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Page 1 of 4 ChemComm

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Pure White-Light and Yellow-to-Blue Emission Tuning in Single Crystal of Dy(III) Metal-Organic Framework

Cite this: DOI: 10.1039/x0xx00000x

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Received 00th January 2012, Accepted 00th January 2012

DOI: 10.1039/x0xx00000x

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Direct white-light emission was first achieved in a single phase material of Dy(III) metal-organic framework, which also shows tunable yellow-to-blue photoluminescence by variation of excitation wavelengths.

White-light-emitting diodes (WLEDs) have wide applications in lasers, indicators, backlights, displays, *etc*. 1-2 Apart from the traditional method of blending multi-LEDs (for example, mixing LEDs with three primary colors red, green and blue, *i.e* RGB), single component (SC) white-light-emitting materials provide an alternative approach to fabricate WLEDs characteristic of uniform and well-balanced color, as well as easy-manipulation and lowcost.3-6 In recent years, lanthanide containing metal-organic frameworks (Ln-MOFs) have aroused special interest in design of SC white-light and color-tunable materials, taking advantages of the intrinsic multiple-colored *f-f* emissions of Ln atoms with efficient Antenna effect, sharp band and long lifetime.

To reach this goal, Ln atoms that could emit pure visible lights, *e.g.* red Eu(III) and green Tb(III) emissions, have been most frequently applied. In principle, if these two primary colors are further combined with blue luminescence, which is often contributed by the ligand-based emission, white-light emitting phosphor can be achieved in a single phase Ln-MOF containing mixed Eu and Tb centers.7-10 So far, most of the white-light emitting Ln-MOFs are constructed on the basis of such trichromatic RGB strategy; however, study on white-light emission of Ln-MOFs with other Ln(III) centers remains quite rare.¹¹ This is probably due to the fact that visible color emissions from other Ln atoms, such as Sm(III) and Dy(III), are usually rather weak. Dy(III) possesses appropriate *f-f* transitions which are able to emit colors in blue (480 nm), yellow (573 nm) and red (661 nm) regions under adequate excitation. Usually yellow emission dominates over the other two emissive colors. Therefore, if a Dy-MOF can be designed to emit yellow (Dy-based) and blue (ligand-based) colors simultaneously, white-light emission can be achieved through a BY dichromatic strategy on the balance of the

blue (B) and yellow (Y) colors. This will provide an alternative way to utilize Ln-MOFs as potential candidates of single-phase white phosphors. Nevertheless, due to the less efficient luminescence in most Dy-MOFs compared with the Eu/Tb counterparts,¹² little attention has been paid to the photoluminescence (PL) of Dy-MOFs, especially SC white-light emitting and color tuning properties.

We herein report a multiple-color photoluminescent Dy-MOF, [Dy(**TETP**)(NO³)3]⋅4H2O (**1**⋅4H2O, **TETP** = 1,1',1''-((2,4,6 triethylbenzene-1,3,5-triyl)tris(methylene))tris(pyridin-4(1H)-one)), in which the ligand **TETP** can play dual functions. On one hand, it can effectively sensitize Dy centers to produce yellow light; on the other hand, it is strongly blue luminescent with its blue emission in Dy-MOF being sufficient to balance yellow emission of Dy to generate white-light, offering the Commission Internationale de l'Eclairage (CIE) chromaticity coordinate of (0.33, 0.35). Moreover, by varying the excitation wavelengths, yellow-to-blue PL colortuning can be readily accomplished depending on the variation of the intensity ratios between the characteristic yellow emission of Dy atoms and the blue luminescence of **TETP** ligand.

solvent molecules, counter anions and H atoms are omitted for clarity (yellow for Dy, blue for N, red for O, and gray for C atoms, respectively).

Complex **1**⋅4H2O was prepared in a simple wet chemical method from reaction of **TETP** ligand and $Dy(NO₃)$ ₃ 6H₂O salt in wateracetone solvent system, affording colorless crystals in reasonable yield (58%) . The phase purity of the bulky sample of 1.4H₂O has been confirmed by powder X-ray diffraction in comparison with its single-crystal data simulation (Fig. S1). Complex **1** crystallizes in the space group $Pna2_1$ with the asymmetric unit consisting of one Dy(III), one **TETP** and three coordinating $NO₃$ ⁻ groups. The Dy(III) centers lie in a tricapped trigonal prismatic geometry, coordinated by three TETP through terminal O and three $NO₃$ ⁻ groups in chelating mode (Figure 1a). Meanwhile, each tripodal **TETP** connects three different Dy(III) with metal and ligand centers as nodes and pyridone arms as linkers, thus generating an intricate three-dimensional (3D) framework in (10, 3)- d (or **utp**) topology¹³ as seen from Fig. 1b.

The UV-adsorption of ligand **TETP** in solution shows a strong $\pi\pi^*$ band around 265 nm, and relatively weak intra-ligand charge transfer $(ILCT)^{10}$ above 300 nm originating from pyridone donors. These ligand transitions are merged and even expanded beyond 400 nm in the solid-state (Fig. S2). The solid-state emission spectrum of **TETP** displays a broad asymmetric emission band around 430 nm with decay lifetimes $\tau = 1.2$ and 2.2 ns (Fig. S3), revealing blue nature of the ligand-centered (LC) luminescence which can only be excited above 300 nm, thereof, dominated by ILCT. Upon excitation at 365 nm at room temperature, the solid sample of **1** displays multiple emission peaks at 430, 480, 573 and 661 nm. The first broad band centered at 430 nm is attributable to the reserved ILCT emission of **TETP**, while the three narrow peaks are originated from $f-f$ transitions between Dy(III) lowest emitting state ${}^{4}F_{9/2}$ and ground levels of ${}^{6}H_{15/2}$, ${}^{6}H_{13/2}$ and ${}^{6}H_{11/2}$. The nature of these emissions has been confirmed by the measurement of their decay dynamics. The 430-nm peak has a decay feature with $\tau = 0.8$ and 3.2 ns, typical of LC emission. And the main peak at 573 nm has a decay lifetime of 31 µs, characteristic of Dy-centered emission (Fig. S4). Remarkably, combination of these four emissions in complex **1** is properly balanced to result in white-light output, offering a CIE coordinate of (0.33, 0.35) which is close to that of pure white light (0.33, 0.33). The collected emissions of **1** (λ_{ex} = 365 nm, detection span: 400-800 nm) give an absolute quantum yield of \sim 7%, which is among the highest value for Dy(III) complex.¹⁴ The white-light can also be directly detected by the naked eye (Fig. 2, inset).

Fig. 2 Solid-state PL spectrum of **1** (λ_{ex} = 365 nm). The emission curve is superimposed on the white-light spectrum. Inset: photograph of white emissive Dy-MOF in a sample vial and CIE chromaticity coordinates.

The PL mechanism and energy transfer (ET) processes involved in this Dy-MOF may be complicated as illustrated in Scheme 1. The

PL spectrum of isostructural Gd-MOF (Fig. S1 and S5) indicates a triplet energy level of **TETP** at \sim 23000 cm⁻¹ (T = 77 K, τ = 0.7 and 6.9 μ s), which is adequate for ET to the ⁴F_{9/2} level of Dy (~21,500) cm⁻¹).¹⁴ Therefore, after **TETP** absorbs the UV light at either $\pi \pi^*$ (singlet transition, $<$ 300 nm) or ILCT (300-400 nm) bands, it can transfer the energy directly¹⁵ or via its triplet state to Dy center to generate the metal-centered (MC) emission; however, the blue LC emission can only be excited in the ILCT spectral region. These ETs are subject to influence of excitation energy. As shown in excitation spectra of **1** (Fig. S6), MC emission (573 nm) can be effectively sensitized via ligand singlet adsorption (< 300 nm), or moderately by ILCT and hypersensitized *f-f* transitions (300-400 nm), while this ILCT excitation in 300-400 nm region also generates LC emission. Monitoring at 430 nm reveals a wide excitation band resembling that of pure **TETP**, and shows structured profile with a few troughs corresponding to *f-f* adsorptions of Dy(III). This implies that ILCT excitation has not efficient Antenna effect as singlet transition of **TETP**, leading to LC emission accompanying with *f-f* emissions. Therefore, fine color-tuning is achievable through a dichromatic approach upon variation of excitations. At proper wavelength, the relative intensity between MC and LC emissions can be balanced to give white-light output directly from the single phase of Dy-MOF.

Scheme 1. Schematic representation of energy absorption, migration, emission (plain arrows) and dissipation (dotted arrows) processes in complex **1**. LC, ligand-centered; MC, metal-centered; ILCT, intraligand charge transfer; S, singlet state; T, triplet state; A, absorption; F, fluorescence; P, phosphorescence; nr, nonradiative; ISC, intersystem crossing; ET, energy transfer.^{10b}

Based on above discussion, this Dy-MOF can be considered to contain two kinds of emissive centers, *i.e.* Dy atoms emitting mainly yellow (573 nm) color and **TETP** components emitting blue color (430 nm), of which the luminescent intensities depend on the ET efficiency at varied excitation wavelengths. As a consequence, the PL of **1** is tunable from yellow to blue according to variation of excitation light. As seen in Fig. 3 and S6, when excited at 265 or 290 nm, the LC blue emission is negligible, while *f-f* emissions of Dy dominate. This denotes an effective ET from ligand Antenna to Dy centers at ligand singlet excitation, resulting in a CIE coordinate of (0.39, 0.44) in yellow region. In contrast, at the excitation of 373 nm, the ligand-to-metal ET becomes less efficient, while LC blue emission predominates over Dy emission. The resulting CIE coordinate is (0.21, 0.20), falling in blue region. In between these two wavelengths, the excitation at 338 nm leads to bluish-white emission with a CIE coordinate of (0.28, 0.29), while the excitation at 365 nm brings pure white-light emission at (0.33, 0.35) as mentioned before. When excited at 77 K, similar emission feature and color-tuning property can be achieved for this Dy-MOF (Fig.

S7), and the short lifetimes (0.9 and 5.5 ns) at 430 nm indicate that the LC emission at this temperature is still in nature of fluorescence as in room temperature, other than phosphorescence as in Gd-MOF.

Fig. 3 Emission spectra of 1 excited at different λ_{ex} . Insets: Chromaticity coordinates of 1 at different excitations (λ_{ex} = 290, 338, 365 and 373 nm; CIE $= (0.39, 0.44), (0.28, 0.29), (0.33, 0.35)$ and $(0.21, 0.20)$ and photographs of the name of our University SYSU written with the white-light emitting **1**.

In summary, we have successfully synthesized a 3D Dy-MOF showing tunable yellow-to-blue PL by variation of excitation light. It is noteworthy that pure white-light emission has been realized by this Dy-MOF in a single phase for the first time. In contrast to the common strategy to utilize mixed-metal Ln-MOFs (e.g. Eu and Tb) for white-light emission, this provides an alternative way to use homogeneous Ln-MOFs as single-phase white-light phosphors, which will make the preparation and manipulation even easier.

We thank the 973 Program of China (2012CB821701), the NSFC projects (91222201, 21373276, 21121061, 21173272), the NSF of Guangdong (S2013030013474), the FRF for the Central Universities, and the RFDP of Higher Education of China for funding.

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Electronic Supplementary Information (ESI) available: [synthetic, photophysical and crystallographic details]. See DOI: 10.1039/c000000x/.

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Table of contents entry

An alternative way to use homogeneous Ln-MOFs as singlephase white-light and color-tunable phosphors has been achieved by assembly of Dy(III)-MOF

