence of the MOF host. It may then be possible to enhance (or quench) the luminescence of the MOF by incorporating specific guest molecules, thereby allowing the MOF to act as a sensor.

12.5.1

Encapsulation of Luminophores

Conceptually, the infiltration of a porous, non-luminescent MOF with a luminescent guest molecule is the simplest way of introducing desired luminescence properties in a MOF material. Müller *et al.* synthesized a selection of well-studied MOFs [MOF-5, MOF-177, UCM-1, and MIL-53(Al)] and investigated the ability to incorporate strongly luminescent guest molecules into these frameworks via the vapor phase [70]. The luminophores they chose were *N,N*-bis(2,6-dimethylphenyl)-3,4,9,10-perylene-tetracarboxylic diimide (dxp) and (2-carboxypyridyl)bis[3,5-difluoro-2(2-pyridyl)phenyl]iridium(III) (F₂-ppy), and in each case, the infiltrated MOF exhibited PL spectra dominated by the luminophore guest. The authors attempted to remove the luminophores by washing the hybrid materials and observed that, for the F₂-ppy compounds, partial to total removal was possible, coupled with decreases in PL intensity. However, removal of dxp was not achieved, possibly indicating stronger interactions with the frameworks. The emissive properties of the dxp-containing MOFs suggest host–guest interactions such as caging effects, aggregate formation, and strong quenching of the MOF host luminescence.

In another example, this time involving infiltration of charged dye molecules, Zhang *et al.* reported the synthesis of nanoscale Ln^{III} MOFs (Ln = Gd, Eu, Yb) that exhibit temperature-controlled guest encapsulation and light harvesting [71]. The MOFs are constructed from highly conjugated linear linker molecules, and when the coordination spheres of the Ln^{III} ions are filled with non-bridging anions such as acetate, the lanthanide clusters are negatively charged. This allows infiltration of the MOFs with cationic dye molecules (*trans*-4-styryl-1-methylpyridinium iodide and methylene blue), upon which the MOF-based luminescence is quenched and FRET emission is observed. The intensity and efficiency of FRET are controllable based on the temperature of the MOF synthesis, which was shown to correlate directly with the percentage loading of the guest luminophore. Interestingly, the dye and linker together in solution (DMF) demonstrated luminescence quenching, but no energy transfer, suggesting that the highly ordered arrangement of the dye guest molecules within the MOF cavities favors light harvesting. This study is significant, not only for the sequestration of a luminophore within the cavities, but also because it demon-

strates that MOF–guest interactions can have profound effects on the resultant PL properties.

It is also possible to use the presence of a guest molecule to quench the luminescence of a MOF efficiently. Park *et al.* reported the structure and luminescence spectra of a mesoporous MOF derived from a superstructure of Tb^{III} ions, triazine-1,3,5-(4,4',4''-trisbenzoate), and dimethylacetamide [72]. The crystal structure reveals large pores (3.9 and 4.7 nm in diameter) and, upon activation, the material shows a Langmuir surface area of $3855 \text{ m}^2 \text{ g}^{-1}$. PL studies demonstrated bright green, sensitized luminescence characteristic of the Tb^{III} ions. To investigate the potential for uptake of guest molecules, vapor-phase ferrocene was introduced to the structure using elevated temperatures and reduced pressure. Figure 12.7 shows several transmission and fluorescence microscopy images, and also luminescence spectra for the as-synthesized and ferrocene-containing framework. A color change of the single crystals from colorless to dark brown, in addition to loss of Tb

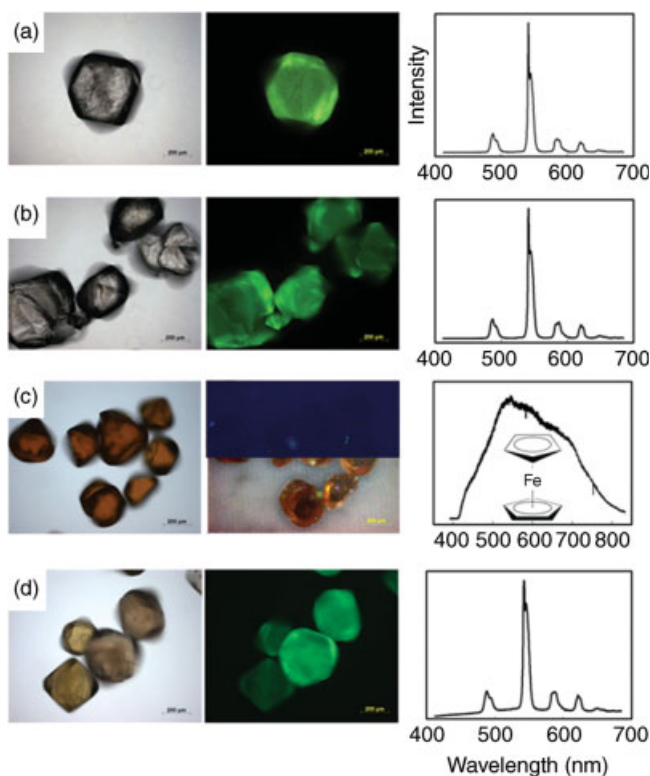


Figure 12.7 Transmission (left) and fluorescence (middle) microscope images and luminescence spectra (right) for a) a single MOF, b) multiple MOFs, c) ferrocene included MOFs, and d) regenerated initial MOFs after removal of ferrocene via evacuation at 50°C for

1 day. In the fluorescence image of c), half of the background color was subtracted to show the crystals. [72] Copyright Wiley-VCH Verlag GmbH & Co. KGaA. Reproduced with permission.

luminescence, are indicative of ferrocene inclusion. Additionally, the authors demonstrated that extraction of the ferrocene moieties is possible by using vacuum and high temperature, and that upon ferrocene removal, the framework luminescence is re-established. Elemental analysis also indicated a high uptake of ferrocene within this MOF.

12.5.2

Guest-Induced Charge Transfer: Excimers and Exciplexes

One way of introducing or modulating the luminescence of a MOF material is to generate an entirely new luminescent species in the form of an excited-state complex. When the two molecules in question are the same, they form a homodimer excited-state complex or *excimer* (excited dimer). In the case where the two molecules are different, the excited-state complex is a heterodimer called an *exciplex* (excited complex)[16]. Formation of exciplexes or excimers in MOFs is possible by two routes: (1) as a result of the framework structure itself or (2) by infiltration with a guest molecule. For the first possibility, the relative orientation of framework linkers allows the formation of an excited-state complex upon absorption of a photon. This could be achieved as a result of the framework topology itself; by “breathing” in a flexible framework that alters the structure and brings the ligands into closer contact (and in the correct orientation); or by framework interpenetration, whereby the two linker molecules are contained on different nets, but located adjacent to one another. For the second possibility, infiltration of the framework with an appropriate guest molecule, and the subsequent interaction of this molecule with the ligands, could result in an excited-state complex. In both the exciplex and the excimer, the effect on the observed luminescence is the formation of typically broad and featureless emission spectra, which are noticeably red shifted in comparison with the individual monomer(s). The exciplex or excimer emission is highly dependent on the orientation of the component molecules, so that formation of these excited-state species and observation of their PL can not only provide information on the presence of a guest molecule (sensor), but can also provide structural information regarding the interaction between the host and guest (structural probe). Cofacial arrangements of the two molecules are generally believed to be favorable to excimer and exciplex formation, and electronic structure modeling suggests that the strength of the interaction is strongly dependent on the separation distance [73].

Exciplexes have been reported in MOFs on only a few occasions. Perhaps the most extensive investigation was conducted by Zaworotko and co-workers, who synthesized several MOFs in which aromatic molecules are intercalated within the framework to allow exciplex formation. In one case, 4,4'-bipyridine (4,4'-bipy) and pyrene form a one-periodic molecular ladder coordination polymer with the unit formula $\{Zn(4,4'\text{-bipy})_{1.5}(NO_3)_2 \cdot CH_3OH \cdot 0.5\text{pyrene}\}_n$, in which the 4,4'-bipy linker assumes a cofacial arrangement with the intercalated pyrene molecule [74]. The pyrene moieties are not coordinated to the zinc ions, but rather trapped within the matrix to form a 2:1 4,4'-bipy–pyrene exciplex. The ratio of relative intensity between the third and first peaks within the vibronic structure of the weak monomer

fluorescence band for pyrene can be used as a probe for the polarity of the micro-environment surrounding the pyrene [16]. The results suggest the possibility of using both excimer formation and the emission of the monomer as an internal probe of adsorbed guest (solvent) molecules. In another example [75], the same group prepared a series of Zn^{II} 4,4'-bipy-MOFs having one-periodic ladder and two-periodic square grid topologies. In these examples, the nature of the aromatic solvent molecule (benzene, chlorobenzene, *o*-dichlorobenzene, *p*-xylene, and toluene) determines the topology of the structure that results. Both λ_{max} and τ_{F} of the fluorescence are influenced by the structure. The emission of the one-periodic structures is red shifted ($\lambda_{\text{max}} \approx 520$ versus 460 nm) in comparison with the two-periodic emission and displays shorter lifetimes ($\tau_{\text{F}} \approx 43$ versus 74 ns). These features of the exciplex luminescence lead to a structural interpretation of face-to-face π - π stacking in the one-periodic structures, but for an edge-to-face C-H $\cdots \pi$ interactions in the two-periodic structure. This work is the most detailed analysis to date of the exciplex phenomenon in MOFs and demonstrates the way in which MOFs can be implemented as a platform to perturb systematically the identity and orientation of exciplex components. Conversely, the study also reveals how subtle changes in the MOF can lead to dramatic changes in the luminescence properties, demonstrating their potential use as good diagnostic handles for MOF structure.

Separately, another research group has demonstrated a case of guest-influenced luminescence in a MOF that may be related to exciplex formation. Stylianou *et al.* described the synthesis of a highly emissive MOF constructed from a strongly fluorescent linker ligand with an electronically isolated pyrene core [29] (see Section 12.3.2). For this MOF, they described solid-state PL properties in which the emission is solely the result of the ligand, which is largely isolated from other ligands or possible quenching moieties. The desolvated MOF can also be resolvated using other solvents, such as H_2O , dioxane, and *p*-xylene, leading to slightly altered PL. Interestingly, whereas the introduction of H_2O and dioxane produces small, but expected, red shifts in λ_{max} , the introduction of *p*-xylene leads to a blue-shift (~ 19 nm). The authors also presented PXRD data that seem to suggest that some of the *p*-xylene moieties are located in a face-to-face π - π stacking interaction with the pyrene core, indicating possible exciplex formation. However, the hypsochromic nature of the shift, together with the fact that the emission spectrum was observed to be no broader than the emission spectra of either the desolvated MOF or the MOFs resolvated with other solvents (DMF, H_2O , or dioxane), make classification of this luminescence emission as an example of exciplex formation in a MOF questionable and undetermined at present.

12.5.3

Encapsulation of Lanthanide Ion Luminophores

In addition to organic molecules, it is also possible to infiltrate MOFs with metal ions to act as guest luminophores. A classic strategy for the fabrication of optically active materials is to dope a compound with luminescent lanthanide ions. Recently, this strategy was shown to be a viable option for modulating MOF

luminescence. Although there are numerous examples of MOFs based upon lanthanides incorporated into the framework and corresponding characterization of these materials' lanthanide-based luminescence, there are relatively few examples of MOFs infiltrated with lanthanide guests. Luo and Batten reported doping a non-lanthanide MOF with Ln^{III} cations and presented data illustrating the ability to affect the luminescence of the material [76]. In this example, a Zn^{II} MOF constructed from 1,2,4,5-benzenetetracarboxylate (H_4btec) generates permanent one-periodic channels. The overall framework structure, $\{(\text{NH}_4)_2[\text{Zn}(\text{btec})]\cdot 6\text{H}_2\text{O}\}_n$, is anionic and the channels are filled with counterbalancing NH_4^+ cations. Subjecting the MOF to solutions of Tb^{III} [$\text{Tb}(\text{ClO}_4)_3$; 10^{-3} – 10^{-6} mol L^{-1}] or Eu^{III} (EuCl_3 ; 10^{-3} – 10^{-7} mol L^{-1}) results in simple cation exchange. The authors demonstrated that the luminescence of the MOF can be tuned, based on the choice of Ln^{III} dopant utilized. The as-synthesized MOF exhibits blue luminescence characterized as LMCT since the ligand itself was shown to be non-luminescent. Upon infiltration with dopants, the blue emission is no longer observed; however, strong emission characteristic of the Ln^{III} ions is generated. Additionally, it was demonstrated that the extent of infiltration plays a role in the intensity of the emission. The MOF hybrids obtained by infusing with a higher concentration of Eu^{III} are 30 times more intense than those obtained from the lower concentration. In the case of Tb^{III} , the observed intensity difference is only eight times more in the case of higher concentration.

Thirumurugan and Cheetham described a series of MOFs constructed with bismuth (Bi^{III}) in which they introduced Ln^{III} ions to modulate the luminescence of the compounds [77]. Owing to similar charge and ionic radii, bismuth-containing compounds are known to behave as good hosts for Ln^{III} ions, and this knowledge has been useful for the development of doped optical materials. Upon subjecting the MOFs to solutions of Eu^{III} and Tb^{III} , the infiltrated materials (~ 2 mol% Ln^{III}) exhibit PL unique from that of the pristine as-synthesized MOF. The neat MOF material has a broad emission centered at ~ 420 nm that was characterized as ligand-centered $\pi^* \rightarrow n$ or $\pi^* \rightarrow \pi$ emission. After doping with Ln^{III} ions, the emission spectra for the hybrid material exhibit characteristic peaks for Eu^{III} ($^5\text{D}_0 \rightarrow ^7\text{F}_j$) and Tb^{III} ($^5\text{D}_4 \rightarrow ^7\text{F}_j$), while the main ligand-centered emission is almost completely suppressed, resulting in red and green luminescence, respectively. Additionally, the authors reported co-doping of the MOFs, using a combination of both Eu^{III} and Tb^{III} (~ 2 mol% each), which results in an overall orange-colored emission and clearly demonstrates the extent to which the luminescence of these materials can be fine-tuned.

Another example demonstrating the effect that encapsulating lanthanide ion luminophores within a MOF has on the PL was reported by Wang *et al.* [78]. They prepared a lanthanide-based MOF capable of encapsulating hydrated Ln^{III} cations by using a bent 1,2,4-trizole-bridged ligand. The encapsulated cations are confined within small volumes as they are hydrogen bonded to the framework, reducing vibrational movements originating from strong O–H oscillators. These vibrational movements normally lead to quenching of the f–f luminescence, but their reduced contribution in these materials leads to enhanced luminescence

intensity, converting the practically non-luminescent $[\text{Ln}(\text{H}_2\text{O})_8]^{3+}$ into a strongly emissive species.

12.6

Applications of Luminescent MOFs

12.6.1

Chemical Sensors

Efficient, portable sensors are increasingly required to support the development of technologies for many fields, from medicine to homeland security, and the number of reports regarding the use of MOFs for such purposes is steadily growing. The ability to tune the emissive properties of MOFs via alteration of the ligand and/or metal cluster type, along with the potential for high porosity with *accessible*, uniform pores, makes MOFs inherently multifunctional materials. In addition, the precise chemical functionality and size of the pores may be selectively tailored, allowing for specific sensing and separation capabilities. These features set MOFs apart from conventional porous materials such as porous silicas and zeolites, which offer only limited degrees of synthetic tunability. As the synthetic strategies for preparing “designer” MOFs improves, so does the ability to create MOFs with functional pores. A recent review by Chen *et al.* summarized developments in MOF materials for molecular recognition and sensing, including MOFs that can act as molecular sieves and effectively separate gases [79]. Jiang *et al.* demonstrated that MOFs can be utilized as stationary phases for liquid chromatography [40]. Additionally, the use of post-synthetic methodologies to alter MOF structure is of recent interest in this context [28, 80–84].

As was discussed in previous sections, the emission properties of solid-state materials are often highly dependent upon details of their structure, which can potentially be tailored via careful selection of the starting materials and growth conditions. Altering emission properties through “simple” changes in host–guest chemistry, such as solvent exchange, has been demonstrated for a variety of porous structures, as discussed in Section 12.3. This may lead to solvatochromic-like effects (although, of course, the guest molecule does not “dissolve” the MOF) that are caused by altering the dielectric environment of the chromophore, resulting in different emission wavelengths, intensities, or both. Changing the solvent molecule does not just alter the dielectric environment, but may also lead to small structural changes, such as lattice expansion or contraction, changes in ligand orientation, or changes in ligand–ligand distance, which can lead to observable differences in luminescence via LLCT or excimer formation. Such spectral shifts are typically relatively small compared with cases in which changing the solvent results in inhibition of LMCT versus ligand-based emission, for example. However, these early findings suggest that MOFs could be used and tailored for specific luminescent sensing capabilities [21]. Solvent incorporation may also result in major changes to the framework structure. For example, Zhu *et al.* reported a lanthanide MOF structure in which

solvent incorporation or removal leads to reversible structural collapse, whereby luminescence is quenched and restored upon a dehydration–rehydration cycle [85]. A similar observation has been reported for one of the rare examples of luminescent MOFs from which emission is exclusively d-metal based, $[\text{Cu}_4\text{I}_4(\text{dabco})_2]_n$ (dabco = 1,4-diazabicyclo[2.2.2]octane). Two different structures were prepared in water and in acetonitrile [86], and the luminescence was found to vary depending on the solvent used. The MOF prepared in water shows temperature-dependent luminescence and emits at 556 nm, whereas the acetonitrile-incorporated MOF shows little temperature dependence and emits at 580 nm. These structures can be reversibly interconverted by solvent exchange. The emission here is cluster-centered, and is a function of the Cu–Cu distance.

It should be noted that the examples in the previous paragraph were concerned with the effects of incorporated solvent on framework luminescence. There are many examples in the literature where this has been observed, and it may be considered to be an expected feature of significantly porous, luminescent structures. In the following sections, a few such effects will be highlighted, but the focus is on the design and use of MOFs for targeted sensing purposes. Although luminescence is typically considered among the more sensitive and attractive for optical sensing methods, an elegant study by Lu and Hupp demonstrated the first use of MOFs for sensing via changes in refractive index [87]. A Fabry–Pérot device fabricated from thin films of ZIF-8 (a zeolitic imidazolate framework) was found to be sensitive to various chemical vapors and gases, with analyte detection readily achievable through transmission UV–visible measurements.

12.6.1.1 Small-Molecule and Ion Sensors

Sensing of metal ions and small molecules by emission can occur by a variety of mechanisms. Ion exchange has been used to alter sensitization in several lanthanide MOFs. Chen and co-workers reported a $\text{Tb}(1,3,5\text{-btc})$ structure in which anionic guests solvated in methanol were incorporated, yielding anion-filled pores [88]. The luminescence from Tb^{III} was found to be enhanced, particularly in the presence of fluoride anion. This enhancement was attributed to a stronger hydrogen-bonding interaction between F^- and methanol leading to attenuation of the O–H stretching vibration, with concomitant reduction of the ability of this vibration to quench the sensitizing btc ligand. Lu and co-workers demonstrated a MOF that displays sensitivity to cation exchange [89]. Here, a Tb^{III} -based MOF exhibited a significant increase in emission intensity upon exchange of K^+ with Ca^{2+} . This was ascribed to rigidification of the imidazole-4,5-dicarboxylate ligand upon complexation with Ca^{2+} , which leads to more efficient energy transfer to the lanthanide ion. Wong *et al.* reported a $\text{Tb}(\text{mucicate})$ structure, wherein the flexible multidentate ligand is used to differentiate between several anions, with fluorescence enhancement in the case of CO_3^{2-} and CN^- [90]. Lewis basic pyridyl sites within a luminescent porous $\text{Eu}(\text{pyridine-3,5-dicarboxylate})$ MOF were utilized for sensing of Cu^{II} ion [91]. This material shows significant quenching upon binding to Cu^{II} . Finally, luminescent intensity modulation was also shown for two Eu^{III} 1,4,8,11-tetraazacyclotetradecane-1,4,8,11-tetrapropionate-based MOFs, whereby

Ag^I exchange significantly alters the relative transition intensities of the emissive Eu^{III} center [92].

Ion sensing is also possible using transition metal-based frameworks, typically through alteration of LMCT efficiency. For example, a net-type anionic MOF, $\{(\text{NH}_4)_2[\text{Zn}(1,2,4,5\text{-benzenetetracarboxylate})]\cdot 6\text{H}_2\text{O}\}_n$, exhibits broad luminescence centered at 440 nm, as a likely result of LMCT and/or ligand-based luminescence [93]. The open structure allows for exchange between the ammonium groups and aqueous metal cations. Titration with increasing amounts of Cu^{II} leads to quenching as a result of ion exchange, and this effect was shown for a variety of other cations in addition. Upon ion exchange with Ln^{III} ions [76], LMCT character was no longer observed, replaced by emission typical of the lanthanide ion instead. These lanthanide-incorporated structures were subsequently doped with MCl_x solutions (M = Na⁺, K⁺, Zn²⁺, Ni²⁺, Mn²⁺, Co²⁺, Cu²⁺), and significant quenching was seen for Co^{II} and Cu^{II} ions.

MOFs may be prepared in which the metal center remains unsaturated, which leads to accessible, potentially luminescent, open metal sites for the sensing of small molecules. The degree of saturation of lanthanide ions in MOFs can often be manipulated via ligand exchange. For example, a porous Eu(1,3,5-btc) MOF with open metal coordination sites shows sensitivity to a variety of solvent molecules, particularly a marked decrease in emission intensity upon exposure to acetone and an increase with DMF exchange [94]. Exchange of coordinated solvents has been shown to alter emission properties in other lanthanide-based MOFs as well. A Eu^{III} MOF with 4,4'-(hexafluoroisopropylidene)bis(benzoic acid) linkers has been used to sense ethanol in air [95]. The Eu^{III} emission at 619 nm was monitored upon exposure, leading to a rapid decrease in luminescence intensity, which was recovered upon exposure to air. The initial quenching was attributed to an effect of the O–H stretching vibration upon coordination of ethanol to the Eu^{III} ions. Harbuzaru *et al.* also demonstrated similar use of an Eu^{III} MOF containing a 1,10-phenanthroline-2,9-dicarboxylic acid linker for pH sensing applications in the biologically suitable range of pH = 5–7.5 [96].

Interruption of LMCT and MLCT processes by small molecules can lead to striking differences in luminescence properties. Jiang and co-workers prepared Cd^{II} MOFs with the same building blocks, but with differing porosity and ligand–ligand separation [40, 97]. Through alteration of the solvent volume and/or temperature, the framework porosity and LMCT signal strength were found to vary reversibly. Kobayashi *et al.* prepared a Pt/Zn-containing 5,5'-dicarboxy-2,2'-bipyridine-based MOF, which exhibits thermochromism and reversible sensitivity to solvent vapors [98]. The thermochromic effect is a result of dehydration of the Pt^{II} centers and the resultant change in Pt–Pt distance, which in turn affects the triplet-sensitized MLCT process. The authors also noted solvatochromic effects with a mechanism similar to this thermochromic effect, whereby various solvents alter the intermetallic Pt distance.

Manipulation of ligand-based luminescence in MOFs has been shown in a variety of cases. As described in Section 12.3.2, Rosseinsky and co-workers prepared $[\text{In}_2(\text{OH})_2(\text{tbaPy})]_n$, which displays a red-shifted emission spectrum upon desolva-

tion as a result of a reduced intermolecular ligand–ligand distance [29]. However, the spectral position and intensity nearly recover fully within 3 h upon re-exposure to DMF. The intermolecular coupling between ligands is highly distance dependent at short range, and addition of various guests leads to alteration in the position and intensity of the pyrene fluorescence. The authors were able to correlate fluorescence changes directly with structural changes, and noted that the long lifetime and quantum yield of 6.7% make these MOFs comparable to commercial lanthanide sensors. As an additional recent example, $\{[\text{Zn}_3(1,3,5\text{-btc})_2]\cdot 12\text{H}_2\text{O}\}_n$ frameworks were found to be selective for ethylamine over butylamine and propylamine, exhibiting significant emission quenching of btc-based luminescence upon increasing doses of ethylamine [99].

12.6.1.2 Oxygen Sensors

Phosphorescent MOFs functionalized with Ir^{III} -containing ligands have been synthesized as potential oxygen sensors [100]. A variety of Zn MOFs were prepared in which the ligands were Ir(2-phenylpyridine)₃ derivatives (see Figure 12.8). Structures 11 and 12 were nonporous; however, structure 10 exhibits permanent porosity and

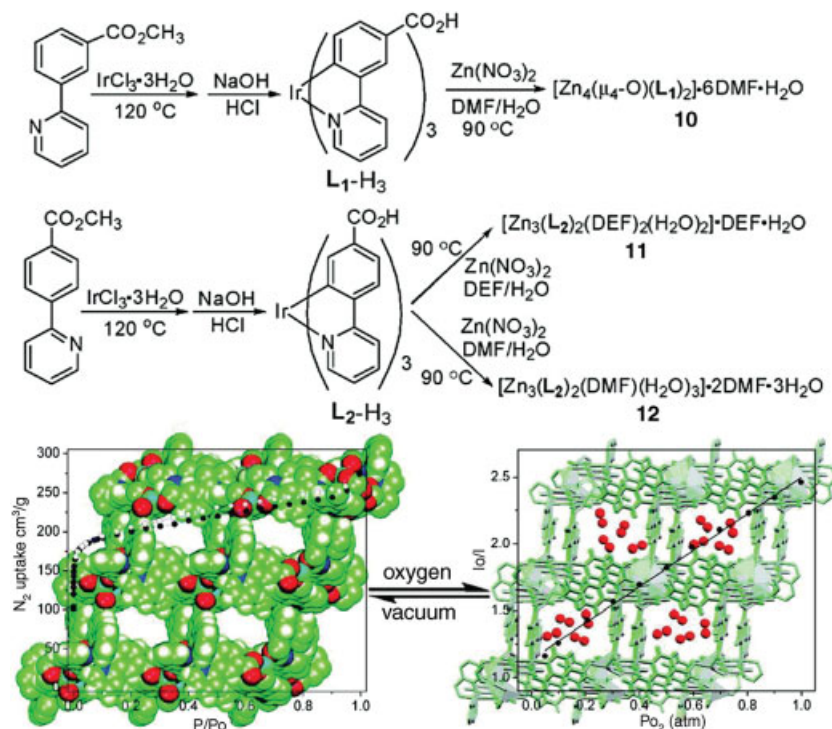


Figure 12.8 (Top) Three structures were synthesized from Ir tris(2-phenylpyridine)-based ligands. 1 was found to be porous, showing rapid and reversible luminescence

quenching by oxygen, as shown schematically (bottom). Reprinted with permission from [100]. Copyright 2010 American Chemical Society.

reversible host–guest chemistry. Triplet-sensitized MLCT was observed from the Ir(2-phenylpyridine)₃ derivatives, at 538 nm for **10** and **12**, and at 565 nm for **11**. It is well known that dioxygen rapidly quenches triplet excited states and, as expected, a decrease in the emission intensity was observed for all three structures in the presence of O₂. Interestingly, upon exposure to O₂, only the porous structure was found to respond rapidly and reversibly. A linear Stern–Volmer plot (see Section 12.2.2; Equation 12.5) was extracted for exposure of this porous MOF to O₂, and the luminescence quenching was shown to be reversible across multiple cycles. The nonporous MOFs have limited accessibility to the interior Ir moieties, and it also is more difficult to remove the O₂ molecules from these structures. Nonetheless, **11** and **12** were also observed to display a degree of reversibility on longer timescales than **10**.

12.6.1.3 Detection of Explosives

Detection of explosives is a topic of significant contemporary relevance, and substantial work has been carried out in this area. Efficient, inexpensive detection of 2,4-dinitrotoluene (DNT) [a volatile indicator of 2,4,6-trinitrotoluene (TNT)] and 2,3-dimethyl-2,3-dinitrobutane (DMNB, an ingredient in plastic explosives), are of high interest in this regard. Typically, luminescent changes upon a redox reaction are used in current detection methods, although this is much more difficult to achieve for DMNB. The first example of a MOF that possesses the ability to detect both markers has been reported recently [101]. The structure [Zn₂(bpdc)₂(bpee)]_n, (bpdc = 4,4'-biphenyldicarboxylate; bpee = 1,2-bipyridylethene) was shown to absorb both molecules of interest quickly (on the order of 10s), reversibly, and efficiently, with their uptake detected through changes in the emission properties of the MOF. Thin layers of the MOFs were exposed to vapors of DMNB and DNT, upon which red-shifted, quenched emission was rapidly observed, as shown in Figure 12.9. Both analytes

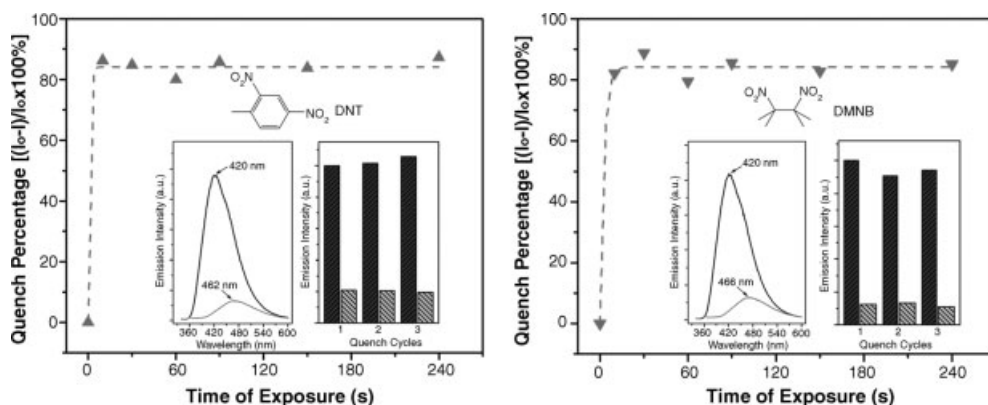


Figure 12.9 Time-dependant fluorescence quenching of MOFs by DNT and DMNB. The red-shifted fluorescence can be seen in the inset, along with three quench/fluorescence regeneration cycles. [101] Copyright Wiley-VCH Verlag GmbH & Co. KGaA. Reproduced with permission.

contain nitro groups, which are known quenchers of emission. The structure maintains a three-periodic, porous geometry in which the ligands are held at appropriate distances to minimize π – π intermolecular interactions between the ligands, and is highly emissive. It is likely that the emission is ligand-based in this MOF. DMF guests incorporated during the synthesis can be removed reversibly, although the structure is slightly distorted upon removal.

12.6.2

Radiation Detection

Radiation detection goes hand-in-hand with the aforementioned interest in explosives detection for homeland security applications. In addition, medical devices and methods in biotechnology rely on radiolabeling. Portable, inexpensive neutron detectors are currently scarce, and are often not as sensitive as required. Interestingly, the same small conjugated molecules that make for suitable ligands in MOFs can also exhibit useful scintillation properties, in particular 4,4'-stilbenedicarboxylic acid. Such molecules are known for their ability to emit light effectively upon exposure to various forms of radiation, yet their solid-state structure remains a challenge for the development of reliable, reproducible sensor devices. The structures of both two-periodic network and three-periodic cubic Zn stilbene MOFs (see Figure 12.2) provide a rigid environments for the stilbene ligands, but the relative spacing and orientation differ, leading to differences in emission wavelengths and lifetimes. The two-periodic network is denser than the porous three-periodic cubic structure. Exposure of both MOFs to high-energy (3 MeV) protons and α -particles leads to radioluminescence, although the materials have different Stokes shifts (Figure 12.10) [102]. The three-periodic stilbene MOF has a larger Stokes shift, likely another consequence of the differences in stilbene–stilbene orientation between the two structures. It was found that these MOFs have scintillation quantum yields that are comparable to those with commercial scintillators. Moreover, the stability to repeated exposure outperforms anthracene, with luminescence degrading only very slowly with integrated doses up 10 MGy ($1 \text{ Gy} = 1 \text{ J kg}^{-1}$). The versatility of MOF syntheses and the commonality between scintillating organic groups and potential as MOF ligands leaves this area of study wide open for development.

12.6.3

Solid-State Lighting

The tunable nature of MOFs and the ability to incorporate multiple luminophores, including guest molecules, ligands, and metals, and the various potential interactions between these groups, can allow for fine-scale adjustment of the emission intensity and color of hybrid MOF materials. Solid-state lighting devices, such as organic light-emitting diodes (OLEDs), represent highly attractive, energy-efficient methods for the generation of light and are the subject of much contemporary research. In particular, white light sources for ambient lighting represent particularly interesting targets [103]. In addition to the tunability of MOF structures and emissive properties

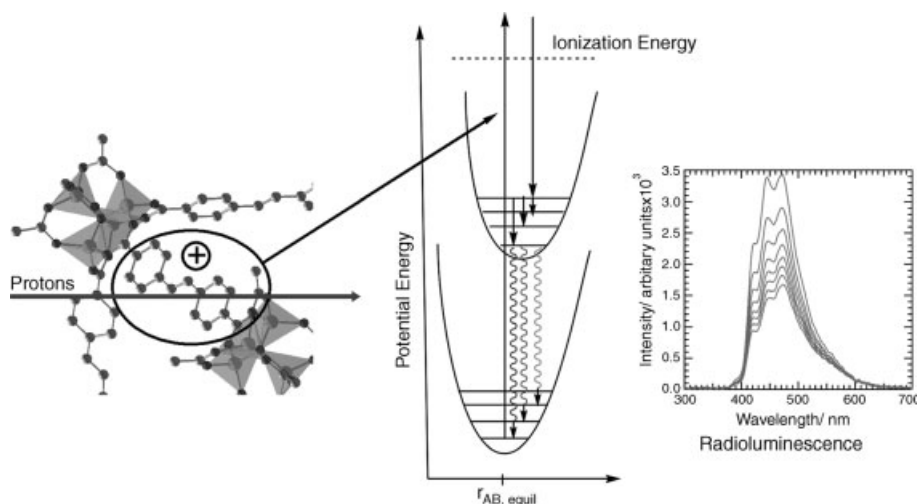


Figure 12.10 Schematic of the fast neutron detection process. Left: Interaction of high energy recoil protons with a rigid stilbene unit (a portion of the cubic 3D stilbene MOF structure is shown with stilbene dicarboxylate linkages between tetrahedral Zn_4O units), resulting in excitation, which decays (center) through

radiationless pathways (solid arrows) to the lowest-lying excited state of the stilbene groups, followed by radiative decay and emission of photons (wavy arrows). The emission profile is shown on the right, revealing diminishing intensity upon repeated doses of high intensity radiation.

discussed previously, their high thermal stability and often intense luminescence are desirable in this context. Current white light-emitting materials typically consist of mixtures of blue- and yellow-emitting materials [104]. Two cases of near white light-emitting MOFs have been reported to date. Ca and Sr frameworks containing 9-fluorenone-2,7-dicarboxylic acid exhibit broad, ligand-based emission that is dependent on the identity of the cation [105]. The rigidification of the ligand both by inclusion in the framework and by cooling to -196°C leads to an enhancement of the quantum yield. The broad emission of the pure ligand is yellow, whereas the Ca structure emits green light ($\lambda_{\text{max}} = 503\text{ nm}$) and the cation-exchanged Sr structure emits orange light ($\lambda_{\text{max}} = 526\text{ nm}$). Both frameworks can be excited between 380 and 460 nm, and the FWHM of the broad emission spectra is 90 nm, comparable to Ce^{III} : YAG phosphors.

The first example of a MOF material in which variable emission processes are harnessed for the purpose of generating white light was reported recently by Wang *et al.* [106]. A multi-process emission profile, highly sensitive to excitation wavelength, was used to manipulate the emission color of the MOF. The authors synthesized an Ag^{I} (4-cyanobenzoate)-based MOF that exhibits variable emission; the color is tunable from yellow to white via variation of the excitation wavelength. The solid-state emission spectrum shows a multitude of peaks, with intensities that vary upon modulation of the excitation wavelength. The maximum emission is at $\lambda = 427\text{ nm}$ when excited by 355 nm light, but shifts to 566 nm upon excitation by

330 nm light. Excitation of the MOF at 350 nm yields a white light-emitting material (to the eye) as, at this excitation wavelength, the 427 and 566 nm bands are of nearly equal intensity. There are several physical causes for the observed emission peaks, as MLCT and ligand-based emission coexist in this structure.

12.6.4

Nonlinear Optics

Nonlinear optics (NLO) properties (including second- and third-order processes) of both organic and inorganic materials are of great interest for a variety of purposes, such as communications, lighting, lasing, and frequency modulation. Second harmonic generation (SHG) is a nonlinear conversion of two photons of a particular frequency to a single photon with twice this frequency (i.e., frequency doubling). A requirement for SHG is non-centrosymmetry, which is challenging in bulk material. Donor–acceptor substituted molecules (i.e., push–pull conjugated) are often used. Current techniques involve assembly at liquid–air surfaces or electric field poling [107, 108]. As MOFs can be tailorable in function and shape, they offer potential as scaffolds amenable to the arrangement of ligands in a non-centrosymmetric manner. Several examples of MOFs exhibiting SHG are known, including Pb^{II} complexes prepared by Yang *et al.*, for which the emission can be assigned to a metal-centered transition involving the s- and p-orbitals of the metal, sensitized by the organic ligands, a relatively rare case of main-group MLCT [41]. A porous, Cd^{II} oxalate-based MOF ionic cluster exhibited among the highest SHG signals recorded for MOF materials, and this structure is also sensitive to cation exchange, illustrating the first NLO MOF with sensing potential [109]. A variety of other examples of engineered, asymmetric MOF crystals that exhibit high SHG responses have been reported [110–114]. In addition, Guo *et al.* prepared a multifunctional ferroelectric, NLO-active MOF [115].

Enhancement of standard one-photon fluorescence often leads to a concomitant increase in two-photon fluorescence, a third-order process. Although this has yet to be measured in a MOF material, the outcome may be of technological interest and warrants investigation. It has been well documented that centrosymmetric π -conjugated molecules with quadrupolar or higher degrees of charge separation have among the highest two-photon cross-sections (δ) [116]. These often have a donor– π -donor or acceptor– π -acceptor geometry and variations thereof. Such molecules are again well suited for MOF chemistry and many such MOF ligands are currently known or may be synthesized by straightforward protocols. In addition, coordination to the metal centers will likely improve the electron-withdrawing capability of the terminal carboxylate groups, and potentially improve the two-photon fluorescence signal strength.

12.6.5

Barcode Labeling

Barcoding is a method that requires reproducible, consistent materials for which readout is rapid. White *et al.* utilized the synthetic flexibility available in MOF

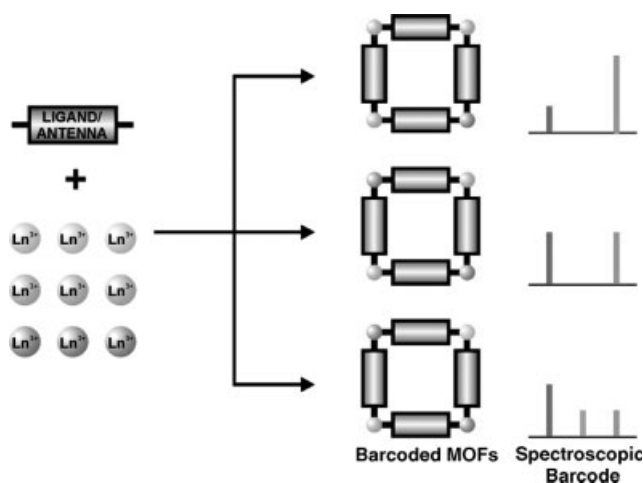


Figure 12.11 Schematic of the development of luminescent barcode materials by controlling the relative amounts of three differing emissive lanthanide ions in the MOF structures. Reprinted with permission from [117]. Copyright 2009 American Chemical Society.

syntheses to introduce multiple metal centers, each of which differs in luminescence color [117]. A mixture of lanthanide ions was used to develop MOFs with differing spectral outputs, which required excitation at a single wavelength as they are all sensitized by the same organic ligand. This allows for “encoding” of the spectral profile, as lanthanide emission is typically sharp, well resolved, and well separated (Figure 12.11). By tuning the ratio of a mixture of Ln^{III} ions (Yb, Er, Nd), a specific “code” results. The authors noted the reproducibility and wide diversity of codes possible. Upon coating in an adhesive matrix, the spectral profile was maintained, demonstrating the versatility of incorporation into other practical morphologies and composites.

12.7

Conclusion

What is so unique about MOFs, aside from the spectacular pore volumes and surface areas that they can exhibit, is that they possess the high local and long-range order characteristic of crystals, the synthetic versatility of organic polymers, the geometric predictability inherent in coordination bonding, *and* often stable nanoporosity. This amazing combination of properties is only beginning to be explored and is likely to lead to many surprises as scientists in fields beyond chemistry consider MOFs from their own perspectives and with their own specialized expertise. The examples cited in this chapter demonstrate that luminescent MOFs are not only a rapidly expanding subset with tremendous potential for a wide variety of applications, but are also an exciting venue for building a new, fundamental understanding of electronic structure, energy transfer, and light emission. It therefore seems

likely that research to date represents only the beginning of a long and interesting scientific story.

Acknowledgments

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