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Highly Stable and Sensitive LnMOF Ratiometric Thermometers Constructed with Mixed Ligands

Yongqin Wei, Rongjian Sa, Qiaohong Li and Kechen Wu*

The mixed-lanthanide metal-organic frameworks (M’LnMOFs) applied for accurate, non-invasive and self-reference temperature measurements have been only recently recognized. It is a great challenge for chemists to fulfil thermostable structure, intense luminescence and high temperature sensitivity on one LnMOF ratiometric thermometer for the thermometric applications. Choosing 2,4-(2,2‘:6′,2″-terpyridin-4′-yl)-benzenesulfonic acid (H2DSTP) as the first ligand and changing ancillary ligand oxalic acid (OA) or 1,4-benzene dicarboxylic acid (BDC), we successfully realized two types of highly stable and sensitive thermometers [Tb1−xEu0.5x(OA)x]·3H2O and [Tb1−xEu0.5x(BDC)x]·2H2O (x=0.01, 0.02) that in addition exhibit brilliant luminescence over a wide temperature range, providing a new strategy to explore luminescence-based M’LnMOF thermometer.

1. Introduction

The rational design and synthesis of metal-organic frameworks (MOFs) has undergone a tremendous development owing to their fascinating structures and functional applications as electronic, magnetic, optical and catalytic materials.1 Solid-state luminescence is particularly targeted owing to the advantage of their designability that allows the fine-tuning of luminescent properties.2 In particular, lanthanide metal-organic frameworks (LnMOFs) offer the advantage of rational choice of light-emissive building blocks for both the lanthanide ions or clusters as nodes and the functional organic ligands as linkers. Moreover, the isostructural LnMOFs make it feasible to maintain the original framework topologies via doping approach while produce novel luminescent properties.3

The search for new materials with temperature-sensing luminescent properties for their application in thermometry is nowadays a very active field.4 The thermal dependence of phosphor is a noninvasive and accurate alternative technique that provides fast response and high spatial resolution, and works remotely by way of an optical detection system even in biological fluids, strong electromagnetic fields and fast-moving objects.5 The study on luminescent thermometers at micro- and nanoscale for the microdevices and potential applications in life sciences is nowadays a very hot research topic.6 On the other hand, the mixed-lanthanide metal-organic frameworks (M’LnMOFs) for thermometers have been only recently realized.7 The developed M’LnMOFs methodology for luminescent thermometers is based on a thermally driven energy transfer from Tb5+ to Eu3+ within the framework solids and the corresponding emission intensity ratio can be strongly temperature-dependent. Such ratiometric thermometer does not require any additional calibration of luminescent intensity and thus overcomes the drawbacks of most luminescent thermometers based on the temperature-dependent intensity of one transition whose accuracy can be heavily affected by extrinsic factors like fluctuations in excitation rate, detection efficiency, probe concentration, optical occlusion, or other local inhomogeneities that alter absolute intensities.8

In theory, we could develop M’LnMOF ratiometric thermometer with high temperature sensitivity by modulating the excited state energy of organic ligand.7b,5 But apart from high sensitivity, a thermometer for application should be process of other properties such as high thermal stability and efficient luminescence detectable over a wide temperature range. It is well known that LnMOFs are prone to involve coordinated water or other solvent molecule due to large ion radius and high coordination number of trivalent lanthanide ions. The high-energy vibrations around inner coordination sphere will lead to low luminescent efficiency and even luminescent quench of lanthanide ions.9 Moreover, the LnMOFs will undergo a subtle structural transformation and even collapse of framework skeleton after the loss of coordinated solvent molecule at high temperature, which further results in the irrecoverable luminescence. Thus, M’LnMOF ratiometric thermometers with thermostable structure, intense luminescence and high sensitivity have not been reported. How to fulfil these terms on one M’LnMOF thermometer is a great challenge for chemists.

Cite this: DOI: 10.1039/x0xx00000x

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We have noticed that introduction of ancillary ligands into LnMOFs often results in the higher luminescent intensity due to the multiple beneficial effects: not only are high-energy vibrations removed from inner coordination sphere, increasing observed lifetime, but a better positioning of the excited state with one of the first ligand increases the efficiency of energy transfer.\(^\text{10,12}\) Furthermore, extruding the coordinated solvent molecule with ancillary ligand will probably enhance the thermal stabilities of LnMOFs. Nowadays there has no systematic study of ancillary ligand effect on the temperature-dependent luminescence of M’LnMOF thermometer. Choosing a designed 2,4-(2,2’-6,2’-terpyridin-4’-yl)-benzenedisulfonlic acid (H\(_2\)DSTP) as the first ligand and changing ancillary ligand, we successfully synthesized two types of LnMOFs, [Ln(OA)\(_0.5\)(DSTP)]•3H\(_2\)O (Ln-OA-DSTP) and [Ln(BDC)\(_0.5\)•3H\(_2\)O (Ln-BDC-DSTP) (Ln = Tb, Eu, OA = oxalic acid, BDC = 1,4-benzene dicarboxylic acid). The ratiometric temperature-sensing properties of the corresponding thermometers Tb\(_{1-x}\)Eu\(_x\)-OA-DSTP and Tb\(_{1-x}\)Eu\(_x\)-BDC-DSTP (x = 0.01, 0.02) distinctly change with the added ancillary ligand. Moreover, the removing of coordinated water molecule from the inner coordination sphere by ancillary ligand results in the high luminescent efficiencies of the thermometers and the tight-bonding polymeric structure leads to their application in the wide temperature-sensing range. The introduction of ancillary ligands into M’LnMOFs provides a new strategy to explore luminescence-based thermometer.

2. Experiment section

Materials and measurements
All commercially available chemicals were of analytical reagent grade and used as received without further purification. Elemental analyses for C, H, N and O were carried out on a Vario MICRO CHNOS Elemental Analyzer. Inductively coupled plasma spectroscopy (ICP) was performed on an Ultima2 Inductively Coupled Plasma OES spectrometer. The Tb/Eu ratios were determined by the ICP analysis and ICP samples were prepared by digesting the dry samples into concentrated HCl (30%). Powder X-ray diffraction (PXRD) patterns were collected in the 2\(\theta\) = 5–50° range on a D/MAX2500 Powder X-ray diffractometer using Cu-K\(_\alpha\) radiation (\(\lambda = 1.5406\) Å) at room temperature. Thermogravimetric analyses (TGA) were performed on a NETZSCH STA 449C instrument with a heating rate of 10°C/min in nitrogen atmosphere. Synthesis of 2,4-(2,2’-6,2’-terpyridin-4’-yl)-benzenedisulfonlic acid (H\(_2\)DSTP). Sodium 4-formylbenzene-1,3-disulfonate (7.8g, 25mmol) and 2-acetylpyridine (5.6mL, 50mmol) were dissolved in 100mL ethanol. To this was added, 30% NH\(_3\) solution (5mL) and NaOH (2g, 50mmol). The solution was stirred vigorously for 0-5°C for 3h, after which an yellow precipitate appeared. The stirring was continued at room temperature for two day and more precipitate was obtained. The precipitate was collected by filtration, washed with methanol. The product was dissolved in 30mL ice water. Hydrochloric acid was added to the solution and the PH value was adjusted to 5, after which a pale-yellow precipitate appeared. The precipitate was collected by filtration, washed with acetone, and oven-dried at 60°C (4.2g, yield: 32% based on Sodium 4-formylbenzene-1,3-disulfonate). Anal. Calcd for H\(_2\)STP•3H\(_2\)O: C 48.18, H 4.04, N 8.03, O 27.50. Found: C 47.93, H 4.12, N 8.24, O 27.15. \(^1\)H NMR (DMSO, Fig. S1, ESI†): 8.71 d 2H, 8.66 d 2H, 8.52 s 2H, 8.29 s 1H, 8.03 t 1H, 7.68 d 2H, 7.49 t 2H, 7.23 d 2H.

Syntheses of MOFs [Ln(OA)\(_0.5\)(DSTP)]•3H\(_2\)O (Ln-OA-DSTP). Taking Tb-OA-DSTP as an example, oxalic acid (OA, 0.15mmol), 2,4-(2,2’-6,2’-terpyridin-4’-yl)-benzenedisulfonlic acid (H\(_2\)DSTP, 0.3mmol), NaOH (0.9mmol), Tb(NO\(_3\))\(_3\)•6H\(_2\)O (0.35mmol), and water (25mL) were sealed into a Teflon cup. The solution was heated at 180°C for 3 days and then cooled to room temperature. The yellow crystals were collected by filtration and washed with DMF and water (88.6mg, yield: 41% based on H\(_2\)DSTP). Anal. Calcd for [Tb(OA)\(_0.5\)(DSTP)]•3H\(_2\)O: C 36.48, H 2.64, N 5.80, O 24.29. Found: C 36.52, H 2.83, N 5.75, O 24.35.

Syntheses of MOFs [Ln(BDC)\(_0.5\)(DSTP)]•2H\(_2\)O (Ln-BDC-DSTP). The MOFs Ln-BDC-DSTP were synthesized similarly to Ln-OA-DSTP except for the use of 1,4-benzene dicarboxylic acid (BDC) to replace oxalic acid.

Single-crystal structure determination

Suitable single crystals of Ln-OA-DSTP and Ln-BDC-DSTP (Ln = Eu, Tb) were carefully selected and glued to thin glass fibers with epoxy resin. Intensity data for single crystal were collected on a Rigaku Mercury CCD diffractometer with graphite-monochromatized Mo K\(_\alpha\) radiation (\(\lambda = 0.71073\) Å). The empirical absorption corrections were performed using the CrystalClear program.\(^{11}\) The structures were solved and refined on \(F^2\) by full-matrix least-squares technique using the SHELX-97 program package.\(^{12}\) Anisotropic thermal parameters were applied to all non-hydrogen atoms. The hydrogen atoms bonded to the carbon atoms of ligands were generated geometrically and the hydrogen atoms of water molecule were located in a difference Fourier map. In the structures of compounds Ln-OA-DSTP, there were extensive areas of residual electron density that could not sensibly be modeled as water molecules.\(^{13}\) Therefore, they were removed via application of the SQUEEZE function in PLATON. The final formulas of the two compounds were determined by combining with elemental analysis. Crystallographic data for the four compounds are listed in table S1 (ESI†).

Photoluminescence (PL) Measurements

Microcrystalline samples in the solid state were used for all photoluminescence measurements. The temperature-dependent emission spectra and the decay curves were recorded on an Edinburgh FLS980 fluorescence spectrometer equipped with a
450W ozone-free Xenon Arc lamp as light source, triple-grated monochromators for the excitation and emission pathways, and a R928P photomultiplier tube (PMT) for signal detection in the single-photon counting mode. The temperature measurements (77–400K) were performed on a Linkam THMSE600 controller. The overall quantum yields at room temperature were determined by the absolute method using a BaSO$_4$-coated integrating sphere as sample chamber on an Edinburgh FLS920 fluorescence spectrometer.

3. Results and discussion

Structural characterization

Two types of LnMOFs, Ln-OA-DSTP and Ln-BDC-DSTP (Ln = Tb, Eu) were synthesized under the hydrothermal reaction of Ln(NO$_3$)$_3$, H$_2$DSTP and corresponding ancillary ligand. All structures were determined by using single crystal X-ray crystallography. In the two types of isostructural LnMOFs, ligand DSTP adopts the same coordination mode and uses three chelating nitrogen atoms of terpyridine, bidentate bridging and monodentate sulfonic groups to link four lanthanide ions, forming a 2D Ln-DSTP structure. Ligand OA using two carboxylate groups to link two metal centers is inserted to the 2D Ln-DSTP, resulting in the final 2D polymeric structure of Ln-OA-DSTP (Fig. 1a). The topology can be viewed as a (4,5)-connected net with short Schläfl (point) symbol {3.4$^2$.5$^2$.6$^2$.} (Fig. 1b). Ligand BDC using two bidentate bridging carboxylate groups to link four metal centers connects the layered Ln-DSTP, resulting in the final 3D polymeric structure of Ln-BDC-DSTP (Fig. 2a). In the compounds Ln-BDC-DSTP, two metal centers are linked by two bridging carboxylate groups and two bridging sulfonic groups, forming a dimer building block. As shown in Fig. 2b and Fig. 2c, ligand DSTP is simplified as a 3-connected node to link the dimer building blocks and the topology of Ln-BDC-DSTP is a (3,8)-connected net with short Schläfl (point) symbol {4$^4$.6$^{18}.8^4$}. The metal centers in Ln-OA-DSTP and Ln-BDC-DSTP adopt a dodecahedron 8-coordinate environment which has no coordinated water and is totally occupied with nitrogen and oxygen atoms of the first ligand DSTP and the ancillary ligand OA or BDC. Such tight-bonding structure is expected to enhance the luminescent efficiency and thermal stability of the corresponding M’LnMOF thermometers.

Ratiometric temperature-sensing properties

Because of the tight polymeric structures of as-synthesized materials, they are virtually insoluble in most solvents such as H$_2$O, CH$_3$OH, CH$_3$CN, THF, DMF, and DMSO. Microcrystalline samples in the solid state were used for all photoluminescence measurements.

The temperature-dependent emission spectra of Ln-OA-DSTP and Ln-BDC-DSTP (Ln = Tb, Eu) from 77 to 400K are illustrated in Fig. 3. As expected, the emission intensity of Tb$^{3+}$ in Tb-OA-DSTP and Tb-BDC-DSTP decreases dramatically with increasing temperature. However, the emission intensity of Eu$^{3+}$ in Eu-OA-DSTP and Eu-BDC-DSTP nearly has no change with increasing temperature. Such significantly different emissions of Tb$^{3+}$ and Eu$^{3+}$ will enable us to develop new M’LnMOF thermometers Tb$_{1-x}$Eu$_x$-OA-DSTP and Tb$_{1-x}$Eu$_x$-BDC-DSTP based on the temperature-dependent Tb$^{3+}$/Eu$^{3+}$ emission intensity ratio of high sensitivity. Moreover, the successful syntheses of Ln-OA-DSTP and Ln-BDC-DSTP should enable us to investigate the ancillary ligand effect.
on the ratiometric temperature-sensing properties. To accomplish the research aims, we synthesized Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP and Tb$_{0.98}$Eu$_{0.02}$-BDC-DSTP with the same molar ratio Tb/Eu, which was confirmed by inductively coupled plasma (ICP) analysis and powder X-ray diffraction (PXRD) measurement (Table S2 and Fig. S2, ESI†). As shown in their temperature-dependent spectra (Fig. 4a and Fig. 5a), the emission intensity of Tb$^{3+}$ decreases dramatically with increasing temperature, while that of Eu$^{3+}$ nearly remains unchanged for Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP and even increases gradually for Tb$_{0.98}$Eu$_{0.02}$-BDC-DSTP. The two thermometers exhibit bright photoluminescence to the naked eyes and show different colors (green, yellow, orange, and red) with increasing temperature (Fig. S3, ESI†), allowing us to visualize the temperature change instantly and straightforwardly.

In the temperature-dependent emission spectra of Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP and Tb$_{0.98}$Eu$_{0.02}$-BDC-DSTP, $^5D_0\rightarrow^7F_2$ transitions of Eu$^{3+}$ overlap distinctly with $^5D_2\rightarrow^7F_3$ transitions of Tb$^{3+}$ at the cryogenic region (77-225K). Thus, the luminescent intensity of $^5D_0\rightarrow^7F_2$ transition is unfit for being chosen as an experimental parameter to reflect luminescent change of Eu$^{3+}$. It is interesting that the $^5D_0\rightarrow^7F_4$ transitions in the red emissions of Eu-OA-DSTP and Eu-BDC-DSTP have comparable intensity with that of $^5D_2\rightarrow^7F_4$ transitions (Fig. 3). It is well known that in most of Eu(III) compounds, the $^5D_0\rightarrow^7F_2$ transition has by far the largest intensity and the $^5D_0\rightarrow^7F_4$ transition is rather weak. The enhanced intensity of the $^5D_0\rightarrow^7F_4$ transition is also observed for Na$_9$[EuW$_{10}$O$_{36}$]$\cdot$14H$_2$O$^{14}$ and [EuCl$_3$(1,3-Pb(CN)$_2$)$_{15}$, which was explained as a consequence of the intensity parameters $\Omega_4$ in a highly polarizable environment.$^{14}$ In view of no overlap of the $^5D_0\rightarrow^7F_4$ transition with any transition of Tb$^{3+}$, intensity ratio of $^5D_4\rightarrow^7F_2$ transition of Tb$^{3+}$ and $^5D_0\rightarrow^7F_4$ transition of Eu$^{3+}$ was chosen as the thermometric parameter ($\Delta=I_{\text{Tb}}/I_{\text{Eu}}$).
Fig. 5 The ratiometric temperature-sensing properties of Tb$_{0.98}$Eu$_{0.02}$-BDC-DSTP: (a) emission spectra for the solid sample recorded between 77 and 400K upon excitation at 360nm, inset: normalized emission intensities of $^5D_4 \rightarrow ^7F_5$ transition of Tb$^{3+}$ (546nm) and $^5D_0 \rightarrow ^7F_4$ transition of Eu$^{3+}$ (702nm); (b) temperature-dependent intensity ratio ($\Delta/\Delta_77K$) and linearly fitted curve; (c) the natural logarithm of the thermometric parameter $\Delta$ changing with the inverse of temperature and the linearly fitted curve.

Fig. 4b presents the temperature-dependent intensity ratio of the thermometer Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP. The $\Delta$ values were normalized to the 77K value (the lowest measured temperature). It is exciting that the temperature measurement can be linearly correlated to experimental parameter in a wide temperature range of 77-275K. Moreover, the sensitivity was calculated based on the equation 1,

$$S = \frac{\Delta \Delta/\Delta}{\Delta}$$  \hspace{1cm} (1)

which showed that the thermometer Tb$_{0.98}$Eu$_{0.02}$-BDC-DSTP has a maximum sensitivity of 2.4% K$^{-1}$ at 275K (Fig. S6, ESI$^\dagger$). The first luminescent M$'$LnMOF thermometer [(Tb$_{0.9931}$Eu$_{0.0069}$)$_2$-(DMBDC)$_3$(H$_2$O)$_4$]·DMF·H$_2$O (DMBDC = 2,5-dimethoxy-1,4-benzenedicarboxylate) reported by Cui et al exhibits a linear correlation between emission intensity ratio and the temperature from 50 to 200K with a maximum sensitivity of 1.3% K$^{-1}$ at 200K.$^{7a}$ The high sensitivity of thermometer Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP in the wide temperature range is really remarkable, which will allow us to highly sensitize the temperature change. In fact, another thermometer Tb$_{0.99}$Eu$_{0.01}$-OA-DSTP is still process of the excellent linear correlation between the temperature and luminescence intensity ratio from 77 to 225K (Fig. S4b, ESI$^\dagger$), indicating that Tb$_{1-x}$Eu$_x$-OA-DSTP is a type of very useful luminescent thermometer.

Another notable character of thermometer Tb$_{1-x}$Eu$_x$-OA-DSTP is the higher thermal stability compared with the most of LnMOFs that involve the coordinated water in the inner coordination sphere. As mentioned above, the loss of coordinated water at high temperature will result in irrecoverable luminescence and even collapse of framework skeleton. In result, the most of M$'$LnMOF thermometers are not fit for being used at the high temperature. In order to confirm the practicability of the two thermometers Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP and Tb$_{0.99}$Eu$_{0.02}$-BDC-DSTP at high temperature, we investigate their thermal stabilities. Thermogravimetric analyses (TGA) showed that the two thermometers have decomposition points of 380 and 515°C (Fig. 6a), respectively. The higher stability of Tb$_{0.98}$Eu$_{0.02}$-BDC-DSTP could be attributed to the tight 3D polymeric structure. The first weight

Fig. 6 The thermal stabilities of the two thermometers of Tb$_{0.98}$Eu$_{0.02}$-OA-DSTP and Tb$_{0.99}$Eu$_{0.02}$-BDC-DSTP: (a) thermogravimetric analyses (TGA); (b) the reversible changes of emission intensity ratio 1/Δ by the alternative thermo-cycles in the range of 300 and 400 K.
loss of the two thermometers below about 150°C is related to
dissociative water molecules in the crystals. But it could not
change the framework skeletons. The measured emission
intensities and values of the emission intensity ratio remain
unchanged, which was verified by the reversible luminescence
under thermo-cycles. As shown in Fig. 6b, the reversible
changes of the emission intensity ratios of $I_{\text{Tb}}/I_{\text{Eu}}$ were observed
by alternative thermo-cycles in the temperature range of 300
and 400K. It is notable that the verge of temperature-dependent
measurement at 400K is far apart from the decomposition
points of the two thermometers. But the emission intensity of
$\text{Tb}^{3+}$ decreases dramatically with increasing temperature and
the measurement at higher temperature will be unreliable.

The encountered problem is no longer linear correlation of the
intensity ratio at elevated temperature because of the low
measured value ($\Delta<1$). It is necessary to find a new function for
the use at high temperature region. We chose the plot of the
natural logarithm of the thermometric parameter $\Delta$ vs the
inverse of temperature, which has been adopted by Carlos et al. As shown Fig. 4c, for temperatures above 225K the
temperature dependence of $\ln(\Delta)$ of thermometer $\text{Tb}_{0.98}\text{Eu}_{0.02}\text{-OA-DSTP}$ follows the equation: $\ln(\Delta)=1520.5/T–5.363$
($R^2=0.994$). The slope value of 1520.5 is larger than the
reported value by a factor of ca. 10. Thus, we successfully
developed a new type of thermostable thermometer $\text{Tb}_{1.0}\text{Eu}_{0.0}$
-OA-DSTP being process of not only the high sensitivity but
also the wide temperature-sensing range.

As shown in Fig. 5 and Fig. S5 (ESI†), the ratiometric
temperature-sensing properties distinctly change with the added
ancillary ligand. $\text{Tb}_{1.0}\text{Eu}_{0.0}\text{-BDC-DSTP}$ ($x=0.01, 0.02$) exhibit the remarkably higher maximum sensitivities (2.8% K⁻¹ at
225K for $\text{Tb}_{0.98}\text{Eu}_{0.02}\text{-BDC-DSTP}$ and 3.9% K⁻¹ at 200K for
$\text{Tb}_{0.99}\text{Eu}_{0.01}\text{-BDC-DSTP}$, larger than that of the corresponding
$\text{Tb}_{1.0}\text{Eu}_{0.0}\text{-OA-DSTP}$ at the same temperature). Although the
thermometric parameters of $\text{Tb}_{1.0}\text{Eu}_{0.0}\text{-BDC-DSTP}$ are linearly
related in a narrower temperature range, the natural logarithm
of the thermometric parameters $\Delta$ are linearly related to the
inverse of temperature in a wide temperature range of 200-
400K. The slope values of 1803.3 and 1655.1 are larger than
that of the corresponding thermometers $\text{Tb}_{1.0}\text{Eu}_{0.0}\text{-OA-DSTP}$.

Energy transfer mechanism
The measurement for lifetimes of $\text{Tb}^{3+}$ and $\text{Eu}^{3+}$ in the pure
MOFs and the $\text{Tb}/\text{Eu}$-codoping MOFs (Fig. S7, ESI†) showed that the lifetime of $\text{Tb}^{3+}$ in the $\text{Tb}/\text{Eu}$-codoping MOF becomes
shorter compared with that in the pure MOF, whereas the
lifetime of $\text{Eu}^{3+}$ increases. This information indicates the $\text{Tb}^{3+}$
to-$\text{Eu}^{3+}$ energy transfer, which is verified by the emission
spectra of $\text{Tb}_{0.98}\text{Eu}_{0.02}\text{-OA-DSTP}$ and $\text{Tb}_{0.99}\text{Eu}_{0.01}\text{-BDC-DSTP}$
on the excitation of 488nm (Fig. S8, ESI†). The higher sensitivities of $\text{Tb}_{1.0}\text{Eu}_{0.0}\text{-BDC-DSTP}$ compared with $\text{Tb}_{1.0}\text{Eu}_{0.0}$
-OA-DSTP ($x=0.01, 0.02$) are related to the efficient energy
transfer from $\text{Tb}^{3+}$ to $\text{Eu}^{3+}$, which is estimated using the equation $\ref{eq:2}$,
$$\eta_{\text{Tb-Eu}} = 1 - \frac{\tau_{\text{Eu}}}{\tau_{\text{Tb}}}$$
where $\tau_{\text{Tb}}$ and $\tau_{\text{Eu}}$ are the emission lifetimes monitoring $^5\text{D}_4$
emission of $\text{Tb}^{3+}$ at 546nm of the pure $\text{Tb}$($\text{III}$) compound and
the corresponding $\text{Tb}/\text{Eu}$-codoping compound, respectively. As shown in Fig. 7, the energy transfer efficiency of $\text{Tb}_{0.98}\text{Eu}_{0.02}$
-OA-DSTP is relatively lower in all temperature range, while
$\text{Tb}_{0.98}\text{Eu}_{0.02}\text{-BDC-DSTP}$ shows different temperature-
dependence of the energy transfer efficiency, which enhances
quickly with increasing temperature, thus resulting in the
quicker enhancement of the emission intensity of $\text{Eu}^{3+}$ and the
higher temperature sensitivity.

The difference of temperature sensitivity is essentially related
to the triplet-state energy location of ligands. In order to
investigate energy transfer between ligands and lanthanide ions,
the emission spectrum of Gd-OA-DSTP was measured at 77K (Fig. S9, ESI†), which showed that the first triplet-state energy of DSTP (19050cm⁻¹) estimated from the phosphorescence peak at about 525nm) is slightly higher than 3D₃ of the Eu³⁺ (17200cm⁻¹) and lower than 3D₄ of the Tb³⁺ (20400cm⁻¹). We could deduce that energy transfer to appropriate level of Tb³⁺ probably came from the first triplet-state of ancillary ligands OA or BDC. The lower triplet-state energy of BDC compared with that of OA will result in the more efficient energy back transfer and further makes energy transfer from Tb³⁺ to Eu³⁺ easier. We estimated the energy back-transfer rate (ν_back) of Tb-OA-DSTP and Tb-BDC-DSTP using kinetic analysis. ν_back is assumed to obey the following Arrhenius-type equation 3,18

$$\ln(\frac{1}{\tau_{obs}} - \frac{1}{\tau_{77K}}) = \ln A - \frac{E_a}{k_B T}$$

where τ_obs is the emission lifetime, τ_{77K} is the emission lifetime at 77K, E_a is the activation energy, k_B is the Boltzmann constant and T is the absolute temperature. The activation energies obtained for Tb-OA-DSTP and Tb-BDC-DSTP (Fig. 8) are 1620±26cm⁻¹ and 1316±36cm⁻¹, respectively, in agreement with the above suggestion that the energy back transfer of Tb³⁺ involves the excited state of ancillary ligand.

4. Conclusion

In summary, we successfully target two types of M’LnMOF ratiometric thermometers Tb₁₋ₓEuₓ-OA-DSTP and Tb₁₋ₓEuₓ-BDC-DSTP (x = 0.01, 0.02) for application by introducing ancillary ligands into the framework solids. The thermometers exhibit brilliant luminescence in the all temperature range of 77-400K. The absolute luminescence quantum yields of Tb₀.₉₈Eu₀.₀₂-OA-DSTP and Tb₀.₉₈Eu₀.₀₂-BDC-DSTP measured at room temperature are 16.2% and 24.5% respectively. Notably, the overall emission of Tb₁₋ₓEuₓ-BDC-DSTP is even more intense at the elevated temperature. The intense luminescence of the thermometers can be explained as a consequence of excellent light-harvesting linker DSTP with large π-conjugated system, efficient energy transfer from Tb³⁺ to Eu³⁺ and high-energy vibrations removed from the inner coordination sphere by the ancillary ligand OA or BDC. Furthermore, the tight-bonding structure as a result of the introduction of the ancillary ligands enhances the thermal stabilities of the thermometers and leads to their application in the wide temperature-sensing range, which has been seldom reported for M’LnMOF thermometers. It is notable that the ratiometric temperature-sensing properties distinctly change with the added ancillary ligand. Although the introduction of ancillary ligands brings difficulty for study on energy transfer mechanism of M’LnMOF thermometers, improving their temperature-sensing properties with ancillary ligands provides a new strategy to explore luminescence-based M’LnMOF thermometers.

Acknowledgements

We gratefully acknowledge the financial support of Natural Science Foundation of China (no. 21171165, 21201165 and 91122015).

Notes and references

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† Electronic supplementary information (ESI) available: Information of supplementary data and figures. CCDC reference numbers 1016955-1016958.


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By introducing ancillary ligands into M’LnMOFs and removing coordinated water molecule from the inner coordination sphere, we successfully realized two types of highly stable and sensitive thermometers that in addition exhibit brilliant luminescence over a wide temperature range, providing a new strategy to explore luminescence-based M’LnMOF thermometer.