A stoichiometric terbium-europium dyad molecular thermometer: energy transfer properties

Guochen Bao1,2,3, Ka-Leung Wong1, Dayong Jin2,3 and Peter A. Tanner1

Abstract

The optical thermometer has shown great promise for use in the fields of aeronautical engineering, environmental monitoring and medical diagnosis. Self-referencing lanthanide thermo-probes distinguish themselves because of their accuracy, calibration, photostability, and temporal dimension of signal. However, the use of conventional lanthanide-doped materials is limited by their poor reproducibility, random distance between energy transfer pairs and interference by energy migration, thereby restricting their utility. Herein, a strategy for synthesizing hetero-dinuclear complexes that comprise chemically similar lanthanides is introduced in which a pair of thermosensitive dinuclear complexes, cyCTb-phEu and cYcEu-phTb, were synthesized. Their structures were geometrically optimized with an internuclear distance of approximately 10.6 Å. The sensitive linear temperature-dependent luminescent intensity ratios of europium and terbium emission over a wide temperature range (50–298 K and 10–200 K, respectively) and their temporal dimension responses indicate that both dinuclear complexes can act as excellent self-referencing thermometers. The energy transfer from Tb3+ to Eu3+ is thermally activated, with the most important pathway involving the F1, J-multiplet at room temperature. The energy transfer from the antenna to Eu3+ was simulated, and it was found that the most important ligand contributions to the rate come from transfers to the Eu3+ upper states rather than direct ligand–metal transfer to 5D1 or 5D0. As the first molecular-based thermometer with clear validation of the metal ratio and a fixed distance between the metal pairs, these dinuclear complexes can be used as new materials for temperature sensing and can provide a new platform for understanding the energy transfer between lanthanide ions.

Introduction

Luminescent physical sensors for monitoring temperature have shown great promise for use in the fields of aeronautical engineering, environmental engineering, and industrial processes1–4. They have distinguished advantages over traditional thermometers in terms of a fast response, a high sensitivity, and a tolerance to extreme atmospheres5–7. In particular, self-referencing optical thermometers do not require additional calibration of the emission intensity and are more accurate due to non-invasive operation8,9.

Materials containing two different lanthanide ions are attractive for the construction of a self-referencing thermometer10. Lanthanide luminescent materials have sharp emission bands11,12 and a large energy shift between the antenna absorption and lanthanide emission 13, which distinguishes them from other luminescent materials, such as organic dyes or quantum dots14. In addition to these properties, lanthanide luminescence has a long lifetime, allowing time-gated techniques to increase the signal-to-noise ratio15. The temperature-dependent quenching and energy transfer between the dopant ions...
or between the host and lanthanide ion allow selective luminescent responses for different lanthanide ions with a change in temperature. Many lanthanide materials have been developed for temperature monitoring, such as lanthanide-doped inorganic nanocrystals, metal-organic frameworks (MOFs), polymers, and co-doped complexes. The reported maximum temperature sensitivities of recent systems involving energy transfer between two different lanthanide ions for ratiometric emission measurements are collected in Table S13. The stoichiometry of these systems is a critical parameter, and the sensitivities usually vary widely with temperature.

However, it is difficult to obtain exactly the same concentrations and qualities from different prepared batches of doped materials; the distance between the donor ion and acceptor ion, which is a crucial parameter for energy transfer, is random and can only be estimated by statistical treatment, and a mixed system with these materials allows donor–donor and acceptor–acceptor energy migration, making the kinetics of transfer more complicated. Hence, the hetero-dinuclear lanthanide complex, with its fixed distance between donor and acceptor, its definite structure and stoichiometric ratio of donor and acceptor, has great potential for development as a luminescent self-referencing thermosensitive indicator. The donor and acceptor in the dinuclear complex are fixed at a certain distance so that the pairwise energy transfer rates can be obtained without statistical treatment. With a stoichiometric arrangement of different lanthanide ions at each binding center, the intramolecular energy transfer between the ions is scarcely interfered with by donor–donor and acceptor–acceptor energy migration. The antenna effect of the chelating chromophore, which possesses a large absorption cross section, allows more incident light to be absorbed and transferred to a lanthanide ion, making such complexes brighter than most lanthanide-doped inorganic nanocrystals. Unsurprisingly, few literature reports include an investigation of the energy transfer processes using lanthanide hetero-dinuclear complexes - and even then with the co-presence of homo-dinuclear species in the complex mixture. Even though great effort has been expended to synthesize pure hetero-polynuclear complexes with different chemically similar lanthanide ions, there remains a considerable need for a simple strategy for controlling the formation of hetero-dinuclear complexes. This is a severe challenge for lanthanide chemists.

Herein, we report a pair of hetero-dinuclear lanthanide complexes, cycTb-phEu and cycEu-phTb, as molecular-based luminescent temperature sensors. In order to avoid mixing the two chemically similar lanthanide ions, europium and terbium, they are situated in two distinct binding sites using two different synthesis steps. The clear validation of the metal ratio and the fixed distance between the energy donor and acceptor make the complexes outweigh other materials, such as doped crystals, MOFs and polymers, both in terms of the understanding energy transfer and in temperature sensing performance. The 1,10-phenanthroline (phen) moiety serves as the chromophore to sensitize europium and terbium, giving an increased temperature-dependent luminescence emission ratio for europium over terbium.

**Fig. 1** The structures of complexes. (a) cycTb-phEu and (b) cycEu-phTb, and (c) optimized geometry of the simplified 100-atom structure of cycTb-phEu as determined using the PBeh-3c functional with basis def2-SVP and effective core potentials for Tb and Eu in ORCA (refer to the SI for details). The charge-transfer nature of the (d) HOMO and (e) LUMO are shown.
The luminescence from terbium and from europium is reduced in intensity with different rates as the temperature increases, and it involves the process of energy transfer from the chromophore to each ion and that from terbium to europium. Complexes cycTb-phEu and cycEu-phTb (Fig. 1) exhibit remarkably different photophysics due to the uniqueness of the two metal binding centers, and their energy transfer processes have been studied in detail.

Results

Structure

Refer to Fig. S1 for the structures and names of the complexes. The 166-atom structure of cycTb-phEu was modeled by MOPAC28,29 using the semi-empirical RM1 software in the LUMPAC program30,31, and the structure is shown in Fig. S13. Both Tb and Eu are 8-coordinated. Eu is coordinated to six S and to two N, whereas Tb is coordinated to four O and four N. A simplified 100-atom structure with aliphatic rings replaced by CH2 groups was employed for the optimization using the PBeh-3c functional in ORCA24–27 (Fig. 1). A comparison of the bond distances for these two different optimizations is provided in Table S10.

Synthesis

Scheme S1 shows the synthesis of the ligands and complexes. Ligand 2 (cyclen 1,10-phenanthroline) was prepared by a substitution reaction of 2-bromoacetyl bromide with 1,10-phenanthrolin-5-amine in the presence of K2CO3 in DCM for 22 h to give 1, which was followed by coupling with a triarmed cyclen in MeCN in the presence of NaHCO3 at room temperature, giving 2 in a 95.8% yield. The triarmed cyclen 1 was prepared via two-step substitution reactions according to the literature13. Complexes 3 (cycLn1-phen) were formed by coordinating lanthanide chlorides with one equivalent of ligand 2 in a mixed solution of MeOH and H2O at room temperature, followed by precipitation with diethyl ether from methanol. Taking advantage of fast coordination to Ln(III) ions and less interference with the energy transfer from the 1,10-phenanthroline chromophore to the Ln(III) ions, pyrrolidine dithiocarbamate was chosen as the ligand for coordination with the second lanthanide ion. The synthesis of complexes 4 (cycLn1-phLn2) was achieved by coordinating lanthanide chloride with one equivalent of complex 3 and three equivalents of ammonium pyrrolidine-1-carbodithioate at room temperature, followed by precipitation with diethyl ether from methanol. The proton nuclear magnetic resonance (1H NMR) spectra of the La motif structure showed that three pyrrolidine dithiocarbamates were coordinated to the complex, which is consistent with a previous report32. This synthetic strategy provides a platform to obtain heterodinuclear complexes that consist of chemically similar lanthanide ions, one of which is not present in both ligand structures.

Singlet and triplet states

The singlet and triplet energy levels were experimentally studied by using the La motif complex, cycLa-phLa, since La3+ does not exhibit 4f-4f electron transitions. The large La3+ ion generates a heavy atom effect towards the surrounding organic ligand and induces spin-orbit coupling, which accommodates intersystem crossing from singlets to triplets. Thus, phosphorescence occurs. The fluorescence from the solid La3+ complex was measured at room temperature, while the phosphorescence was recorded at 77 K (Fig. 2). The room temperature emission under broad band 273 nm excitation was observed for cycLa-phLa with a maximum at 398 nm (25121 cm−1) (Fig. 2b), which is attributed to a π-π* singlet transition of the ligand. At lower temperature, the fluorescence intensity decreases, the band redshifts, and phosphorescence from

Fig. 2 Optical spectra. The 77 K phosphorescence a and 298 K fluorescence b of cycLa-phLa (λexc = 273 nm) in the solid state. c Room temperature absorption spectrum of cycLa-phEu in buffer solution at pH 7.4 at two concentrations: black, 5 μM, red, 10 μM. The insert d shows the simulated absorption spectrum in the gas phase between 300 and 700 nm as determined from LUMPAC30 with the triplet levels calculated to low energy of 316 nm marked by vertical lines. The singlet transitions were broadened with widths of 2000 cm−1.
T1 dominates at longer wavelengths (Fig. 2a). The zero phonon line of the T1 → S0 transition is at 497 nm (20,124 cm⁻¹), and there is a progression of the totally symmetric ring C = N mode at 1471 ± 6 cm⁻¹ to lower energy (compare with the Fourier transform infrared (FT-IR) spectrum, Fig. S2). The excitation spectrum of the phosphorescence (Fig. S3a) demonstrates the population of T1 from a singlet state at ~400 nm. The excitation spectrum of the singlet fluorescent (Fig. S3b) shows a further singlet state at 273 nm. A strong band at or near this wavelength is observed for all of the cyclen-ligand systems in solution, for example, cycLa-phEu (Fig. 2c). In fact, the room temperature excitation spectra of cycEu-phLa (Fig. S4) and cycLa-phEu (Fig. S5) demonstrate the participation of other singlet states in the energy transfer routes from the ligand to Eu3+. Notably, by comparing the intensity of the ligand and Eu3+ absorption bands in the excitation spectra, the ligand–metal energy transfer is more efficient when Eu3+ is bound to the phen rather than to the cyclen moiety (Figs. S4 and S5).

The triplet lifetime was determined to be 0.26 s at 77 K (Fig. S6a), whereas the singlet lifetime was measured at room temperature to be 1.6 ns (Fig. S6b).

**Lanthanide luminescence at low temperature**

The emission spectra of complexes cycTb-phEu and cycEu-phTb were measured in the solid state at 10 K with excitation at 355 nm into the ligand absorption band (Fig. 3). These emission spectra exhibit the characteristic Eu3+ and Tb3+ emission bands: 5D0 → 7F2 (J = 0–6) for Eu3+ and 5D4 → 7F2 (J = 6–4) for Tb3+. In each case, the emission from Eu3+ dominates with the forced electric dipole transitions 5D0 → 7F2 (J = 2, 4) being the strongest and more intense than the magnetic dipole-allowed transition 5D0 → 7F1. The 5D0 → 7F0 transition, which is active for Eu3+ situated at sites with C5v, C2v, and C₅ᵥ symmetry, was observed as a single sharp band in each spectrum, indicating that only one type of complex is present. The spectral bands are listed in Tables S1 and S2 together with the derived Eu3+ and Tb3+ energy level data. For Eu3+, the major difference between the two spectra is the stronger relative intensity and number of resolved bands of the 5D0 → 7F4 transition for cycEu-phTb. Interestingly, the emission bands from either Eu3+ or Tb3+ situated in the more rigid cyclen binding site are sharper than when Eu3+ or Tb3+ is coordinated in the phenLn(pdtc)₃-binding site in both dinuclear complexes.

The hypersensitive transition 5D0 → 7F2 is more influenced by the coordinative environment than other forced ED transitions and serves as an indicator of the local symmetry of a coordination site33. The area ratio of the 5D0 → 7F0 to that of 5D0 → 7F1 is 3.93 for cycTb-phEu and 2.33 for cycEu-phTb. The forced electric dipole transition, 5D0 → 7F0, exhibits a weak emission intensity in the luminescence spectrum of each binuclear complex.

The 10 K emission spectrum of cycTb-phen is displayed in Fig. 3c. The near-environment of Tb3+ is the same as in cycTb-phEu (Fig. 3a); however, the (pdtc)₃Eu moiety is absent. The Tb3+-Eu3+ distance in cycTb-phEu is 10.6 ± 0.1 Å. The crystal field levels of Tb3+, as deduced in Table S1 from the Tb3+ emission spectra in Fig. 3a, c, are the same within experimental error for cycTb-phen and cycTb-phEu. For example, the splittings of the J-multiplets 7F₆ and 7F₅ are 407 ± 5 cm⁻¹ and 494 cm⁻¹, respectively, in each case. This gives an indication of the “spectroscopic vision” of Tb3+ and shows the atomic core-like nature of 4f orbitals so that the crystal field experienced by Tb3+ is effectively the same for both systems.

**Emission decay lifetimes of Tb3+ and Eu3+ and temperature dependence**

The intensity of emission was monitored as a function of time after pulsed excitation for the 5D0 (Eu3+) and 5D4 (Tb3+) states by measuring the emission decays for the 5D0 → 7F4 transition of Eu3+ and for the 5D4 → 7F₅ transition of Tb3+ because the spectral bands of these
transitions are well-separated from those of other transitions. The excitation wavelength, 355 nm, overwhelmingly excites the phen antenna, which may then transfer energy to Eu$^{3+}$ and Tb$^{3+}$. A further energy transfer from Tb$^{3+}$ to Eu$^{3+}$ is well-documented in the literature; however, the use of excitation at 490 nm produced very weak emission in the present study due to weak absorption. Thus, we were unable to directly study this energy transfer process using this excitation wavelength. Fig. 4a–d display the Eu$^{3+}$ $^5D_0$ and Tb$^{3+}$ $^5D_4$ emission decays for cycTb-phen and cycEu-phen at various temperatures, whereas Fig. 4e shows the Tb$^{3+}$ decay for cycTb-phen. An analysis of the decay curves is presented in Tables S3–S7 using mono- and bi-exponential functions and direct integration of the decay curves. Generally, the decays are not mono-exponential due to the initial faster decay, and the approach to mono-exponential behavior increases with time, as exemplified in the fits to the decay curves after ~0.2 ms. The Eu$^{3+}$ decay in cycEu-phen, as shown in Fig. 4a, comes closest to the behavior of mono-exponential decay. Taking the values measured after ~0.2 ms following the decay pulse, the lifetimes of the Eu$^{3+}$ $^5D_0$ excited state in complexes cycTb-phen and cycEu-phen decrease slightly from 0.34 to 0.25 ms and
from 0.64 to 0.58 ms, respectively, as the temperature increases from 10 to 298 K due to multiphonon relaxation and back-transfer to $^5D_1$. The $^3Tb^{3+}$-$^5D_4$ lifetimes exhibit greater changes: in cycTb-phEu, from 0.42 to 0.11 ms (from 10 K to 298 K) and in cycEu-phTb, from 0.65 to 0.13 ms (from 10 K to 200 K). The changes follow the same patterns as above but are different in magnitude upon considering the values of the “steady-state” lifetime, $\tau_{ST}$ (Tables S4 and S7). Additional processes besides multiphonon relaxation are thus involved, such as energy transfer from $^3Tb^{3+}$ to $Eu^{3+}$.

### Energy transfer from $^3Tb^{3+}$ to $Eu^{3+}$

There have been many previous literature studies that have investigated this energy transfer with varying concentrations of $^3Tb^{3+}$ and $Eu^{3+}$ (Table S9). Ideally, this energy transfer can be assessed from a comparison of the luminescence decays of cycTb-phEu and cycTb-phLa; however, since we did not synthesize the latter complex, we considered the decays of cycTb-phEu and cycTb-phen as an alternative estimation. The likely energy transfer pathways are from $^3Tb^{3+}$-$^5D_4$ (20,482 cm$^{-1}$) to $^3Tb^{3+}$-$^5D_1$ (with the lowest crystal field level at 19,048 cm$^{-1}$ as shown in Fig. S5 (literature value at ~19,100 cm$^{-1}$)) and to $^3Tb^{3+}$-$^5D_0$ (17,233 cm$^{-1}$, Table S1), with energy differences of 1434 and 3249 cm$^{-1}$, respectively. The participation of $^5D_1$ in Fig. 3a and b is not evident, particularly when compared with the spectrum of cycLa-phEu (Fig. S5), supporting the direct energy transfer from $^3Tb^{3+}$ to $Eu^{3+}$-$^5D_0$ rather than to $^5D_1$. From first order selection rules, the pathways of $^5D_4$-$^7F_6$, $^7F_5$ involve an electric quadrupole and/or forced electric dipole nonradiative transition. The transition $^7F_0$-$^5D_0$ is dipole and quadrupole forbidden to first order, although the thermally assisted $^7F_1$-$^5D_0$ transition is both forced electric dipole and electric quadrupole allowed. Both $^5D_4$-$^7F_6$ and $^7F_0$-$^5D_0$ are forbidden by the exchange selection rule, $|\text{J'}-\text{J}|=0$, 1 with $\text{J}=0 \leftrightarrow \text{J}=0$ forbidden. The transfer via the charge-transfer state $Eu^{2+}$-$^5D_0$ was not considered.

The long-term (> 0.2 ms) luminescence decay constants (i.e., reciprocal lifetimes) of cycTb-phEu ($k_{Tb\text{Eu}}$) and cycTb-phen ($k_{Tb\text{Eu}}$) are listed in Table 1 together with the difference in $k_{Tb\text{Eu}}$-$k_{Tb\text{Eu}}$, taken to indicate the $^3Tb^{3+}$-$^5D_0$ energy transfer rate. Alternative descriptions of the energy transfer rate by employing $k_{ET}$ and $k_{ET''}$ are also defined in the table. The magnitude of $k_{Tb\text{Eu}}$ does not change markedly with temperature (by 10% from 10 K to room temperature). In contrast, $k_{Tb\text{Eu}}$ considerably increases by 270% with temperature, and $k_{ET'}$ can be described by an exponential growth model (Fig. S8), clearly indicating the importance of temperature in the energy transfer process, which is understood as the thermal population and involvement of $^1F_1$. Considering only the $Eu^{3+}$-$^7F_0$ and $^1F_1$-$^7F_6$ multiplets, with the mean energy of the latter equal to 369 cm$^{-1}$ (from Table S1), and following the description in ref. 44, the energy transfer rate can be expressed as:

\[
k_{ET}(Tb \rightarrow Eu) = aN(7F_1)\times T + bN(7F_0)
\]

\[
k_{ET}(Tb \rightarrow Eu)/N(7F_0) = a \times 3 \exp(-369/kT) x T + b
\]

where $a$ and $b$ are constants; the partition function is given by $[1 + 3 \exp(-369/kT)] = P$, where $k$ is the Boltzmann constant; the average $^7F_1$ energy is 369 cm$^{-1}$; and the populations of $^7F_1$ and $^7F_0$ are $N(7F_1) = n_1 = 3 \exp(-369/kT)/P$ and $N(7F_0) = n_0 = 1/P$, respectively. Taking the results from Table 1, Fig. 5a shows a fit of the $k_{ET'}$ values from 10 to 298 K with the use of Eq. 2.

The energy transfer pathway at low temperature, therefore, involves $^7F_0$ and phonon(s) emission:

$^5D_4^{3+}Tb^{3+}(20482) + ^7F_0^{Eu^{3+}}(0) \rightarrow ^7F_6^{Eu^{3+}}(2019 - 2513) + ^5D_0^{3+}Tb^{3+}(17233) + \text{phonon}(736 - 1230)$

and/or

$^5D_4^{3+}Tb^{3+}(20482) + ^7F_0^{Eu^{3+}}(0) \rightarrow ^7F_6^{Eu^{3+}}(0 - 405) + ^5D_1^{3+}Tb^{3+}(19048) + \text{phonon}(1029 - 1434)$

where representative ranges of energies (in cm$^{-1}$) are given in parentheses; or it involves the participation of $^7F_4^{3+}$ $Tb^{3+}$ with the absorption of a low energy phonon. The energy transfer rate is higher at room temperature because the forced electric dipole transition, $^7F_1 \rightarrow ^5D_0$, is

### Table 1 Temperature variation of long-term decay constants, $k_{Tb\text{Eu}}$ and $k_{Tb\text{Eu}}$, for $^5D_4^{3+}Tb^{3+}$ emission in cycTb-phEu and cycTb-phen

| Temp (K) | Rate constant (s$^{-1}$) |
|---------|-------------------------|
|         | $k_{Tb\text{Eu}}$ | $k_{Tb}$ | $k_{ET}$ | $k_{ET'}$ | $k_{ET''}$ | $k_1$ |
| 10      | 2398           | 1555       | 843          | 943          | 35,507         | 32,062       |
| 50      | 2421           | 1545       | 876          | 1103         | 29,165         | 27,863       |
| 100     | 2740           | 1523       | 1217         | 1493         | 34,131         | 49,826       |
| 150     | 3086           | 1583       | 1503         | 2118         | 51,931         | 64,851       |
| 200     | 3751           | 1563       | 2188         | 3494         | 117,128        | 84,531       |
| 250     | 5618           | 1687       | 3931         | 5858         | 132,653        | 89,047       |
| 298     | 8811           | 1714       | 7096         | 21678        | 159,544        | 123,854      |

Temperature variation of Tb–Eu energy transfer rates deduced from the long-term lifetimes ($k_{Tb\text{Eu}}$-$k_{Tb\text{Eu}}$). Direct subtraction of decay curves ($1/r_1 = k_{ET}$, with $r_1$ from Table S8), and steady-state lifetimes ($1/r_2$ (cycTb-phEu)-1/$r_{ST}$ (cycTb-phen)) are $k_{ET}$, with $r_{ST}$ from Tables S4 and S7). The steady-state lifetime is determined by integration of the decay curves: $r_{ST} = \frac{1}{\int_0^\infty \frac{1}{r} \, dt}$, refer to the discussion above Table S3. The fitted parameter $k_1 = 1/r_2$ is from Table S7.
involved:

\[ ^{5}\text{D}_4\text{Tb}^{3+}(20482) + ^{7}\text{F}_1\text{Eu}^{3+}(300 - 442) \rightarrow ^{7}\text{F}_1\text{Tb}^{3+}
\]

\[ (3289 - 3908) + \text{Eu}^{3+} (17,233) \pm \text{phonon} \]

Alternatively, the decay curves of cycTb-phEu and cycTb-phen were directly subtracted (Fig. S9) and the resulting plots were well-fitted by a bi-exponential function (Table S8). The lifetime \( \tau_1 \) represents the reciprocal of the energy transfer rate from \( ^{5}\text{D}_4 \text{Tb}^{3+} \rightarrow ^{7}\text{F}_1 \text{Eu}^{3+} \), \( k_{ET} \) (listed in Table 1). The values are comparable with those for \( k_{ET}' \), except that the room temperature value is much higher. Figure 5b shows a plot of the experimental values of \( k_{ET} \) against inverse temperature and displays a fit using Eq. 2. Figure 5a. The fits of the subtracted curves (Fig. S9) produce a rise time, \( r_2 \), in each case, which is equal to the reciprocal of the rate constant, \( k_r \). This rate constant gives a linear plot against temperature, as shown in Fig. 5d, which is expected for the phonon occupation number of a one phonon process. Finally, the plot included in Fig. 5c uses another alternative description of the energy transfer rate, \( k_{ET}'' \), which was derived from steady-state decay lifetimes. Although the numerical values differ, the three fittings serve to confirm the importance of the participation of \( ^{7}\text{F}_1 \) in the energy transfer process at room temperature.

The energy transfer efficiency from \( ^{5}\text{D}_4 \text{Tb}^{3+} \rightarrow ^{7}\text{F}_1 \text{Eu}^{3+} \) (\( \eta_{ET} = 1 - k_{TB}/k_{TB\text{Eu}} \)) has very different values when calculated using the three alternative sets of rate constants. The value is above 0.9 and is reasonably independent of temperature when using steady-state lifetimes.

**Energy transfer from antenna to Eu\(^{3+}\) ion**

The ligand triplet state energy of cycLa-phLa was experimentally determined to be 20,124 cm\(^{-1}\) (Fig. 2), which is below the \( ^{5}\text{D}_4 \) energy (20,482 cm\(^{-1}\)) of Tb\(^{3+}\) in cycTb-phEu. Nevertheless, strong emission is observed from cycTb-phen at room temperature upon excitation into the ligand singlet state to populate the \( ^{5}\text{D}_4 \) state (Fig. S7). The \( ^{5}\text{D}_0 \) level (17,233 cm\(^{-1}\)) of Eu\(^{3+}\) lies sufficiently below the triplet state (\( \sim 3300 \text{ cm}^{-1} \)) to avoid backtransfer. In addition to triplet \( ^{7}\text{T}_1 \) - metal transfer, there are also opportunities for transfer to the lanthanide ions from higher ligand levels upon 355 nm excitation.

The calculation of singlet and triplet levels of cycTb-phEu was performed by first optimizing the structure using the Sparkle/RM1 method in MOPAC\(^{28}\). Then, the optimized geometry was used in LUMPAC with semi-empirical method\(^{30}\). Very different triplet state energies resulted when the number of states in the calculation was varied. Figure 2d displays the result with 25 states, where the lowest triplet state energy was calculated at 20,501 cm\(^{-1}\) (487.8 nm). In this calculation, there are 25 triplet states and 19 singlet states with
transition wavelengths to $S_0$ that are longer than 315 nm, with the lowest energy singlets calculated at 391, 416, and 433 nm. The schematic relative energies of the ligand Tb$^{3+}$ and Eu$^{3+}$ levels in cycTb-phEu are shown in Fig. 6 with ligand levels from the above calculation and with the metal ion levels determined experimentally or from other systems. The ability to assign definitive energy transfer pathways is therefore complex.

The 355 nm excitation into the dyad system can populate the ligand level at ~360 nm (Fig. S4), which we associate with the calculated level (Fig. 6) at 28,550 cm$^{-1}$. The gap below this singlet level is spanned by 1 phonon ($\sim 770$ cm$^{-1}$) so that internal conversion can be fast. In fact, all of the singlet-singlet gaps down to the lowest singlet state calculated at 23,084 cm$^{-1}$ (433 nm) are spanned by one phonon. The latter ligand singlet level is identified with the weak band at 437 nm in Fig. S4. From this figure, it is demonstrated that the rate of internal conversion is slower than the rate of antenna-metal ion energy transfer for these two singlet states. The energy transfer rates from not only these two states, but all of those potentially populated by 355 nm excitation, to the Eu$^{3+}$ levels were investigated using the LUMPAC program, and the results from three different optimizations and calculations are displayed in Tables S11 and S12. The calculated energies represent those from vertical Franck-Condor transitions to unrelaxed excited states. The LUMPAC program neglects intra-ligand intersystem crossing and internal conversion processes. However, it represents the most informative analysis of antenna-metal energy transfer processes currently available. It is evident from Tables S11 and S12 that the major energy transfer pathways from the ligand to Eu$^{3+}$ involve upper Eu$^{3+}$ $J$-multiplets, followed by internal nonradiative relaxation to $^5D_0$.

**Thermometric properties**

To investigate their potential application for luminescence thermometry, the temperature-dependent photophysical properties of complexes cycTb-phEu and cycEu-phTb were examined in terms of both the luminescence intensity and lifetime, at intervals between 10 and 298 K (Fig. 7). Three selected temperatures for the emission spectra of these compounds are employed in Figs. S10a, b to show the trends in emission intensity. Moreover, the changes in intensity of the Tb$^{3+}$ $^5D_4 \rightarrow ^7F_5$ and Eu$^{3+}$ $^5D_0 \rightarrow ^7F_4$ transitions clearly demonstrate the trends. For cycTb-phEu, the major decrease in intensity occurs above ~50 K for Tb$^{3+}$, and it is above ~100 K for Eu$^{3+}$, as shown in Fig. S11a, whereas the decreases begin to occur at lower temperatures for cycEu-phTb (Fig. S11b). The changes in the long-term lifetimes of the $^5D_4$ and $^7F_0$ states for these complexes are shown in Fig. 7c, d. These changes are greater for Tb$^{3+}$ than for Eu$^{3+}$.

The different temperature-dependent luminescent emission of the $^5D_4 \rightarrow ^7F_5$ transition of Tb$^{3+}$ and the $^5D_0 \rightarrow ^7F_4$ transition of Eu$^{3+}$ indicates that these dinuclear complexes are potential self-referencing thermosensitive probes that do not require any cumbersome calibration of the emission intensity. This makes a luminescent thermometer more reliable. The two target emission intensity measurements lie in clear spectral windows so that background interference is absent. The ratio of the emission intensities, $R$, follows an exponential growth curve with temperature (Fig. S12), and the natural logarithm of the ratio is plotted against temperature in Fig. 7e, f with cycEu-phTb providing the better fit. The relative
sensitivity is $S = (dR/dT)/R$, i.e., the change in ratio with temperature divided by the value of the ratio. Since, from Fig. S12b:

$$R = A \times \exp(BxT),$$

where $A = 3.96$ and $B = 0.01857K^{-1}$, the relative sensitivity is constant at 1.86% K$^{-1}$. The temperature resolution, $dT$, is given by:

$$dT = (1/S) \times (dR/R)$$

where $dR$ is the error (standard deviation) in $R$ value at temperature $T$ and is < 1 °C at $T > 200$ K (Fig. S12c).

The dyad cycEu-phTb is therefore suitable for use as a luminescent self-referencing thermo-monitor with temperature responsive luminescence and lifetimes.

**Discussion**

A strategy to synthesize hetero-dinuclear complexes that consist of chemically similar lanthanides has been introduced by which cycTb-phEu and cycEu-phTb were produced. The different energy gaps between the ligand triplet state and the acceptor lanthanide ion states, as well as the Tb$^{3+}$ to Eu$^{3+}$ energy transfer, result in different luminescence performances for each metal ion, giving an increased temperature-dependent luminescence emission ratio for europium over terbium. Both dinuclear complexes illustrated excellent temperature sensitivity over a wide temperature range, with cycEu-phTb having the best potential as an optical thermometer using an excitation wavelength of 355 nm. The temperature-dependent energy transfer between the two lanthanide ions that quenches Tb$^{3+}$ emission has been rationalized by observing the importance of the $^{7}F_{1}$ state at room temperature. LUMPAC calculations point to the energy transfer to Eu$^{3+}$ from higher ligand states, rather than from the lowest triplet state.

**Materials and methods**

All chemicals were purchased and received without further purification. NMR spectra were measured on a Bruker400 (400 Hz) magnetic resonance spectrometer with chemical shifts expressed as parts per million (ppm) and coupling constants, $J$, as Hertz (Hz). Mass spectrometry was taken on an ABI QSTAR Elite quadrupole-time-of-flight mass spectrometer using electrospray ionization (ESI) as the ion source. The HPLC measurements were conducted on an Agilent 1200 series HPLC (Column: Vision HT C18 HL 5u, length 250 mm, Serial No. 5151920 ID 4.6 mm). Fourier transform-infrared spectroscopy was performed on a PerkinElmer Paragon 1000 PC FT-IR spectrometer with KBr tablets. The emission spectra and decay lifetime measurements were recorded on a Horiba fluorescence spectrometer with a xenon lamp and a SpectreLED as the excitation sources, on an iHR550 spectrometer with a Nd$^{3+}$:YAG laser as the excitation source, and on a Mini-tau from Edinburgh Instruments.
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Author details
1Department of Chemistry, Hong Kong Baptist University, Kowloon Tong, Hong Kong SAR, People’s Republic of China. 2Institute for Biomedical Materials and Devices (IBMD), Faculty of Science, University of Technology Sydney, Sydney, NSW 2007, Australia. 3School of Mathematical and Physical Sciences, Faculty of Science, University of Technology Sydney, Sydney, NSW 2007, Australia.

Authors’ contributions
The experimental work was performed by G.B.; the manuscript was written by G.B. and P.A.T.; calculations were performed by P.A.T.; overall supervision of the project was performed by K.L.W. and D.J.

Conflict of interest
The authors declare that they have no conflict of interest.

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